

# Parallel local search for the Costas Array Problem

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**Abstract**—The Costas Array Problem is a highly combinatorial problem linked to radar applications. We present in this paper its detailed modeling and solving by Adaptive Search, a constraint-based local search method. Experiments have been done on both sequential and parallel hardware up to several hundreds of cores. Performance evaluation of the sequential version shows results outperforming previous implementations, while the parallel version shows nearly linear speedups up to 4,096 cores.

## I. INTRODUCTION

During the last decade, the family of Local Search methods and Metaheuristics has been quite successful in solving large real-life problems [22], [23], [24]. Applying Local Search to Constraint Satisfaction Problems (CSP) has also been attracting some interest [9], [18], [23], [25] as it can tackle CSPs instances far beyond the reach of classical propagation-based solvers.

A generic domain-independent Local Search method named Adaptive Search was proposed in [9], [10]. It is a metaheuristic that takes advantage of the structure of the problem to guide the search and that can be applied to a large class of constraints (*e.g.*, linear and non-linear arithmetic constraints and symbolic constraints). Moreover, it intrinsically copes with over-constrained problems.

We will describe in this paper how the Costas Array Problem can be modeled easily in the Adaptive Search framework and how this model can be refined progressively in order to achieve very good performances on sequential machines. The Costas Arrays Problem (CAP) has been introduced to the constraint programming community in [25]. It is an abstract problem that was motivated by a sonar application in the 1960's but still has practical interest in radar and software-defined radio applications [3]. A whole community is active around the CAP (<http://www.costasarrays.org>) as several problems are not yet solved, *e.g.*, no constructive algorithm exists for larger values of  $n$  and the question of the existence of a Costas array of size  $n$  for  $n = 32$  is still open. From a constraint programming point of view, it is an interesting problem because CAP is conceptually related

to three well-known CSPs: the perennial *Nqueens* problem, the *All-Interval Series* problem (`prob007` of CSPLib [19]), and the *Golomb rulers* problem (`prob006` of CSPLib), but it is much more difficult to solve, mainly because of a tricky, bi-dimensional `alldifferent` constraint.

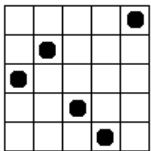
When attacking a highly combinatorial problem, an interesting approach is also to consider parallel implementations in order to boost performance. In the last decade, with desktop computers turning into parallel machines with 2, 4 or even 8 core CPUs, designing and implementing efficient parallel constraint solvers has become an increasingly developing research field. However, only a few efficient parallel implementations of constraint solvers have been reported, *e.g.*, the seminal work of [34], [35] on quad-core machine and the Comet system [23] which has been parallelized for small clusters of PCs, both for its local search solver [28] and its propagation-based constraint solver [29]. Recent experiments have been done up to 12 processors [30], and speedups tend to level off after 10 processors. It has to be noted that it is not obvious how to scale such implementations up to a few hundreds of processors, either because of the shared-memory nature of the solver or because of the intrinsic sequentiality of the solving method or of the problem itself.

In [13], [7], [6], a parallel version of a constraint solver based on local search is defined with a multi-start approach requiring no communication between processes (also known as *Pleasantly Parallel*). On classical CSP benchmarks from CSPLib, this simple parallelization scheme gives good results, with a factor 50-70 speedup for 256 cores, but this is far from ideal speedup (*e.g.*, factor 256 speedup for 256 processors), even for large problem instances. It is thus an open question to know whether this is due to the classical (structured) CSP benchmarks used or if this is a limitation of the method. In this paper we address the problem of modeling a very combinatorial problem, with a low density of solutions (the CAP) in the sequential version and we further investigate if it scales up to a large number of processors and exhibits better speedups. For this we

used three platforms: (1) the HA8000 machine, an Hitachi supercomputer with 15,232 cores installed at University of Tokyo, (2) the GRID'5000 infrastructure, the French national Grid for the research, which contains a total of 8,596 cores deployed on 9 sites distributed in France and (3) the IBM Blue Gene/P machine JUGENE with 294,912 cores at the Jülich Supercomputing Center in Germany. Experiments on both HA8000 and GRID'5000 show nearly linear speedups w.r.t. the sequential version, for instance 120 for 128 cores and 230 for 256 cores. On the Blue Gene/P, we obtained linear speed-ups up to 4,096 cores.

The rest of this paper is organized as follows. Section II presents the Costas Array Problem and Section III the Adaptive Search method. Section IV presents the modeling of the Costas Array Problem within the Adaptive Search formalism, and the performance of the sequential implementation. Section V details the experiments on three different parallel hardware: the HA8000 supercomputer, the GRID'5000 platform and the Blue Gene/P supercomputer. Section VI concludes the paper and briefly discusses about future work.

## II. THE COSTAS ARRAY PROBLEM



A Costas array is an  $n \times n$  grid containing  $n$  marks such that there is exactly one mark per row and per column and the  $n(n-1)/2$  vectors joining the marks are all different.

We give here an example of Costas array of size 5. It is convenient to see the Costas Array Problem (CAP) as a permutation problem by considering an array of  $n$  variables  $(V_1, \dots, V_n)$  which forms a permutation of  $\{1, 2, \dots, n\}$ . The Costas array above can thus be represented by the array  $[3, 4, 2, 1, 5]$ .

