

A Genetic-Algorithm-Based Approach to the Design of DCT Hardware Accelerators

MARIO BARBARESCHI and SALVATORE BARONE, Department of Electrical Engineering and Information Technologies, University of Naples Federico II, Italy

ALBERTO BOSIO and MARCELLO TRAIOLA, Institut des Nanotechnologies de Lyon, École Centrale de Lyon, France

JIE HAN, Department of Electrical and Computer Engineering, University of Alberta, Canada

As modern applications demand an unprecedented level of computational resources, traditional computing system design paradigms are no longer adequate to guarantee significant performance enhancement at an affordable cost. Approximate Computing (AxC) has been introduced as a potential candidate to achieve better computational performances by relaxing non-critical functional system specifications. In this paper, we propose a systematic and high-abstraction-level approach allowing the automatic generation of near Pareto-optimal approximate configurations for a Discrete Cosine Transform (DCT) hardware accelerator. We obtain the approximate variants by using approximate operations, having configurable approximation degree, rather than full-precise ones. We use a genetic searching algorithm to find the appropriate tuning of the approximation degree leading to optimal trade-offs between accuracy and gains. Finally, to evaluate the actual HW gains, we synthesize non-dominated approximate DCT variants for two different target technologies, namely Field Programmable Gate Arrays (FPGAs) and Application Specific Integrated Circuits (ASICs). Experimental results show that the proposed approach allows performing a meaningful exploration of the design space to find the best trade-offs in a reasonable time. Indeed, compared to the state-of-the-art work on approximate DCT, the proposed approach allows an 18% average energy improvement while providing at the same time image quality improvement.

CCS Concepts: • **Hardware** → **Circuit optimization**; **Circuits power issues**; *Switching devices power issues*; *Application specific integrated circuits*; *Full-custom circuits*; • **Applied computing** → **Multi-criterion optimization and decision-making**.

Additional Key Words and Phrases: Code Mutation, Generic Algorithm, Approximate Computing Techniques, Design Space Exploration, JPEG, Discrete Cosine Transform.

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1 INTRODUCTION

With the increasingly fast growing amount of information processed by modern computing systems, energy efficiency has become a stringent requirement. Therefore, design activities are becoming more challenging and traditional techniques seem more and more unsuitable. An increasingly popular solution is the **Approximate Computing (AxC)** design paradigm. **AxC** exploits the gap between the high accuracy level provided by a computer system and the moderate

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53 accuracy required by a given application to achieve performance gains or energy savings by carefully reducing accuracy.
54 In this perspective, AxC is most effective when applied to applications dealing with redundant data, end-user perceptual
55 limitations, or error resilient algorithms[12, 13, 43].

56 AxC has raised many design challenges due to the variety of **Approximate Computing Techniques (AxCTs)** [26].
57 Each AxCT can introduce different approximation degrees. This leads to a huge amount of different approximate
58 system configurations. Evaluating the approximation impact on the output quality at the application level is often
59 quite time-consuming and not trivial. Indeed, it is usually achieved through simulations or executions of the whole
60 approximate application [12] to ensure that the quality specifications are met. An additional challenge is related to the
61 lack of a general automation tool and a methodology for the **Design Space Exploration (DSE)**.
62

63 One of the main fields of application for AxC is image processing: imperceptible reduction of image quality can
64 lead to important computational resources savings. Most of the research work focuses on the JPEG compression,
65 either considering the algorithms as a whole or its individual computational steps. Concerning the design of hardware
66 accelerators, researchers focused on the approximation of **Discrete Cosine Transform (DCT)** accelerators, mainly
67 targeting figures of merit such as circuit complexity, delay, area and power dissipation [1, 2, 11, 21].
68

69 Unfortunately, the effect of the different approximation techniques and relative configurations (i.e., approximation
70 degrees) are only analyzed individually and without a supporting methodology.
71

72 Conversely, in this work, we assess the impact of approximation – for different approximation degrees – on the DCT
73 computation by performing a fully automated DSE. Starting from the DCT algorithm, we first perform an **Abstract Syntax
74 Tree (AST)** analysis to gather information on the operations suitable for approximation. Then, we generate parametric
75 approximate versions. Through the approximation parameters, we can tune the approximation degree. Finally, we build
76 a **Multi-objective Optimization Problem (MOP)** to find the Pareto-optimal values for the aforementioned approximation
77 parameters. To converge towards a Pareto front, we use a **Genetic Algorithm (GA)**. The fitness functions driving the GA
78 are the *quality reduction minimization* and the *gain maximization*. To perform the DSE in an acceptable time, we operate
79 at high-abstraction level, by modeling the hardware approximation gains with an estimation function. We measure the
80 quality reduction by evaluating the JPEG execution over a large image dataset. Finally, after the DSE, we synthesize the
81 obtained approximate DCT configurations on both **Field Programmable Gate Array (FPGA)** and **Application Specific
82 Integrated Circuits (ASICs)** target technologies and demonstrate the effectiveness of the approach w.r.t. the state of the
83 art.
84

85 The remainder of this paper is structured as follows: Section 2 reviews the existing related work; Section 3 provides
86 preliminary technical background, while Section 4 describes in detail the proposed workflow. In order to evaluate the
87 proposed approach, Section 5 describes the experimental setup, reports result and compares our approach with previous
88 studies. Finally, Section 6 draws the conclusions.
89

90 2 RELATED WORK

91 The effectiveness of imprecise computation for error-resilient applications has been demonstrated both for software
92 and hardware components implementing inexact algorithms [12, 43]. The use of approximation is sometimes intrinsic,
93 e.g. in digital signal processing, where analog signals are discretized and quantized. In many scenarios, the voluntary
94 introduction of approximation turns out to be beneficial. For instance, the perceptual limitations of human senses can
95 be leveraged to reduce the data precision, thus to reduce the storage requirements [32] or to improve performances
96 of multimedia and signal processing applications [35]. Likewise, executing iterative-refinement algorithms with a
97 reduced precision of intermediate computation can improve performances, with little or even no effects on the quality
98

of results [31]. Different AxCTs have been proposed in the scientific literature [26]. Some examples use bit-width optimization [19] and loop-perforation [37].

Employing the AxC full potential requires dealing with several challenges. (i) Approximation error assessment: since compliance with output quality constraints has to be guaranteed, approximation error assessment is crucial. The actual error assessment may require the simulation of both exact and approximate applications, which is very costly. Nevertheless, in order to reduce the required effort, an estimation can be obtained by resorting to a representative workload. On the other hand, a different approach, based on Bayesian inference [38, 39], analytical models [10], machine-learning [27], and even neural networks [46] have been proposed in order to avoid simulations. (ii) The choice of metrics for error estimation: they may strongly depend on the particular application. Thus, choosing the right metric, or the right combination of metrics, is of major concern and it is not a trivial task. (iii) Approximate configuration choice: a given application can be approximated in many different ways. Among all the different approximate configurations, choosing those providing the best trade-off between accuracy loss and resource gains is one of the major challenges in AxC. Indeed, low error and high benefits are conflicting goals.

Concerning hardware design, initial approaches – although not efficient nor scalable – addressed the aforementioned issues by manually identifying approximable sub-parts of a given application. Afterwards, there have been several attempts to define systematic and automated approaches. Ranjan et al. [34], for instance, identify combinational sub-circuits – such as arithmetic units – by looking at the Register Transfer Level (RTL) description of the circuit to be approximated, selecting the optimal quality/energy operating-point using stochastic gradient-descent, in order to maximize energy savings. Nepal et al. [28] perform AST manipulations to generate several approximate versions of a given circuit, starting from either behavioral or RTL descriptions. Variant providing the optimal trade-off between accuracy and gains is selected by means of a stochastic greedy algorithm.

The synergetic effects of multiple approximation techniques, acting at different levels, have been also investigated in the scientific literature. In [23, 46], for instance, the precision-scaling technique is exploited in conjunction with voltage-scaling, since the former usually leads to lower circuit delay, paving the way for aggressive voltage scaling, which provides higher impacts on energy savings.

However, all the aforementioned approaches either combine multiple design objectives in a single-objective optimization problem or optimize a single parameter while keeping the others fixed. Therefore, the resulting solutions are centered around a few dominant design alternatives and do not cover the whole Pareto front [42]. Researches in [36, 41] addressed the problem by using Cartesian genetic programming to search for Pareto-optimal approximate circuit implementations requiring progressively fewer hardware resources. Unfortunately, such approaches did not focus on complex systems, rather on arithmetic components, such as adders and multipliers, which are used as building blocks for hardware accelerators.

In [27], circuits from a library of approximate components are selected to generate an approximate accelerator for a given application. On the basis of contributions from single components, machine learning techniques are adopted to estimate the overall quality and hardware cost of the accelerator, without requiring simulations and synthesis. A similar approach has been presented in [10]. A set of analytical models of quality and resource requirements are derived for a library of approximate components. Then these are used to estimate resource needed and accuracy of accelerator designed through high-level synthesis of C language description.

In this work, we focus on a complex system, performing image processing. Specifically, we focus on the JPEG compression. In [2], a framework relying on inexact computing to perform the DCT computation for the JPEG has been proposed. The framework acts on three levels: (i) at the application level, it exploits human insensitivity to high-frequency

variation to use a filter and discard high-frequency components; (ii) at the algorithmic level, multiplier-less fast algorithms are employed for the actual DCT computation on integer coefficients; (iii) at hardware level, rather than using a simple truncation for adder circuits, authors used *Inexact-Adder Cells (IACs)* to compute less significant bits instead of the *Full-Adder Cells (FACs)*. Therefore, firstly the JPEG quantization step is performed only low-frequency components of an image block; thus the high frequency filter implementation comes down to simply setting some DCT coefficients to zero. Then, at algorithmic level, since the DCT is the most effort-demanding step in JPEG, fast DCT algorithms have been used [6–9, 14, 29, 30]. Those algorithms reduce the DCT complexity from $O(N^2)$ to $O(N)$ and require only integer additions. Finally, at the hardware level, three different IAC families are considered in [2], i.e. the *Approximate Mirror Adder (AMA)* [21], the *Approximate XOR-based Adder (AXA)* [45] and the *IneXact Adder (InXA)* [1]. The framework in [2] mainly aims at assessing the joint impact of those three levels of approximation. However, it presents two main weaknesses: (i) approximation is introduced by manually tuning the individual approximation parameters and (ii) output quality is assessed over only four images. In this paper, we propose a methodology to overcome the highlighted problems by defining a systematic and automatic approach for approximate hardware design.

