# Power-aware Allocation of Fault-tolerant AUTOSAR Software Applications with End-to-end Constraints via Hybrid Particle Swarm Optimization

#### Abstract

Software-to-hardware allocation plays an important role in the development of resource-constrained automotive embedded systems that are required to meet timing, reliability and power requirements. This paper proposes an Integer Linear Programming optimization approach for the allocation of fault-tolerant embedded software applications that are developed using the AUTOSAR standard. The allocation takes into account the timing and reliability requirements of the multirate software applications and the heterogeneity of their execution platforms. The optimization objective is to minimize the total power consumption of the applications that are distributed over multiple computing units. The proposed approach is evaluated using a range of different software applications from the automotive domain, which are generated using the real-world automotive benchmark. The evaluation results indicate that our proposed allocation approach is effective while meeting the timing, reliability, and power requirements of the considered automotive software applications.

### 1. Introduction

The software-to-hardware allocation is a very important step during the development of automotive embedded systems. Basically, it allows the designer to explore system-level solutions that meet functional and extra-functional software requirements together with resource availability on the execution platform. Software allocation is a well-researched area in the domain of embedded systems, including in hardware/software co-design [1], platform-based system design [2] and the Ychart design approach [3]. It is a type of bin-packing problem, and therefore finding an optimal solution, in the general case, is NP-hard [4]. The methods to solve such problems can be exact [5], which means solutions are guaranteed to be optimal, or heuristic, which deliver near-optimal solutions [6][7]. Exact methods such as Integer Linear Programming (ILP) [8] have been used widely in several resource optimization problems. In contrast to heuristic methods, ILP returns optimal solutions faster for relatively small problems [9]. However, many problems in real-time systems are nonlinear by nature [10], e.g., response time of cause-effect actions, system reliability, etc. To benefit from linear optimization techniques, non-linear functions are approximated using Linearization - a widelyused technique in the optimization of non-linear problems.

In case of fault-tolerance with replication [11], the search space to find the optimal allocation is increased due to the replicas. The search space becomes even larger if we assume that the real-time system executes over different sampling rates, known as multirate [12], in which case the feasible (timed) paths that pass through the different sampling points (or activation patterns) increase exponentially with the number of activation patterns increase. Furthermore, due to the different sampling rates that result in oversampling and undersampling effects, the timing analysis of signals propagation is complex [13]. Existing methods of software allocation lack exact results for the timing analysis of multirate systems.

In this paper, we propose an allocation scheme based on ILP for relatively small- and medium-sized fault-tolerant distributed applications, with the number of allocatable components not exceeding 15, operating-system tasks less than 100, and cause-effect chains in the range of 30 to 60. These parameters are deducted from the real-world automotive benchmark [14], and from previous experience in developing automotive systems and experiments. The applications are distributed over heterogeneous computing units that share a single network. The allocation aims for minimizing the total power consumption of the system while meeting timing and reliability requirements. Our proposed solution targets the automotive domain, in particular systems that conform to the AUTomotive Open System ARchitecture (AUTOSAR) standard. In comparison to related work [9][15][5], we consider a fault-tolerant and multirate system model. Furthermore, we follow a highly integrated approach in the allocation process, which includes response-time analysis (RTA), and utilization bound checking (UB), as well as bounding the level of fault tolerance via the imposed reliability requirement on the application. The main contributions of our work are: i) an ILP model for the allocation of a fault-tolerant multirate application on heterogeneous nodes with the objective of minimizing the total power consumption, and ii) an approach for reducing overhead of replications and cause-effect chains on the allocation of such applications.

Our approach is evaluated on synthetic automotive applications that are generated according to the real-world automotive benchmark proposed by Kramer et al. [14]. In the evaluation, we show the performance of our proposed approach in terms of allocation time and resource efficiency with respect to the size of applications. The tool and the synthetic applications used in this experiment are publicly available from BitBucket<sup>1</sup>.