Historically, these arrays have been developed in the 1960's to compute a set of sonar and radar frequencies avoiding noise [11]. A very complete survey on Costas arrays can be found in [14]. The problem to find a Costas array of size  $n$  is very complex since the required time grows exponentially with  $n$ . In the 1980's, several algorithms have been proposed to build a Costas array given  $n$ , such as the Welch construction [20] and the Golomb construction [21], but these methods cannot built Costas arrays of size 32 and some higher non-prime sizes. Nowadays, after many decades of research, it remains unknown if there exist any Costas arrays of size 32 or 33. Another difficult problem is to enumerate all Costas arrays for a given size. Using the Golomb and Welch constructions, Drakakis *et. al* present in [16] all Costas arrays for  $n = 29$ . They show that among the  $29!$  permutations, there are only 164 Costas arrays, and 23 unique Costas arrays up to rotation and reflection. There are constructive methods known to produce Costas arrays of order 24 to 29 [4], [38], [15], [16].

The Costas Array Problem has been proposed as a challenging combinatorial problem by Kadioglu and Sellmann in

[25]. They propose a local search metaheuristic, *Dialectic Search*, for constraint satisfaction and optimization, and show its performance for several problems. Clearly this problem is too difficult for propagation-based solvers, even for medium size instances (*i.e.*, with  $n$  around 18 – 20). Let us finally note that we do not pretend that using local search is better than constructive methods in order to solve the CAP. We rather consider the CAP as a very good benchmark for testing local search and constraint-based systems and to investigate how they scale up for large instances and parallel execution.

In [37], Rickard and Healy studied a stochastic search method for CAP and concluded that such methods are unlikely to succeed for  $n > 26$ . Although their conclusion is true for their stochastic method, it cannot be extended to all stochastic searches: their method uses a restart policy which is too simple and they also used an approximation of the Hamming distance between configurations in order to guide the search which they recognized themselves not to be a very good indicator. However, they studied in this paper the distribution of solutions in the search space and have shown that clusters of solutions tend to spread out from  $n > 17$ , which justify our multi-walk approach presented in Section V to reach linear speedup for high values of  $n$ .

## III. THE ADAPTIVE SEARCH METHOD

Adaptive Search (AS) was proposed in [9], [10] as a generic, domain-independent constraint-based local search method. This metaheuristic takes advantage of the structure of the problem in terms of constraints and variables and can guide the search more precisely than a single global cost function to optimize, such as for instance the number of violated constraints. The algorithm also uses a short-term adaptive memory in the spirit of Tabu Search in order to prevent stagnation in local minima and loops.

### A. Algorithm

The input of the method is a problem in CSP format: a set of variables with their (finite) domains of possible values and a set of constraints over these variables. For each constraint, an *error function* needs to be defined: it gives, for each tuple of variable values, an indication of how much the constraint is violated. Consider an  $n$ -ary constraint  $c(X_1, \dots, X_n)$  and associated domains  $D_1, \dots, D_n$  for each variable. An error function  $f_c$  associated to the constraint  $c$  is a real-valued function from  $D_1 \times \dots \times D_n$  such that  $f_c(X_1, \dots, X_n)$  has value zero iff  $c(X_1, \dots, X_n)$  is satisfied. The error function will in fact be used as a heuristic to represent the degree of satisfaction of a constraint and will thus give an indication on how much the constraint is violated. This idea has also been proposed independently by [18], where it is called "penalty functions", and then reused in the Comet system [23], where it is called "violations". For example, the error function associated with an arithmetic constraint  $|X - Y| < c$ , for

a given constant  $c \geq 0$ , can be  $\max(0, |X - Y| - c)$ . AS relies on iterative repair, based on variable and constraint error information, seeking to reduce the error on the worst variable so far. The basic idea is to compute the error function for each constraint, then combine for each variable the errors of all constraints in which it appears, thereby projecting constraint errors onto the relevant variables. This combination of errors is problem-dependent, see [9] for details and examples, and represents the heuristic distance to a solution. It is usually a simple sum of absolute values, *i.e.*, Manhattan distance, although it might also be any other distance or maybe a weighted sum if constraints are given different priorities. Finally, the variable with the highest error is designated as the “culprit” and its value is modified. In this second step, the well known min-conflict heuristic [31] is used to select the value in the variable domain which is the most promising, that is, the value for which the total error in the next configuration is minimal. In order to prevent being trapped in local minima, the AS method also includes a short-term memory mechanism to store configurations to avoid. More precisely, variables can be marked Tabu and “frozen” for a given number of iterations. It also integrates reset transitions to escape stagnation around local minima, see Section III-B. As in every local search method, it is also possible to restart from scratch when the number of iterations becomes too large. The AS base algorithm is described in Figure 1.