3 PRELIMINARY TECHNICAL BACKGROUND

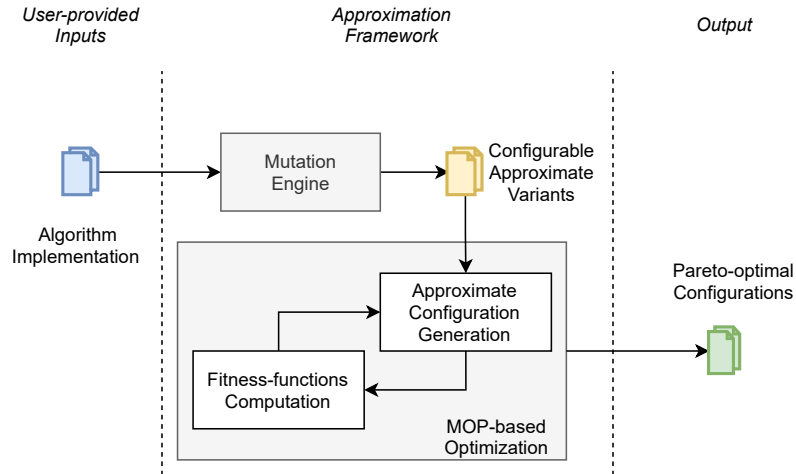


Fig. 1. The workflow of our approach.

In this Section we discuss the proposed workflow, which is depicted in Figure 1, while providing the required technical background. In particular, we firstly detail the proposed generation process of approximate variants. Secondly, we describe how we perform a DSE to converge towards the Pareto-optimal approximate variants.

3.1 Approximate variants generation

Given an algorithm implementation, in order to automatically generate approximate variants while having control on the error, gathering information on the operations suitable for approximation is necessary. We obtain this information through the analysis of the algorithm implementation's AST. Hence, we employ *mutators* [4, 5], to make systematic modifications to the AST, producing altered implementations. Mutators are defined as a set of rules to search and

209 modify the **AST**. The rule definition is generally application independent and does not require the user to know the
 210 algorithm or its specific implementation. In particular, we consider, as an approximation technique, the replacement of
 211 exact sums with approximate ones. We produce a mutator that automatically replaces exact sums with the approximate
 212 counterparts, without knowing the particular algorithm or its implementation. In this way, we obtain *approximate*
 213 *variants* of the original algorithm. Moreover, the approximate sums are configurable in their degree of approximation.
 214

215 The number of variants and the number of configurations grow quickly with the number of operations suitable for
 216 approximation. Consider, for instance, an algorithm implementation with n approximable operations, each allowing k
 217 different degrees of approximation: $\binom{n}{j}$ different approximate variants can be defined by simultaneously approximating
 218 j operations, and k^j different approximate configurations can be defined for each of the variants. Therefore, the total
 219 number of approximate configurations is $\sum_{i=1}^n k^i \times \binom{n}{i}$. At this point, the main challenge is to find values for the
 220 approximation parameters leading to the Pareto-optimal trade-offs between performance gains and accuracy losses, i.e.
 221 to perform a **DSE**.
 222
 223

224 3.2 Design space exploration

225 As mentioned in the introduction, we model the **DSE** as a **MOP**. Basically, a **MOP** consists of a set of *fitness-functions* to
 226 minimize/maximize at the same time and a set of *constraints* to be met, as reported in (1),
 227
 228

$$\begin{aligned}
 229 \quad \Gamma &= \{\gamma_i : A \rightarrow \mathbb{R}, i = 1 \dots k\} \\
 230 \quad \Psi &= \{\psi_j : A \rightarrow \{0, 1\}, j = 1 \dots l\} \\
 231 \quad A &\subseteq \mathbb{R}^n
 \end{aligned} \tag{1}$$

232 where Γ and Ψ are the set of fitness-functions and the set of constraint-functions, respectively. While the functions
 233 of the former set assume values in \mathbb{R} , or its subset, the constraint functions assume either the value 1 or 0 to indicate
 234 that the constraint is or is not met, respectively.
 235

236 Equation (2) describes the set of solutions for (1). For non-trivial **MOPs**, $|X| > 1$, where $|\cdot|$ expresses the size of the
 237 set, i.e. the number of elements it contains.
 238

$$239 \quad X = \{x \in A : \gamma_i(x) \leq \gamma_i(x'), x \vdash \psi_j, x' \neq x, i \in [1, k], j \in [1, l]\} \tag{2}$$

240 Indeed, since different objectives (i.e., fitness functions) often represent conflicting goals, the **DSE** goal is to seek for
 241 a set of equally good solutions being close to the *Pareto-front* (2). Let us consider two solutions, $x, y \in X : x \neq y$, x is
 242 said to *dominate* y i.f.f. (3) holds, i.e. x shows better or equally good objective values than y in all objectives and at least
 243 better in one objective. If a solution is not dominated by any others, it is called a *Pareto-optimal* solution.
 244

$$245 \quad x < y \iff \gamma_i(x) \leq \gamma_i(y) \forall i \in [1, k] \wedge \exists j \in [1, k] : \gamma_j(x) < \gamma_j(y) \tag{3}$$

246 Due to the rapid growth of the size of the solution space as the number of decision variables, fitness functions and
 247 constraints increases, using exact solving algorithms turns out to be very computation-intensive. Consequently, a
 248 variety of heuristics aiming at producing an approximation of the Pareto-front have been proposed in the scientific
 249 literature. Some well known heuristics are hill-climbing, ant-colony [18] and **Evolutionary Algorithms (EAs)**.
 250

251 **GAs**, a subclass of **EAs**, have been largely used in the literature to find Pareto-fronts for **MOPs** [15]. **GAs** are inspired
 252 by and also borrows terminology from the evolutionary theory: *mutation*, *crossover* and *selection* mechanisms cause the
 253 extinction of weak and unfit species while strong ones have greater opportunities to survive and pass their genes to
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future generations. In particular, we resort to **Multi-Objective Evolutionary Algorithms (MOEAs)**. MOEAs operate on a set of *individuals*, called *population*, that evolves and, eventually, converges to a set of Pareto-dominant solutions. Each individual is represented as a *chromosome*, i.e. a data structure encoding the search space. During the evolution process, new offspring is generated either through or in combination of *crossover* and *mutation* [17]. A *crossover* takes two parent chromosomes to produce a new chromosome. Thence, we resort to the **Non-dominated Sorted Genetic Algorithm-II (NSGA-II)**, which is widely employed as a standard approach [16].

It follows an elitist principle, i.e. the elites of a population are given the opportunity to be carried to the next generation. The evolution process starts from an *initial population*, which is typically randomly generated or seeded in areas where optimal solutions are likely to be found, and it goes through the following steps.

At the beginning, the NSGA-II builds a random initial population P_0 of size N , and sorts the latter using *non-dominationsorting*, which will be presented below. Thus, each individual is assigned a *rank* equal to its non-domination level. At the first iteration, usual crossover, mutation and binary tournament selection operator are used to build the Q_0 offspring population.

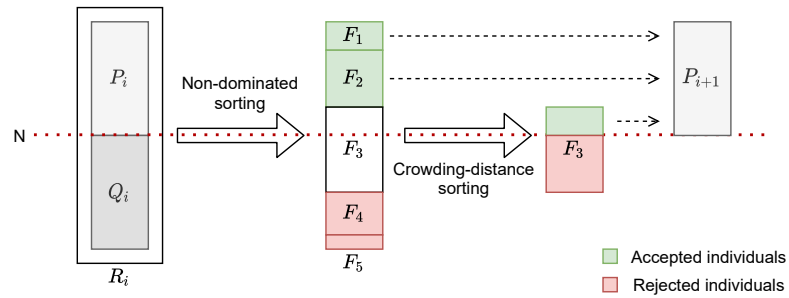


Fig. 2. The NSGA-II selection strategy.

The behavior of the algorithm during a generic i -th iteration is summarized in Figure 2: (i) a combined population $R_i = P_i \cup Q_i$ is formed exploiting crossover and mutation, then sorted on the basis of non-domination, and finally split in subsets $\{F_1, \dots, F_m\}$; (ii) individuals from the highest rank subsets F_1 are preserved in the next iteration; if the amount of surviving individuals from F_1 is less than N , individuals from lower rank subsets $F_2 \dots F_l$ are also preserved, and they survive for the next iteration; (iii) in order to select exactly N individuals from R_i , individuals from the last non-rejected non-domination subset F_l are sorted on the basis of the *crowdingdistance*, and selected using the *crowded selectionoperator* (4).

In order to perform non-domination sorting, for each individual $p \in P$, the NSGA-II computes (i) the non-domination count $n_p = |D_p|$, $D_p = \{q \in P : q < p\}$, i.e. the amount of individuals q which dominate p , and (ii) the set $S_p = \{q \in P : p < q\}$, i.e. the set of individuals q which are dominated by p . Individuals in the highest rank subset of R_i , i.e. F_1 , have $n_p = 0$. Then, for each $p : n_p = 0$, each $q \in S_p$ is visited and n_q is decremented by one. If the term n_q becomes zero, q is placed in the second-highest rank subset of R_i , i.e. F_2 . The procedure is iterated until each subset is fully identified.