The rest of the paper is organized as follows. Section 2 provides a brief overview of AUTOSAR-based software development, emphasizing the role of software allocation. Section 3 describes the system model, and Section ?? describes the extra-functional models including the timing, reliability, and power consumption models. Section 4.7 presents the proposed allocation scheme, and in Section 6, we provide an evaluation of the proposed approach using the automotive benchmark. In Section 8, we compare to related work. Finally,

 $<sup>^{1} \</sup>verb|https://bitbucket.org/nasmdh/archsynapp/src/master/$ 

we conclude the paper in Section 9, and outline the possible future work.

#### 2. AUTOSAR

The AUTomotive Open System ARchitecture (AUTOSAR) partnership has defined the open standard AUTOSAR for automotive software architecture that enables manufacturers, suppliers, and tool developers to adopt shared development specifications, while allowing sufficient space for competitiveness. The specifications state standards and development methodologies on how to manage the growing complexity of Electronic/Electrical (E/E) systems, which take into account the flexibility of software development, portability of software applications, dependability, efficiency, etc., of automotive solutions. The conceptual separation of software applications from their infrastructure (or execution platform) is an important attribute of AUTOSAR and is realized through different functional abstractions [16].

# 2.1. Software Application

According to AUTOSAR, software applications are realized on different functional abstractions. The top-most functional abstraction, that is the Virtual Function Bus (VFB), defines a software application over a virtual communication bus using software components that communicate with each other via standard interfaces of various communication semantics. The behavior of a software component is realized by one or more atomic programs known as *Runnables*, which are entities that are scheduled for execution by the operating system and provide abstraction to operating system tasks, essentially enabling behavioral analysis of a software application at the VFB level. The Runtime Time Environment (RTE), which is the lower-level abstraction, realizes the communication between Runnables via RTE Application Programming Interface (API) calls that respond to events, e.g., timing. Furthermore, the RTE implementation provides software components with the access to basic software services, e.g., communication, micro-controller and ECU abstractions, etc., which are defined in the Basic Software (BSW) abstraction [16].

#### 2.2. Timing and Reliability of Applications

The timing information of applications is a crucial input to the software allocation process. Among other extensions, the AUTOSAR Timing Extension specification [17] states the timing descriptions and constraints that can be imposed at the system-level via the *SystemTiming* element. The timing constraints realize the timing requirements on the observable occurrence of events of type *Timing Events*, e.g., Runnables execution time, and *Event Chains*, also referred to as *Cause-effect Chains* that denote the causal nature of the chain. In this work, we consider periodic events and cause-effect chains with different rates of execution (or activation patterns).

Although the importance of reliability is indicated in various AUTOSAR specifications via best practices, the lack of a comprehensive reliability design

recommendations has opened an opportunity for flexible yet not standardized development approaches. In this paper, we consider application reliability as a user requirement and, in the allocation process, we aim at meeting the requirement via optimal placement and replication of software components.

### 3. System Model

The system model shown in Figure 1 is captured as 5-tuple  $\langle A, N, \mapsto, Req, Pro \rangle$ , where  $A = \{A_k : k = 1, \dots, n_A\}$  denotes software applications,  $N = \{n_i : i = 1, \dots, n_N\}$  heterogenous computation (or computing nodes)  $\mapsto: A \to N$  maps the software applications to at least a single computing node, and Req, Pro are assignment functions, respectively defines the requirements of the applications and resource provisions of the computing nodes. The function  $Req: A \to (RL, EE, CL): \mathbb{R}^+ \times [0,1] \times \mathbb{I}^+$  a, assigns the reliability, end-to-end timing and criticality level of applications. The function  $Pro: N \to (HZ, PW, FR): \mathbb{R}^+ \times \mathbb{R}^+$  assigns, respectively the processor speed, power consumption and failure rate of the computing nodes.

The end-to-end timing requirements are the timing constraints over the end-to-end functional behaviors of the applications. These requirements are often specified on the *cause-effect chains* consisting of software components and (potentially network messages) within the application. The reliability requirement is the expected reliability goal of an application which is discussed further in Subsection 4.3, and the criticality level signifies the importance of an application over other applications that have lower criticality levels, thus prioritizing the application during resource contention. The criticality levels are defined systematically, e.g., following the hazard analysis according to the ISO 26262 standard for functional safety in road vehicles [18].

