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# Software Analysis for Heterogeneous Computing Architectures

A research automation framework towards more efficient  
HW/SW co-design

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I certify that except where due acknowledgement has been given, the work presented in this thesis is that of the author alone; the work has not been submitted previously, in whole or in part, to qualify for any other academic award; and the content of the thesis is the result of work which has been carried out since the official commencement date of the approved research program.

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Georgios Zacharopoulos  
Lugano, 21 October 2019

*To my parents Areti and Dimitris.  
For always being there for me.*



Men give me credit for some genius. All the genius I have lies in this; when I have a subject in hand, I study it profoundly. Day and night it is before me. My mind becomes pervaded with it. Then the effort that I have made is what people are pleased to call the fruit of genius. It is the fruit of labor and thought.

Alexander Hamilton





# Abstract

**NOTE: The abstract will be written in the end of the thesis.**

Performance increase, in terms of faster execution and higher energy efficiency, is a never-ending research endeavor and does not come for free. The breakdown of Dennard scaling, along with the seemingly inevitable end of Moore's law economic aspect, present a new challenge to computer architects striving to achieve better performance in the modern computer systems. Heterogeneous computing is emerging as one of the solutions to overcome these limitations in order to keep the performance trend rising. This is achieved by turning the focus to specialized Hardware (HW) that can accelerate the execution of a Software (SW) application or a part of that application. The goal is to design efficient HW/SW computer architectures, where a general purpose CPU is coupled with a number of specialized HW accelerators.

The choice of which parts of an application to be accelerated, though, as well as the type of accelerators to be used, while taking into account the underlying memory system, are all non-trivial research questions and depend heavily on the SW applications characteristics that are going to be accelerated. Therefore, an in-depth SW analysis can be crucial, prior to designing a heterogeneous system, as it can provide valuable information and subsequently highly benefit performance. My initial research is revolving around various ways that SW analysis, by extending the compiler frameworks and, hence, their potential, can offer this type of information and move one step closer towards optimizing and automating the design of hybrid HW/SW systems.



# Acknowledgements

This is where I acknowledge people.



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# Introduction

Performance increase, in terms of faster execution and higher energy efficiency, is a never-ending research domain and does not come for free. Living in an era where there is an immense amount of data, the demand for performance by modern computing systems rises even more. Technological giants, such as Google and Facebook, gather and compute loads of data, for instance during Machine Learning related applications and lengthy simulations. This large amount of data processing requires immense computational power and ends up in lengthier and lengthier execution time.

Moore's law [63], an observation made by the co-founder of Intel Gordon Moore, predicts that the number of transistors that can be used in the same area of an integrated circuit will double roughly every 18 months. Complimentary to that, Dennard scaling [21], also known as MOSFET scaling, states that voltage and current are proportional to the size of a transistor. Therefore, as long as the same chip area is retained, power stays constant and, at the same time, more transistors of smaller size can fit onto it. Unfortunately, this is no longer the case. The transistor size has decreased over the years, but the amount of power per transistor has, recently, stopped decreasing accordingly, resulting in current leakage, a phenomenon also known as the Breakdown of Dennard scaling [23].

The breakdown of Dennard scaling [23], along with the seemingly inevitable end of Moore's law economic aspect [68], present a new challenge to computer architects striving to achieve better performance in the modern computer systems. Heterogeneous computing is emerging as one of the solutions in order to keep the performance trend rising. This is achieved by turning the focus to specialized Hardware (HW) that can accelerate the execution of a Software (SW) application or a part of that application. Specialized HW accelerators are implemented in platforms where they can be either reprogrammable, thus allowing for a large degree of flexibility as various implementations may take place utilizing the HW resources of the platform (e.g. an FPGA board), or hardwired, such as an Application-Specific Integrated Circuit (ASIC). The first type of HW implementation sacrifices part of the potential performance achieved by allowing for flexible

designs, as the same HW resources can be reprogrammed. The latter offer no flexibility but can provide better performance in comparison to FPGAs. Under the scope of this research both HW implementations were considered.

Since the performance of a general purpose CPU is becoming limited, due to physical and technological constraints, alternative computer architectures are required. Homogeneous parallel CPUs are used in order to expose parallelism of computation in SW applications, but performance is still restricted by the parts of computation that cannot be parallelized, a fact known also as Amdahl's law. Instead of a general purpose CPU – or homogeneous parallel CPUs – managing the execution of SW applications, specialized pieces of HW, namely accelerators, can be used alongside with a general purpose CPU and execute the most demanding parts of an application in terms of computation. Consequently, the need for a powerful single CPU is no more that critical, as the execution can be offloaded to other parts of HW as well. As a result, we achieve both a more balanced execution with the use of different HW resources, and we offload the execution of specific, more demanding parts of the computation to specialized HW accelerators.

One example of a widely spread heterogeneous architecture is the addition of a GPU to a CPU on the same chip, in order to exploit the parallelism and computing power that a GPU has to offer, when it comes to image processing and 3D graphics rendering. Other examples are general purpose CPUs coupled with dedicated HW that execute specific kernels or even full applications. The latter architecture could come in a number of variations, with one or more HW accelerators, and different types of coupling, tightly or loosely [19]. The design of the first option, tightly or co-processor model, is done by using the accelerator as an Instruction Set Extension in the default pipeline of the CPU. The latter implements the connection between CPU and accelerator loosely, without any knowledge of the underlying CPU micro-architecture.

The goal of HW/SW co-design research is to design efficient heterogeneous computer architectures, so that the time latency and energy requirements are ever decreasing. The heterogeneous system that I considered during my research comprises a general purpose CPU, loosely coupled with a number of specialized HW accelerators, dedicated to the acceleration of specific parts of an application.

The choice of which parts of an application to be accelerated, though, as well as the type of accelerators to be used, while taking into account the underlying memory system, are all non-trivial research questions and depend heavily on the SW applications characteristics that are going to be accelerated. In addition to the accelerator selection problem, every HW accelerator can be synthesized with a number of optimizations embedded onto it, according to the characteristics of the task that is targeted for acceleration. For instance, in case a loop is included



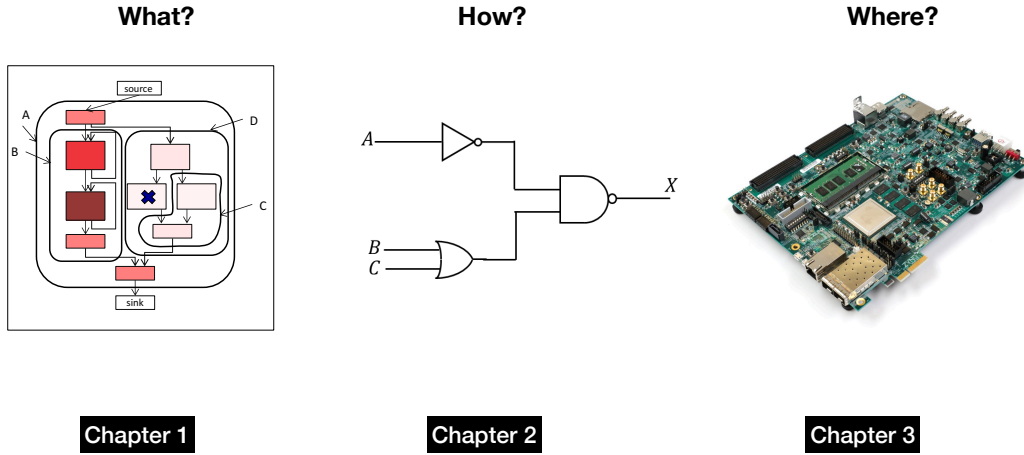


Figure 1. Overview of the research that has been conducted during my PhD and the respective chapters of the PhD thesis.

in the execution, there could be a loop unrolling factor taken into account during the synthesis of the accelerator that may dramatically affect execution time. Another example would be the addition of a memory buffer, e.g. a scratchpad memory, to reduce the memory latency of the execution. Furthermore, the underlying memory system, as in every computer architecture, can significantly affect the overall performance, due to communication latency, and should be taken into account during the selection of the accelerators to be implemented, along with their respective potential optimizations.

Therefore, an in-depth SW analysis can be crucial, prior to designing a heterogeneous system, as it can provide valuable information and subsequently highly benefit performance. Furthermore, such an analysis can be performed in short time (typically within a few seconds) and can be portable to other target applications or platforms. The research during my PhD has revolved around various ways that SW analysis, by extending the LLVM compiler framework [38] and, hence, its potential, can guide a HW engineer by making informed decisions early in the development cycle.

An overview of the research conducted during my PhD is depicted in Figure 1. This can be viewed as a map of this PhD thesis in order to navigate throughout my research time-line and present a high level view of how each piece is connected to each other.

Chapter 1 answers the question of *what* should be accelerated, namely which parts of computation, given a constraint on HW area resources. Under the scope of this chapter the RegionSeeker tool-chain is presented [87]. RegionSeeker is an

LLVM based framework that, given a SW application provided as input, identifies and selects, in a fully automatic fashion, HW accelerators under the constraint of an area (HW resources) budget. The granularity of the candidates for acceleration considered is that of a subgraph of the control flow graph of a function, with a single control input and a single control output. These candidates are called regions. After identification takes place, a selection algorithm solves the problem of finding the subset of the initial regions list that, under a given area constraint, maximizes the collective speedup obtained. The evaluation of RegionSeeker took place by using both an industrial tool, such as Xilinx Vivado HLS [79], and a research HW accelerator simulator, such as Aladdin [66]. Experiments carried out with these tools revealed an improvement of performance compared to the state-of-the-art and a speedup gain of up to 4.6x.

In Chapter 2, the analysis that is presented attempts to answer the research question of *how* the identified and selected HW accelerators should be implemented in order to achieve improved performance. Under that scope, Data Reuse analysis, during the execution of a specific domain of applications, reveals the effectiveness of private local memory structures [85]. This is achieved with the aid of compiler polyhedral analysis that detects the amount of data reuse on a specific domain of applications. The analysis provides automatically the appropriate dimensions of a memory buffer attached to the HW accelerator that would carry out the execution of the applications and minimize the communication between the accelerator and the main memory. Furthermore, for HW accelerators that contain loops, an optimal Loop Unrolling factor can be predicted for each of the included loops [86]. The most suitable Loop Unrolling factor for each loop is defined according to the target of optimization, which can be either less use of HW resources or better speedup. With the aid of a prior LLVM based analysis of the loops and Machine Learning classification, predictions can be performed on a set of loops and the respective Loop Unrolling factors may be subsequently applied during the synthesis phase of the accelerators.

Finally, Chapter 3 tackles the research question of what should be accelerated but at the same time taking into account *where* the specialized HW is hosted. An analysis of the system at hand and its memory hierarchy can affect vastly the selection of HW accelerators and subsequently the performance achieved. Latency due to data exchange between the HW accelerators and main memory could add a significant overhead to the overall computation time. In this chapter AccelSeeker, an LLVM based tool-chain, is presented. AccelSeeker performs thorough analysis of applications and estimates memory latency along with computational latency of candidates for acceleration. The granularity of the candidates for acceleration is that of a subgraph of the entire call graph of the application.

HW accelerators are selected by an algorithm that maximizes speedup, or energy efficiency under a given area budget. The evaluation of AccelSeeker took place on Zynq UltraScale platform by Xilinx, considering a demanding and complex application such as H.264. With respect to methodologies based solely on profiling information AccelSeeker attained an improved performance with an up to 2x speedup over a general purpose SW processor.

Automating the design and implementation of heterogeneous systems, while improving their performance is the broad goal of this PhD thesis. All chapters of this document attempt to provide a step closer on attaining this goal and expanding the state-of-the-art, as well as opening new paths to future work.



# Chapter 1

## Automatic Identification and Selection of Accelerators

Moving towards a heterogeneous era, HW accelerators, dedicated to a specific task, can improve both speedup of execution and energy efficiency in comparison to a general purpose CPU or a set of homogeneous CPUs. Nonetheless, the identification and selection of which parts of the computation are to be implemented in HW is a complex and demanding task. A thorough understanding of the application to be accelerated is necessary, the HW resources (area) budget is often tight and the granularity of the candidates for acceleration can dramatically affect the overall execution time. Furthermore, optimizations may be applied to a given, identified HW accelerator and this can produce multiple versions of equivalent computation instances, that in turn can result in various heterogeneous architectures with different characteristics and different performance gains. In order to address these issues I present an automated methodology that receives as input the source code of a given application and outputs a number of HW accelerators to be considered for acceleration. Among these candidates a selection takes place that maximizes collective speedup, given an area constraint. Finally, multiple versions of the same candidate can be considered during the selection phase as well.

### 1.1 Motivation

What is the rationale behind designer choices, when manually choosing application parts to be accelerated in HW, and how can those choices be replicated by an automated tool instead? Although it is possible, perhaps, that *all* of a designer's rationale cannot be replicated automatically — potentially because it requires

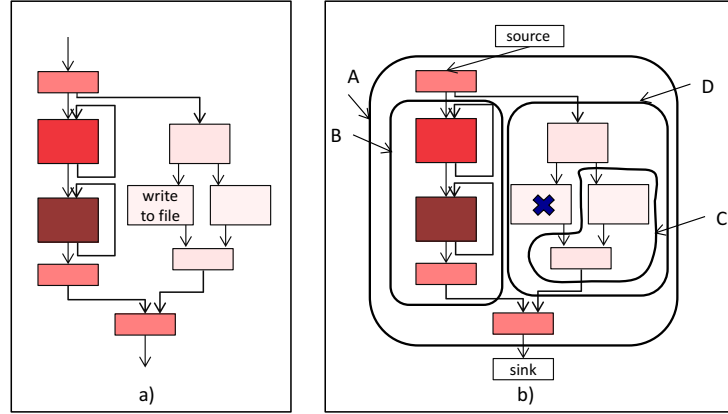


Figure 1.1. a) Example Control Flow Graph of a function, color-coded with frequency of execution (the darker the basic block, the more frequent). b) B and C are Valid Subgraphs; A and D are not Valid Subgraphs because they contain a forbidden node. B is also a CFG region, because it has a single control flow input and output.

a deep knowledge of the application at hand — it is certainly still desirable to identify at least a subset of the actions that can be automated.

Typically the designer aim will be: given an available accelerator area, extract as much as possible of the computation, under the constraint to require no more than that area, in order to maximize the resulting speedup.

Under the scope of this research I identify subgraphs of the control flow graph that have a single input control point and a single output control point, which herein will be called *regions*, as good candidates for acceleration. The rationale is that these subgraphs have a single entry point, and this corresponds to the moment of execution when the accelerator is called, and a single exit point, hence duly returning to a single location in software when the accelerator is done. Note that this type of control flow subgraph has been previously proposed and explored in compiler research — under the name of *SESE* (Single Entry Single Exit) in [2], [34], and under the name of *Simple Region* in an LLVM implementation [38] — with the aim of improving the quality of *SW code generation*, and as a scope for applying compiler optimizations and parallelization. The idea of identifying the same type of subgraph is borrowed and applied here in a novel way and to a different scenario and aim: that of automatically selecting HW accelerators.