Along with convergence to the Pareto-front, it is also desired that an EA maintains a good spread in the obtained set of solutions. In order to preserve diversity, the NSGA-II adopts the *crowdingdistance* as a metric for the density of solutions surrounding a particular solution. Such distance is the average distance of two points on either side of the considered solution, along each of the fitness functions. The computation requires sorting the population M times,

313 according to each fitness function, in ascending order. Each time the population is sorted, the boundary solutions –
 314 i.e. solutions with the smallest and the largest fitness – are assigned an infinite distance, while all other solutions are
 315 assigned with a crowding distance equal to the absolute normalized difference of fitness of the two adjacent solutions.
 316 The overall crowding distance is the sum of individual distances corresponding to each fitness function. Using crowded
 317 distance, the usual definition of Pareto dominance (3) is slightly modified as in (4), where $x \approx y$ denotes x and y do
 318 not dominate each other and $d(\cdot)$ is the crowding distance of a given solution. The *crowdedselectionoperator* preserves
 319 diversity by promoting solutions located in less crowded areas of the solution space.
 320
 321

$$322 \quad x \prec_n y \iff x \prec y \cup (x \approx y \wedge d(x) > d(y)) \quad (4)$$

323
 324 Modeling a specific optimization problem so that it can be resolved by a MOEA is not trivial and there are no general
 325 rules. In our case, as mentioned in Section 3.1, we approximate a number of operations in the DCT algorithm. According
 326 to their approximation degree, each of the operations contributes to error and gains differently. On the one hand, it
 327 may be easy to analytically define a relationship between the degree of approximation and the corresponding gains, by
 328 using properties of the specific approximation. On the other hand, doing the same with the error is not trivial, due to its
 329 propagation through the data-flow. Therefore, fitness functions should be defined case by case. Finally, regarding the
 330 constraints, identifying acceptable variation ranges for one or more genes or identify an error threshold is generally
 331 feasible. For instance, for an n-bits adder the variation interval of the gene could be $\{0, 1, \dots, n\}$ to cover the degree
 332 that goes from “no-approximation” to “totally approximate”.
 333

334 Finally, from a technical standpoint, approximate configurations can be represented by using a vector – i.e. a
 335 chromosome, in the MOEA context – having as many elements – i.e. genes – as operations. The value of each element
 336 represents the approximation degree for the corresponding operation.
 337

338 Being the goal of the DSE to find the Pareto-optimal approximation degree for all the operations, the NSGA-II
 339 provides a suitable tool to perform the exploration of such a large design space, in a reasonable time.
 340
 341

342 4 TOWARDS APPROXIMATE DCT

343
 344 In this section, firstly we discuss the mathematical steps leading to a set of linear equations to compute DCT coefficients
 345 using only additions. These equations are general and independent of the particular DCT algorithm. Secondly, we
 346 present the approximate DCT variants generation and how we model the DSE as a MOP so that it can be resolved by
 347 using the NSGA-II.
 348
 349

350 4.1 Addition-based computation of DCT coefficients

351 As we already mentioned, the DCT computation is known to have $O(N^2)$ complexity and requires resource-intensive
 352 functional units, such as floating-point arithmetic modules. The algorithm proposed in [25] requires 11 multiplications
 353 and 29 additions to compute the one-dimensional eight-point DCT needed by the JPEG compression. It is considered
 354 the most efficient exact algorithm, since the lower bound on the number of multiplications required for such DCT
 355 computation has been proven to be 11 [20]. In order to achieve an additional reduction in resource requirements, authors
 356 of [3] moved parts of the DCT computation to the JPEG quantization step. Furthermore, transformed coefficients can be
 357 scaled and rounded such that floating-point operations can be superseded by integer ones: the resulting algorithms are
 358 significantly faster and they find extensive use in practical applications. However, integer multiplication is still complex
 359 and resource intensive; thus, many low-complexity multiplier-less algorithms have been proposed [6–9, 14, 29, 30].
 360 As in [3], all of these algorithms split the DCT computation into two consecutive steps: the first one is referred to as
 361
 362
 363
 364

Table 1. Comparison among **DCT** algorithms in terms of number of operations

Method	Additions	Multiplications	Shifts	Total operations
DFT (definition)	432	192	0	624
FFT	58	6	0	64
DCT (definition)	56	64	0	120
Arai algorithm	29	5	0	34
BAS08 [7]	18	0	2	20
BAS09 [8]	18	0	0	18
BAS11 [9] (a=0)	16	0	0	16
BAS11 [9] (a=1)	18	0	0	18
BAS11 [9] (a=2)	18	0	2	20
CB11 [14]	22	0	0	22
BC12 [6]	14	0	0	14
PEA12 [29]	24	0	6	30
PEA14 [30]	14	0	0	14

approximate-**DCT**, which involves only integer operations, while the second step is embedded into the quantization and takes advantages of floating-point operations the latter requires. Moreover, they all avoid computing **DCT** coefficients separately or iteratively. Instead, they extensively use matrix algebra and its properties. To show how the above mentioned algorithms work, let X be an input image tile, which is a 8×8 matrix; its two-dimensional **DCT** transform, from now on simply **DCT**, is described by the following equation:

$$F = C \cdot X \cdot C', \quad (5)$$

where C is referred to as **DCT matrix**. C contains the cosine function values at the needed frequencies. The X and the F matrices have the same dimensions. The elements in F represent the **DCT** coefficients as the frequency progressively increases: low-frequency components are closer to the top-left corner, while high-frequency ones are placed close to the bottom-right corner. The C matrix can be split into two matrices, T and D , as reported in Equation (6).

$$F = C \cdot X \cdot C' = D \cdot (T \cdot X \cdot T') \cdot D \quad (6)$$

Different algorithms define T and D in different ways, so the number of computation operations may vary from algorithm to algorithm, as reported in Table 1. Splitting C allows integers-only matrix multiplications. Indeed, T contains only the values $\{0, \pm \frac{1}{2}, \pm 1, \pm 2\}$ and it is orthogonal, i.e. $T' = T^{-1} \Rightarrow TT' = T'T = I$, where I is the identity matrix. Note that multiplying by $\frac{1}{2}$ or 2 comes down to shifting to the right or to the left, respectively; this, at hardware level, is reduced to simple wiring. This means T allows computing the **DCT** using only additions. Nevertheless, the multiplication of D in (6) still requires floating-point operations. In fact, D is a diagonal matrix consisting of values in the $[-1, 1]$ range, with $\{\frac{1}{2}, \frac{1}{\sqrt{2}}, \frac{1}{\sqrt{8}}\}$ being typical values. For this reason, resorting to properties of diagonal matrices allows obtaining the following equation:

$$F = T \cdot X \cdot T' \circ (\text{diag}(D) \cdot \text{diag}(D)'), \quad (7)$$

where \circ is the Hadamard product, i.e. an element-wise multiplication. Thus, the integer null-multiplicative part $T \cdot X \cdot T'$ can be isolated from floating-point operations required by D . Afterwards, floating-point operations can be performed outside the **DCT**, and embedded into the JPEG quantization step, as shown in the following equation.

$$F_Q = [F \oslash Q] = [T \cdot X \cdot T' \circ (\text{diag}(D) \cdot \text{diag}(D)') \oslash Q] = [T \cdot X \cdot T' \circ \hat{Q}] = [(T \cdot (T \cdot X)') \circ \hat{Q}] \quad (8)$$

$$\hat{Q} = (\text{diag}(D) \cdot \text{diag}(D)') \oslash Q, \quad (9)$$

where \hat{Q} in (9) is the *complete quantization matrix* and the \oslash operator is the Hadamard division, i.e. an element-wise division. From (8), it follows $F = (T \cdot (T \cdot X)')$, which means that the approximate two-dimensional DCT transform can be computed using the one-dimensional DCT transform twice, reducing the complexity from quadratic to linear.

The only substantial difference between the different multiplier-less DCT algorithms is the T matrix. Hence, it is straightforward to derive a set of equations to calculate the one-dimensional DCT coefficients. Equations in (10), for instance, refers to the BC12 algorithm [6].

$$\begin{aligned}
 f_0 &= x_0 + x_1 + x_2 + x_3 + x_4 + x_5 + x_6 + x_7 & f_1 &= x_0 - x_7 \\
 f_2 &= x_0 - x_1 - x_2 + x_3 + x_4 - x_5 - x_6 + x_7 & f_3 &= x_4 - x_3 \\
 f_4 &= x_0 - x_3 - x_4 + x_7 & f_5 &= x_5 - x_2 \\
 f_6 &= x_2 - x_1 + x_5 - x_6 & f_7 &= x_6 - x_1
 \end{aligned} \tag{10}$$

Furthermore, some terms – for instance $(x_0 + x_7)$ – are involved in the computation of multiple f_i coefficients, which allows to further reduce the amount of operations.

4.2 Approximate variants generation and MOP modeling

Once the addition-based equations for the DCT coefficients are defined, simple implementations for the DCT computation algorithm can be derived, such as BAS08 [7], BAS09 [8], BAS11 [9], BC12 [6], CB11 [14], PEA12 [29] and PEA14 [30]. Within those, we introduce further approximation by replacing exact sums by configurable approximate ones. Such approximate sums allow setting two parameters, i.e., the **Number of Approximate Bits (NAB)** and the type of adder hardware cell to use (namely, a classic **FAC** or an **IAC**). This was the same approach adopted in [2]. However, while in [2] the approximation was manually introduced, we propose to automate the replacement process by considering the **AST** of the algorithm implementation. Moreover, in [2], approximation parameters were tuned manually and one at a time, keeping the others at fixed values. Unfortunately, this does not necessarily lead to Pareto-optimal solutions. Conversely, we propose to find the optimal value for the approximation parameters through an automatic **DSE**. We model the **DSE** as a **MOP**, so that it can be resolved by using a **MOEA**. In the next subsections, we detail the **MOP** modeling.