Adaptive Search is a simple algorithm but it turns out to be quite efficient in practice. As a simple comparison, [6] provides timings between AS and the Comet 2.1.1 system on a few classical benchmarks. AS is about 40 times faster than Comet on the N-queen problem (for  $N=10000$  to  $50000$ ) and 100 to 500 times faster than Comet for the Magic Square problem (prob019 in CSPLib), for  $N=30$  to  $50$ . Of course, it should be noted that Comet is a complete and very versatile system while AS is just a C-based library. In [25] a new metaheuristics named Dialectic Search is compared with the 2001 version of AS [9], showing that both methods have similar results for the Magic Square problem (instances from  $N=20$  to  $50$ ). However when compared with the 2003 version of AS [10], AS is about 15 to 40 times faster than Dialectic Search (depending on the instances), on the same reference machine. Obviously, performance comparisons are always tricky, but one or two orders of magnitude can be considered as significant, especially as speedups increase with problem size. The basic code for permutation problems and examples are available as free software at the URL: <http://cri-dist.univ-paris1.fr/diaz/adaptive/>

### B. Tuning Adaptive Search

1) *Plateaux*: In [10], a simple but very effective improvement of the original algorithm was proposed. In AS, when a variable is selected and all alternative values for this variable give a global cost worse than the current one, this variable is

tagged as “Tabu” for a given number of iterations. However, what should be done when there is no improvement, but only equal-valued moves? In that case, a “plateau” is found in the global cost function landscape, and the question is to either follow this plateau or not. A simple idea is to introduce a probability  $p$  for this. With a good tuning (*e.g.*, probability of 90% to 95% of following a plateau) this boosts the performance of the algorithm by an order of magnitude on some problems such as Magic Square. Indeed the current sequential AS version of Magic Square problem can solve instances as big as  $400 \times 400$  (*i.e.*, 160,000 variables with domains of size 160,000) in about one hour on average.

2) *Reset*: When too many variables become Tabu, there is a risk of “freezing” the configuration and of getting trapped around a local minimum. The AS method thus needs a diversification operator to avoid such cases. This is done by performing a (partial) reset, *i.e.*, by assigning fresh values to a given percentage of the problem variables (parameter *RP* of the algorithm). A reset is triggered by the total number of variables being marked Tabu at a given iteration (parameter *RL* of the algorithm). This is the generic default reset mechanism of AS, but the reset procedure can also be customized if needed to become dedicated to a given problem. We use this ability for solving efficiently the CAP.

3) *Random*: Using a reliable pseudo-random number generator is essential for local search algorithms. For sequential methods, using generic random functions provided by standard libraries is often good enough. However, the need for better random functions (*i.e.*, more uniform) already appeared in stochastic optimization methods such as Particle Swarm Optimization (PSO) [40]. Therefore, when designing a massively parallel method with several hundreds or thousands of stochastic processes running at the same time, one has to choose carefully their random seed and to be assured of the correctness of the distribution. At this point, using a generic random function can turn out to be insufficient.

To ensure equity, we choose to generate the seed used by each process via a pseudo-random number generator based on a linear chaotic map. This method shows robust properties of distribution and has been implemented for cryptographic systems, like Trident [32].

## IV. SOLVING THE CAP WITH ADAPTIVE SEARCH

### A. Basic Model

The CAP can be modeled as a permutation problem by considering an array of  $n$  variables ( $V_1, \dots, V_n$ ) which forms a permutation of  $\{1, 2, \dots, n\}$  (*i.e.*, implicit `alldifferent` constraint on variables  $V_i$ ).  $V_i = j$  iff there is a mark at column  $i$  and row  $j$ . To take into account constraints on vectors between marks (which must be different) it is convenient to use the so-called *difference triangle*.

This triangle contains  $n-1$  rows, each row corresponding to a distance  $d$ . The  $d$ th row of the triangle contains the

## Algorithm 1 Adaptive Search Base Algorithm

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<b>Input:</b> problem given in CSP format: <ul style="list-style-type: none"> <li>• variables <math>X_i</math> with their domains</li> <li>• constraints <math>C_j</math> w/error functions</li> <li>• function to project errors on vars</li> <li>• cost function to minimize</li> </ul>	<b>some tuning parameters:</b> <ul style="list-style-type: none"> <li>• <math>TT</math>: # iterations a variable is frozen</li> <li>• <math>RL</math>: # frozen variables triggering a reset</li> <li>• <math>RP</math>: % of variables to reset</li> <li>• <math>MI</math>: max. # iterations before restart</li> <li>• <math>MR</math>: maximal # of restarts</li> </ul>
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**Output:** a solution if the CSP is satisfied or a quasi-solution of minimal cost otherwise.