A motivational example is provided in Figure 1.1a, which depicts the CFG of an example function, color-coded with frequency of execution (the darker the

basic block, the more frequent). A possible choice, when *manually* identifying accelerators, is to work at the granularity of functions: implement, in HW, the function most frequently executed. However, this choice might not be ideal, as the downside can be twofold: 1) a part of a function might be less frequently executed than other parts (the right side of the CFG, in the example in Figure 1.1a), therefore effectively wasting accelerator real estate. 2) a part of a function might contain non-synthesizable constructs — such as the “write to file” system call in Figure 1.1a or a function call that cannot be inlined. On the other side of the spectrum, choosing simply within the scope of single basic blocks — therefore, the body of the frequently executed loop in the picture — may not be ideal either, as the accelerator will be called once in every iteration of the loop, which may result in a large overhead. Furthermore, some speedup potential might be missed, as larger CFG regions might expose better synthesis optimizations.

CFG regions are proposed therefore as candidates for accelerators considering a granularity that can go from a single loop to an entire function, and anything in between. The main body of my research for this work is the consideration of CFG regions as candidates and a method to automatically identify and select these regions.

## 1.2 Problem Formulation

The suggested methodology identifies and investigates the performance of regions by analyzing, at the Intermediate Representation (IR) level, the Control Flow Graphs of the functions comprising a target application. A CFG represents the flow of control through a program.

**Definition: CFG.** A CFG is a Directed Cyclic Graph  $G$ , where  $V(G)$  is the set of nodes and  $E(G)$  is the set of edges. Each node in a CFG corresponds to a basic block in a function, and each edge to the control flow within that function.

A source node is added, connected only to the entry basic block of the function, and a sink node, connected only to the exit of the function. Figure 1.1b shows an example of CFG. A node in the CFG is marked as forbidden if it corresponds to a basic block containing instructions that cannot be synthesized in HW — for example operating system calls.

**Definition: Valid Subgraph.** A Valid Subgraph is any subgraph of the CFG that does not contain a forbidden node. In Figure 1.1b: B, and C are Valid Subgraphs; A and D are not Valid Subgraphs because they contain a forbidden node.

**Definition: Region.** A region  $R$  of a CFG  $G$  is a Valid Subgraph such that there exists a *single* edge going from a node in  $V(G) \setminus V(R)$  to a node in  $V(R)$  and a

single edge going from a node in  $V(R)$  to a node in  $V(G) \setminus V(R)$ . In Figure 1.1b: B is a region, while C is not.

Under the scope of this chapter, all and only regions are considered as candidates for identification of accelerators. Given a merit  $M()$  and cost  $C()$  function for each region we can formulate the problem of selecting accelerators as follows:

**Problem: Region Selection**

Let  $\mathcal{R} = \{R_1, R_2, \dots, R_n\}$  be a set of regions, with associated cost and merit functions  $C$  and  $M$ . For any subset  $X \subseteq \{1, 2, \dots, n\}$  of regions, we denote by  $M(X) = \sum_{i \in X} M(R_i)$  the sum of the merits of its regions, and we denote by  $C(X) = \sum_{i \in X} C(R_i)$  the sum of the costs of its regions.

We want to select a subset  $X$  of regions such that

1. No two regions belonging to the same CFG overlap, i.e.,  $V(R_i) \cap V(R_j) = \emptyset$ , for all  $1 \leq i, j \leq n$
2. The cost  $C(X)$  is within a user-given cost budget  $C_{\max}$
3. The merit  $M(X)$  is maximized

This problem definition maps to what we have identified in the previous section as the designer aim: given an available accelerator area, extract as much as possible of the computation, under the constraint to require no more than that area, in order to maximize the resulting speedup.

### 1.3 Region Selection Algorithms

The *Region Selection* Problem requires a previously identified set of regions as input. To identify regions, and hence gather such set, an existing LLVM pass is reused, which in turn is based on an algorithm of linear time complexity published in [34]. Then, given the available set of regions, the more computationally expensive *Region Selection* Problem must be tackled, and algorithms to solve it are explained in the following.

Firstly an exponential, exact branch-and-bound method based on a binary-tree search is provided; secondly, a fast (polynomial) non-exact, greedy method; thirdly, a meet-in-the-middle approach, still exponential but scaling faster than exact, that we call exact-on-cropped.

Before delving into the algorithm explanation, a running example is provided in Figure 1.2. The Figure depicts the CFGs of two functions, and highlights five regions identified within them (labeled A,B,C,D,E in the picture).





Figure 1.2. a) Running example, used to explain the selection algorithms: the CFGs of two functions are depicted, with five regions identified in them — labelled from A to E. b) The overlap graph for the five regions.

In the following, we denote by  $O$  the set of region overlaps, i.e., the set

$$\{(i, j) \mid V(R_i) \cap V(R_j) \neq \emptyset\}.$$

The edges of the graph represented in Figure 1.2b correspond to the set  $O$  of region overlaps, for the running example.

### 1.3.1 Exact Method

In the exact method the Problem *Region Selection* is reduced to the *independent set problem*. In particular, we construct an undirected graph  $G$  where  $V(G) = \{1, 2, \dots, n\}$ , i.e., there is one node for each region, and  $E(G) = O$ , i.e., two nodes are connected if the corresponding regions overlap. It is easy to see that a set  $\{i_1, i_2, \dots, i_r\}$  satisfies condition 1 of the *Region Selection* Problem if it is an independent set of  $G$ . Figure 1.2b shows the overlap graph  $G$  corresponding to the running example: region A overlaps with all regions but E, hence edges are added linking A with B, C and D, and so on. Examples of independent sets in this graph are  $\{A\}$ ,  $\{B, D\}$ ,  $\{B, C, E\}$ .

The algorithm recursively explores the independent sets of  $G$ , similarly to the Bron-Kerbosch algorithm [10], and its steps will be followed with the aid of the running example, and with its corresponding tree exploration, shown in Figure 1.3. The algorithm maintains a set  $X$ , which is the active independent set (initialized to  $\emptyset$ ) and a set  $P$  of available nodes (initialized to  $V(G)$ ). At each iteration, the algorithm chooses a node  $u$  in  $P$  such that  $C(X \cup \{u\}) \leq C_{\max}$ , i.e.,

it satisfies condition 2 of the *Region Selection Problem*, and recursively explores the configurations

1.  $X' = X \cup \{u\}$ ,  $P' = P \setminus (\{u\} \cup N(u))$
2.  $X' = X$ ,  $P' = P \setminus \{u\}$

where  $N(u) = \{v \mid (u, v) \in E(G)\}$  is the set of neighbours of  $u$  in the overlap graph. Configuration 1 traverses all the independent sets that contain  $X$  and  $u$ . The choice of  $P'$  maintains this invariant, as all the neighbours of  $u$  are removed from  $P$ . Instead, configuration 2 traverses all the independent sets that contain  $X$  but not  $u$ . Note that any independent set is visited *once only*.

This process can be exemplified through Figure 1.3: the root of the tree represents the empty set, and set  $P$  at this point contains all regions. Then, inclusion of region  $A$  is first explored, and the set  $P$  is updated by removing all regions overlapping with  $A$ :  $P = \{E\}$ . According to the merit and costs of all regions in this example, shown in the table within the picture, the merit (60) and cost (35) of the solution currently explored is also updated.

At every point of the exploration, a new node  $u$  is considered for addition in the current independent set. If there is no node  $u$  satisfying condition 2 of the *Region Selection Problem*, the algorithm records the set  $X$  and backtracks, as  $X$  is maximal with respect to condition 2. For the running example in Figure 1.3, the cost budget  $C_{\max}$  is equal to 35. Hence, exploration stops at  $X = \{A\}$  because the cost budget has been reached, and backtracks. The next region chosen is  $B$ , sets  $X$  and  $P$  are again updated accordingly, to  $X = \{B\}$  and  $P = \{C, D, E\}$ , and exploration continues.

*Optimization 1:* To speed up the search, the algorithm maintains the maximum merit  $M_{\max}$  of the independent sets explored so far. In this way, if  $M(X \cup P) < M_{\max}$  the algorithm can backtrack, as no superset of  $X$  has a merit larger than the maximum one found so far. This optimization can be seen at work, among others, in the tree-node where  $X = \{B\}$  and  $P = \{D, E\}$ . In fact,  $M_{\max}$  is 75 at that point in the exploration (it was reached by set  $X = \{B, C, E\}$ ), while the current merit  $M(\{B\})$  is 30, and the remaining potential gain of  $P = \{D, E\}$  is 40.  $M_{\max}$  cannot be reached, and the algorithm can backtrack.

*Optimization 2:* To make the above exact pruning strategy effective, the algorithm adopts the strategy of choosing the node  $u$  with maximum merit among the ones which satisfy condition 2 of the *Region Selection Problem*. In practice, this means that candidate regions are considered in order of decreasing merit. In Subsection 1.5.2, it is shown that these two optimizations greatly increase the scalability of the exact method.

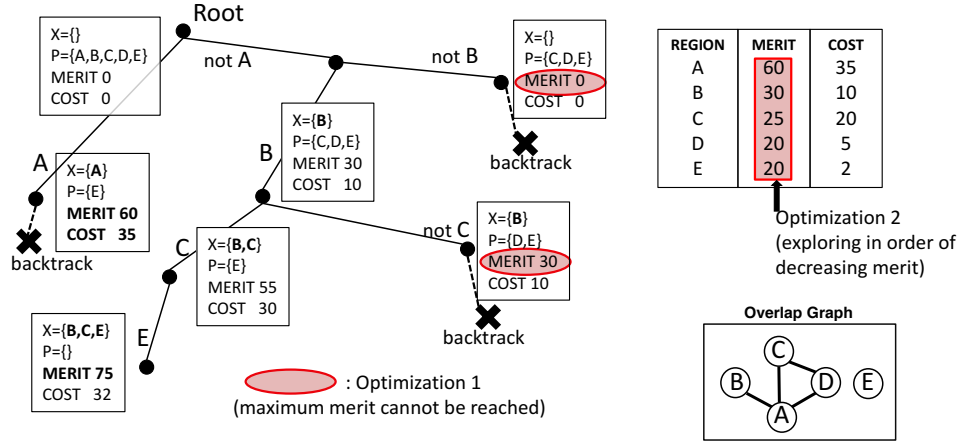


Figure 1.3. Tree exploration performed by **exact**, for the running example of Figure 1.2, and for a cost budget of 35.

At the end of the exploration, the algorithm reports the set(s) recorded with merit equal to  $M_{\max}$ , i.e., satisfying condition 3 of the *Region Selection Problem*. In the running example, this corresponds to set  $X = \{B, C, E\}$ .

### 1.3.2 Greedy Method

The algorithm implementing a greedy selection maintains a set  $X$  (initialized to  $\emptyset$ ), which is the current partial solution, and a set  $P$  of available regions (initialized to  $\{1, 2, \dots, n\}$ ). At each iteration, the algorithm selects the region  $u$  in  $P$  with largest merit such that  $C(X \cup \{u\}) \leq C_{\max}$ , and continues to the next iteration with  $X' = X \cup \{u\}$  and  $P' = P \setminus (\{u\} \cup \{v \mid (u, v) \in O\})$ . The choice of  $P'$  guarantees that the set  $X$  satisfies condition 1 in each iteration. If there is no region  $u$  satisfying condition 2 the algorithm terminates and reports  $X$ . In the running example of Figure 1.3, this corresponds to simply stopping exploration at set  $X = \{A\}$ .

Since the greedy method never backtracks, it is often trapped in local minima, and therefore cannot guarantee optimality. On the other hand, it converges to a solution very fast, and is used as a naive solution for generating comparative results (Subsection 1.5.2).

### 1.3.3 Exact-on-cropped Method

The previous two algorithms represent two ends of the spectrum: exact and exponential on one side; non-exact, fast and naive on the other. A third solution,

which strikes a balance between them, comes from the observation that while the list of regions identified in an application is long — potentially too long to be processed exactly — the list of *meaningful* regions is short, where by meaningful is mean contributing tangibly to the overall speedup. In other words, the distribution of regions with respect to speedup provided is very skewed.

The third algorithm alternative is therefore to apply the *exact* algorithm (Section 1.3.1), but only to a *cropped* list of regions in input. This in practice corresponds to ignoring a number of low-speedup regions in the selection problem. In Subsection 1.5.2 it is showed that such approach, while of course still of exponential complexity, can greatly improve the scalability of the exact algorithm, still retrieving high-quality solutions.

## 1.4 The RegionSeeker Framework

The RegionSeeker framework is an automated methodology that identifies candidates for HW acceleration from application source code. An extensive SW analysis, based on the LLVM compiler infrastructure, performs, apart from the identification, an estimation of the performance gain (merit), along with the HW resources (cost), of each candidate. Subsequently given a HW resources constraint, a selection of the identified HW accelerators takes place that maximizes the cumulative performance gain, as detailed in Section 1.3. First the LLVM toolchain built for this purpose is analyzed, then the employed platform model and the benchmarks used for a comparative evaluation are detailed.

### 1.4.1 LLVM Toolchain

The analysis passes of RegionSeeker were built within the latest version (3.8) of the *LLVM Compiler and Toolchain* [38]. The LLVM infrastructure provided the compiler ground in order to develop my own analysis passes, as well as the tools used for profiling. *Region Identification* pass, as depicted in Algorithm 1, was developed to identify and provide an initial estimated evaluation to the identified regions. The pass receives as input applications developed in C or C++ and performs the analysis in the Intermediate Representation (IR) level.

*Region Identification* pass iterates over every function of the provided input applications and, using the existing *RegionInfo* LLVM pass [74], identifies regions within every function. Subsequently, forbidden nodes within regions are identified and labeled, such as system calls or calls to functions that are not inlined. The regions containing these nodes are marked as invalid. Conversely, the valid

---

**Algorithm 1** LLVM Analysis Pass - Region Identification

---

**Input:** Application written in C/C++**Output:** List of Identified and Profiled Regions

```

1: function RunOnFunction()
2:   Region_List = NULL
3:   RI = getRegionInfoAnalysis()
4:   for Region in Function do
5:     if RegionIsValid() then
6:       EvaluateRegion(Region)
7:       Region_List.Add(Region)
8:   return Region_List
9:
10:  /* Estimate Merit for Region */
11:  function EvaluateRegion(Region)
12:    for Basic Block in Region do
13:      getProfilingInfo(Basic Block)

```

---

regions are evaluated by a profiling via instrumentation routine. Profiling via instrumentation requires generating an instrumented version of the code, which gives more detailed results than a sampling profiler. Using this information, the basic blocks are annotated in each function with their respective execution frequency, along with the aid of *ClrFreqCFGPrinter* LLVM pass [88].

By exploiting the execution frequency of every basic block, the respective evaluation for every region is computed, which is being associated to the region as estimated merit, and the region is being added to the list of identified regions. The final output of our analysis pass is a list of valid regions, or else accelerator candidates, with an estimated merit attached to them.

The region list output is saved in a file, which is in turn processed by the selection algorithms exact, greedy and exact-on-cropped, which are implemented as standalone programs in C++.

#### 1.4.2 Platform Model and Performance Metrics

The performance benefit achievable by application-specific acceleration is dependent on multiple target-specific parameters, including the adopted memory hierarchy, the employed bus protocol, the interconnect strategy and the number of considered processors, accelerators and memories.