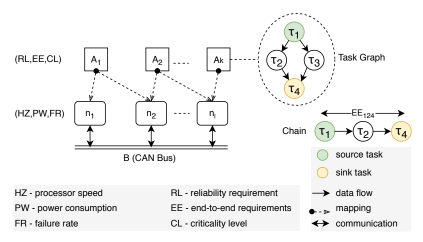


Figure 1: System Model.

Notation	Description
/* Application related */	
$A = \{A_k : k = 1, \dots, n_A\}$	software applications*
$g^{(A_k)}$ denotes the directed acyclic graph	of $A_k$ , where $V(g)$ is the set
of node/vertices and $E(g)$ is the set of ea	
$C^{(k)} = \{c_i^{(k)} : i = 1, \dots, n_C\}$	software-component types of $A_k^{**}$
$Q_i^{(k)} = \{q_{i,j}^{(k)} : j = 1, \dots, n_{Q_i}\}$	component replicas of type $c_i^{(k)}$
/* Execution platform related */	
$N = \{n_i : i = 1, \dots, n_N\}$	computation (or computing) nodes
B	shared CAN bus
$M = \{m_i : i = 1, \dots, n_M\}$	messages on the CAN bus
$\tau, c, m, \gamma$ denote iterator variables, respec	ctively for task, component,
chain and node, e.g., $\forall \tau \in T^{(A_k)}$ .***	
/* Mapping related */	
$\mathbf{x} = {\mathbf{x}_k : k = 1, \dots, n_{\mathbf{x}}} : \bigcup Q_i^{(A_k)} \mapsto M$	a mapping vector from $Q^{(A_k)}$ to $M$
k, i, j denote iterator index-variables, re	spectively for the mapping
vector $\mathbf{x}$ , and rows and columns of the r	$\text{matrix } \mathbf{x}^{(k)}, \text{ e.g., } x_{ij}^{(k)}.***$
$g_{ au}^{(A_k)}(\mathbf{x})$	directed acyclic graph of task nodes for $A_k$ .
$\Gamma^{(k)} = \{\Gamma_i^{(k)} : i = 1, \dots, n_{\Gamma}\}$	end-to-end chains
$\Gamma_{i}^{(k)} = (e_{i})_{i=1}^{Z}$	a chain of $e \in V(g_{\tau}^{(A_k)}) \cup M$
/* Functions related */	- (3, )
$Power(\mathbf{x})$	total power consumption of $A$ in $\mathbf{x}$
$Reliability_a(\mathbf{x})$	application reliability of $a \in A$ in $\mathbf{x}$
$ResponseTime_{\tau}(\mathbf{x})$	response time of $\tau \in V(g_{\tau}^{(A_k)})(\mathbf{x})$
$Delay_{\gamma}(\mathbf{x})$	age delay of $\gamma \in \Gamma^{(A_k)}$ in $\mathbf{x}$

\*Note: the total elements in a set S is denoted by  $n_S,$  e.g.,  $n_A$  denotes the number of

#### 3.1. Software Applications

The software application represents an independent and self-contained user-defined software functionality, e.g., x-by-wire, electronic throttle control, flight control, etc. The application is assumed to be developed using the principles of model-based development and component-based software engineering [19][20].

**Definition 1 (Software Application Model).** It is modeled as a set of communicating software components. Note that a software component is a design-time concept, representing the lowest-level hierarchical element in software architecture of the application. The set of software components in the application is modeled as a directed acyclic vertex-weighted graph  $\langle V_{\tau}, L_{\tau}, w \rangle$  of periodic task nodes  $V_{\tau}$ , where  $a_{ij} \in L_{\tau}$  refers to the data-flow link from node  $\tau_i$  to node  $\tau_j$  and  $i \neq j$ . The function  $w: V_{\tau} \to (e_{n_i}, D, P)$  assigns computation cost to each task node, where  $e_m$  is worst-case execution tim (WCET) on the computing node  $n_i$ , D deadline and P period.