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1:  $Restart \leftarrow 0$ 
2: repeat
3:    $Restart \leftarrow Restart + 1$ 
4:    $Iteration \leftarrow 0$ 
5:   Compute a random assignment  $A$  of variables in  $V$ 
6:    $Opt\_Sol \leftarrow A$ 
7:    $Opt\_Cost \leftarrow cost(A)$ 
8:   repeat
9:      $Iteration \leftarrow Iteration + 1$ 
10:    Compute errors of all constraints in  $C$  and combine errors on each variable
11:    ▷ (by considering only the constraints in which a variable appears)
12:    select the variable  $X$  (not marked Tabu) with highest error
13:    evaluate costs of possible moves from  $X$ 
14:    if no improvement move exists then
15:      mark  $X$  as Tabu until iteration number:  $Iteration + TT$ 
16:      if the number of variables marked Tabu  $\geq RL$  then
17:        randomly reset  $RP$  % variables in  $V$  (and unmark those Tabu)
18:      end if
19:    else
20:      select best move and change  $X$ , yielding the next configuration  $A'$ 
21:      if  $cost(A') < Opt\_Cost$  then
22:         $Opt\_Sol \leftarrow A \leftarrow A'$ 
23:         $Opt\_Cost \leftarrow cost(A')$ 
24:      end if
25:    end if
26:  until  $Opt\_Cost = 0$  (a solution is found) or  $Iteration \geq MI$ 
27: until  $Opt\_Cost = 0$  (a solution is found) or  $Restart \geq MR$ 
28: output( $Opt\_Sol, Opt\_Cost$ )
  
```

Figure 1. Adaptive Search Base Algorithm

differences  $V_{i+d} - V_i$  for all  $i = 1, \dots, n - d$  (i.e., the difference of values at a distance  $d$ ). Ensuring all vectors are different comes down to ensure the triangle contains no repeated values on any given row (i.e., alldifferent constraint on each row). Here is the difference triangle for the Costas array given as example in Section II.

	3	4	2	1	5
$d = 1$	1	-2	-1	4	
$d = 2$		-1	-3	3	
$d = 3$			-2	1	
$d = 4$				2	

In AS, the way to define a constraint is done via error

functions. At each new configuration, the difference triangle is checked to compute the global cost and the cost of each variable  $V_i$ . Each row  $d$  of the triangle is checked one by one. Inside a row  $d$ , if a pair  $(V_i, V_{i+d})$  presents a difference which has been already encountered in the row, the error is reported as follows: increment the global cost and the cost of both variables  $V_i$  and  $V_{i+d}$  by  $ERR(d)$  (a strictly positive function). For a basic model we can use  $ERR(d) = 1$  (to simply count the number of errors). Obviously a solution is found when the global cost equals 0. Otherwise AS selects the most erroneous<sup>1</sup> variable and will try to improve it.

<sup>1</sup>i.e., the variable with the highest total error.

## B. Optimization

In the basic model, the function  $ERR(d)$  can be a constant (e.g.,  $ERR(d) = 1$ ) but a better function is  $ERR(d) = n^2 - d^2$  which “penalizes” more errors occurring in the first rows (those containing more differences). The use of this function instead of  $ERR(d) = 1$  improves the computation time (around 17 %).

Moreover, a remark from Chang [8] makes it possible to focus only on distances  $d \leq \lfloor (n-1)/2 \rfloor$ . In our example, it is only necessary to check the 2 first rows of the triangle (i.e.,  $d = 1$  and  $d = 2$ ). This represents a further gain in computation time (around 30 %).

Another source of optimization concerns the *reset* phase. Recall that AS maintains a Tabu list to avoid to be trapped in local minima and, when too many variables become Tabu, the current configuration is perturbed to escape the current local minimum. By default AS resets a certain percentage of variables (see parameters  $RL$  and  $RP$  in the previous algorithm). We found good results with  $RL = 1$  and  $RP = 5\%$  (as soon as 1 variable is marked Tabu, reset 5% of the variables). Whereas this default behavior of AS is general enough to escape any local minimum, it sometimes “breaks” some important parts of the current configuration (but conversely, if we want to preserve too many variables, we can be trapped in the local minimum). AS allows the user to define his own reset procedure: when a reset is needed this procedure is called to propose a pertinent alternative configuration. Our customized reset procedure tries 3 different perturbations from the current configuration:

- 1) Select the most erroneous variable  $V_m$ . Consider each sub-array starting or ending by  $V_m$  and shift it (circularly) from 1 cell to the left and to the right.
- 2) Add a constant “circularly” (i.e., modulo  $n$  to maintain the permutation) to each variable. The current implementation tries the following 4 constants: 1, 2,  $n-2$ ,  $n-3$  (but middle values  $n/2$ ,  $n/2-1$ ,  $n/2+1$  are also pertinent).