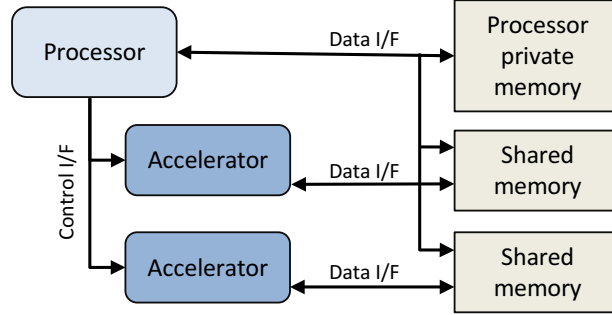


Figure 1.4. Target ASIP model, featuring a host processor interfacing special-function accelerators through control interfaces and shared data memories.

In order to assess the performance of RegionSeeker, a system comprising a single processor and multiple accelerators was assumed, exchanging shared data with scratchpad memories (Figure 1.4). The processor activates the accelerators via a memory-mapped interface, thus requiring a transaction on the system bus. When activated, accelerators read and write data to and from the scratchpads, computing their outputs, which can then be accessed by the processor. In this way, no data movement surrounding the execution of accelerations is required. Accelerators are interfaced to scratchpad memories with ports having a latency of one clock cycle. The control interface between the processor and the accelerators has a latency of 10 clock cycles. These parameters correspond to the ones employed by the `ap_memory` and `s_axilite` interfaces, respectively, provided by Xilinx Vivado.

The run-times of the non-accelerated part of the considered benchmarks are measured using the Gem5 simulator [8], modeling an ARMv8-A processor with an issue width of 1. The processor model is atomic, with in-order execution. It is interfaced with separate instruction and data memories with an access latency of one clock cycle.

Hardware execution times are retrieved using *two different HLS frameworks*: the Aladdin simulator and the Xilinx Vivado HLS commercial tool-suite. Aladdin targets ASIC implementations. It allows a fast evaluation, but does not produce a synthesizable netlist as output; nonetheless, the estimations offered by this tool are within 1% of the ones derived from an RTL implementation [66]. Hardware instances generated with Vivado HLS are instead intended for FPGA designs. Synthesis-runs within this framework are more time-consuming, but provide exact cost (HW resources) and merit (speedup) figures of each accelerator, as well as a direct path to its realization.

The cost  $C()$  of regions (and, for comparison, basic blocks and functions) was computed as the amount of required resources. We expressed them in terms of IC area in the case of Aladdin ( $mM^2$ ), and the maximum between their required flip-flops and look-up tables, on a Virtex7 FPGA, in the case of Vivado. The merit  $M()$  of a region was set as the difference between its hardware and software run time, across all its invocations in an application.

### 1.4.3 Benchmarks

Real-world applications of varying size from the CHStone embedded applications benchmark suite [33] were considered. `adpcm` performs an encoding routine, whereas `sha` is a secure hash encryption algorithm, widely used for generating digital signatures and the exchange of cryptographic keys. `aes` is a symmetric-key encryption algorithm. `gsm` performs a linear predictive coding analysis, used for mobile communication. `dfmul` and `dfsine` are smaller kernels that perform double-precision floating-point multiplication and sine functions employing integer arithmetics. `jpeg` and `mpeg2` are larger applications, implementing JPEG and MPEG-2 compression, respectively.

## 1.5 Experimental Results

This section investigates and quantitatively assesses the results and contributions of RegionSeeker from multiple perspectives. First, the speedup deriving from considering regions as targets for acceleration is evaluated, with respect to state-of-the-art solutions based on functions and basic blocks. Then, the performance of the algorithms proposed to solve the *Region Selection* Problem is analyzed. Finally, the robustness of region-based acceleration when varying architectural-specific parameters is explored.

### 1.5.1 Regions as a Choice for Accelerators

In order to evaluate the benefits of RegionSeeker, we comparatively assess it against two state-of-the-art alternatives. The first is to identify accelerators *automatically*, but only within the scope of data flow — which means within the scope of single basic blocks — as done by state-of-the-art approaches such as [84], [56], [16], [61], and [26] to name only a few. In particular, the state-of-the-art algorithm proposed in [76] and used in [54] and in [26] was implemented, that identifies maximum convex subgraphs within basic blocks. These



Figure 1.5. Comparison of speedups obtained on eight CHStone benchmarks by selecting regions, only basic blocks and only functions, varying the area constraint, using Aladdin and Gem5 for merit and cost evaluation.





Figure 1.6. Comparison of speedups obtained on eight CHStone benchmarks by selecting regions, only basic blocks and only functions, varying the area constraint, using Vivado\_HLS and Gem5 for merit and cost evaluation.

methods identify the largest part that can be synthesized and accelerated within a basic block, and hence represent an upper-bound on the speedup that can be achieved by identification methods that work at the data-flow (basic block) level. The second is to mimic the *manual* approach of selecting entire functions, which is also the scope supported by high-level synthesis tools [14] [77] [79].

In the experiments, RegionSeeker with the exact -on- cropped selection method was used, discarding regions that provide less than 10% of the maximum merit. Two sets of experiments were performed: first Aladdin, and then Vivado HLS were used to estimate merit and cost, highlighting that the RegionSeeker methodology can be used across different high-level synthesis tools, and more importantly verifying that the regions selected are largely the same, independently of the cost and merit estimation model used.

Figure 1.5 showcases the achieved speedup, when employing Aladdin, by the accelerators selected by RegionSeeker (labeled regions in the figure), with respect to the entire run-time of the applications and for different area constraints. For small-to-medium size applications such as *adpcm*, *aes*, *gsm* and *sha* speedup gains for RegionSeeker vary from 1.6x up to 3.2x. For smaller kernels, larger variations can be observed, as for *dfmul* and *dfsin* the speedup reaches 1.12x and 3.9x respectively. Finally, for larger benchmarks such as *jpeg* and *mpeg2* speedup is fairly significant: 2.5x for the former and up to 4.3x for the latter can be reached using RegionSeeker.

Similar trends are observed when Vivado HLS is instead used for the accelerator synthesis, as reported in Figure 1.6: RegionSeeker consistently outperforms state-of-the-art approaches which target either single basic blocks or entire functions, across all benchmarks. These results highlight that the achievable speedups are highly influenced by which segments of applications are selected for accelerations, and that such choice is only marginally influenced by the adopted merit and cost estimation tool. In fact, it was verified that across the two sets of experiments, the regions chosen were *the same* in 80% of the cases. As an example, out of 10 regions selected to achieve a 2.2x speedup for the *jpeg* benchmark, 8 are the same when using either Aladdin or Vivado HLS for merit and cost estimation, and the ones that differ contribute to less than 14% of the provided gain.

The speedup that can be obtained by accelerating basic blocks is hampered by their small granularity and, consequently, the high number of switches between software and hardware execution. Moreover, in this setting many optimization opportunities during the hardware implementation of the accelerators are missed, because they only arise when control flow is considered, as is instead the case for regions. On the other hand, the speedup derived by selecting whole

functions trails the one corresponding to regions, because of two reasons. First, function selection is limited to the ones which do not present forbidden nodes, and this might rule out promising regions within them. Second and more importantly, it is also inflexible from an area viewpoint, which is especially visible when few hardware resources are available for acceleration. In those cases, the selection of functions often detects only few feasible candidates, with a small merit (e.g., in jpeg and mpeg2, for an area of less than  $0.5 \text{ mM}^2$ ).

This limitation is not present for regions, as only the part pertaining to individual hotspots inside a function can be selected. Indeed, the performance of RegionSeeker stems from the high flexibility of the selection approach, as it allows the consideration of the entire spectrum of granularity ranging from whole functions to single loops, ultimately enabling a better exploitation of speedup for a given area budget.

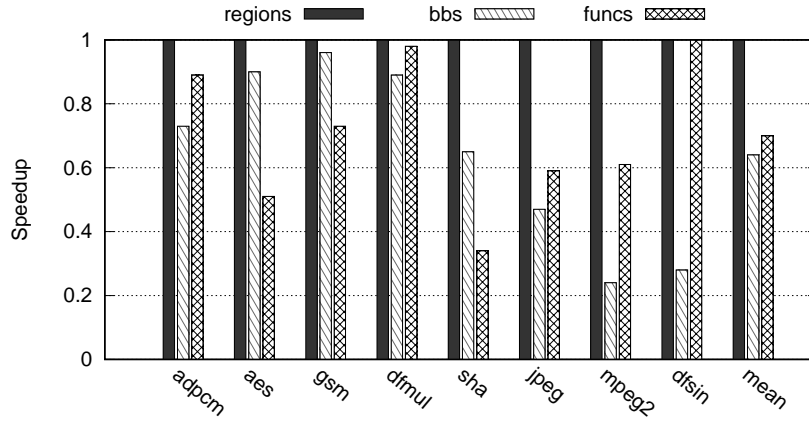


Figure 1.7. Normalized Speedup of RegionSeeker with respect to function and basic block selection, considering, for each benchmark, a fixed area constraint. Synthesis performed with Vivado HLS.

A summary of the performed experimental exploration is presented in Figure 1.7. It reports the normalized speedups obtained by RegionSeeker compared to basic block and function identification, when the maximum considered area budget and Vivado\_HLS are employed. The rightmost column set illustrates that, on average, RegionSeeker harnesses approximately 30% higher speedups with respect to the two baseline methods. Moreover, while in some cases the baselines match the performance of RegionSeeker (e.g.: gsm for basic blocks, dfsin for functions), neither of them can do that consistently across different area constraints and across applications, showcasing the suitability of control-flow regions as accelerator candidates.

### 1.5.2 Performance of Selection Algorithms

In Section 1.3, three selection algorithms were presented: an exact algorithm that might not scale for large benchmarks, a naive greedy, and a meet-in-the-middle approach where the exact algorithm is applied only to a cropped list of regions, as opposed to all regions of a benchmark. In this section, the performance of these algorithms is evaluated, in terms of scalability and of goodness of the solution found.

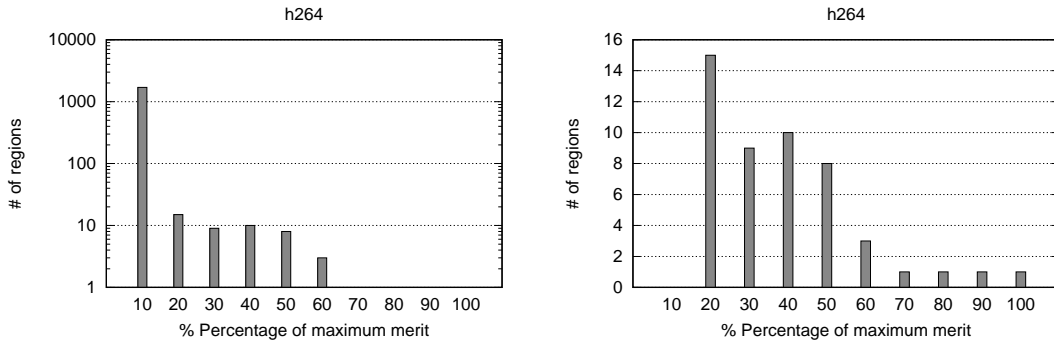


Figure 1.8. Left: The 1745 regions of H.264 are partitioned here in ten bins, according to their merit  $M()$ . Regions providing up to 10% of the maximum merit fall in the first bin, from 10% to 20% in the second bin etc. Notice that the distribution is extremely skewed. Right: The nine most profitable bins are here shown in linear scale, for clarity. The skewness of this distribution is leveraged by the **exact-on-cropped** selection algorithm.

To this end, a complex benchmark is targeted (namely, H.264 [70]), which has a code size of more than ten thousand lines of code and contains thousands of regions. Note that, while Aladdin provides merit and cost estimation for a region very quickly (a matter of milliseconds per region) and therefore could have been employed for this experiment of algorithm scalability, it does, however, require the outlining of the selected accelerator specifically *as a function call* within the application source code — and this currently needs to be done manually. This technical limitation means that Aladdin estimations cannot be used for the scalability experiments in this section. Hence, a more abstract model was employed, which could be implemented directly within the LLVM toolchain, relying on the LLVM intermediate representation and without needing manual intervention on benchmark source code. The cost of a region was estimated as the area required to implement its DFG nodes, and its merit as the cycles saved between SW and



Figure 1.9. Left: Speedup achieved on the H.264 benchmark by **exact-on-cropped**, cropped by considering only regions providing at least 5%, 10% and 20% of the maximum merit, and by **greedy**. Right: Corresponding algorithms execution time.

HW execution, where the latter is the delay of the nodes on the DFG critical paths, each multiplied by their respective frequency.

Firstly, in Figure 1.8 it is shown, for the H.264 benchmark, the distribution of all regions with respect to their merit  $M()$ . It becomes apparent in this figure that the distribution is extremely skewed, with very few regions having high merit and the majority providing a negligible one. This is to be expected, as it is well known that a large percentage of time, in running a software application, is typically spent on a small percentage of code. And of course a region merit is proportional to the frequency of execution of its corresponding code segment.

For a large benchmark such as H.264, the exact algorithm does not scale if it is fed all 1745 regions. However, it is reasonable to expect that, given the distribution seen, the goodness of the solution found should decrease only slightly when discarding a large number of low-potential regions, while scalability could grow tangibly. This assumption is confirmed by the data in Figure 1.9, where the comparison of performance and scalability of the exact-on-cropped selection algorithm is performed, at three different levels of cropping — that is, considering regions providing at least 5%, 10% and 20% of the maximum merit, respectively.

Figure 1.9 (left) plots the estimated speedup achieved by implementing a set of regions in hardware, selected by the three exact-on-cropped levels, and by greedy, for different area constraints. It can be seen that, when changing the level of cropping, the goodness of the solutions found by exact-on-cropped differs only slightly (as expected, little is lost when some low-potential regions



Figure 1.10. Execution time of the **exact-on-cropped** algorithm (cropping level at 10%), when the two optimizations described in subsection 1.3.1 are removed.

are ignored) and it is altogether largely superior to fast but naive greedy.

However, as a second issue worth observing, the exploration space for the three different levels of cropping differs greatly, and hence the time spent by each of these algorithms to terminate. This can be seen in Figure 1.9 (right) : orders of magnitude separate the time spent by each, with **exact-on-cropped** at 5% taking hours, and at 10% taking seconds.

Last, the effect of the optimizations that were devised for improving search tree exploration is shown. These optimizations were described in Section 1.3.1 and exemplified in Figure 1.3. They are 1: pruning the exploration tree when a certain best merit, found so far, cannot be reached, and 2: processing regions in order of decreasing merit. In Figure 1.10 it can be seen how the two optimizations affect the algorithm run time. When pruning is turned off, more than three orders of magnitude are lost, in time. When the list of regions is processed in an order different than that of decreasing merit (in this experiment, an order of decreasing *density* is considered, i.e. *merit divided by cost*) more than two orders of magnitude are lost.

### 1.5.3 Impact of the Interface Overhead

The initiation of an accelerated routine on a dedicated hardware block always entails a timing penalty  $T_{\text{Overhead}}$ . Such overhead is highly dependent on the interface protocol between the processor and the application-specific accelerators. While the definition of such protocol is outside the scope of this work, the

impact of adopting different values for this parameter is worthy to investigate, when different selection methods are employed.

Two observations can be made by analyzing the results of Figure 1.11, reported for the sha benchmark. Firstly, the speedup obtained by RegionSeeker is not affected in any significant way by the variation of the value of  $T_{Overhead}$  among one, ten and twenty cycles, while the speedup for basic blocks is indeed affected. This is to be expected, since basic block level accelerators require a higher number of invocations (e.g.: for each iteration of an intensive loop) than region-level accelerators. Secondly, while by decreasing the value of  $T_{Overhead}$  the speedup of basic block increases, it does not increase in a significant way and is still less tangible than the speedup achieved by RegionSeeker.