Multiple applications can be executed on the same computating node(s) and can share the on-board network, e.g., the CAN bus. Since the applications can have different criticality requirements, the computing nodes and the network should provide an isolation mechanism in order to avoid interference of lower-criticality applications on higher-criticality applications. The isolation of different criticality applications on the same execution platforms is important in the so-called mixed-criticality systems [21], which can be already found in the avionics domain and also trending in other domains such as the automotive domain where safety-critical applications, such as x-by-wire and electronic throttle control systems, are required to be consolidated with the infotainment system on the same ECUs [?].

#### 3.2. Scheduling Software Applications

The applications are scheduled on the heterogeneous execution platform by considering their respective requirements such as the criticality levels, reliability requirements, and end-to-end timing requirements. There are several techniques in the literature that deal with the scheduling of mixed-criticality applications on *uniprocessor* systems [21]. In our problem, though distributed applications, each task is mapped to a single computing node and the mapping is static. In this case, the schedulability of tasks can be performed per computing node, following the uniprocessor scheduling. Therefore, in the context of this work, the distributed applications are schedulable if the tasks, messages and the cause-effect chains meet their respective timing requirements, but also meet reliability requirements under the limited resources of the execution platform.

In this work, we consider the partitioned criticality (PA) technique to schedule the mixed-criticality applications, which basically prioritizes higher critical applications over their lower-criticality counterparts. In contrast to other techniques, PA does not require a runtime monitoring of tasks, e.g., using servers [22, 23, 24], though less efficient. Note: other scheduling techniques together with the PA technique can be used with our approach.

#### 3.2.1. Scheduling Tasks and Messages

We assume tasks are scheduled using the fixed-priority preemptive scheduling polity (FPPS) [?]. Initally, applications are prioritized based on their criticality levels following the PA technique, and within each application the tasks are prioritized according to the deadline monotonic (DM) priorities assignment.

$$\forall \tau_1 \in V(g_{\tau}^{(A_i)}) \tau_2 \in V(g_{\tau}^{(A_j)}) \ [cri(A_i) > cri(A_j) \implies Pri(\tau_1) > Pri(\tau_2)],$$

where cri, pri are predicates which determine the critiality and priority of tasks  $\tau_1, \tau_2$ , respectively;  $V(g_{\tau}^{(A_i)}), V(g_{\tau}^{(A_j)})$  are the set of tasks which implement the applications  $A_i, A_j$ , respectively.

The schedulability of tasks is checked by the classical response-time analysis shown in Equation (1) [25, 25], which computes the worst-case response time of each task, denoted by  $R_{\tau}$ . According to the analysis, if the response time of each task is less than or equal to its deadline, that is  $R_{\tau} \leq Deadline_{\tau}$ , the taskset is schedulable otherwise it is not.

$$R_{\tau} = c_{\tau} + \sum_{j \in hp(\tau)} \left\lceil \frac{R_{\tau}}{P_j} \right\rceil c_j, \tag{1}$$

where  $c_{\tau}, c_{\tau_h}$  are exection times of the lower and higher tasks, respectivel;  $hp(\tau)$  is the predicate that returns the higher-priority tasks of task  $c_{\tau}$ .

The messages in the CAN bus are scheduled using a fixed, non-preemptive scheduling policy. Similar to the tasks, the priority of messages followes the PA techniques to achieve the mixed-criticality requirement. This can easily be achived by inheriting the priority of sender task, that is  $pri(m) = pri(\tau)|\tau = pre(m)$ , where pre(m) finds the sender task. The schdulability of messages is checked using the classical response-time analysis of the CAN network, presented by Rob Davis et. al [?] as shown in Equation (2). Thus, the worst-case response time of a message is computed as the summation of its *jitter* time (that is, the time taken by the sender task to queue for transmittion)  $J_m$ , the *interferance* time (that is, the message delay in the queue)  $w_m$ , and its *transmission* time (that is, the longest time for a signal or data to be transmitted)  $c_m$ .