4.2.1 Chromosome encoding. To enable the **MOEA** to resolve the MOP, each approximate configuration is modeled as a chromosome. Chromosome's genes represent the two aforementioned approximation parameters, i.e., the **NAB** value and the type of adder hardware cell to use. Thus, if N_{op} is the number of addition required, each chromosome is composed of $2 \cdot N_{op}$ different genes. Chromosomes are provided with an additional gene representing the approximation degree for the high frequency filter. Thus, each chromosome is composed of $2 \cdot N_{op} + 1$ genes.

Let us consider an approximate configuration from the population and its corresponding chromosome. Suppose a mutation occurs: depending on the particular gene being altered, a different approximate configuration will be generated. The latter will differ from the origin one in (i) the number of approximate bits for a certain sum operation, (ii) the adder cell to be used for a certain sum, or (iii) the amount of frequencies discarded by the high frequency filter. Despite the fact that it involves multiple genes and, hence multiple characteristics of a configuration, the same reasoning also applies to crossover. The **NSGA-II** selection operator will determine which of the individuals will survive, on the basis of fitness. Fitness functions driving the **DSE** are detailed in the following sections.

469 4.2.2 *Error fitness-function.* For the MOEA to be able to evaluate the error entailed by the approximations, we need to
 470 define and error fitness-function to minimize.

471 In [1], the authors computed, through exhaustive simulations, the error-rate for different numbers of erroneous bits,
 472 for IAC adders. Unfortunately, this kind of measurements is not suitable for complex algorithms, such as the DCT, as it
 473 does not take into account the error propagation. Therefore, we resort to the whole JPEG compression, performed on a
 474 representative data set, to estimate the error. Section 5 discusses the experimental setup in more detail.

475 Regarding error metrics, in [2] the Maximum Difference (MD), the Average Difference (AD), the Mean Square Error
 476 (MSE) and the Peak Signal-to-Noise Ratio (PSNR) have been considered. Unfortunately, since all of them consider single
 477 pixels, these metrics turn out to be too sensible to noise [33]. Therefore, they are not particularly suited to evaluate the
 478 effectiveness of approximations on image processing algorithms. To overcome this issue, we resort to the Structural
 479 SIMilarity (SSIM) [44] to evaluate differences among images. Its formal definition is reported in Equation (11), where X
 480 and Y are two sets of data (i.e., the images), μ_X and μ_Y are their mean values, σ_X^2 and σ_Y^2 are their variances, σ_{XY} is their
 481 co-variance, L is the value range in which elements of X and Y can vary, and k_1 and k_2 are tuning parameters (typically
 482 equal to 0.01 and 0.03, respectively). Values of $SSIM(X, Y)$ span in the range $[-1, 1]$. Values of $SSIM(X, Y) \approx 1$ mean
 483 that X and Y are structurally similar, while values of $SSIM(X, Y) \approx 0$ mean that there is no similarity between the two
 484 images. Values smaller than zero are meaningless [44].

$$485 \quad SSIM(X, Y) = \frac{(2\mu_X\mu_Y + k_1) \cdot (2\sigma_{XY} + L \cdot k_2)}{(\mu_X + \mu_Y + k_1) \cdot (\sigma_X^2 + \sigma_Y^2 + L \cdot k_2)} \quad (11)$$

486 The effectiveness of image quality assessment increases if (11) is applied locally rather than globally since (i) image
 487 statistical features are usually highly spatially non-stationary, (ii) image distortions, which may or may not depend on
 488 the local image statistics, may also be space-variant, and (iii) only a local area in the image can be perceived with high
 489 resolution by human observers [44]. Since, in practice, a single overall quality measure of the entire image is required,
 490 the Mean SSIM (MSSIM) from Equation (12) is adopted. There, X and Y are the reference and the distorted images,
 491 respectively, x_j and y_j are the image contents at the j -th local window; and M is the number of local windows in the
 492 image. Typically, the MSSIM index is computed considering 11×11 Gaussian weighted circular windows rather than
 493 on 8×8 square tiles [44].

$$494 \quad MSSIM(X, Y) = \frac{1}{M} \sum_{j=1}^M SSIM(x_j, y_j) \quad (12)$$

495 As SSIM, the lower the MSSIM index the lower the similarity between X and Y sets; thus, in order to define a suitable
 496 fitness-function for the MOEA to minimize error, we adopt the Structural DISSIMilarity (DSSIM) – $DSSIM(X, Y) =$
 497 $1 - MSSIM(X, Y)$. In particular, we compute the DSSIM between a standard JPEG compressed image X and an image
 498 Y which is obtained by using a certain approximate configuration of a given approximate algorithm. Both X and Y
 499 originate from the same non-compressed source image. We perform this operation for several images and use the
 500 average DSSIM as final error fitness-function.

501 4.2.3 *The reward fitness function.* In order to accurately assess resource savings, area, power consumption and maximum
 502 clock speed should be measured. Unfortunately this would require the synthesis and simulation of large hardware
 503 designs. Thus, performing an hardware synthesis for each design explored in the DSE is a very time-consuming process.
 504 Therefore, we resort to a gain estimation to drive the DSE. In particular, we estimate the gain (from now on ‘reward’)

from the number of transistors required to implement an inaccurate cell, using the data from [2]. This constitutes a good predictor for area and power gains, as experimentally evaluated in Subsection 5.2, for both ASIC and FPGA. For convenience, in Table 2 we report, from [2], the number of transistors required to implement inaccurate cells of the mentioned IACs. Concerning the operating frequency, in this case the approximation does not entail any change, as further explained in Subsection 5.1.

Table 2. Transistor count from [2] for inexact-adder cells

Cell	Full Adder	AMA1	AMA2	AMA3	AMA4	AXA1	AXA2	AXA3	InXA1	InXA2	InXA3
Transistors	58	20	14	11	14	8	6	8	6	8	6

Let us detail the reward function. Let N_{op} be the number of operations required to compute the single-dimensional DCT and let nab_i be the NAB for the i -th addition. We compute the total number of saved transistors as

$$\sum_{i=0}^{N_{op}-1} nab_i \cdot (T_{FA} - T_{IAi}), \quad (13)$$

where T_{FA} and T_{IAi} are the number of transistors required by the FAC and the i -th IAC, respectively. Finally, since the number of additions required by each algorithm varies, we use a normalized measure, as reported in the following equation:

$$\rho = \frac{1}{2 \cdot N_{bits} \cdot N_{op} \cdot T_{FA}} \sum_{i=0}^{N_{op}-1} nab_i \cdot (T_{FA} - T_{IAi}) \quad (14)$$

where N_{bits} and N_{op} are the number of bits on which each of the sums is expressed and the number of sums required for the DCT computation, respectively.

Thanks to the so-defined reward function estimation, we are able to execute the DSE without performing any time-consuming syntheses of the approximate designs explored. This allows us to explore a lot more approximate designs, which would be infeasible otherwise. As a result of the DSE, we obtain the Pareto-optimal approximate designs in terms of accuracy and gains. Hence, we actually synthesize these designs to a particular technology.

5 EVALUATION AND EXPERIMENTAL RESULT

In this section, we firstly describe the DCT algorithm hardware implementation. Secondly, we validate the suitability of the chosen reward fitness function. Thirdly, we show our experimental setup and related result. Finally, we perform a comparison with previous work.

5.1 DCT algorithm hardware implementation

In order to be able – at the end of the DSE – to measure the final gains, we encoded all the above mentioned DCT algorithms in VHDL. Such implementations guarantee high flexibility: they handle the configuration of both the type of adder cells to use for each addition and the number of bits to approximate (NABs). This allows the synthesis of any solution eventually found in the DSE process. VHDL implementations follow Equation (8), which allows splitting the two-dimensional DCT into two consecutive one-dimensional DCTs, separated by a transposition block, which transposes the signals. The transposition block implementation in hardware comes down to being just wiring.

A block schema of the two consecutive one-dimensional DCTs is depicted in Figure 3: X_1, \dots, X_7 represent the rows of the image tile being transformed, while F_1, \dots, F_7 represent the rows of the transformed block. A RTL schema of the

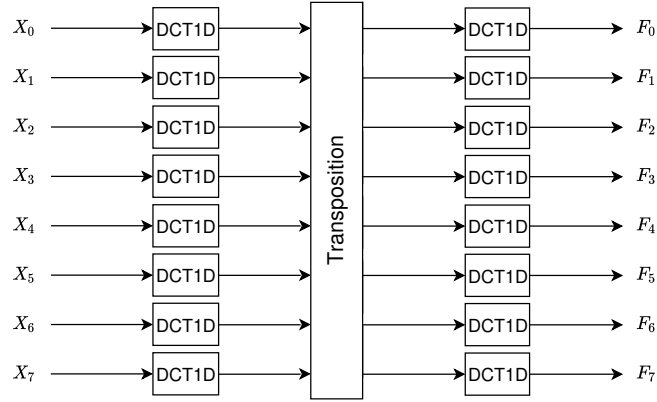


Fig. 3. RTL block schema for the BC12-2D hardware implementation

single dimensional **DCT** computation block (DCT1D) is shown in Figure 4; without loss of generality, the schema refers to BC12 [6], since the differences between different algorithms are negligible.

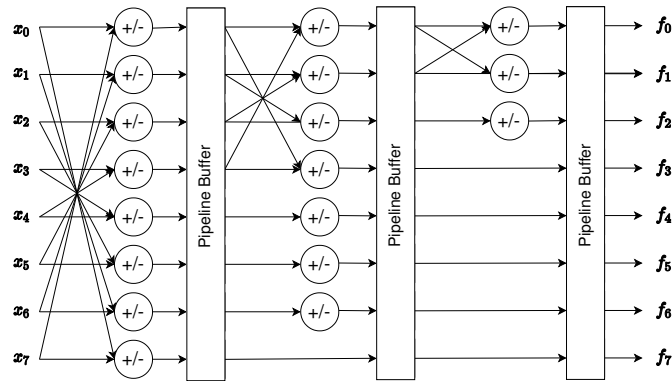


Fig. 4. RTL block schema for the BC12-1D hardware implementation

The architecture of the one-dimensional **DCT** computing block is pipelined, with pipe registers separating the adders needed for the partial-sums computation. The one-dimensional **DCT** has three clock cycles latency, thus the whole two-dimensional **DCT** block is six clock cycles latency. Each of the partial sums is performed using a configurable approximate adder. The scheme of a configurable approximate adder is depicted in Figure 5: it is a ripple-carry adder whose least significant bits are computed by **IACs**, while the most significant ones are computed by classical **FACs**. The number of approximate sums, i.e. **IACs**, is configurable by means of the **NAB** parameter.