Figure 1.11. Impact of the initiation overhead for regions and basic blocks selection strategies, considering  $T_{Overhead}$  values of 1, 10, and 20 clock cycles.

## 1.6 RegionSeeker MuLTiVersioning

High Level Synthesis (HLS) tools, such as Vivado HLS by Xilinx, may employ optimizations to HW accelerators design in order to increase performance, i.e. obtain faster execution. These HLS optimizations were not taken into account by RegionSeeker framework in the previous section. Default, non optimized versions of HW accelerators were identified and selected instead. In this section an extended RegionSeeker framework is presented, which performs the selection not only among possible CFG subgraphs, but also among different versions of each identified subgraph, namely different versions of the regions identified. This extension is referred to in the rest of the document as RegionSeeker: the MuLTiVersioning approach.

### 1.6.1 Methodology

The rationale, supporting the extension of RegionSeeker framework, is to achieve improved speedup that can be provided by exploiting a more varied set of HW accelerators to select from, with different optimizations implemented onto them. This is being achieved by instantiating different versions of each HW accelerator with the same functionality, yet different speedup gains and different area (HW resources) requirements. The set of optimizations that were considered in order to design different HW implementations of the same accelerators are:

1. The Loop Unrolling (LU) factor, in accelerators that contain loops.
2. The loop pipelining option, being either on or off.
3. The array partition factor, which is the number of input and output ports of the memory buffer (scratchpad) attached to the accelerator.

Loop unrolling optimization is an HLS directive that, in the context of High Level Synthesis instantiates multiple copies of the logic implementing the functionality defined in a loop body, drastically impacting the performance of HW accelerators [37] [36]. This directive can be applied in HW accelerators containing loops whose trip count can be statically defined. It should nonetheless be applied in a careful manner, as it entails a high area cost for the duplicated logic. Furthermore, the resulting benefits can be hampered by loop-carried dependencies and frequent memory accesses.

Loop pipelining is an additional HLS directive applied in loops that allows the pipelining of the operations contained in a single body of a loop and across consecutive iterations. Restrictions regarding loop-carried dependencies across con-



secutive iterations can limit the application of the loop pipelining optimization as the result of the output of a loop iteration would be required in the following one, thus not allowing the pipelining of the loop body operations.

Given an initial set of HW accelerators, i.e., a set of regions that is derived by the RegionSeeker framework, multiple versions for each region can be generated that maintain the same functionality. Each version may employ one of the optimizations listed above, or a combination of them.

All versions of the HW accelerators were evaluated by the Aladdin HW accelerator simulator. Aladdin targets ASIC implementations. It provides a fast evaluation, but does not generate a synthesizable netlist, as opposed to Vivado HLS. Nonetheless, the estimations provided are within 1% of the ones derived from a Register-transfer level (RTL) implementation, according to the developers of Aladdin [66]. For all simulated versions of the selected regions (or HW accelerators), the number of Cycles and number of Functional Units (FU) Area were retrieved. For the SW execution time the gem5 simulator [8] was used with two CPU settings: a) TimingCPU (a simple and slow CPU with only two pipeline stages) and b) O3CPU (a complex and fast CPU with five pipeline stages and other resources such as a branch predictor, reorder buffer etc).

The exact selection algorithm, as detailed in Subsection 1.3.1, was used subsequently to perform the subset selection that maximizes speedup, given an initial set of HW accelerators along with their respective versions, as well as a specific area (HW resources) budget. An important note is that no more than one version of each candidate can be selected, as only one realization of the respective SW execution is required. To ensure that, each version is marked with the same set of basic block indexes and the selection takes place by considering the overlapping graph presented in Figure 1.3. As a result multiple versions of the same region cannot be selected during the selection phase as this would violate the first condition of the *Region Selection* problem as defined in Subsection 1.2.

### 1.6.2 Experimental Results

The experimental setup was the same as in the RegionSeeker framework, with a system comprising a single SW processor and multiple loosely coupled HW accelerators, exchanging shared data with private local memories. The processor invokes the accelerators via a memory-mapped interface, thus requiring a transaction on the system bus and as soon as the HW accelerators execution is complete, control returns to the SW processor.

Experiments were run in jpeg benchmark from the CHStone embedded applications benchmark suite. Four different approaches were compared: a) the

*min* where the regions with the least amount of area are included in the set and hence can be selected, b) the *base* where the regions with median values of area are selected, c) the *max* where only the maximum area regions can be selected and finally d) the MuLTiVersioning approach where any possible version of the regions can be selected. The speedup achieved on jpeg over the whole run time of the application (Figure 1.12 Top), as well as over solely the SW time of the selected regions Figure 1.12 Bottom) is showcased.

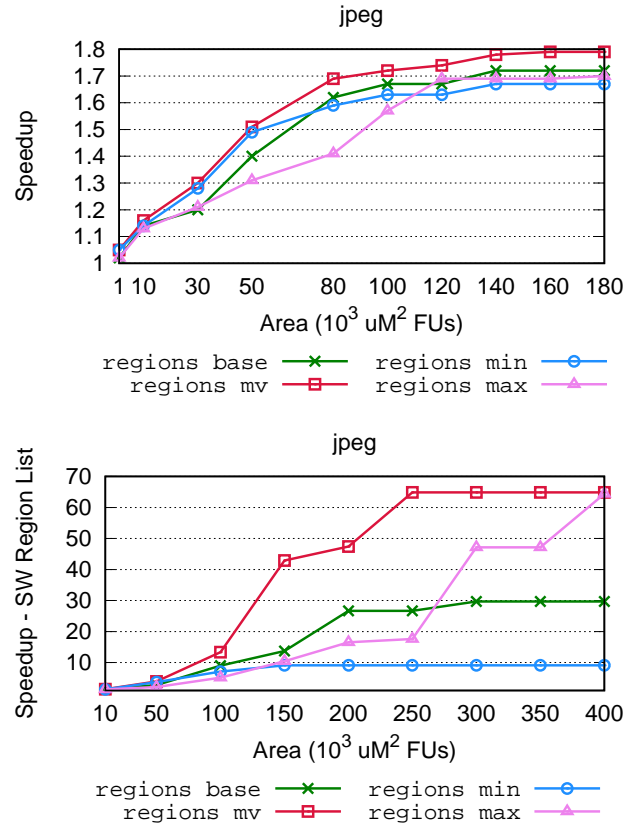


Figure 1.12. Comparison of the speedup obtained on jpeg benchmark, over the Top: Total run time of the application and Bottom: SW time of only the equivalent kernels (regions), varying the area constraint, using Aladdin and gem5 for Speedup and Area evaluation. Four approaches are compared: The min where the regions with least amount of area are selected, the base where the regions with median values of area are selected, the max where only the maximum area regions can be selected and finally the MuLTiVersioning approach where any version of the regions can be selected.

The strength of the MuLTiVersioning approach and the benefit of having a

variety of potential candidates to select from is demonstrated by the experimental outcome of the jpeg application (kernels run-time). In Figure 1.12 for any given area point, the speedup obtained is higher than any other methodology. For a medium area point ( $200 * 10^3 uM^2$ ), the speedup achieved with MuLTiVersioning is 1.7x more than the second best, *base* approach. For a large area constraint ( $400 * 10^3 uM^2$ ) the MuLTiVersioning speedup is more than 2x compared to *base* and more than 6x compared to *min*.

## 1.7 Related Work

Automatically identifying parts of computation to be accelerated is often called, in literature, Instruction Set Extension identification, or also HW/SW Partitioning. The distinction that is most relevant, for this research work, is the *scope* at which the suggested techniques perform identification: identifying accelerators or custom instructions at the data flow or the control flow level.

*Data Flow Level.* State-of-the-art methods have been published in literature to automatically identify, *within a single basic block*, the subgraph of data flow that maximize speedup when implemented in HW as a custom instruction according to varying architectural constraints. A non-extensive list include works [84], [56], [16], [61], [26] and [43], where the problem of identifying subgraphs under convexity, I/O constraint, and/or area is tackled; in [76] and [54] the I/O constraint is relaxed, to be regained via I/O serialization [57], [76], [6], [3]. In [17] the focus of the identification process is also on DFG nodes within single basic blocks, and the constraints that are taken into account are a limited number of read and write ports, and area. The methodology proposed in [25] is not limited by I/O in the selection process, but clusters MAXMISOs [4] in order to form MIMOs (Multiple Input Multiple Output instructions) that can be executed as a single instruction.

In none of the above pieces of research, though, the inclusion of the control flow of the application is considered during the identification process. The technique proposed in this chapter, instead, pushes identification *beyond* the basic block level and identifies entire regions of the Control Flow Graph of the application as candidates for acceleration. Compiler transformations such as if-conversion and loop-unrolling can be, and are, used by several of the techniques mentioned above in order to enlarge the scope of within-basic-block identification, by enlarging basic blocks. Nevertheless, the scope remains limited to those techniques and cannot include *all* kinds of control flow.

*Control Flow Level.* A smaller amount of research has looked into identifica-

tion within CFGs. In [89] it is proposed to implement CFG regions with multiple control exits as accelerators. However, the presence of multiple control outputs significantly complicates the processor-coprocessor interface, as opposed to a single-entry single-exit approach. Another paper proposing HW/SW partitioning [7] presents a clustering methodology that operates on a control-data network compiled from an Extended Finite State Machine (EFSM) model. While it targets control flow to a certain extent, their methodology is limited to applications that can be modeled using EFSMs, therefore considering a much more limited scope than that of generic Control Data Flow Graphs compiled from source code, as does the methodology proposed in this chapter.

Finally, the authors of a recent work [2] consider Single Entry Single Exit regions but their target is to identify strictly parallelizable loop regions and offload them to an MPSoC target platform. This approach is limited in terms of excluding non-parallel regions from being potential candidates to be accelerated, and also in terms of not being cost-efficient, in case a designer needs to set a specific area constraint for the accelerators.

*Compiler Transformations.* Within compiler research, it is fairly common to identify CFG subgraphs for code optimization reasons. For example, trace scheduling, superblock and hyperblock scheduling [32], identify regions of the CFG in order to perform global code scheduling and improve code generation. *SESE* (Single Entry Single Exit) regions have been proposed in [34], and their identification was reimplemented in the LLVM framework in an analysis pass called *RegionInfo*, for the purpose of improving the performance of code generation. For my SW analysis, the idea of CFG region identification was borrowed from compiler research and was applied to automatically identify and select HW accelerators.

*Application Specific Instruction set Processor (ASIP) architectures and design practices.* HW Accelerators that are embedded in an Application Specific Processor can be either developed as hardwired Integrated Circuits (ICs), or mapped onto reprogrammable systems. In the first scenario, examples of Application-Specific Integrated Circuit (ASIC) platforms exist, such as the Tensilica Xtensa from Cadence [13] and the ARC processor from Synopsys [71]. These tools can be extended with accelerators and complex instructions. The CPUs can be configured during the design process to maximize performance and efficiency, without enduring the overhead of reconfiguration. An alternative, not as performing yet more flexible, is offered by FPGA-based Systems-on-Chips (SoCs), such as the Arria10 family [5] by Altera and the Zynq SoCs [80] by Xilinx.

The instances mentioned above support the generation of HW circuits, but do not provide implementation paths for differentiating the execution between HW

and SW. Conversely, High Level Synthesis (HLS) tools allow designers to move parts of applications, written in C or C++, between processors and accelerators. Research endeavors in this domain include LegUp [14] and ROCCC [28], while commercial applications comprise the Vivado HLS [79] suite from Xilinx (for FPGAs) and StratusHLS [12] from Cadence (for ASIC development). However, these HLS frameworks place the responsibility of partitioning a SW application on the application developer.

## 1.8 Conclusions

The RegionSeeker framework, along with its MuLTiVersioning extension, are methodologies that extend the state-of-the-art in the HW/SW co-design domain. They provide efficient solutions to the problem of automatically deciding which parts of an application should be synthesized to HW, under a given area budget. The accelerators identified by RegionSeeker consistently outperform the ones derived by data flow level algorithms and strictly function level candidates, across applications of widely different sizes and for varied area constraints. As an example, RegionSeeker offers up to 4.5x speedup for the mpeg2 benchmark compared to as SW execution. This work was published in IEEE Transactions on Computer-Aided Design of Integrated Circuits and Systems (TCAD) journal [87]. The MuLTiVersioning approach extends the initial selection pool of candidates and, compared to default HW accelerators configurations, offers enhanced speedup on the jpeg application of up to 1.8 on the entire application and up to 65x speedup on the relative kernels that are synthesized into HW, meaning the computation time that is synthesized in HW excluding the time that is left to SW.



## Chapter 2

# Automatic Optimizations for Accelerators

Identifying good candidates for HW acceleration is the first step to realize heterogeneous computing system designs that offer increased performance compared to a homogeneous system restricted to general purpose SW CPU(s). However, a set of optimizations applied on HW accelerators can decrease even more the computation times, thus leading to an improved performance compared to default non-optimized HW accelerator implementations. Modern High Level Synthesis (HLS) tools can apply such optimizations to HW accelerators and increase the performance of their implementations, as well as the overall performance of the entire heterogeneous system. HLS tools such as Vivado HLS [79], however efficient, though, are far from optimal since they require a lot of manual decisions from the programmer's part when it comes to the choice of *how* these accelerators can be synthesized. Furthermore, the resolution of which optimizations may be applied to which HW accelerators can be a complex problem, as it depends heavily on each HW accelerator characteristics.

In order to bring automation one step forward in HW/SW co-design, and under the scope of this part of my research, I tackled the problem of automating the decision making process of which optimizations should be applied to candidates for HW acceleration within a certain context. These optimizations include memory management of the data consumed and produced by the HW accelerators, a set of optimizations targeted to loops (e.g. loop pipelining, loop unrolling, loop flattening etc), pipelining consecutive pieces of computation such as subsequent function calls or loop bodies of consecutive iterations and array optimizations, such as array partitioning in blocks of the same size. Among the various optimizations available, I have focused on two major categories: a) Data Reuse analysis

and b) Loop Unrolling factor prediction. Both of these instances are explained in more detail in the following two sections.

## 2.1 Data reuse Analysis

### 2.1.1 Motivation



Figure 2.1. At each iteration, sliding window applications process a subset of the input data (a). The managed set of subsequent iterations present a high degree of overlap, both in the horizontal (b) and vertical (c) dimensions. The framework in 2.1.3 automatically leverages both, maximizing data reuse (d).

Loops are ideal candidates for acceleration. In almost every application, there is a number of them that contain a large number of iterations and there is a sufficient amount of computation taking place in their bodies. In addition to that, there are nested loops which commonly show a high level of data reuse. An effective exploitation of data reuse across consecutive iterations of loops can significantly lower the required amount of data exchange between HW accelerators and the main memory, thus reducing the respective bandwidth to and from accelerators, and increasing their performance.

An example of such high data reuse can be observed in sliding window applications, where there is typically a window of accesses scanning a wider domain, such as a two-dimensional array. Given that the level and pattern of data reuse



is known a priori, it is feasible to design specific memory structures, also known as memory buffers, attached to the HW accelerators. These memory buffers can exploit data reuse by keeping data locally and, hence, minimize the memory latency due to communication with the main memory.