- 3) Left-shift from 1 cell the sub-array from the beginning to a (randomly chosen) erroneous variable different from  $V_m$ . In the current implementation we test at most 3 erroneous variables.

As soon as the global cost of a perturbation is strictly inferior to the entry global cost the local minimum is considered as *escaped* and AS continues with this (perturbed) configuration. This works, on average, in 32% of the cases (independently from  $n$ ). Otherwise, all perturbations are tested exhaustively and the best (i.e., whose global cost is minimal) is selected. This dedicated reset procedure provides a speedup factor of about 3.7 and is thus very effective.

## C. Evaluation of the sequential implementation

In this section we study the performances of the sequential AS implementation. Since AS is a stochastic method and

uses randomness for initial configurations and during its execution, each benchmark has been executed 100 times. Classically, there are three interesting ways for aggregating those results: considering the best case, the worst case and the average case (i.e., minimum, maximum and average of 100 execution times). The experiments were run on a Dell Precision T7500 (Intel Xeon W5580, 3.20 GHz and 24 GB of 1333MHz SDRAM) running Linux (the AS code is compiled using GCC 4.4.5, with the -O3 flag).

Size		Time	Iterations	Local min	ratio
16	avg	<b>0.08</b>	12665	6853	60
	min	0.00	212	117	
	max	0.45	69894	37904	
17	avg	<b>0.59</b>	73430	38982	30
	min	0.02	2591	1361	
	max	2.39	294580	156154	
18	avg	<b>3.49</b>	395838	207067	116
	min	0.03	2789	1538	
	max	19.81	2254001	1178875	
19	avg	<b>29.46</b>	2694319	1372671	95
	min	0.31	28911	14798	
	max	127.78	11619940	5922204	
20	avg	<b>250.68</b>	20536809	10278723	66
	min	3.89	319368	159127	
	max	1097.06	89791761	44945485	

Table I  
EVALUATION OF THE SEQUENTIAL IMPLEMENTATION (TIMES IN SEC.)

Table I presents the results of the evaluation. For each instance it details the execution time (in seconds), the number of iterations, the number of local minima encountered and the ratio between the average time and the minimum time (when the minimum time is zero we used the numbers of iterations). For  $n = 18$  and above, an exponential behavior seems to appear: the computation time for solving instance  $n$  is an order of magnitude greater than  $n - 1$ . More interestingly, for all instances, *the best case is much faster than the average case* (see last column of Table I). This important property convinced us to experiment on parallel machines as we will see later.

In [25], Kadioglu and Sellmann propose a novel and interesting local search meta-heuristic called *Dialectic Search* (DS). The authors show how DS performs well on the CAP and compare DS with a tabu search algorithm using the quadratic neighborhood implemented in Comet. The comparison was done for instances 13 to 18 and revealed that DS is between 2 and 3 times faster than Comet on CAP. It is thus very interesting to compare our AS implementation with DS. The comparison was however difficult because Kadioglu and Sellmann used an outdated machine and no rigorous scaling information is available to compare their Pentium-III 733 MHz with modern processors. Fortunately we managed to find an exactly identical machine. The following table presents the comparison with DS considering for both systems the average of 100 executions. Times are given in seconds measured on a Pentium-III 733 MHz for AS and taken from [25] for DS.

Size	DS	AS	DS / AS
13	0.05	0.01	5.00
14	0.26	0.05	5.20
15	1.31	0.24	5.46
16	7.74	0.97	7.98
17	53.40	7.58	7.04
18	370.00	44.49	8.32

Table II  
AS SPEED-UPS W.R.T DS

Unfortunately, the paper about DS does not give other data than time executions, therefore we are not in measure to compare the number of iterations, local minima, etc. This table clearly shows that AS outperforms DS on the CAP: for small instances AS is five times faster but the speedup seems to grow with the size of the problem, reaching a factor 8.3 for  $n = 18$ .

CAP has also been used as a benchmark in the Constraint Programming community and we can compare with a CP Comet program made by Laurent Michel and based on the modeling in MiniZinc by Barry O’Sullivan<sup>2</sup>. As could be expected, CP is much less efficient than local search, and this Comet program is about 400 times slower than Adaptive Search for CAP19.