The **DCT** is computed on 8×8 image tiles, each one made of three different color channels. Each element's value spans from 0 to 255. For this reason, each one of the single-dimensional **DCT** output terms can be expressed, at most, as the sum of eight elements. Therefore, the maximum value for the single-dimensional **DCT** terms is $8 \times 255 = 2040 < 2048 = 2^{11}$. As a consequence, the two-dimensional **DCT** output terms can have a maximum value of $8 \times 2040 = 16320 < 16384 = 2^{14}$. As a result, 14 bits turn out to be sufficient to represent the **DCT** frequency coefficients. It is also worth highlighting that replacing **FACs** with **IACs** leaves the overall structure unchanged. As anticipated in Subsection 4.2.3, this means that the

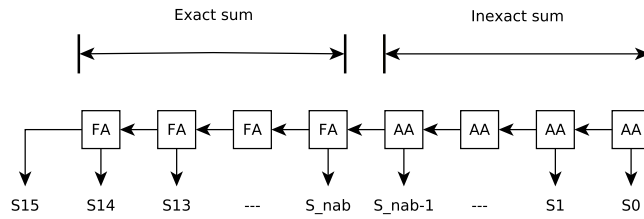


Fig. 5. Inexact ripple-carry adder

approximation does not affect the latency nor the operating frequency of the device. Therefore, we can conclude that the maximum frequency of operation depends entirely on the target technology. In order to compute the maximum operating frequency for our hardware designs, we performed a preliminary **FPGA** and **ASIC** synthesis, varying **NABs** and **IACs**, while targeting a Xilinx Zynq-7020 **FPGA** and a 65nm **Fin Field-Effect Transistor (FinFET)** technology. Preliminary synthesis confirmed our aforementioned observation: approximation does not impact on the operating frequency, which depend only on the target technology. Synthesis tools reported a maximum operating frequency in the 250MHz and 770MHz range for **FPGA** and **ASIC**, respectively.

5.2 Reward fitness function evaluation

As mentioned in Section 4.2.3, the reward fitness function described in Equation (14) performs an estimation of the approximate designs gains. As main advantage, such an approach enables an immediate gain estimation, without the need to resort to highly time-consuming circuit syntheses. This paves the way to a much wider exploration, thus to a more thorough **DSE**.

Despite of such an advantage, it is not trivial to reveal the correlation between transistor reduction and **FPGA** resource savings. In order to validate Equation (14) as suitable gain estimator for **FPGA**, we conducted two preliminary experiments: (i) we synthesized several configurations of approximate adder by varying the **IAC** to be used and **NABs** in order to appreciate programmable resources overhead; (ii) we further synthesized several approximate configuration of **DCT** hardware accelerators on both a Xilinx Zynq-7020 **FPGA** and 65nm **FinFET** [22] technologies. In these experiments, we did not measure the accuracy, since we are only interested in resource savings.

As reported in Figure 6, there is an appreciable overhead reduction, in terms of occupied **Look-Up Tables (LUTs)**, for any adder using **IAC** as **NABs** increase. Indeed, Boolean minimization process, performed during synthesis, gets more possibilities to reduce the size of circuits in terms of literals, fan-in and number of **LUTs** since truth tables of **IACs** are designed to simplify Boolean expressions [2].

Furthermore, we used different **DCT** algorithm hardware accelerators from the literature (**BAS08** [7], **BAS09** [8], **BAS11** [9], **BC12** [6], **CB11** [14], **PEA12** [29] and **PEA14** [30]). For each accelerator, we gradually increased the **NABs** to observe the corresponding hardware resources trend.

Figures 7 and 8 report the amount of **LUTs** and a comparison between reward estimation given by Equation (14) and the actual reward on **FPGA**, respectively, for each of the considered **DCT** algorithms. As for the former, an appreciable resource overhead reduction is observable; as for the latter, the actual reward trend is sufficiently close to the predicted one. As one can notice, the predicted reward from Equation (14) has the same slope as actual reward trend. Furthermore,

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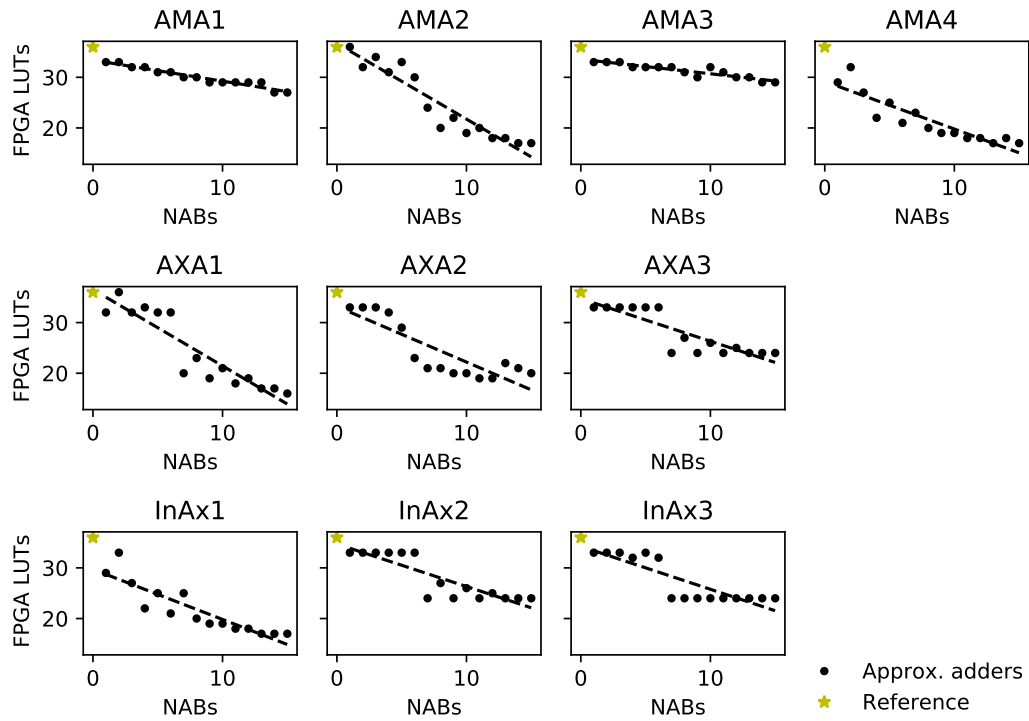


Fig. 6. Area requirements of 16 bits adders on FPGA, varying the NABs

the predicted reward is almost a lower bound for the actual reward, which is significantly relevant in order to achieve a fair DSE.

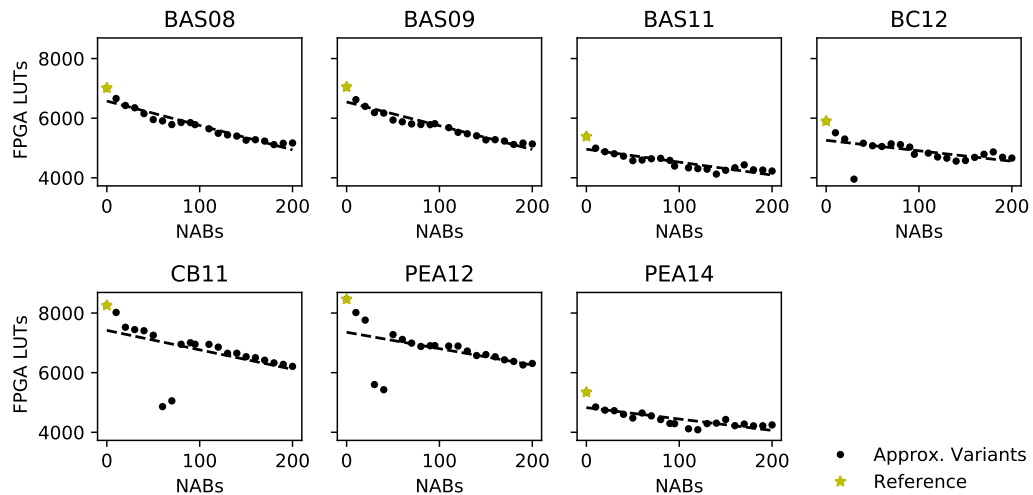


Fig. 7. Area requirements of DCT algorithms on FPGA, varying the NABs

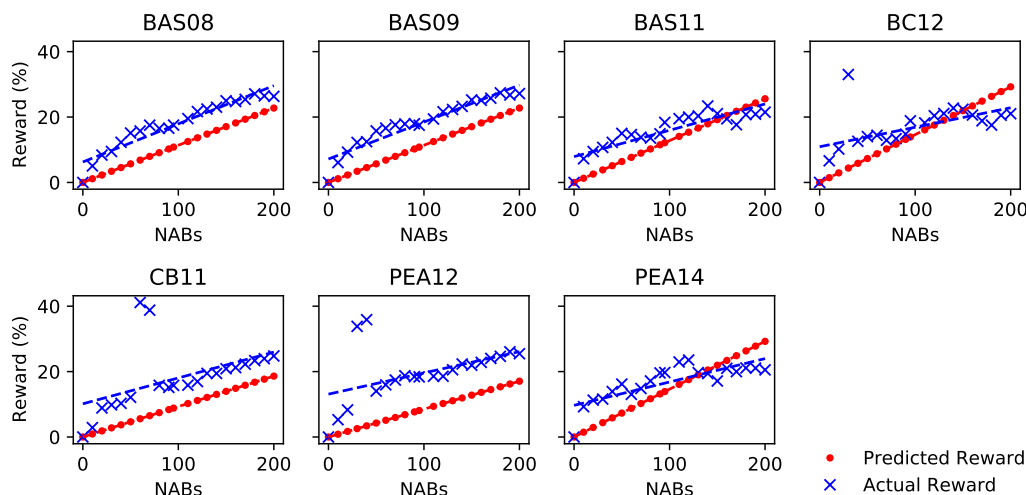


Fig. 8. Expected vs. actual reward for DCT algorithms on FPGA, varying the NABs

Finally, we measured the Mean Absolute Percentage Error (MAPE) between the reward values predicted by Equation (14) and the actual ones from the syntheses, for both FPGA and ASIC. The measured MAPE ranges between [0.33%, 0.65%] and [0.82%, 2.44%], respectively for ASIC and for FPGA. Such low values of MAPE reveal that Equation (14) is an accurate estimator for the involved reward. Therefore, we concluded that Equation (14) is a suitable estimation to forecast approximation gains.