Data reuse exploitation in High Level Synthesis (HLS) is still in premature stage. State-of-the-art methods [75] either rely on manually rewriting the source code, preceding HLS, or on source-to-source translation [55] [65], and are therefore poorly integrated in HLS tool-chains.

The methodology detailed in Subsection 2.1.3 attempts to bridge this gap. It presents a compiler-driven framework, based on the LLVM Polly [74] library, able to identify automatically data reuse potential in computational kernels in order to guide the synthesis of complex HW accelerators. As seen in Figure 2.1 these accelerators exploit timely unrolling and pipelining, by embedding a local storage holding elements which are re-used across iterations.

Sliding window applications, common in the image processing field, are targeted for acceleration. In such domain, a transformation is applied to each element of a large two-dimensional input array (a frame) according to the values in a smaller domain of accesses (a window). Large, yet constrained, input/output links are considered as the main architectural constraint in the design of this type of HW accelerators. Such arrangement is usually supported by commercial application specific platforms, such as the Tensilica Xtensa processor [73].

### 2.1.2 Related Work

In the domain of identifying automatically accelerators, research has so far focused mostly on accelerating data-flow [26] [30], not taking into equal account the potential for optimization by memory accesses. Exceptions are provided by papers [9] [31] where the authors support the claim that accelerators with custom storage can provide better speedup compared to the ones that accelerate data-flow only. However, these papers focus on the identification of the accelerators, and do not present a methodology to automatically identify the optimization potential, as well as synthesize them accordingly.

In sliding window applications, there are research endeavors both by academia and industry to exploit data reuse. The smart buffers [29] generated by the ROCCC compiler [77] allow for automatic detection of data reuse opportunities, but cannot be interfaced with interconnects of varying width. The methodology described in [44] employs reuse buffers spanning multiple frame columns, which pose a significant area overhead. Both [29] and [44] are not able to combine Hardware unrolling and pipelining, which are instead jointly supported the

methodology detailed in 2.1.3. An alternative approach, described in [22], requires a great deal of Hardware resources as well, as it requires the storage of large parts of a frame being processed inside the custom hardware. In [83], the authors propose an analytical method to gather microarchitectural parameters for sliding-window applications on FPGAs. Their design however ultimately needs to be manually implemented and hence the work neglects high level synthesis aspects.

The commercial Vivado HLS tool requires extensive manual rewrite of the source code, in order to instantiate a reuse memory buffer. On the other hand, the approach presented here relies on automated code analysis to derive the characteristics of the target application.

### 2.1.3 Methodology

In order to generate the custom-storage HW accelerators, a two-steps methodology is carried out. First the data reuse analysis of the application takes place and then the synthesis of the part of computation to be implemented in HW. The first is performed with the utilization of compiler static source code analysis while in the latter details regarding the Hardware implementations are provided. The phases of analysis and synthesis lead to the design and implementation of custom-storage accelerators that manage to minimize the latency due to data transfer between main memory and the HW accelerators.

#### Data Reuse Analysis

In order to identify the level of data reuse that takes place throughout the execution of every window application, there are three pieces of information that are vital: a) the size of the window, b) the stride and c) the frame size. An example of data reuse, accounting for roughly 66% the size of the window between consecutive iterations, can be seen in Figure 2.2. The window size is the access pattern within the innermost body of the loop. The innermost and outermost loop stride is the value of the induction variable increase for the innermost and the outermost loop respectively. Finally, the frame size is the iteration space within which the sliding window is moving. To extract this information I have developed an analysis pass, based on the LLVM Polly framework [74]. Application-specific parameters are then considered in conjunction with architecture constraints (input/output width) to automate the synthesis of efficient HW accelerators.

The analysis pass, as seen in Algorithm 2, iterates over regions of the application functions and identifies Static Control Parts (SCoPs). The SCoPs are sub-



Figure 2.2. Data Reuse of memory accesses (66.6%) across consecutive iterations of a loop, in a sliding window application.

graphs of the control flow graph of a function where the flow of control is known statically. For each SCoP, loop and scalar evolution information is collected from the body of the loop.

Loop information supports methods that can provide the loop depth, so as to identify the innermost loop. Scalar evolution information can be used to extract the loop trip count, which is the iteration space of each loop, and thus compute the frame size. The stride value, both vertical and horizontal, is obtained by a function that was developed based on existing methods of the Loop analysis LLVM pass. Lastly, the read memory accesses of the innermost body of the loop are identified by using `isl` functions, which compute the distance (or delta) of each of these read accesses with respect to the first one. Given the access pattern, the window size is computed as the minimum enclosing rectangle. After having identified the necessary information, the implementation of a local buffer that fits these needs can be carried out.

### Hardware Implementation

The parameters retrieved with the analysis pass (frame size, stride, horizontal and vertical window size) and the characteristics of the interconnect (input and output width) are employed to derive efficient HW accelerators implementation with local storage and data reuse.

As showcased in Figure 2.2, accelerators implementations embed multiple

---

**Algorithm 2** LLVM Analysis Pass - SCoP Identification and Data Reuse Analysis
 

---

**Input:** Application written in C/C++

**Output:** List of Identified SCoPs and their respective Frame, Window, Stride sizes/values.

```

1: function RunOnRegion()
2:   getAnalysis(ScopInfo)
3:   scop = getScop()
4:   RunOnScop(scop)
5:
6: function RunOnScop(scop)
7:   LI = getLoopInfo()
8:   SE = getSE()
9:   if L == OutermostLoop then
10:    getTripCountForLoop()
11:    getStrideForLoop()
12:   else if L == InnerMostLoop then
13:    getReadMemoryAccesses()
14:    ComputeDistancesForReadAccesses()
15:    ComputeWindowSize()

```

---

combinatorial datapaths, each executing one iteration of the loop body of the target application. The input interface embeds a local storage, whose horizontal size corresponds to the available input data width of  $IN_w$  data elements, while its vertical size is equal to the vertical size of the application window  $h$ . It is implemented as a  $IN_w * h$  shift register, operating in the vertical (top-down) direction. During execution, the first row of the shift register is filled with input data in each clock cycle. A subset of the elements stored in the shift register is connected to each of the different datapaths according to their managed sets, e.g.: the first one having inputs corresponding to the buffer columns ranging from 0 to  $w - 1$  (the horizontal size of the window) and the second one corresponding to the buffer columns ranging from 1 to  $w$ . Figure 2.2 illustrates such scheme for the simple case of  $IN_w = w + 1$ .

At the beginning of the execution,  $h$  rows are stored in the shift register before activating the datapaths logic. Afterwards, this activation is performed for each new row, discarding the last (topmost) line and storing a new one in the first (bottom) position of the shift register. At the completion of a vertical slide of the window through the frame, a new one is started, increasing the horizontal

displacement of the buffer by  $IN_w - w + 1$  elements.

Finally, since no reuse opportunities are present for outputs, the output interface simply concatenates the values generated by the datapaths, and transfers them as a single and wide memory access.

#### 2.1.4 Experimental Results

The evaluation of this approach is carried out in three benchmarks of varying window sizes. Sobel is an edge detection algorithm with an access pattern of a 3x3 window. BlockSAD is a kernel in H.264 and is used to detect the similarity among 4x4 blocks. Finally, Maximum Filter computes the brightest pixel among neighbors in 8x8 blocks. Three different configurations were considered, spanning from a single datapath and minimum input width (Conf.1) to multiple datapaths and increased input width (Conf.2 and Conf.3) as seen in Figure 2.2. Multiple datapaths translate to more parallel windows executing and, hence, increased demand in area resources.

The comparison of the approach introduced above is carried out against two state-of-the-art HLS tools: ROCCC and Vivado HLS. Vivado HLS is compared in two modes, one being the default (Vivado\_norew) and the other one after extensive manual rewrite of the source code (Vivado\_rew) in order to obtain increased data reuse.

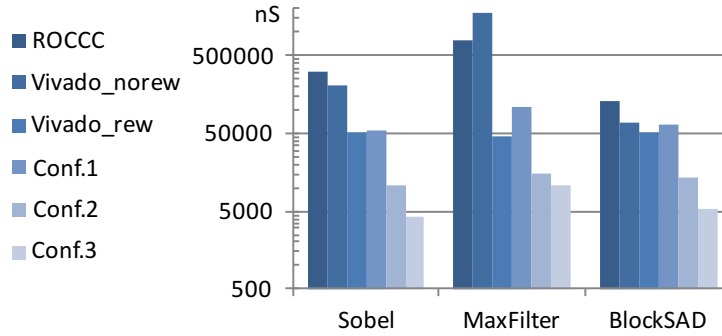


Figure 2.3. Execution time to process a 100x100 frame.

Execution time, as seen in Figure 2.3, is extracted from a targeted Xilinx Virtex7 FPGA platform. It can be observed that ROCCC systems have similar performance with respect to Vivado\_norew ones. Conf.1 accelerators — even though they do not require code modifications — are as efficient as Vivado\_rew ones. Conf.2 and Conf. 3 that are supported only by the framework presented in 2.1.3,



Figure 2.4. Comparison of required resources for our generated systems and for baseline approaches: LUTs (a) and Flip-Flops (b).

dramatically decrease run-times, with an order-of-magnitude speed up on average between Conf.1 and Conf.3. The other state-of-the-art tools *fail to provide an equivalent solution with such low execution time*. Figure 2.4 reports the amount of area resources required by ROCCC, Vivado HLS and our own generated accelerators. Unsurprisingly, accelerators featuring a high number of datapaths (Conf.3) require more resources than single-datapaths approaches (Conf.1, Vivado). Nevertheless, the area increase in terms of Flip-Flops is comparable to the other to state-of-the-art tools, as the size of the buffer only is increased slightly to support a high degree of parallelism. On the other hand, the results highlight that complex accelerators require an increased amount of combinatorial logic (LUTs), with respect to ROCCC and Vivado HLS.

### 2.1.5 Conclusions

It has been demonstrated that static source code analysis can be crucial, when it comes to automatically optimizing the synthesis of accelerators that are dedicated to sliding window applications. My SW analysis identifies data reuse, as well as data locality, and subsequently allows to exploit these characteristics by making use of appropriate memory buffers. The experimental results reveal an order-of-magnitude performance improvement with respect to state-of-the-art methodologies. This work was published in HiPEAC IMPACT 2017 Seventh International Workshop on Polyhedral Compilation Techniques [85].

## 2.2 Machine Learning Approach for Loop Unrolling Factor Prediction

### 2.2.1 Motivation

High Level Synthesis tools, utilized to synthesize accelerators, require manual decisions to be made, so as to build efficient accelerators. These decisions regard the choice of high-level optimizations and transformations to be applied, therefore a good understanding of the SW parts to be accelerated is essential. Optimizations applied during synthesis can highly improve the performance achieved, as well as allocate less HW resources for a given computer architecture. Nonetheless, the selection of optimizations is a challenging task due to two main reasons. First, hardware synthesis is a time-consuming process, limiting in practice the amount of possible implementations that can be evaluated. Second, the effect of assigning different values to directives is difficult to foresee, due to low-level application characteristics.

Simulation tools such as Aladdin [66] have been developed in order to rapidly estimate the performance and cost (area) of HLS-defined designs. Nonetheless, even when employing estimation tools, an exhaustive evaluation of all directives settings for each candidate accelerator in a heterogeneous system is still unfeasible beyond simple cases. Addressing this challenge, a machine learning framework is proposed that is able to infer the proper implementation of an HLS design based on its characteristics, automatically derived from a source code analysis pass, based within the LLVM compiler framework [38].

Within the scope of this piece of research, the focus is placed on *loop unrolling*, an already well known optimization from the compiler domain, as well as the HW domain. The loop unrolling optimization replicates the body of a loop a given number of times in order to expose parallelism, which especially in a HW implementation can lead to substantial speedup gains [37]. This directive should nonetheless be applied sensibly, because it entails a high area cost for the duplicated logic; in addition, its ensuing benefits can be hampered by loop-carried dependencies and frequent memory accesses.

It is thus clear that there is a trade-off between execution time and the area budget that a computer architect has at hand, as well as the level of complexity and more potential side effects that need to be taken into consideration. Since HW realizations are targeted, the goal of this work is to reach a *sweet spot* between performance and HW resources.

Within the sphere of this research work, the following contributions are made.

First, a novel *Machine Learning* approach is introduced, based on Random Forest classification, instead of estimation-based models, to predict accurately the optimal loop unrolling factor of loops in applications to be synthesized in HW. The use of this methodology can provide results with better prediction score and in much less time, compared to the state-of-the-art. Second, the whole process is fully automated, from the analysis of the input applications, using the LLVM compiler infrastructure [38], up to the training of the Random Forest Classifier. Finally, the trained Random Forest classifier can be used to generate accurate loop unrolling predictions for any given application – a piece of information that can directly be used by an HLS tool such as Vivado HLS, in order to synthesize parts of these applications to HW.

### 2.2.2 Related Work

Research papers have explored the applicability of machine learning to apply compiler optimizations. In Software compilers, it has been employed by Agakov et al. [1] to speed up iterative compilation, by Monsifrot et al. [47] to produce compiler heuristics and by Kulkarni et al. [36] to select the order in which optimization passes should be performed. Stephenson et al. [69] have made use of Supervised Classification, such as near neighbor (NN) classification and Support Vector Machines (SVM) methods, to produce accurate predictions in optimal unrolling factors. In all above-mentioned research works the authors targeted Software compilation; in Subsection 2.2.4, a comparative performance evaluation of the framework presented in Subsection 2.2.3 to the methodology proposed by Stephenson et al. is carried out, showcasing the benefit of the choice of loop features and classification strategy in the HLS scenario.

Liu et al. [40] used a Random Forest classification model in the context of HLS, extending the Iterative Refinement framework proposed in [42] [51] [82] and [90]. They address a different problem with respect to the one tackled in this section: that of retrieving the set of Pareto-optimal implementations of a given design by navigating its configuration space. A similar stance, addressing system-level design, is illustrated by Ozisikyilmaz et al. [50]. As opposed to these works, my aim is to perform a predictive assignment of synthesis directives, based on a training performed on an independent input set. This problem was also investigated by Kurra et al. [37]. Contrary to their methodology, my methodology does not depend on a detailed estimation delay model of the loop body so as to predict loop unrolling factors in HLS instances.





Figure 2.5. Overview of the Loop Unrolling Prediction methodology.

### 2.2.3 Methodology

In this section, first the employed objective function that determines the optimal loop unrolling factor is presented. Then, the LLVM analysis pass that was developed in order to automatically extract relevant loop features and the approach followed to retrieve the area and run-time performance of HLS designs is detailed. Lastly, the supervised learning classifier method is demonstrated, which, during the training phase, gathers the data from the previous steps to produce a loop unrolling predictor, and, during the test phases, assigns loop unrolling factors based on loop features.

#### Optimal Loop Unrolling Factor – Objective Function

The *optimal loop unrolling factor* is defined as follows. Given a defined set  $S$  of unrolling factors, e.g.  $S: \langle 1, 2, 4, 8, 16, 32, 64 \rangle$ , there is one for every loop that maximizes the **Impact (I)**, given by the following formula:

$$I(L, A) = \alpha \cdot \frac{(L_1 - L)}{L_1} + \beta \cdot \frac{(A_1 - A)}{A_1}, \quad \alpha + \beta = 1$$

Where  $L_1$  is the latency of the of the function containing a loop and being synthesized as HW accelerator, for Loop Unrolling Factor (LUF) that is equal to one, i.e., a fully rolled loop.  $L$  is the latency of the HW accelerator for any possible LUF from the defined set. Respectively,  $A_1$  is the area resources requirements of the HW accelerator with LUF equal to one and  $A$  the area for any possible LUF

from the defined set.