$$R_m = J_m + w_m + c_m \tag{2}$$

$$w_m = B_m + \sum_{\forall k \in hp(m)} \left\lceil \frac{w_m + \tau_{bit}}{P_k} \right\rceil c_k \tag{3}$$

$$B_m = \max_{\forall k \in I_n(m)} (c_k),\tag{4}$$

Note: we assume no jitter, therefore, the interference formula is reduced as shown in Equation (3), where  $B_m$  is the blocking time caused by the lower-priority messages using the CAN bus (since it is non-preemptive) and is computed by Equation (4); hp(m) finds the higher-priority messages, which delay the trasmission of the message m in the queue as well as in the transmission.

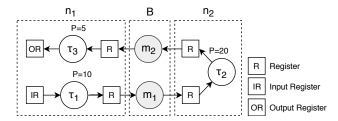


Figure 2: A Cause-effect Chain, mapped on nodes  $n_1$  and  $n_2$ .

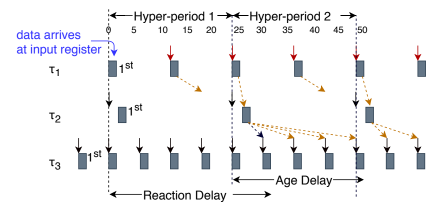


Figure 3: Reaction and Age Delays of the Cause-effect Chain, Shown in Figure 2.

## 3.2.2. Scheduling Cause-effect Chains

The software application can be considered as a set of cause-effect chains  $\Gamma^{(k)} = \{\Gamma_i^{(k)} : i=1,\ldots,n_\Gamma\}$ , which are directed paths in the graph, annotated by end-to-end timing constraints. They represent sequences of actions triggered usually by external events (or causal actions or stimuli) and produce corresponding effects (or responses), e.g., pressing a rotary-wheel to activate a cruise control system, pressing a brake pedal to slow down a car, etc. The end-to-end requirements put upper-bounds on the duration of the stimuli-response elapse time. An example of a cause-effect chain is shown in Figure 2, which consists of three independently clocked tasks  $\tau_1, \tau_2, \tau_3$ , and messages  $m_1, m_2$ . It uses single-register buffers for communication, which is a common practice in control systems design, e.g., automotive software applications [26].

The end-to-end delay in a chain is the duration between the reading of data from the input register by the source task  $Source(\Gamma_i^{(A_k)})$  to the writing of same data to the output register by the last (or sink) task  $Sink(\Gamma_i^{(A_k)})$ . Since we assume the chains consist of independently clocked tasks, the delay usually varies due to the undersampling and oversampling effects in the chains. In this work, we are particularly interested in the two types of delays that are widely used in the automotive and similar systems, namely the age delay and the reaction delay.

The difference between the two types of delays is demonstrated in Figure ??. The tasks  $\tau_1$  and  $\tau_2$  execute on node  $n_1$ , whereas task  $\tau_3$  executes on node  $n_2$ . Note:  $\tau_2$  communicates with  $\tau_3$  via a CAN bus, which is not shown in the figure for simplicity. The red inverted arrows in the figure represent the reading of data from the input register, whereas the dashed-curve arrows represent the timed paths through which the data propagates from the input to the output of the chain. Thus, the age delay is the time elapsed between a stimulus and its corresponding latest non-overwritten response, i.e., between the  $3^{rd}$  instance of  $\tau_1$  and the  $10^{th}$  instance of  $\tau_3$ . It is frequently used in the control systems applications where freshness of data is paramount, e.g., braking a car over a bounded time. And, the reaction delay is the earliest time the system takes to respond to a stimulus that "just missed" the read access at the input of the chain. Assume that data arrives just after the start of the  $1^{st}$  instance of  $\tau_1$  execution. The data corresponding to this event is not read by the current instance of  $\tau_1$ . In fact, the data will be read by the  $2^{nd}$  instance of  $\tau_1$ . The earliest effect of this data at the output of the chain will appear at the  $7^{th}$ instance of  $\tau_3$ , which represents the reaction delay. This delay is useful in the body-electronics domain where first reaction to events is important, e.g., in the button-to-reaction applications. For detailed discussion of the different delay semantics, we direct the reader to check research work by Mubeen et al. [13]. The age delay is computed using Equation (5) and Equation (6) for a single node and multiple nodes, respectively.