5.3 Experimental Setup

In this section we describe the experimental setup that we used in order to evaluate our approach. We considered seven different DCT algorithms and ten types of IACs. As for the DCT algorithms, we considered BAS08 [7], BAS09 [8], BAS11 [9], BC12 [6], CB11 [14], PEA12 [29] and PEA14 [30]. As for the IACs families, we considered AMA [21], AXA [45] and InXA [1]. The considered DCT algorithms and approximate adders are the same taken into account by the authors of [2].

Figure 9 sketches our workflow as a whole: to speed-up the simulation process, we modeled each of the above mentioned multiplier-less DCT algorithms by using C/C++ implementations straightly derived from equations (8) and (10). Starting from such implementations, approximate variants generation is performed using the Clang-Chimera tool, which is an Clang/LLVM-based C/C++ source-to-source mutation engine part of the IDEA framework [4, 5]. For each DCT algorithm, the Clang-Chimera tool produces mutated sources which allow to configure, for each of the sums, both the NABs and type of adder hardware cell to use (i.e. either FAC or IAC). Furthermore, as mentioned in subsection 4.2, we modeled the DSE as a MOP. The MOP resolution is performed by using the ParadisEO framework, a template-based evolutionary computation library [24]. In order to perform the hardware synthesis, non-dominated solutions from the DSE are employed to configure the VHDL implementation discussed in Section 5.1. Finally, we measure actual gains.

To find a suitable MOEA configuration, we conducted several DSE campaigns with different MOEA parameters. As a result, we deduced two things: (i) to obtain a populous frontier and avoid local sub-optimum, we need to increase

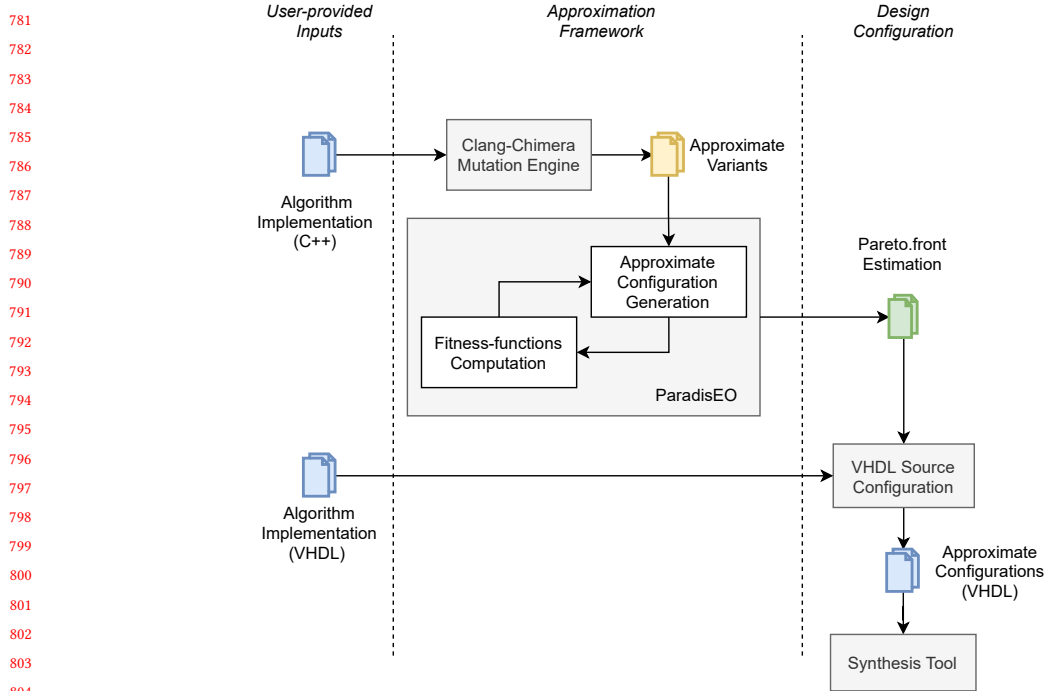


Fig. 9. Actual workflow of our approach.

the initial population size as much as possible; (ii) in order to avoid long-run exploration around local sub-optimum, mutations have to take place frequently. Hence, we set our MOEA parameters as follows: initial population equals to 2000 individuals, mutation and crossover probabilities set to 0.7 and 0.9, respectively. We did not set any maximum error threshold.

Details on the fitness-functions employed for the DSE have been provided in Section 4.2.2 and Section 4.2.3. As mentioned in Section 4.2.2, we resort to the whole JPEG compression, performed on a representative data set, to estimate the error. The considered data set [40], consists of 44 different images, covering a wide set of common features, including among others a flat gray scale, foreground subject with a messy background, and high contrast images.

5.4 Experimental Result

In this Section, we report experimental results. Firstly, we discuss the DSE results. Secondly, we discuss both silicon die area and power consumption reductions we achieved for designs resulting from DSE, synthesized on both FPGA and ASIC.

5.4.1 DSE results. Figure 10 reports the Pareto front provided by the MOEA for all the considered algorithms. The reference is, for each algorithm, its non-approximate implementation, depicted as a gold star. It is important to bear in mind that such algorithms are non-exact DCT versions (see Section 4) and that the JPEG implemented with a non-exact DCT algorithm produces lower-quality images compared to its exact version. For this reason, the reported non-approximate solutions exhibit already some error. Their reward value is zero, since they do not use any IACs, so

they do not achieve any approximation gain according to Equation 14. As envisioned, the graphs highlight increasing expected rewards as the error increases.

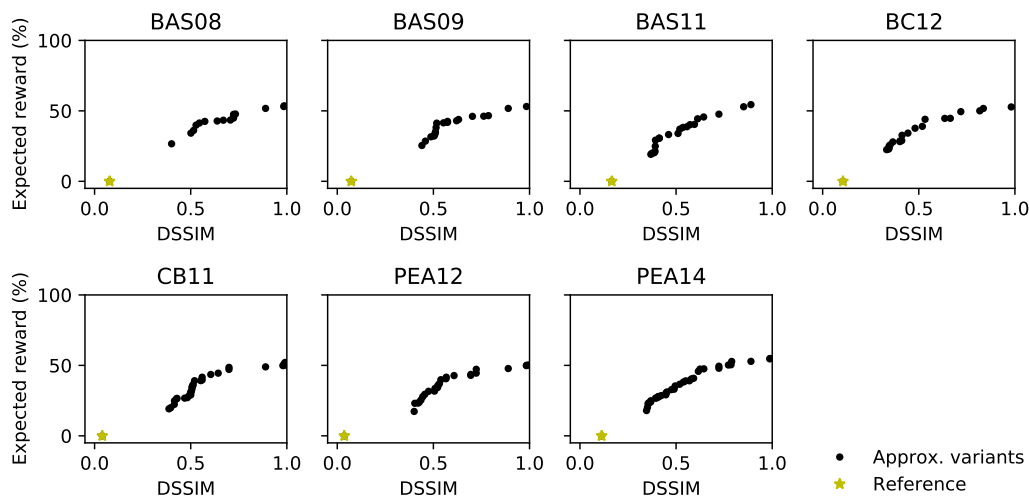


Fig. 10. Pareto-front estimation provided by the MOEA

Concerning the exploration time, the DSE has been conducted on a host PC equipped with 16 GB of RAM and an Intel i7-3770 CPU running at 3.9 GHz. On that hardware platform, the DSE for each algorithm took about 20 hours to complete. Therefore, thanks to the proposed fitness-function (Equation 14), we only needed few days to complete the exploration for all the algorithms. Indeed, by simply evaluating the proposed fitness-function we avoided performing a circuit synthesis to compute the reward. In fact, on the same hardware platform, the ASIC synthesis tool requires, on average, of 22 minutes to accomplish a single synthesis, while the FPGA tool needs 30 minutes (17 for synthesis and 13 for technology mapping), on average. Considering that the MOEA configurations had a population of 2000 individuals (i.e., 4000 circuit variants to synthesize at each iteration), performing the first of the seven iterations alone would have required $22 \times 4000 = 88000$ minutes – i.e. about 60 days – for ASIC technology.

It is worth noting that an exhaustive DSE is undoubtedly unfeasible, even in the case evaluating a single solution requires negligible time, since the size of solution spaces ranges between $2.66 \times 10^{49} \approx 2^{164}$ and $1.081 \times 10^{80} \approx 2^{265}$.

After the DSE, to correctly evaluate the final gains, we synthesized the obtained approximate configurations to both ASIC and FPGA technologies. Over all the algorithms, the total number of obtained non-dominated approximate configurations to synthesize was 164, i.e. ≈ 24 per each algorithm, on average. For the reader convenience, in the following figures we plotted the experimental result data along with the corresponding first-order interpolation to highlight the trend.