Subsequently, the optimal LUF is defined as the one that maximizes the Impact function above. Note that, when  $LUF = 1$ , then  $I(L, A) = 0$  which corresponds to a baseline implementation.  $I(L, A)$  may also be negative for suboptimal LUF choices (where unrolling might increase area without decreasing latency), but will always be  $\geq 0$  for optimal unrolling factors.

For the evaluation presented in Subsection 2.2.4 three different strategies were considered: a) Optimize for both latency and area ( $\alpha = \beta = 0.5$ ). In this configuration a balance is maintained between decreasing the number of execution cycles and keeping low the usage of HW area resources in a given implementation. b) Optimize for latency ( $\alpha = 0.7, \beta = 0.3$ ). Minimizing latency is favored by this approach, thus focusing on increasing the speedup of an application, and finally c) Optimize for area ( $\alpha = 0.3, \beta = 0.7$ ). This approach aims at decreasing the area budget of the implementation, therefore achieving an average speedup, but maintain low usage of HW area resources. All three configurations fulfill different architectural needs and explore realistic alternative scenarios.

#### LLVM Analysis Pass – Loop Features Extraction

Loop features are automatically identified by an analysis pass (depicted as point 1 in Figure 2.5) that was developed within the LLVM compiler infrastructure [38]. Features are retrieved starting from applications written in C or C++, operating on their Intermediate Representation, provided by the LLVM front-end passes.

My *LLVM Loop Unrolling Prediction Analysis Pass* iterates over functions of the applications and identifies loops. On each of them, it performs loop, scalar evolution and dependence analysis to extract their features, summarized in Table 2.1: the critical path, the trip count, the presence of loop carried dependencies and the required memory accesses (load and stores).

The choice of features is based on the factors that influence the cost and the achievable speedup of Hardware unrolled loops: a loop with a long critical path may be expensive to duplicate, while loop carried dependencies and memory accesses may force a serialization of execution irrespectively of the degree of unrolling. These considerations lead us to consider a markedly different feature list with respect to works focusing on software targets, such as the one of Stephenson et al. (Table 2.2).

Features - X Vector
<i>Critical Path</i>
<i>Loop Trip Count</i>
<i>Has Loop Carried Dependencies</i>
<i># Load Instructions</i>
<i># Store Instructions</i>

Table 2.1. Features extracted by LLVM LU Analysis Pass.

Features - X Vector 1	Features - X Vector 2
<i># Operands</i>	<i># Floating Point Operations</i>
<i>Range Size</i>	<i>Loop Nest Level</i>
<i>Critical Path</i>	<i># Operands</i>
<i># Operations</i>	<i># Branches</i>
<i>Loop Trip Count</i>	<i># Memory Operations</i>

Table 2.2. Feature vectors selected by Stephenson et al. [69].

### Latency and Area Estimation

To establish a link between LUFs and performance/cost of implementations, latency and area values must be extracted both for the loops in the training set (in order to optimize the classifier) and the ones in the test set (to measure its accuracy). Aladdin [66] (point 3 in Figure 2.5), a pre-RTL power-performance simulator for Hardware accelerators was utilized in order to retrieve latency and area information. All functions in the considered benchmarks were simulated by employing each feasible unrolling factor in the  $S$  set defined above on every loop contained. Latency is reported by Aladdin in clock cycles, while area is expressed in  $\mu m^2$  in a 45nm technology. The result is shown as point 4 in Figure 2.5.

The Impact ( $I$ ) was computed afterwards for the different  $\alpha$  and  $\beta$  values, to retrieve the optimal loop unrolling factor for every loop of a function, which is the index of the LUF that maximizes  $I$ . The result is three vectors  $\{Y1, Y2, Y3\}$  (point 5 in Figure 2.5) that contain the target values for the classification algorithm. The  $Y1$  vector includes the optimal loop unrolling factor that balances the Hardware implementation of the accelerators in terms of low latency and low area. Values in the  $Y2$  vector favors low-latency implementations, applying more aggressive loop unrolling, whereas  $Y3$  favors low-area ones.

---

**Algorithm 3** LIVM Analysis Pass - Loop Unrolling Prediction Analysis

---

**Input:** Application written in C, C++**Output:** X (Feature Vector)

```

1: function RunOnFunction( )
2:   for BB in Function do
3:     if L=getLoopForBB() then
4:       LoopUnrollingPredictionAnalysis(BB,L)
5:
6: function LoopUnrollingPredictionAnalysis(Basic Block BB, Loop L)
7:   LI=getLoopInfoAnalysis()
8:   SE=getScalarEvolutionAnalysis()
9:   DA=getDependenceAnalysis()
10:  /* Gather Features for X Vector */
11:  x1=getCriticalPath(BB)
12:  x2=getTripCountForLoop(L)
13:  x3=getLoopCarriedDependencies(BB)
14:  x4=getNumberOfLoadInstructions(BB)
15:  x5=getNumberOfStoreInstructions(BB)

```

---

## Random Forest Classification

This information (X and Y vectors) extracted as described above are used as input to a Random Forest (RF) classifier (point 6 in Figure 2.5). Supervised learning is performed by detecting the correlation between the input, the compiler extracted information, used as the X feature vector, and the output, which is the optimal loop unrolling factor for each loop.

Random Forest was used as the supervised learning model, which has been shown by Liu et al. [40] to outperform alternatives such as Multilayer Neural Networks and Support Vector Machines classification in the context of HLS design space exploration. Random Forest algorithms follow a decision tree methodology, combining many weak classifiers to derive a strong one, allowing the generation of low-complexity and robust classifiers.

The algorithm employed, as presented in Algorithm 4, follows an approach similar to a *k-fold cross validation* strategy. The whole data set (X and Y vectors, see points 2 and 5 in Figure 2.5) is divided randomly between a training set and a test set, where the training set is equal to 80% of the whole data set and the remaining 20% is the test set. Then, the Random Forest model is used for the

---

**Algorithm 4** Random Forest Classification - Training and Test

---

**Input:** X and Y Vectors**Output:** Trained Random Forest Classifier

---

```

1: for  $i$  in  $NumberOfTrainingSessions$  do
2:    $X\_train, X\_test, Y\_train, Y\_test = train\_test\_split(X, Y)$ 
3:   /* Training Phase */
4:    $M = RandomForestLearningModel$ 
5:    $M.train(X\_train, Y\_train)$ 
6:   /* Evaluation Phase */
7:    $Pred = M.predict(X\_test)$ 
8:    $Error = abs(Pred - Y\_test)$ 
9:    $Score = M.score(X\_test - Y\_test)$ 

```

---

training process on the training set and out-of-sample predictions are carried out for each element of the test set. After all predictions on the test set have been computed, the prediction score and the average error (as defined in Subsection 2.2.4) are computed for the current training session.

#### 2.2.4 Experimental Results

To evaluate the classification performance of a trained classifier, two different metrics were adopted. The *Prediction Score* states the percentage of optimal (according to  $I(L, A)$ ) LUFs that were correctly identified on the out-of-sample test set. The *Average Error* instead measures the average distance between the indexes in  $S$  of the correct and the predicted LUF.

In order to comparatively evaluate the proposed methodology, combining Random Forest classification and LLVM-based loop features extraction, benchmarks of different complexity were considered. Small and medium-sized ones are `adpcm`, an audio encoding kernel, `stencil`, an implementation of an iterative algorithm that updates array elements according to a given pattern, and `sha`, a secure hash encryption method used in the information security domain. `jpeg` and `mpeg2` are instead larger benchmarks, which perform image and video compression, respectively. Applications were drawn from the CHStone [33] and the Scalable Heterogeneous Computing (SHOC) benchmark suites [20]. In total, they comprise 87 different loops.

Random Forest classification was implemented using the Scikit-learn suite [52], that includes state-of-the-art implementations of Machine Learning models

in python. Scikit-learn was also employed to re-implement the two methods proposed by Stephenson et al. [69], that are consider as baselines.

Giving an initial proof of concept for the strategy proposed, Figure 2.6 reports the difference between the indexes of the predicted optimal (according to impact value) loop unrolling factors and the ones retrieved with an exhaustive exploration, considering 18.000 out-of-sample predictions on all the benchmark loops. Results are highly concentrated on zero, indicating a high rate of correct predictions.

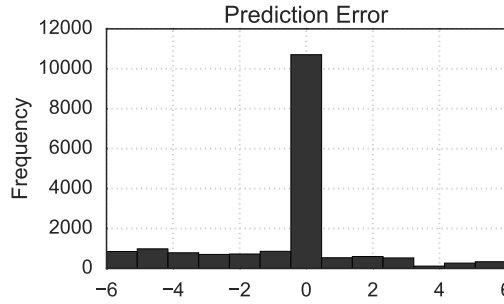


Figure 2.6. Distribution of Loop Unrolling Factor Prediction Errors over 18.000 out-of-sample predictions.

### Classification Models and Features

The evaluation of the choice of features ( $X$  vector in Table 2.1) and training model Random Forest (RF), takes place against two state-of-the-art methodologies proposed by Stephenson et al. [69]. The latter present different classification strategies: Support Vector Machines (SVM) and Near Neighbor (NN) and a different choice of investigated loop features, reported in Table 2.2. Figure 2.7 shows, for a choice of  $\alpha = 0.5$ , the prediction score and the average error of the nine strategies resulting from different feature vectors and classification strategies. Experimental results highlight that the presented framework ( $X$  feature vector and RF classification) outperform other choices, reaching a prediction score above 60% and an average error of less than 1.4. Similar results were obtained for impact functions favoring area or latency ( $Y2$  and  $Y3$  vectors).

### Iterative Refinement

In the second round of experiments, the comparison of the proposed method was carried out against an Iterative Refinement approach, used in [42] [51] [82] [90].



Figure 2.7. Left: Comparison of the Prediction Score across Random Forest, Nearest Neighbor, Support Vector Machines models and the respective feature selection: X vector, Stephenson et al. X1 and X2 vectors [69]. Right: Features extracted by my LLVM Loop Unrolling Analysis Pass.

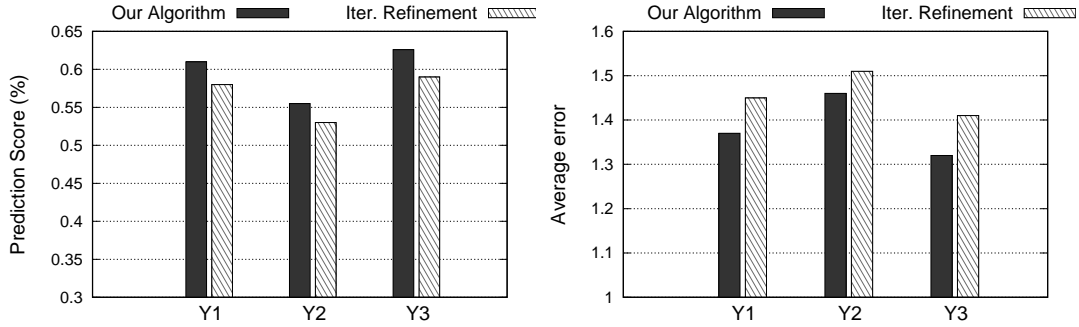


Figure 2.8. Left: Comparison of the Prediction Score across Random Forest, Nearest Neighbor, Support Vector Machines models and the respective feature selection: X vector, Stephenson et al. X1 and X2 vectors [69]. Right: Features extracted by my LLVM Loop Unrolling Analysis Pass.

Iterative Refinement uses part of the training data set to obtain a first version of the classifier, whose performance is then improved by using a second, disjoint set of input and outputs.

For this evaluation, three different settings of Y target vectors  $\{Y1, Y2, Y3\}$  were considered, as described in Subsection 2.2.3. The employed data, the features (X vector) and the training model (Random Forest) were the same both for Algorithm 4 and the one using Iterative Refinement. For Iterative Refinement, 75% of the training set is allocated for the initial training phase, and the remaining 25% for the refinement phase.

The prediction score, as seen in Figure 2.8, ranges from 53% to 63% across the three Y vectors. Nevertheless, our methodology consistently outperforms

the Iterative Refinement approach, while achieving the highest prediction score (63%) for the setting that favors low area resources (Y3). A similar observation can be made for the average error values, where the suggested approach keeps a lower average error for all predictions, across all vectors, with the one related to the Y3 vector being the lowest (1.32).

### 2.2.5 Conclusions

A novel methodology based on LLVM analysis and Random Forest classification that performs loop unrolling factor prediction for HLS designs was performed. This approach achieves better prediction score in comparison to state-of-the-art Machine Learning methods. Experimental evidence showcases that, by carrying out accurate predictions of loop unrolling factors, high performance accelerator implementations can be realized, while avoiding time-consuming exhaustive explorations. This work was published in the 2018 International Conference on High Performance Computing & Simulation (HPCS) [86].



## Chapter 3

# System Aware Accelerators Identification

In the final Chapter of the dissertation the notion of automatic selection of HW accelerators is extended by taking into account the underlying system where the specialized accelerators are hosted. The knowledge of the characteristics of the system that is targeted can be critical for the choice of the HW accelerators to be implemented. The memory system of a platform for instance can vastly affect the latency due to data exchange between the main memory of the system and the HW accelerators. AccelSeeker, an LLVM-based tool-chain, is presented as a framework that performs automatic identification and selection of HW accelerators targeted to a specific System-on-Chip (SoC). AccelSeeker performs thorough analysis of applications source code and estimates memory latency along with computational latency of candidates for acceleration. The granularity of the candidates for acceleration is expanded to that of a subgraph of the entire call graph of an application to accommodate for the communication overhead between main memory and the HW accelerators. **NOTE:** Write for EnergySeeker.

### 3.1 AccelSeeker: Accelerators for Speedup

#### 3.1.1 Motivation

System-level design, and heterogeneous computing as a whole, is witnessing a breakthrough. Emerging best practices based on High Level Synthesis (HLS) allow unprecedented productivity levels. HLS dramatically shortens development cycles by employing C/C++ descriptions as entry points for the development of both Software (SW) and Hardware (HW) greatly facilitating the task of migrating

functionalities between the two.

However, the design of heterogeneous systems comprising SW processors and HW accelerators is still a demanding endeavor, during which key decisions are left solely to manual effort and designer expertise [11] [48]. Furthermore, the long time required for HW synthesis, coupled with the huge space of alternative implementations exposed by real-world applications, limits in practice the number of accelerator choices that can be considered manually by a designer before HW/SW partitioning is settled.

In order to limit the entailed design effort, it is therefore crucial to identify the set of viable acceleration options quickly, and also early in the design process, before performing later and more detailed estimations. This key step is currently poorly supported by design automation tools. Indeed, state-of-the-art early partitioning strategies are solely based on profiling information [81] [38] which, as was also shown by the authors of [72], may often be misleading.