## V. PARALLEL IMPLEMENTATION AND PERFORMANCE ANALYSIS

Parallel implementation of local search metaheuristics have been studied since the early 90’s, when multiprocessor machines started to become widely available, see [39] for a general survey and concepts, or [33] for basic parallel versions of Tabu search, simulated annealing, GRASP and genetic algorithms. With the availability of clusters in the early 2000’s, this domain became active again [12], [2]. Apart from domain-decomposition methods and population-based method (such as genetic algorithms), one usually distinguishes between single-walk and multiple-walk methods for Local Search. Single-walk methods consist in using parallelism inside a single search process, *e.g.*, for parallelizing the exploration of the neighborhood, see for instance [26] for such a method making use of GPUs for the parallel phase. Multiple-walk methods (also called multi-start methods) consist in developing concurrent explorations of the search space, either independently or cooperatively with some communication between concurrent processes. A key point is that independent multiple-walk methods are the easiest to implement on parallel computers and can lead to linear speed-up if solutions are uniformly distributed in the search space and if the method is able to diversify correctly [39]. However, previous experiments [6] showed that on structured problems such as (non-random) CSPs, independent multiple-walk parallelization does not achieve

ideal speedups, reaching only a factor 50-70 speedup for 256 cores. It might be because solutions are not uniformly distributed in the search space, and are for instance regrouped in ”clusters”, as was shown for solutions of SAT problems near the phase transition in [27]. Therefore some sequential computation is needed to get to the vicinity of such clusters.

### A. Motivation and Implementation

As remarked in Section IV-C, it can be seen than in 100 sequential runs of the algorithm starting from different (random) initial configurations, the minimal one is much faster than the average (sometimes more than 100 times). Then if we can run in parallel many AS engines we can hope for a global execution time equal to the minimal one, and therefore a good speedup.

We implemented a parallel version of AS using Open-MPI, an implementation of the MPI standard [17]. Preliminary experiments and performance results on classical CSP benchmarks are described in [6]. The parallelization is straightforward and based on the idea of multi-start and independent multiple-walk: fork a sequential AS method on every available cores. But on the opposite of the classical fork-join paradigm, parallel AS shall terminate as soon as a solution is found, not wait until all the processes have finished (since some searches initialized with ”bad” initial configurations can take some time). Thus, some non-blocking tests are involved every  $c$  iterations to check if there is a message indicating that some other process has found a solution; in which case it terminates the execution properly. This results in a high number of independent work units, a high CPU to I/O ratio, and no inter-process communication for computation.

Four different testbeds were used on three platforms:

- **HA8000**, the Hitachi HA8000 supercomputer of the University of Tokyo with a total number of 15,232 cores. This machine is composed of 952 nodes, each of which is composed of 4 AMD Opteron 8356 (Quad core, 2.3 GHz) with 32 GB of memory. Nodes are interconnected with a Myrinet-10G network attaining 5 GB/s in both directions. HA8000 can theoretically achieve a performance of 147 Tflops, but we only accessed to a subset of its nodes as users can only have a maximum of 64 nodes (1,024 cores) in normal service.
- **GRID’5000** [5], the French national Grid for research, which consists of 8,596 cores deployed on 9 sites in France. We used two subsets of the computing resources of the Sophia-Antipolis node: **Suno**, composed of 45 Dell PowerEdge R410 with 8 cores each, thus a total of 360 cores, and **Helios**, composed of 56 Sun Fire X4100 with 4 cores each, thus a total of 224 cores.
- **JUGENE**, the IBM Blue Gene/P supercomputer at the Jülich Supercomputing Center containing 294,912 cores. This machine is composed of 73,728 nodes, each

<sup>2</sup>[http://www.g12.cs.mu.oz.au/mzn/costas\\_array/CostasArray.mzn](http://www.g12.cs.mu.oz.au/mzn/costas_array/CostasArray.mzn)

of which is composed of 4 Power PC 450 32-bits at 850Mhz with 2 GB of memory. The network is a 3-dimensional torus reaching a bandwidth of 5.1GB/s. The overall peak performance is 1 Pflops.