5.4.2 ASIC Synthesis. We synthesized all the obtained non-dominated approximate configurations to ASIC, by using the 65nm FinFET [22] technology and the Cadence Genus Synthesis Solution tool. We resorted to the synthesis reports for the silicon die area of the approximate configurations. In Figure 11, we report the result. Concerning the power consumption, to determine whether the synthesis power report provides a satisfying accuracy, we simulated the whole workload for two algorithms (BAS08 and BAS09) and collected the resulting power consumption. As a result, we realized

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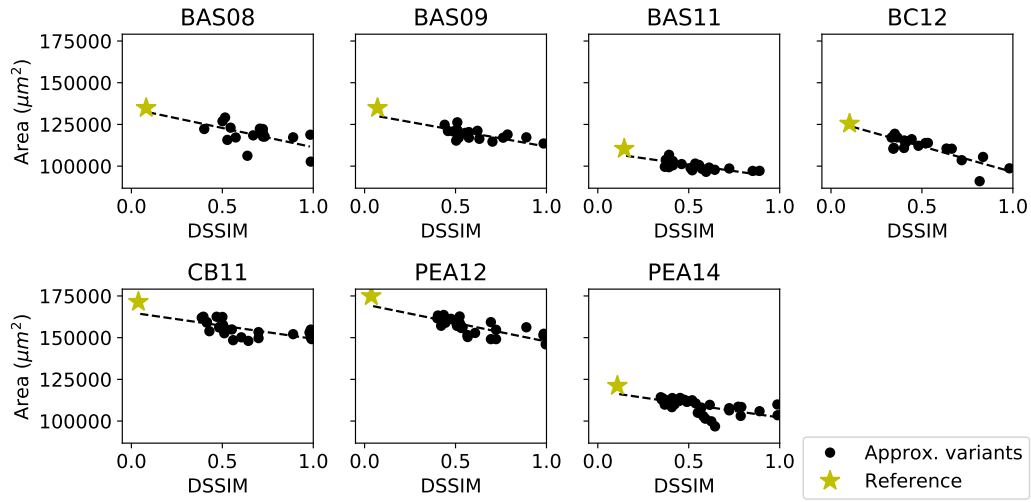


Fig. 11. ASIC silicon die area (μm^2)

that the difference between the power consumption resulted from the workload simulation and that estimated by the synthesis tool only differed by 5%, on average. We considered the synthesis report accuracy sufficient, thus in Figure 12 we show the power results from the synthesis report. Please note that the scale on the left axis (static power) is different from the scale on the right axis (dynamic power).

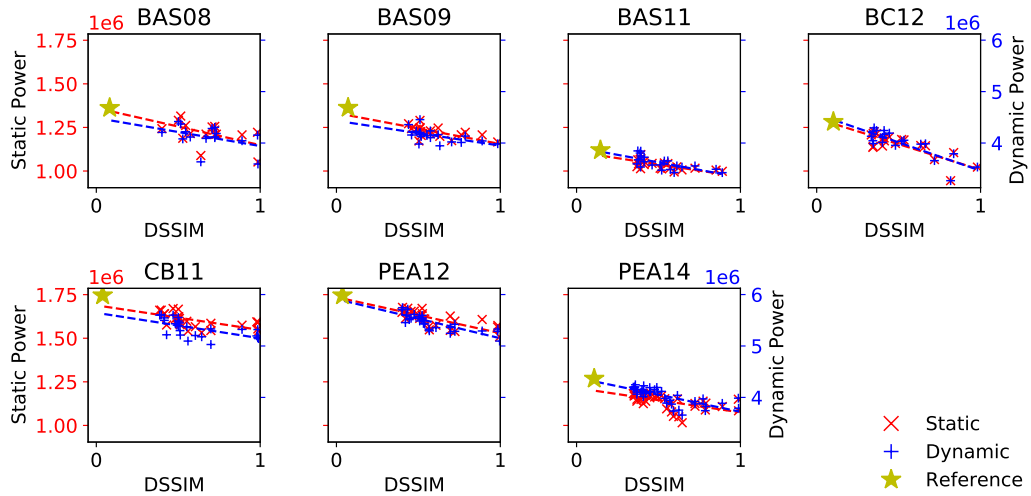


Fig. 12. Power consumption estimation for ASIC (nW)

Power savings are achieved due to both the reduced area and the lower switching activity that IACs exhibit w.r.t FACs, as also reported in [2].

It is worth highlighting that the trends shown in Figures 11 and 12 are perfectly in line with the trend predicted by our approach (see Figure 10). Indeed, higher reward in Figure 10 corresponds to lower area/power in Figures 11 and 12.

Table 3. Minimum and maximum savings while targeting ASIC

Algorithm	Area Savings (%)		Power Savings (%)	
	min	max	min	max
BAS08	9	25	5	20
BAS09	5	15	5	15
BAS11	5	12	9	13
BC12	6	27	6	25
CB11	5	17	3	10
PEA12	7	17	5	15
PEA14	5	23	4	18

For the reader convenience, Table 3 reports a summary of the minimum and maximum area/power savings we achieved during the experimental campaign while targeting ASIC.

5.4.3 *FPGA Synthesis.* We synthesized all the obtained non-dominated approximate configurations to a Xilinx Zynq-7020 MPSoC. To get a fair estimation of hardware requirements, we used only its embedded FPGA and inhibited Digital Signal Processors (DSPs) usage.

Figure 13 reports synthesis result in terms of number of LUTs for all the considered algorithms. As expected,

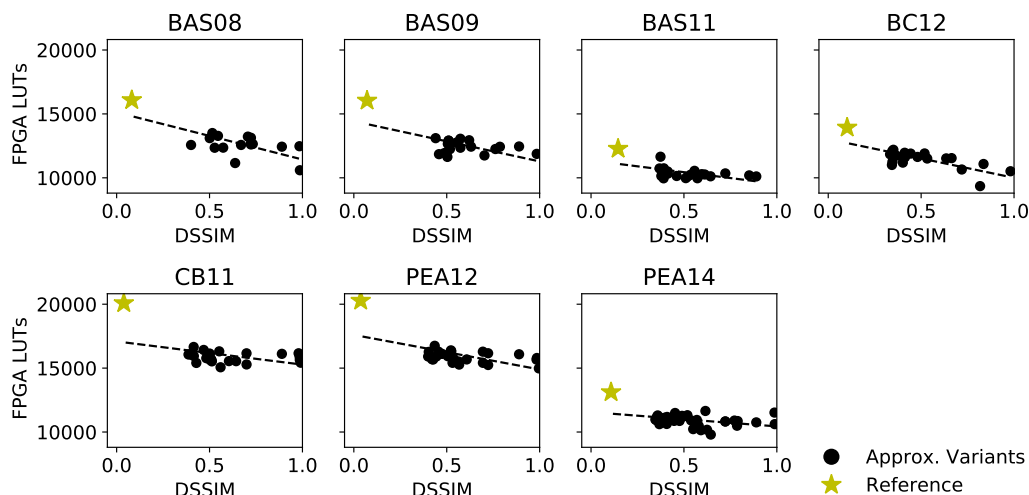


Fig. 13. FPGA resource requirements

approximate solutions require less resources than the precise implementation, as highlighted by the decreasing general trend.

In order to correctly evaluate energy savings, we performed a post-synthesis timing simulation, using the Dynamic Power Analysis tool provided by the Xilinx Vivado. In this case, since the synthesis report has a very low confidence level for power consumption estimation, we resorted to a workload simulation for all the solutions the DSE provided, for all the algorithms. In this way, we achieved a high confidence level power estimation.

Figure 14 shows static and dynamic power consumption for all the algorithms.

The static power of the FPGA is largely caused by the fabric of the device and does not directly depend on used resources, while dynamic one is directly linked to the user design, due to the input data pattern and the design internal

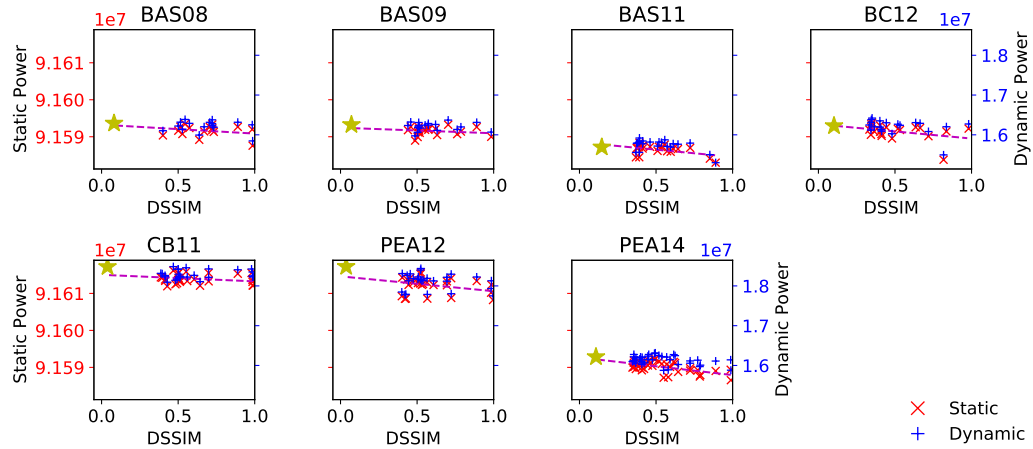


Fig. 14. Power consumption estimation for FPGA (nW)

Table 4. Minimum and maximum savings for FPGA synthesized approximate configurations

Algorithm	LUTs Savings (%)		Dynamic Power Savings (%)	
	min	max	min	max
BAS08	27.5	48.2	0.1	3.4
BAS09	32.6	42.5	0.3	1.8
BAS11	30.6	40.4	0	2.5
BC12	31.1	50.8	0	5.2
CB11	30.9	42.7	0.5	2.6
PEA12	31.3	42.7	0.7	4.4
PEA14	29.2	44.5	0.1	2.8

activity. Being our hardware implementations of approximate **DCT** characterized by low overhead, i.e. device resources usage falls between 6 and 13 %, it is necessary to split power consumption in static and dynamic since the former turned out to be about an order of magnitude greater than the latter one for the target **FPGA** device.