To fill this gap and offer an efficient solution to the problem stated above, AccelSeeker is presented. AccelSeeker offers a methodology to identify and select the suitable acceleration candidates in an application, from its software source code. Being implemented within the LLVM [38] compiler infrastructure, it first models an estimation of the cost (required resources) and merit (potential speedup) of all candidate accelerators in a single, quick pass, and then selects the set that maximizes the estimated speedup, while not exceeding a resource target. The use of AccelSeeker can therefore guide Integrated Circuit architects in the early design phases, highlighting which segments of a computing flow should be targeted with HLS, and hence where to focus the process of applying optimizations to.

On the other hand, it indicates which parts are not likely to yield tangible benefits if realized in HW — either because they present a low computational footprint, or because their characteristics hamper their potential for HW acceleration, e.g. they require an excessive amount of data transfers while performing limited computations.

The approach of AccelSeeker is markedly different from that of performance estimators, as the most promising candidates are identified in a single, high-level exploration, reducing the scope of further, and more detailed, estimations. It also differs from approaches based solely on profiling information, as profilers do not offer a measure of costs and run-times of HW implementations. Furthermore, they do not account for invocation overheads – potentially leading to the selection of frequently called, but small, candidates. Finally, data transfers is an important factor that state-of-the-art profilers do not take into account – hence potentially suggesting candidates requiring an excessive amount of communi-

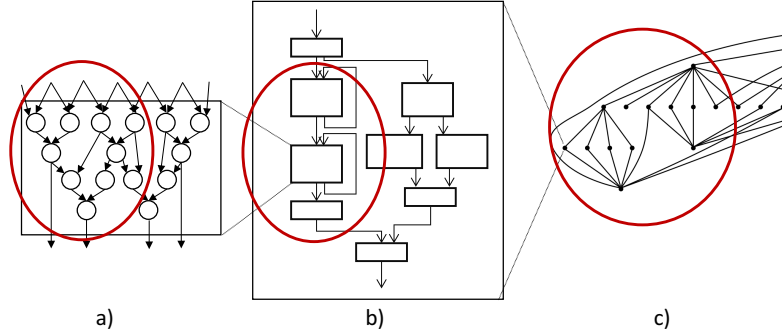


Figure 3.1. Evolution of the SoA in automatic selection of custom instructions/accelerators: (a) from data-flow level [56] [26], (b) to control-flow level [87] [49], (c) to function-call graph level (this work).

cation, that in turn can significantly weaken any potential performance gained. AccelSeeker, instead, takes into account both the communication overhead and all the characteristics of the platform that are required in order to generate an estimation model which leads to high-performance HW accelerators choices.

### 3.1.2 Related Work

High Level Synthesis tools have considerably matured in recent years [45]. Nowadays, available commercial tools (e.g.: Xilinx Vivado HLS [79], Cadence Stratus HLS [12]), as well as academic alternatives (e.g.: Legup [14], Bambu [53]) support the design of very large accelerators from C/C++ code. They reach performance levels comparable to hand-crafted implementations specified in low-level Hardware Description Languages (HDL) such as VHDL or Verilog [41].

Nonetheless, the automated selection of the application parts most amenable to HW acceleration is still an open research topic. Selection approaches based on synthesis results [15] scale poorly to complex applications, as these are only available late in the design process. Estimation frameworks offer a detailed analysis on the performance and resource requirements of a HW-accelerated system while avoiding full synthesis runs, either by interfacing software and HW simulators (e.g., gem5 [8] and Aladdin [66] in [67]), or by adopting a hybrid stance, in which HW execution times are estimated, while software ones are measured on a target platform (as in Xilinx SdSoC [35]). However, in both cases, estimations are performed *after* the partitioning of HW and software, which is left to a trial-and-error basis. A methodical solution for partitioning is instead proposed

in this chapter.

The downside of poor partitioning choices, and consequently the importance of automated tools such AccelSeeker that guide the selection of high-quality accelerator sets, is even more prominent when considering the high effort required to optimize the implementation of HLS-defined accelerators. Design optimization entails the specification of multiple directives to steer designs towards the desired area-performance trade-off. The link between directives and the performance of implementations is not straightforward, hence requiring the evaluation of multiple alternatives to reach the intended results, as exemplified in [62] [90] [24] [40]. It is therefore key to focus up-front this optimization effort only on those candidate accelerators which can lead, from an application perspective, to tangible speedups.

To this end, the approach introduced here is inspired by previous works on automatic identification of instruction set extensions. Most techniques in this field target customizable processors augmented with application-specific functional units, within the processor pipelines. Hence, these techniques usually constrain their search to the scope of single basic blocks [56] [26], as depicted in Figure 3.1a. Recently, the authors of [87] and [49] have instead targeted the identification of larger code segments, including control-flow structures belonging to single functions (depicted in Figure 3.1b). However, such scope still falls short of the one employed in HLS tools, which are devoted to the implementation of dedicated accelerators interfaced on a system bus [19]. In this setting, the cost of data movement becomes so prominent that even control-flow structures inside functions fail to deliver performance. Suitable accelerator candidates must then encompass entire functions, including in turn all functions called within their call tree. AccelSeeker considers this same granularity (Figure 3.1c), advancing the state-of-the-art in automatic accelerators identification.

The methodology embedded in AccelSeeker, whose high-level scheme is depicted in Figure 3.2, is detailed. First, the definition of a candidate for acceleration is provided, and then how the selection of a subset of potential candidates to be implemented in hardware is being performed (A and C in the figure). Subsequently, the approach employed to estimate the candidates performance and resource requirements (B in the figure) is presented.

### 3.1.3 Candidate Identification

In order to discover which parts of an application can be most profitably accelerated in hardware, the investigation of its function-call graph takes place, i.e., a Directed Acyclic Graph  $G(N, E)$ , where every node  $n \in N$  corresponds to a

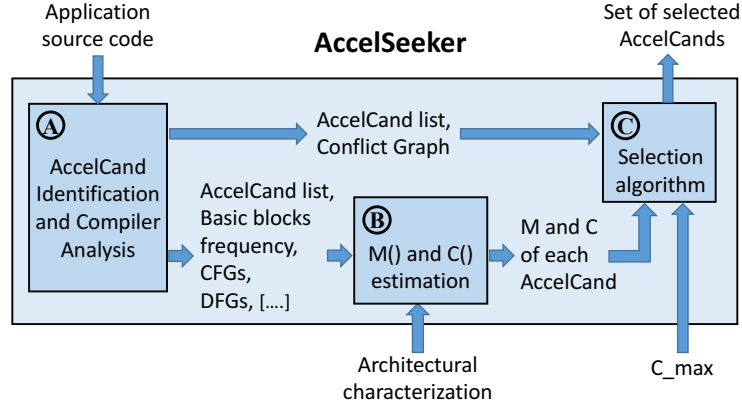


Figure 3.2. The phases of the AccelSeeker approach. A) Candidates for acceleration are identified, from source code analysis. B) Given an estimation of merit and cost for each candidate, and C) given a maximum available cost, AccelSeeker performs selection of a subset of such candidates.

function and every edge  $e = (u, v) \in E$  corresponds to a call from function  $u$  to function  $v$ . A *root* is a node that reaches all other nodes of the graph, i.e., for every other node  $n \in N$ , there exists a path from the root to it. The function-call graph  $G$  has a root, which represents the top-level function of the application. Figure 3.3a shows an example of a call graph (note that edge directions are not shown here, for picture clarity; they are, however, intended from top to bottom).

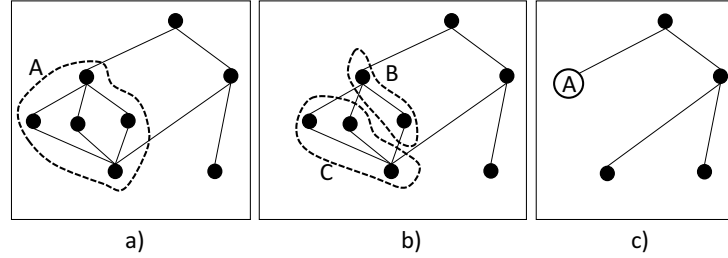


Figure 3.3. a) An example call graph. Black nodes are the functions present in a SW application, and edges represent function calls. Subgraph A is an AccelCand. b) Subgraphs B and C are not AccelCands (B has outgoing edges, C has no root). c) Call graph resulting from selection of A as accelerator.

A candidate accelerator is defined, and called **AccelCand**, as a subgraph  $S(N_s, E_s)$  of graph  $G$ , exhibiting the following two characteristics: the subgraph has a root; the subgraph has zero outgoing edges. The former means that the subgraph has a node that reaches all other nodes in the subgraph; the latter means that, for every node  $n_s \in N_s$ , no edge  $(n_s, m_s)$  exists in  $G$  such that  $m_s \notin N_s$ .

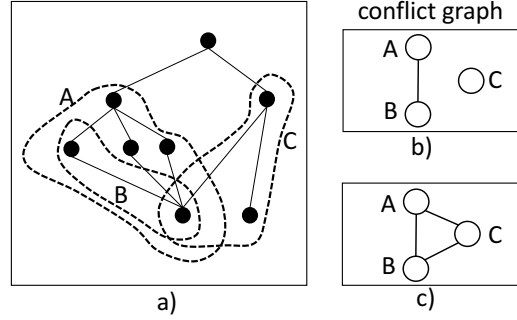


Figure 3.4. a) Three overlapping AccelCands: A, B, C. b) Conflict graph considered in the problem formulation: complete overlap is a conflict, while partial overlap is allowed c) Conflict graph adopted in [87] instead, where any kind of overlap is considered a conflict.

Figure 3.3a and 3.3b show three example subgraphs, labeled A, B, C. While subgraph A is an AccelCand, subgraph B is not, because it does not have zero outgoing edges, and subgraph C also is not an AccelCand, because it does not have a root. The call graph resulting from selection of AccelCand A as accelerator is shown in Figure 3.3c: the whole subgraph is subsumed to a single (accelerator) call.

The methodology is limited to considering call graphs that are acyclic, and hence constructs such as recursion cannot be dealt with. This is in line with the limitations of HLS tools.

#### 3.1.4 Problem Statement and Candidate Selection

Given a call graph  $G(N, E)$ , there exist  $|N|$  AccelCands in it; every node of  $G$  is, in fact, the root of one and only AccelCand. The problem of *Selection* is that of choosing, among all of the  $|N|$  AccelCands, the subset to be realized as accelerators.

Each AccelCand is associated with a merit  $M()$  – an estimation of the number of cycles saved by it when implemented in HW as opposed to SW – and a cost  $C()$  – an estimation of the area needed by it when implemented in HW. Note that the methodology here proposed is agnostic to the way cost and merit are defined. Of course, their definition should correctly reflect the SW and HW architectures that the methodology is targeting, and the details of how functions  $M()$  and  $C()$  are defined for the experiments in this paper are given in Subsection 3.1.5.

Given a set of AccelCands, defined and identified as described in the previous

subsection, and given a cost and merit associated to each of them, the problem that is tackled is the following.

**Problem: *Accel Selection***

Let  $A = \{A_1, A_2, \dots, A_n\}$  be a set of AccelCands, with associated cost and merit functions  $C$  and  $M$ . For any subset  $S \subseteq A$ , it is denoted by  $M(S) = \sum_{i \in S} M(A_i)$  the *sum* of the candidate merits, and by  $C(S) = \sum_{i \in S} C(A_i)$  the *sum* of their costs.

A subset  $S$  of AccelCands is selected such that:

1. The merit  $M(S)$  is maximized
2. The cost  $C(S)$  is within a user-given cost budget  $C_{\max}$
3. No two candidates in set  $S$  are in *conflict*

The concept of *conflict* is defined in the following way: two candidates  $A_i$  and  $A_j$  in  $S$  are in conflict if and only if  $A_i \subset A_j \vee A_j \subset A_i$ , i.e., if one completely includes the other.

The concept of conflict that is considered in the problem formulation is exemplified with the help of Figure 3.4. A call graph is shown, where 3 AccelCands (out of the possible 8, as there are 8 nodes in the call graph) are highlighted, and they are labeled A, B, and C. A *conflict graph* is presented as well, in Figure 3.4b, which is an undirected graph where nodes represent AccelCands, and an edge is added between two nodes if the two candidates are in conflict. A and B are in conflict because B is completely contained in A.

The reason behind the concept of conflict is that, in the problem formulation (and in the implementation to solve it) the merit of a set of candidates is defined as *additive*: it is the sum of the individual merits of candidates selected. Since the merit of B is already completely counted within the merit of A, the two candidates should not be both selected (and their merit should not be counted twice). Note that, if the overlap is instead only partial – as is the case for candidates A and C, which in this example share a call to a single function – the two merits are correctly modeled separately, as it can be identified how many calls to the “shared” functions come from within candidate A, and how many come from within candidate B. Hence, partial overlap does not constitute a conflict.

The problem formulation of *Accel Selection* borrows from that found in Chapter 1. It has however an important difference. Since in Section 1.2 Regions are targeted within a the control-flow graph of a single functions as candidates for acceleration, no overlaps are allowed within the same selection. Conversely, in the current section, subgraphs of the function-call graph are considered, hence allowing solutions including partially overlapping AccelCands.

### 3.1.5 Cost and Merit Estimation

Herein, the abstract cost and merit, which are automatically computed from source code, are detailed (Figure 3.2, B). As the goal of the presented framework is to select the most performing candidates *in advance of their optimization*, AccelSeeker considers their default implementations, e.g., ones where no function is inlined and no loop is unrolled. High-performance implementations will likely have greater resource requirements, in turn potentially requiring to discard some of the selected AccelCands. Nonetheless, these additional design decisions will be performed within the limited scope of the candidate set retrieved by AccelSeeker (as opposed to the whole design) thereby easing the ensuing effort.

Architectural characterization.

AccelSeeker bases its estimations on few parameters characterizing the target platform. Being them only related to the modeled architecture, but independent from the application, the characterization represents a one-time effort for a given hardware target. For the experiments in Section 3.1.8, this task was performed by employing a series of micro-benchmarks, synthesized on a Zynq Programmable System-on-Chip (PSoC). The methodology, however, is not limited to this target. On the contrary, it can be adapted to different computing architectures (e.g.: ASIC implementations) by measuring 1) the area and critical path of single operators (adders, multipliers, etc.), 2) the overhead entailed by initiating an acceleration invocation, 3) the time required to transfer inputs and outputs and 4) the resources employed to realize accelerator-memory links (realized by default as master axi ports in Zynq systems).

Cost Estimation.