$$\Delta_s(\Gamma) = \alpha_{\tau_j} - \alpha_{\tau_i} + R_{\tau_j}$$
 single node (5)

$$\Delta_m(\Gamma) = \sum_{q \in subch(\Gamma)} \Delta_s(q) + \sum_{m \in msg(\Gamma)} R_m, \quad \text{muliple nodes}$$
 (6)

where  $\tau_i = sink(\Gamma)$ ,  $\tau_i = source(\Gamma)$ ,  $\alpha_\tau$  returns the activation time of  $\tau$ .

Assume  $\Gamma \in \Gamma_i^{(A_k)}$  is a chain, if the chain is mapped on a single node, the age delay  $\Delta_s$  is a mere difference between the activation of the sink task  $\alpha_{\tau_j}$  and the activation of the source task  $\alpha_{\tau_i}$  plus the worst-case response time of the sink task  $R_{\tau_j}$  in the longest timed path that complies with the definition of the age delay. On the other hand, if the chain is mapped to multiple nodes, the age delay  $\Delta_m$  can be compositionally computed [27] as follows: the chain is partitioned into a set of sub-chains per node, indicated by the predicate  $subch(\Gamma)$ , and for each sub-chain  $a \in subch(\Gamma)$ , the age delay is computed recursively using the same method to used to compute the age delay for a single node, and the result is added to the response-times of the messages involved in the chain,  $msq(\Gamma)$ .

### 3.3. AUTOSAR Runnables

The AUTOSAR standard introduced the notion of *Runnables* to facilitate early analysis, that is at the VBF level, and to support interoperability of automotive applications across different execution platforms. Basically, runnables are schedulable pieces of codes similar to tasks. In this work, we assume

$r_{i,j}^{(k)}$	$m_h$	$(e_h, P)$	$\prec r_{i,j}^{(k)}$
1,1	1	(1, 10)	2,1
2,1	1	(1,5)	
2,2	1	(1, 15)	
3,1	2	(1,20)	
4,1	3	(1, 10)	
5,1	3	(1,20)	

$ au_i^{(k)}$	$\bigcup r_{i,j}^{(A_k)}$	$(e_h, P)$
1	1,2;2,1	(2,5)
2	2,2	(1, 15)
3	3,1	(1, 20)
4	4,1;5,1	(1, 10)

Table 2: Tasks-Runnables Mappings.

Table 1: Runnables Timing Specifications.

periodically activated runnables with support for multiple worst-case executions that correspond to the different computation processor types. Unlike tasks, runnables' functional and extra-functional properties, e.g., timing, memory requirements, are part of the AUTOSAR software component specification. Therefore, the software application model defined in Definition 1 is extended to accommodate the notion of runnables, using the following simplified formal definition.

**Definition 2 (AUTOSAR Software Application Model).** It is modeled as directed acyclic vertex-weighted graph  $g_r = \langle V_r, L_r, w_r, v \rangle$  of runnable nodes  $V_r$ , where  $a_{ij} \in L_r$  represents either a triggering or data-flow link from the runnable  $r_i$  to runnable  $r_j$  and  $i \neq j$ . The cost  $w_r = w$  defines the timing model of the runnable nodes, which is equivalently defined as the cost of the task nodes explained in Definition 1.

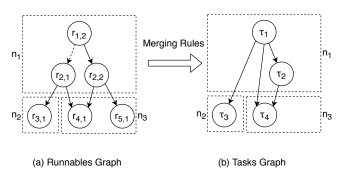


Figure 4: Example of a Software Application, Modeled as Directed Acyclic Graph, where --->, ---> denote triggering link, data-flow link, respectively.