### B. Experiments on parallel machines

Let us now present the result of the parallel experiments. The following tables show the execution times on HA8000 (Table III), JUGENE (Table IV) and GRID'5000 (Table V). Execution times do not include the deployment time, negligible on big benchmarks. Behaviors on both platforms are similar and exhibit good speedups. We give the timings in seconds for 50 executions of each benchmark with the average time, median time, minimal time and maximal time. We can see that with more cores, the maximal time decreases a lot and thus the difference between minimal and maximal times decreases. Moreover, in our case the median time is always below the average time, meaning we have more fast runs rather than slow ones. The median time presents here a speedup at least as good as the average time. For instance, for  $n = 20$  and using 256 cores on HA8000, speedup obtained w.r.t sequential runs is a factor 170 concerning the average time and a factor 210 for the median time.

Size	1 core	32 cores	64 cores	128 cores	256 cores	
18	avg	<b>6.76</b>	<b>0.25</b>	<b>0.23</b>	<b>0.24</b>	<b>0.26</b>
	med	4.25	0.18	0.18	0.20	0.23
	min	0.23	0.00	0.00	0.00	0.00
	max	22.81	1.07	0.90	0.94	0.78
19	avg	<b>54.54</b>	<b>1.84</b>	<b>1.00</b>	<b>0.72</b>	<b>0.55</b>
	med	43.74	1.45	0.76	0.57	0.44
	min	0.51	0.0	0.03	0.02	0.01
	max	212.96	6.62	5.24	3.48	2.22
20	avg	<b>367.24</b>	<b>13.82</b>	<b>8.66</b>	<b>3.74</b>	<b>2.18</b>
	med	305.79	11.53	5.06	2.36	1.44
	min	9.51	0.05	0.03	0.03	0.06
	max	1807.78	54.26	36.98	23.87	9.21
21	avg	-	<b>160.42</b>	<b>81.72</b>	<b>38.56</b>	<b>16.01</b>
	med	-	114.06	53.04	30.68	10.12
	min	-	1.63	2.13	1.49	0.73
	max	-	654.79	335.66	145.59	93.13
22	avg	-	<b>501.23</b>	<b>249.73</b>	<b>128.47</b>	<b>60.80</b>
	med	-	450.45	178.85	99.62	55.90
	min	-	0.23	0.35	0.26	1.58
	max	-	1550.25	935.51	406.15	196.26

Table III  
EXECUTION TIMES (IN SEC.) ON HA8000

Size	256 cores	512 cores	1,024 cores	2,048 cores	4,096 cores	
21	avg	<b>84.41</b>	<b>43.66</b>	<b>27.86</b>	<b>10.21</b>	<b>5.97</b>
	med	60.22	30.31	23.67	5.56	4.47
	min	2.43	0.85	1.46	0.27	0.13
	max	346.03	274.69	108.14	93.89	21.98
22	avg	-	<b>265.12</b>	<b>148.80</b>	<b>76.24</b>	<b>36.12</b>
	med	-	166.47	79.63	63.24	28.00
	min	-	1.34	1.95	0.81	0.60
	max	-	1831.96	638.34	277.96	154.89

Table IV  
EXECUTION TIMES (IN SEC.) ON JUGENE

For small instances ( $n = 18$ ), since computation times of parallel executions on many cores are below 0.5 second, they are maybe not significant because of interactions with operating system operations.

For medium instances, we have for example a **226 times speedup** w.r.t. sequential execution on Suno for  $n = 19$  on **256 cores**. Indeed, for different CAP instances, speedups w.r.t. sequential version are between 75 and 120 on 128 cores on HA8000, and between 100 and 170 on 256 cores on HA8000, between 120 and 137 on 128 cores on Suno, between 204 and 226 on 256 cores on Suno and between 120 and 130 on 128 cores on Helios.

For bigger instances, such as  $n = 21$  and 22, we do not have timings on HA8000 for the sequential version because a sequential problem resolution takes on average more than one hour, and the maximum resource utilization is currently limited to one hour because of power savings. For  $n = 21$  on Suno we have a **218 times speedup on 256 cores** w.r.t. sequential execution. Concerning  $n = 22$ , as sequential computation takes many hours, we limit our experiments on all machines to executions on 32 cores and above. Therefore in Figure 2, we will only give timings from 32 to 256 cores. On JUGENE, we ran our experiments starting from 256 cores. Indeed, despite HA8000 processors or Grid'5000 processors at 2.4Ghz, Blue Gene/P processors have a low frequency (850 Mhz), and are then significantly slower to solve a given problem. Moreover, JUGENE's job scheduler forces a time-out of 30 minutes for any job using less than 1025 cores, which made difficult on few cores our experiments which require 50 runs each. This is why we started experiments on JUGENE from 256 cores for  $n = 21$  and 512 cores for  $n = 512$ . For these two instances, one can see we reach nearly linear speed-ups up to 4,096 cores: We have a speed-up of **14.13 for n=21 w.r.t executions over 256 cores** (perfectly linear speed-up would be 16) and of **7.33 for n=22 w.r.t executions over 512 cores** (8 for a perfectly linear speed-up). Details about speed-ups on the JUGENE machine are represented by Figure 3. We can see that on all platforms, **execution times are halved when the number of cores is doubled**, thus achieving ideal speed-up.

This is graphically depicted in the next figure on a log-log scale. As a final result, we note that we can now solve  $n = 22$  in about one minute on average with 256 cores on HA8000. We can explain these linear speed-ups due to the good distribution of solution clusters over the search space for  $n > 17$  as shown in [37]. Thus, parallel searches using a multi-start approach greatly increase its chance to start near from a solution.