The cost  $C()$  of an AccelCand is computed as the sum of its estimated logic and memory real estates. Regarding logic, the sum of the required resources (independently for look-up tables and DSP blocks) is performed of the arithmetic operations present in its top function. If function calls are present, then recursively the area of the called functions is taken into account as well. Furthermore, mimicking the default implementation of Xilinx PSoC accelerators, the addition of the cost of the logic required by a master axi port for each array present in the AccelCand parameters list. Then, the memory area is derived from the size of the arrays storing the input/output and intermediate values required by the accelerator. The I/O size is determined by analyzing the elements in the parameters list of the candidate top function, while the memory required for intermediate



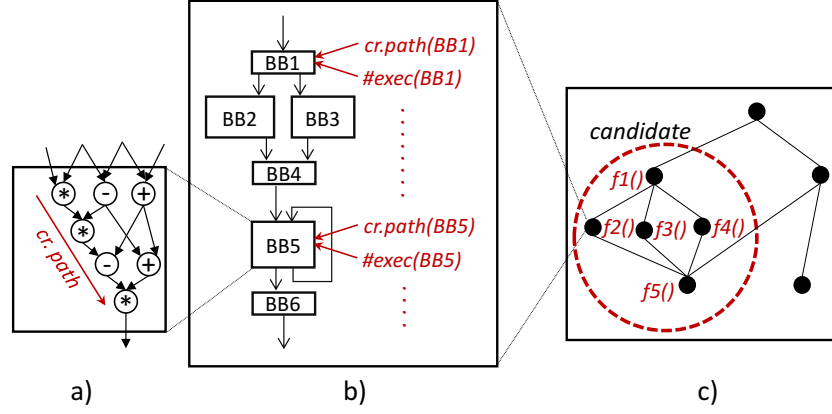


Figure 3.5. Estimation of hardware computation times at the basic block (a), function (b) and AccelCand (c) levels.

values is derived from the variable declarations in each candidate, ultimately determining the number of required BRAM blocks. In line with the limitation of HLS tools, dynamic memory allocations are not supported.

#### Merit Estimation.

The merit  $M()$  of an AccelCand is expressed in terms of the number of clock cycles that are saved by implementing it as a hardware accelerator instead of executing it in software. In turn, the estimation of hardware run times must account both for computation bounds and host-accelerator communication overheads. The latter are retrieved by considering the number of required memory accesses, scaled by an architecture-specific factor.

To assess the computation time of candidates in hardware, a bottom-up approach was carried out, as also exemplified in Figure 3.5. First, the maximum propagation delay of each of the Basic Blocks (BBs) present in an AccelCand (both in the top function and in its callees) is computed. This is achieved by traversing their DFGs and accounting for the operations delays, thus retrieving the longest input-to-output paths (Figure 3.5a). Critical paths of BBs are then expressed in clock cycles, dividing the propagation delays with the period of the system clock. By multiplying the critical paths with the number of executions of each BB, the associated workload is computed. Finally, an estimate of the computation time of an AccelCand is the sum of the workloads of its constituent BBs (Figure 3.5b-c).

Software run-times are estimated in a similar fashion, but instead of comput-

ing critical paths at the BB level, the sum of the latency (in clock cycles) of all its constituent operations is computed, thus modeling that these are processed sequentially in software.

From the gathered data, the merit of an AccelCand  $i$  is computed as follows:

$$M(i) = [T_{sw}(i) - (T_{oh} + \max(T_{Hw}^{comp}(i), T_{Hw}^{comm}(i)))] \times n_{exec}$$

where  $T_{sw}(i)$  is the AccelCand run-time in software,  $T_{oh}$  is the fixed overhead required to configure and start the hardware acceleration,  $T_{Hw}^{comp}(i)$  and  $T_{Hw}^{comm}(i)$  are the run-times when  $i$  is hardware-accelerated, assuming its performance is either computation or communication bound. Finally,  $n_{exec}$  is the number of times the AccelCand is executed in the application.

### 3.1.6 Compiler Analysis

AccelSeeker is implemented as a compiler pass within the LLVM 3.8 [38] infrastructure as seen in Algorithm 5. The resulting implementation comprises methods for the identification and analysis of the AccelCands (Figure 3.2, A), for the estimation of their merit and cost (Figure 3.2, B), and for their selection (Figure 3.2, C). Further details are provided on how the data needed in these phases is retrieved using LLVM IR-level analysis.

*AccelCands identification and analysis.* For the generation of the call graph, every function is annotated with caller/callee relationships; the call graph is subsequently traversed to identify all valid AccelCands as defined in section 3.1.3. At this level information is detected regarding the overlapping of such AccelCands, needed for the creation of the conflict graph, and for subsequent selection. The control flow graph of each candidate, and the data flow graph of each basic block are extracted so that they can be used as input for the cost and merit calculation, according to the method already detailed above.

*Execution Frequency.* The number of invocations of each candidate as well as the execution frequency of each basic block in each candidate is retrieved via LLVM with dynamic profiling. A profiling-via-instrumentation routine is used, which requires the generation of an instrumented version of the code, and then enables the obtained frequencies to be annotated back to the IR level.

*SW Latency Estimation.* The SW latency estimation is being computed by an implemented function within the LLVM analysis pass according to the number of instructions included in a given candidate, as well as the number of instructions included in function calls, if any. The frequency of execution of each Basic Block that the instructions reside is taken into account as well, for a single invocation of the candidate.

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**Algorithm 5** LLVM Analysis Pass - Cost and Merit Estimation
 

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**Input:** Application written in C, C++

**Output:** Candidate List with Estimated Merit and Cost

```

1: function RunOnModule(M)
2:   RunOnFunction(F)
3:
4: function RunOnFunction(F)
5:   /* Merit Estimation */
6:    $T_{SW} = \text{getSWLatencyEstimation}(F)$ 
7:    $T_{HW}^{comp} = \text{getHWLatencyEstimation}(F)$ 
8:    $n_{exec} = \text{getNumberOfInvocations}(F)$ 
9:    $T_{HW}^{comm} = \text{getIORequirements}(F)$ 
10:
11:  /* Cost Estimation */
12:   $A_{HW} = \text{getHWAreaEstimation}(F)$ 
13:   $A_{MAXI} = \text{getMAXIAreaEstimation}(F)$ 

```

---

*HW Latency Computation Estimation.* Conversely, the HW latency estimation takes place by estimating the latency of the string of instructions that have been characterized for the HW implementation that is being targeted. The implemented function estimates the HW latency of each Basic Block as the critical path of the Basic Block multiplied by its respective execution frequency. The total HW latency of a candidate is estimated as the sum of all estimated HW latencies of all Basic Blocks included in the candidate. Both the SW and the HW estimation take place in a bottom-up fashion, by first performing the estimations of the leave candidates and moving onwards to the ones containing calls to others.

*HW Latency Communication Estimation.* In order to take into account the memory latency overhead due to data exchange between the implemented HW accelerators and main memory ( $T_{HW}^{comm}(i)$ ), I/O requirements for each AccelCand are estimated within the LLVM framework by retrieving the parameter list of each candidate and obtaining the data requirements of each candidate type (e.g. size of array of integers, size of a struct etc).

*Area of HW Logic Estimation.* The cost of the candidate is estimated as the total area resources required. The area estimation of logical units is carried out by accounting for the Look Up Tables (LUTs) and the DSPs of characterized operations within a single Basic Block, and subsequently summing up all the resources of all Basic blocks included in a candidate.

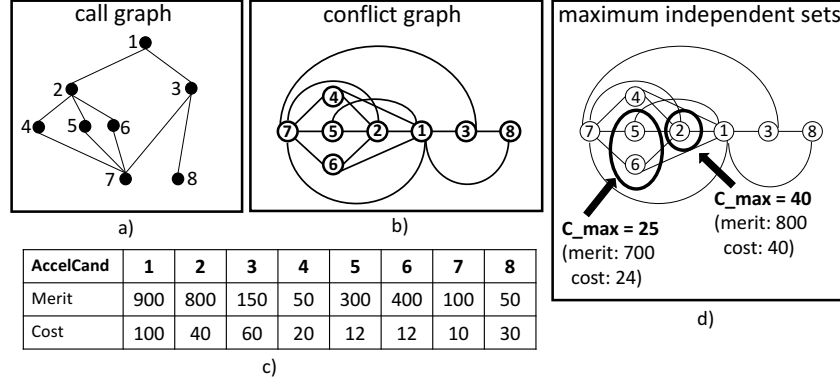


Figure 3.6. a) An example call graph with eight nodes, and hence eight AccelCands, and b) the corresponding conflict graph. c) Given example merits and cost values associated to the eight AccelCands, d) and given a maximum tolerated cost  $C_{max}$ , maximum independent sets that solve problem Accel Selection are shown. A maximum independent set maximises merit, while not exceeding the given cost, and not including conflicts.

*Area of Master AXI ports Estimation.* To account for the HW resources required for a Master AXI port, the parameter list of each candidate is analyzed. Every array identified accounts for extra logical units (LUTs), contributing to the total area.

### 3.1.7 Selection Algorithm

Solving the *Accel Selection* problem on the function-call graph of the application corresponds to solving the *independent set problem* on the resulting conflict graph. The conflict graph, in fact, expresses which pairs of AccelCands are in conflict; thus, an *independent set* of the conflict graph satisfies condition 3 of the *Accel Selection* Problem.

An algorithm is therefore implemented that recursively explores the independent sets of the conflict graph, similarly to the Bron-Kerbosch algorithm [10], and that returns the set  $S$  with the highest merit  $M(S)$  (hence satisfying condition 1 of the problem formulation) and whose cost  $C(S)$  does not exceed a user-given maximum cost  $C_{max}$  (hence satisfying condition 2 of the problem formulation). This returned set represents the optimal solution to the *Accel Selection* Problem.

An algorithm solving an independent set problem is of course one of non-polynomial complexity. Our exact implementation is still able to find the optimal solution for the experiments in this paper in a matter of milliseconds, even for the

considerable dimension of the function-call graph of the application considered (a function-call graph of 63 nodes, as detailed in the experiments section).

An example of selection can be seen in Figure 3.6. First the example call graph is depicted, which has 8 nodes and hence 8 AccelCands, each *rooted* in one of the 8 nodes. The eight corresponding AccelCands are not depicted in this figure, but some of them can be seen in Figure 3.4a: for example, the AccelCand rooted in node 2 is depicted there and labelled A, the AccelCand rooted in node 3 is also depicted in the same Figure and labelled B, etc. Figure 3.6b depicts the complete conflict graph corresponding to this example. As can be seen, candidate 1 (corresponding to the whole graph) is in conflict with all other candidates, candidates 2 and 3 are not in conflict (they only overlap in function 7) etc. Now, given example values of cost and merit for each candidate (in Figure 3.4c), the maximum independent set is found in the conflict graph, which maximises the sum of merits of the candidates selected, does not overcome a maximum sum of cost, and does not include conflicts. Two examples (for  $C_{max} = 25$  and for  $C_{max} = 40$ ) are shown in Figure 3.4d.

### 3.1.8 Experimental Results

### 3.1.9 Conclusions

AccelSeeker was presented, a framework for assisting system architects during the early design phase of hardware-accelerated systems. By automatically assessing the potential speedup of different hardware acceleration choices, in their default implementation, as well as the hardware resources they demand, AccelSeeker allows architects to pinpoint the code sections that are worthy targets for further, more detailed analysis and optimization. AccelSeeker performs the identification of candidate accelerators, as well as their area and speedup estimations, through compiler analysis passes implemented within the LLVM compiler, without requiring lengthy and detailed evaluations of each acceleration candidate individually. It then automatically selects the set of candidates that maximize estimated speedup under a given resource constraint. Experimental evidence highlights that the hardware/software partitioning selected by AccelSeeker vastly outperforms choices that are solely based on profiling information.

## 3.2 EnergySeeker: Accelerators for Energy Efficiency

### 3.2.1 Motivation

eterogeneous Computing calls for specialized Hardware (HW) that can serve the purpose of both accelerating applications and saving significant amounts of energy. Vivado High Level Synthesis (HLS) [79] is one of the state-of-the-art, HLS tools that are used to build accelerators and synthesize parts of software applications to HW. Such HLS and HW Description Language (HDL) tools, however efficient, are far from optimal since they require a lot of manual decisions from the programmer's part. When facing a tight area budget, the decision of which parts of the computation to accelerate or which parts to materialize in low power HW accelerators is not trivial. It requires a deep understanding of the Software (SW) application and its characteristics, as aspects of computation intensity and memory management are complex and hard to identify.

Furthermore, the long time required for hardware synthesis, coupled with the huge space of alternative implementations exposed by real-world applications, limit in practice the number of candidate choices that can be considered manually by a designer before HW/SW partitioning is settled. Simulation tools such as Aladdin [66] have been introduced that allow orders-of-magnitude faster evaluation of the performance of hardware-accelerated applications. These frameworks are not generating a functional hardware implementation, but only provide an representation of the run-time characteristics, the energy requirements and of the required resources of a selected target for acceleration. Nonetheless, these simulation tools still require considerable manual effort to set up experimental environments, a task that must be repeated for every considered candidate.

In order to address these challenges we present EnergySeeker, a methodology to estimate the suitability of acceleration candidates from application source code, subsequently allowing their automatic identification and selection. EnergySeeker, implemented within the LLVM [38] compiler infrastructure, first provides a measure of the cost (required resources) and merit (potential energy saved) of candidate accelerators, and then selects the set that maximizes the estimated energy efficiency gain, within a specified HW resources budget. The use of EnergySeeker could guide HW engineers in the early design phases, highlighting which parts of computation are power-hungry and should be targeted for the synthesis of low-power accelerators, and which parts, instead, are not likely to lead to any significant energy savings. The reason for the latter would be that the computation time in HW could be drastically increased compared to the one in SW (e.g. they have a high memory communication overhead) or that the HW

resources required are so energy costly that they match, or exceed, the respective energy requirements of the SW-only side.

### 3.2.2 Related Work

A large body of research work has been dedicated to energy efficiency in heterogeneous computing systems. In [46] [18] the authors present a clustered many core computing system with tightly coupled OpenRISC 32-bit processing elements. Energy efficiency is achieved through parallelism, yet it is stated that more attention to the memory hierarchy and more specialization in HW realizations would improve both performance and energy efficiency in their proposed system. Applied on small kernels, ultra-low-power RISC-V cores result in a slowdown but offer improvements in energy savings, when long idle periods are present in Internet of Things (IoT) applications [64]. Convolutional Neural Networks (CNNs) are present in IoT devices as well. In [59] a dedicated Hardware realization is suggested to carry out the execution of CNNs on a 65-nm SoC, and thus achieve reduced energy consumption. In none of these instances, though, identification, estimation or selection of the parts of execution to be synthesized in Hardware are challenged.

In [60] an automated methodology that performs estimation of performance and applies optimizations on Hardware accelerators for low-power Deep Neural Networks (DNNs) is presented. Aladdin simulator [66] is used to perform the power requirements estimation and the final implementation is validated on a 40-nm CMOS technology. Furthermore, Machine Learning accelerators: a Support Vector Machine accelerator and an Active Learning Data Selection accelerator, have been coupled with a low-power processor [39] to run medical applications and minimize the power requirements.

HW/SW partitioning has been investigated [78] in order to achieve better energy efficiency in the Advanced Encryption Standard (AES) algorithm by using the OpenSSL [58] library. According to the data blocks size of the encryption an automated method partitions incoming encryption tasks to Hardware implementations or extended Instruction Set implementations of the existing Software processor. These methodologies focus on implementing energy efficient HW realizations on a) specific applications while b) automating some of the processes of identification, estimation and selection of the parts of the execution to be materialized in Hardware. Our methodology instead is general enough to accept any application as input written in C or C++ while fully automating all three processes required to identify and select the most suitable Hardware accelerators that would lead to enhanced energy efficiency.

Finally in [27] an automated methodology that performs estimation on performance and power for SW/HW partitioning is suggested. Given a functional C/C++ description and user defined HW/SW mapping, an estimation is provided and subsequently a design space exploration and optimization phase in order to improve a given configuration. Contrary to our methodology, the selection process of the Hardware candidates is not automated and an area resources budget is not being taken into account.

### 3.2.3 Experimental Results

## 3.3 Conclusions



## Conclusions



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