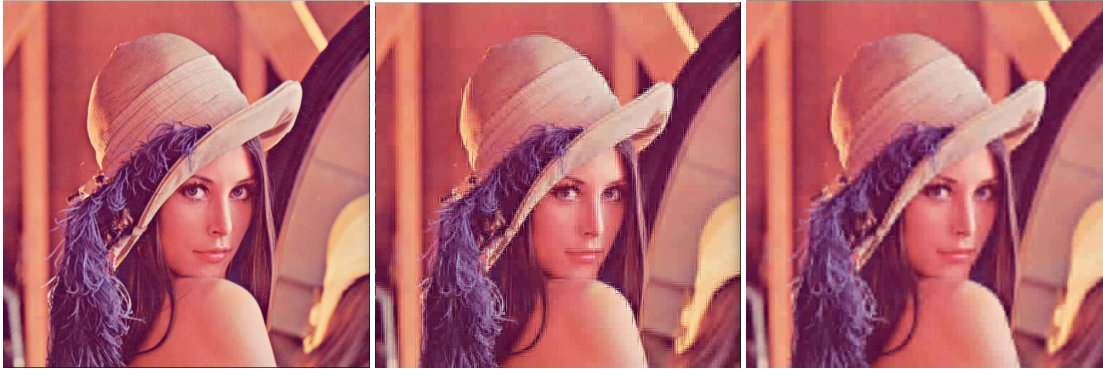
Also in this case, power savings are achieved thanks to both the reduced total area and the logical structure of **IACs**: **FPGA LUTs** implementing **IACs** have a lower switching activity than those implementing **FACs**, as reported in [2].

As in the **ASIC** case, also for **FPGA** the trends shown in Figures 13 and 14 are perfectly in line with the trend predicted by our approach (see Figure 10). Indeed, higher reward in Figure 10 corresponds to lower area in Figures 13.

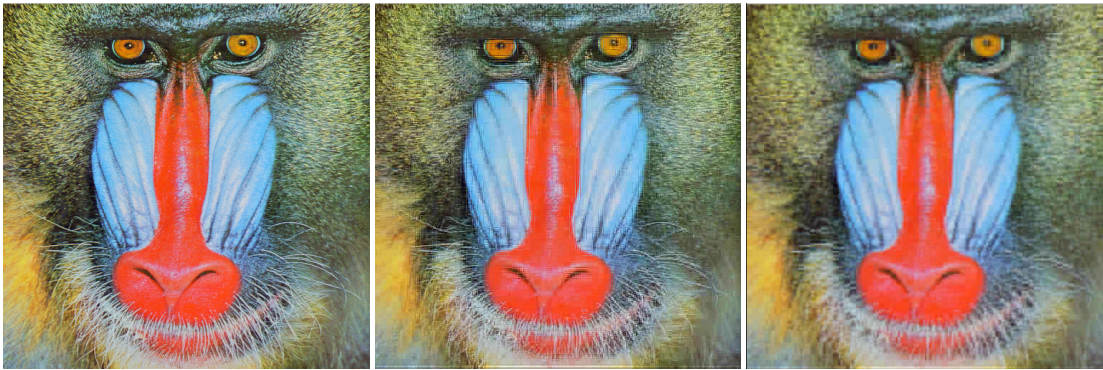
As done for **ASIC**, we reports a summary of the minimum and maximum area/power savings we achieved during the experimental campaign while targeting **FPGA** in Table 4.

5.4.4 Visual Test. Since JPEG belongs to image processing domain, we also provide a visual test: Figure 15 shows, from left to right, the standard JPEG-compressed image of Lena and Baboon, the same images compressed using the exact version of the BC12 algorithm [6] – which exhibit a DSSIM of 0.10, and requires $125473.92 \mu m^2$ and $5691946 \mu W$ when implemented on **ASIC**, or 5902 LUTs and $107933980 \mu W$ while targeting **FPGA** – and, finally, the ones compressed with its approximate variant having 0.33 as DSSIM value and 0.22 of reward, which correspond to $8362.64 \mu m^2$ and $352.711 \mu W$ saved for **ASIC** and 1846 LUTs and $94506.744 \mu W$ saved for **FPGA**. As the reader can easily figure out, the quality differences are barely perceivable.

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(a) Visual test with Lena



(b) Visual test with Baboon

Fig. 15. Visual test

5.5 Comparison with previous work

In this subsection, we compare the results obtained with our approach with those obtained in the work in [2]. Authors of [2] estimated gains G through the following equations:

$$G = \frac{V_i - V_e}{V_e} \quad (15)$$

$$V_i = P_i \cdot nab + (N - nab) \cdot P_e \quad (16)$$

where V_i and V_e represent the average energy required to perform an addition, by an inexact N -bits adder and by an exact N -bits adder respectively, P_i and P_e represent the average energy required by a single IAC and by a full-adder cell, respectively. Values of P_i and P_e used in [2] were measured by using the 45nm Complementary Metal-Oxide Semiconductor (CMOS) technology and are reported in Table 5. Such equations have the same goal as Equation (14), i.e. predicting the gains achieved thanks to the approximation. While Equations (15) and (16) take into account the energy consumption parameters of the individual adder cell, Equation (14) takes into account only the number of transistors.

In [2], authors performed a manual exploration. In particular, firstly they tried different IACs and decided to always resort to InXA2 in their experiments, based on its energy delay product. Then, they tried different NAB values for the

Table 5. Energy consumed by a single adder cell from [2]

Cell		FullAdd	AMA1	AMA2	AMA3	AMA4	AXA	InXA1	InXA2	InXA3
Energy(fJ)	Avg.	0.9267	0.513	0.6631	0.6649	0.478	0.4042	0.1535	0.0563	0.3409
	Max.	2.3668	0.9794	0.7203	0.7116	0.6271	0.8924	0.2096	0.1291	0.4211

InXA2 adder and finally set it to 4 for all the experiments (i.e., for all the **DCT** algorithms). Besides, they used the **PSNR** metric to measure the JPEG error entailed by the approximate **DCT** variants. Conversely, we adopted the **DSSIM** index as error metric – which is more suitable for image processing – and we let the **MOEA** decide which inaccurate cell to use and how many bits to approximate (i.e., the **NAB** parameter) for each of the sums. A minor difference concerns the implementation of the adders: while 32-bit adders were considered in [2], we considered 14-bit adders.

In order to effectively compare the two studies it is necessary to place them under the same conditions. Thus, we executed the JPEG algorithm on the same four images considered in [2] – i.e. Lena, Cameraman, Boat and Pepper – by using the approximate **DCT** variants obtained with our approach and computed the **PSNR** metric. Hence, we computed energy savings according to Equation (15), considering 32-bit adders. Figure 16 shows the obtained results. Concerning

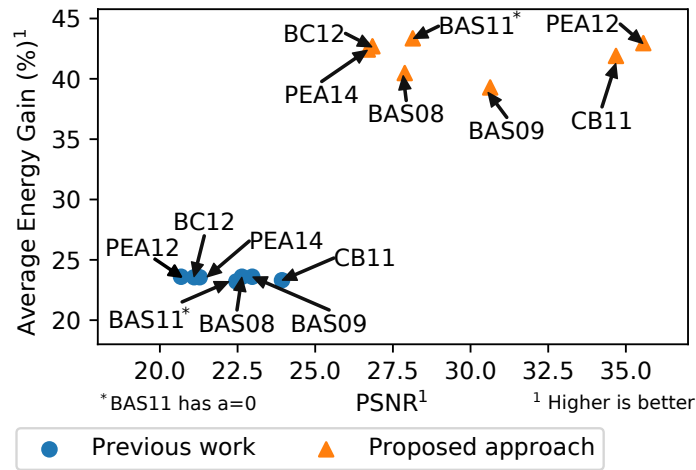


Fig. 16. Comparison with results from [2]

both energy consumption and PSNR, our approach allowed a significant improvement for all the considered algorithms compared to the approach adopted in [2]. In detail, our approach allowed an absolute improvement spanning from 15.69% to 20.15% (average 18.38%) concerning the energy gain and from 5.24 dB to 14.88 dB (average 7.91 dB) concerning the PSNR. Therefore, with our approach we were able to produce higher quality images, i.e. with less error, while consuming less energy. This is the result of the thorough **DSE** made possible by the proposed approach. Indeed, using a **MOEA** allows performing a multi-objective optimization more efficiently and automatically. Moreover, not needing to synthesize each approximate variant allows exploring more extensively the design space in a reduced time.

6 CONCLUSION AND FUTURE WORK

In this manuscript, we presented a novel generic and fully automatic approach for the approximation of DCT hardware accelerators. It enables automatic approximate DCT variant generation and automatic space exploration, by using the GA approach.

We analyzed and modeled several algorithms from the literature to compute a fast and lightweight version of the DCT. For each algorithm, we applied approximation by substituting full-precise adders with several approximate ones from the literature having configurable approximation degree. In this way, we can obtain different approximate configurations of the algorithms, depending on the chosen approximate adders and their approximation degree. Approximate adders introduce inaccuracy in the computation but also achieve gains in terms of area and power consumption. For each algorithm, we performed a DSE to find the non-dominated approximate designs in terms of trade-off between inaccuracy and gains. We modeled the DSE as a MOP and we used a GA to solve it.

After the DSE, we synthesized the obtained designs by targeting both FPGA and ASIC. To do so, we implemented all the algorithms as re-configurable hardware designs. Finally, we evaluated the actual gains in terms of area and power consumption. Experimental results clearly showed that, with the proposed approach, it is possible to perform a meaningful DSE to find the best trade-offs between output accuracy and resource gains in a reasonable time. Finally, the comparison performed with previous work clearly showed the advantages of the proposed approach.

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