According to the AUTOSAR specification [28], runnables are mapped to tasks, and the tasks execute the runnables respecting their timing specifications. In the mapping process, one or more runnables can be merged to optimize the runtime execution by reducing the number of schedulable tasks. Thereore, through the mappings, eventually runnables graphs are refined by tasks graphs as shown in Figure 4 (b). In this work, the following rules are applied in order to merge any runnables a, b, that is the link  $(a, b) \in L_r$  merges to a task node  $v \in V(g_T)$ , if the following rules satisfy:

(i) the runnables are co-hosted in the same computing node, that is

$$a \mapsto m \wedge b \mapsto m$$

(ii) activation periods of the runnables are the same, i.e., a.P = b.P

If the rules are satisfied, the task's timing specifications are set as follows: i) the WCET of the task is set to the sum of the WCET of the runnables,  $v.e_i = a.e_i + b.e_i$ , ii) the period and deadline of the task is set to the least-common multiple (LCM) of the runnables' periods, v.P = v.D = lcm(a.P, b.P). Otherwise, runnables are not merged, instead, each runnable that is not merged is mapped to a task while preserving the timing specifications of runnables.

## 3.4. Reliability of Software Applications

Redundancy is the most common way to increase the reliability of a system. In this context, software application reliability refers to the probability that a software application functions correctly by the time t, or within the time interval [0,t] [29]. Redandancy can be implemented according to different schemes, such as hot stand-by, cold stand-by, etc [30], in this work, we consider the hot-standby scheme, where replicated components maintain the same state. However, only the primary replicas act on the environment, e.g., activating an actuator. The primary software component is the one in operation and is indicated by  $q_{i,1}^{(k)}$ , the secondary software component, which is in the stand-by, by  $q_{i,2}^{(k)}$ , etc., for a software application  $A_k$ . Note: the software components are replicated unless the reliability requirements of applications are satisfied.

In this work the details of the redundancy scheme are abstracted away under the following assumptions: i) Hot stand-by redundancy technique is used for the replacement of failed components, which are identical and are allocated on different nodes, ii) software components need to be replicated if the application's reliability requirement is not met without replication, otherwise they are not replicated, iii) the time needed to detect and replace a faulty component is considered negligible and will not be taken into account in the response time analysis of tasks and delay calculation of cause-effect chains, iv) Because of its simplicity, the mechanism for detection and replacement of faulty components will be considered fault-free, and therefore will not be included in the reliability calculations.

Under this assumptions, the reliability of a software application is equivalent to the reliability of the execuiton platform such as the computating nodes and the communication bus, if any, on wich the application is deployed. The reliability of a computing node or the bus is calculated using  $e^{\lambda t}$ , where  $\lambda$  is an exponentially distributed failure-rate. However, the reliability calcultion of the execution platform that services a software application is not trivial in the case with replication, e.g., the series-parallel reliability approach cannot be applied in the general case, due to the functional interdependency created between computing nodes as the result. To demonstrate the functional interdependency, let us assume a software application  $A_k$ , having component configurations with

$n_1 \qquad n_2 \qquad n_3$	$n_1$	$n_2$ $n_3$	
$q_{1,1} q_{2,1}$	$q_{1,1}$	$q_{2,1}$ $q_{2,2}$	
$q_{3,1}$	$q_{3,1}$	$q_{1,2}  q_{3,2}$	
(a) Without Replication.	(b) With	n Replication	

Figure 5: Deployments of the Software Application  $A_1$ .

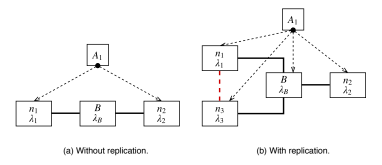


Figure 6: Reliability Block Diagrams of the Software Application.

and without replication as shown Table 5b and 5a, respectively, where  $q_{i,j}^{(A_k)}$  is the  $j^{th}$  software-component replica of software-component type  $c_i^{(A_k)} \in C_i^{(A_k)}$ . Note: for readability of the example, we remove the superscript k.