Up to now, we focused on the *average* execution time in order to measure the performance of the method, but a more detailed analysis could be done. In [1], [36], a method is introduced to represent and compare execution times of stochastic optimization methods by using so-called *time-to-target plots*. Observe that, for the CAP, the target value to

Size	Suno					Helios			
	1 core	32 cores	64 cores	128 cores	256 cores	1 core	32 cores	64 cores	128 cores
18	avg	<b>5.28</b>	<b>0.16</b>	<b>0.083</b>	<b>0.056</b>	<b>8.16</b>	<b>0.24</b>	<b>0.11</b>	<b>0.06</b>
	med		0.11	0.065	0.04		0.19	0.06	0.04
	min	0.01	0.00	0.00	0.00	0.13	0.00	0.00	0.00
	max	20.73	0.64	0.34	0.19	37.5	1.08	0.46	0.26
19	avg	<b>49.5</b>	<b>1.37</b>	<b>0.59</b>	<b>0.41</b>	<b>52</b>	<b>2.3</b>	<b>0.87</b>	<b>0.40</b>
	med		1.09	0.38	0.33		1.27	0.60	0.25
	min	0.67	0.02	0.01	0.00	0.72	0.05	0.0	0.01
	max	279	9.41	2.74	1.82	234.45	10	4.14	2.11
20	avg	<b>372</b>	<b>12.2</b>	<b>5.86</b>	<b>2.67</b>	<b>444</b>	<b>14.3</b>	<b>7.63</b>	<b>4.52</b>
	med		10.6	4.63	2.01		8.28	5.16	2.76
	min	4.45	0.14	0.07	0.0	5.71	0.21	0.01	0.01
	max	1456	50.6	26	19.2	2540	139	41.7	18.7
21	avg	<b>3743</b>	<b>171</b>	<b>51.4</b>	<b>34.9</b>	<b>5391</b>	<b>153</b>	<b>101</b>	<b>36.7</b>
	med		108	38.5	21.8		111	68.6	24.1
	min	265	5.56	0.24	0.27	96.6	2.18	0.45	0.29
	max	10955	893	235	173	18863	657	560	161
22	avg	-	<b>731</b>	<b>381</b>	<b>200</b>	-	<b>1218</b>	<b>520</b>	<b>220</b>
	med	-	428	286	135	-	819	276	133
	min	-	24.7	13.1	5.23	-	78.9	4.12	3.01
	max	-	6357	1482	656	-	4635	3184	1670

Table V  
EXECUTION TIMES (IN SEC.) ON GRID'5000 (SUNO AND HELIOS)

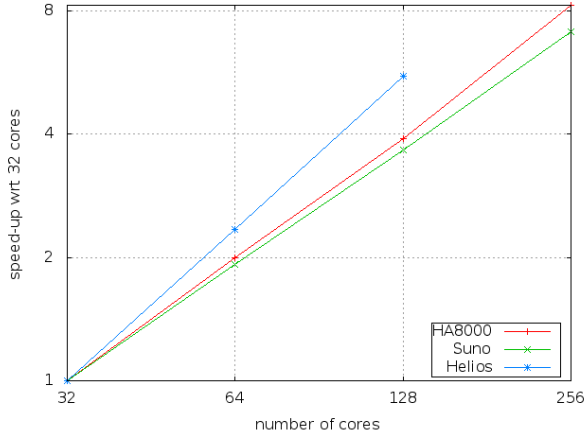


Figure 2. Speed-ups (HA8000 - Grid'5000) for CAP 22 w.r.t. 32 cores

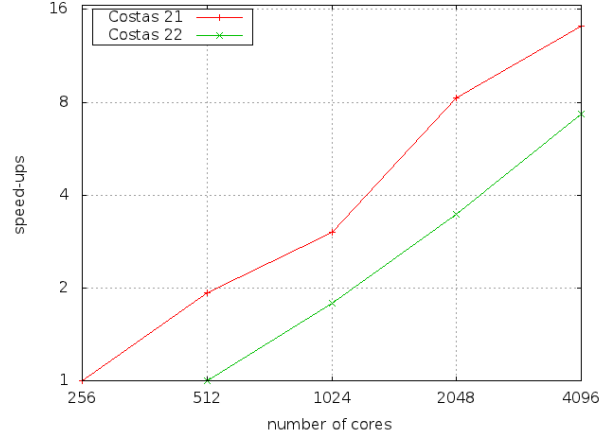


Figure 3. Speed-ups on the JUGENE for CAP 21 and 22

achieve is obviously *zero*, meaning that a solution is found. It is then easy to check if runtime distributions could be approximated by a (shifted) exponential distribution of the form:  $1 - e^{-(x-\mu)/\lambda}$ . Then, according to [39], it is possible to achieve linear speed-ups by multiple independent walks if we have an exponential runtime distribution.

Figure 4 presents time-to-target plots for CAP 21 in order to compare runtime distributions over 32, 64, 128 and 256 cores.

Points represent execution times obtained over 200 runs and lines correspond to the best approximation by an exponential distribution. It can be seen that the actual runtime distributions are very close to exponential distributions. Time-to-target plots also give a clear visual comparison between instances of the same method running on a different number of cores. For instance it can be seen that we have

around 50% chance to find a solution within 100 seconds using 32 cores, and around 75%, 95% and 100% chance respectively with 64, 128 and 256 cores.

## VI. CONCLUSION AND FUTURE WORK

We detailed the modeling and solving of the CAP with the Adaptive Search method. The CAP is a hard combinatorial problem for medium and large instances, too difficult to solve with classical propagation-based solver and we thus used a constraint-based local search solver. We presented a simple modeling and have shown how to further tune the resolution within the Adaptive Search framework. Careful modeling and tuning, together with an efficient parallelization, achieved very good performance. We proposed a parallel version based on the idea of multi-start and independent multiple-walk which naturally provides *Pleasantly Parallel*



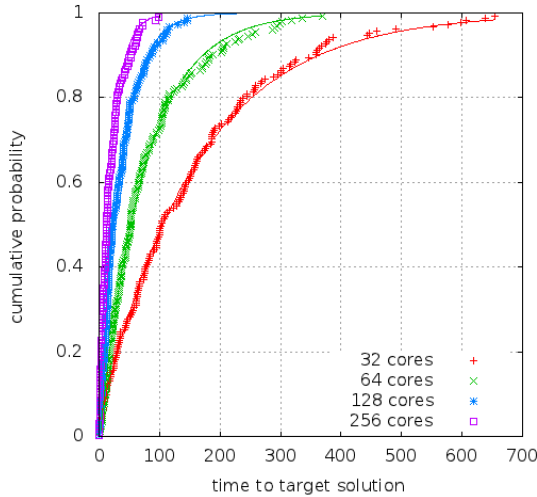


Figure 4. Time-to-Target plots for CAP 21 over 32, 64, 128 and 256 cores

computations and appears viable as it exhibits a nearly linear speed-up behavior. Indeed, we obtained speed-ups w.r.t. sequential implementation of 120 for 128 cores and 226 for 256 cores for medium instances, *i.e.*, nearly linear. For the largest instance studied in this paper (CAP 22), execution times are halved when the number of cores is doubled from 32 cores onward (from 256 cores on the Blue Gene/P), and linear speed-ups can be observed up to 4,096 cores. We are currently continuing our experiments by tackling larger instances and using more cores.

Future work will focus on more complex parallel execution methods with inter-processes communication, *i.e.*, in the dependent multiple-walk scheme, in order to further improve performance. The communication mechanism will be designed with the goals of (1) minimizing data transfers as much as possible, as we aim at massively parallel machines with no hierarchical memory, and (2) re-using some common computations and/or recording previous interesting crossroads in the resolution, from which a restart can be operated.

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