The reliability of the software application without replication forms a series path, indicated by the reliability block diagram (RBD) of Figure 6a, hence is computed as products of the reliability of  $n_1, n_2$  and B. However, with replication, two computing nodes can form as series and parallel to service the sofware application, e.g., due to  $q_{1,1}$  and  $q_{2,2}$  or  $q_{3,2}$ ,  $n_1$  and  $n_3$  make series, and due to  $q_{3,1}$  and  $q_{3,2}$ , the nodes make parallel, to relialize partional functionality of the application. In this case, the series-parallel diagram depicted in Figure 6b does not accurately capture the reliability calculation of the application with replication. Note: the red-dashed line between  $n_1$  and  $n_3$  indicates the possibility of the computing nodes becoming series. To address this problem, we use the exact reliability calculation technique based on the enumeration of the different failure states of the computing nodes, that is the different failure states of the execution platform is enumerated exaustively, and subsequently, the total probability the software application functions is computed, which is discussed in great length in Subsection 4.3.

# 4. Define the Problem of Mapping the Software Applications

The software applications A are partitioned on the execution platform  $\langle N, B \rangle$  by fulfilling the user-defined software applications requirements such as applications reliability  $RL^{(A_k)}$ , end-to-end timing requirements of chains  $EE_i^{(A_k)}$  and software applications criticality  $cl^{(A_k)}$ , and by meeting design constrains such as placement

restrictions committed by the designer. The partitioning is conducted by mapping the respective software-component replicas (or software components in short) of applications effectively on the computing units M' and the communication between software components on the network bus. The partitioning should result in reduced total power-consumption of the applications  $Power(\mathbf{x})$ , which is achieved by selecting lower-power computing units  $N' \subseteq N$ , where  $\mathbf{x}$  is a possible partitioning matrix (or partition in short) with  $x_{ij}^{(k)}$  representing the mapping of the software component  $q_{i,j}^{(k)}$  to the computing node  $n_h$ , where  $h = x_{ij}^{(k)}$ . Note: under this partition, the reliability of applications  $Reliability_{A_k}(\mathbf{x})$ , the timing of chains  $Delay_{\Gamma_i^{(A_k)}}(\mathbf{x})$  should satify the respective requirements.

The power consumption of applications as well as the delay of chains depend on the tasks graphs  $\bigcup g_{\tau}^{(A_k)}$ , as well as the mapping of the tasks nodes to computing units and the links to the network bus. In this case, the tasks graphs has be constructed from the partition  $\mathbf{x}$ . However, if the tasks graphs are apriori to the software partitioning problem, which can be the case if no runnables from different components map to a single task, only the tasks nodes and the links has to be updated with the mapping information.

### 4.1. Task Graph Formulation

The task graph formulation implies the generation of a task graph, with the cost over the tasks nodes and the links updated for the partition  $\mathbf{x}$ . The tasks nodes are updated with the timing information and the mapping to computing units, likewise, the links with the mapping to the network bus. The formulation has two steps: i) for a mapping  $\mathbf{x}$ , we update the runnables with nodes information to which they are mapped by traversing  $\mathbf{x}$ , in linear-time complexity O(n), where  $n = |\mathbf{x}| * n_r$ , and  $n_r$  is the runnables per software component, using Equation (7), ii) by applying the merging rules, we traverse the runnables graphs in the system and subsequently generate the tasks graphs using Equation (8).

$$\forall k, ij \ \forall r \in V(g^{(A_k)}) \ node_r = n_h$$
 , where  $h = x_{ij}^{(k)}$  (7)

$$\forall k \ g_r^{(A_k)}(\mathbf{x}) \xrightarrow{\text{Eqn. (13);Merging Rules}} g_{\tau}^{(A_k)}(\mathbf{x}), \tag{8}$$

where  $V(g^{(A_k)})$  is the nodes of the runnables graph  $g^{(A_k)}$ .

# 4.2. Total Power Consumption

Power consumption refers to the energy usage of electronic components in an integrated circuit, e.g., processor, memory, I/O devices, etc., per time unit. There are several models (or techniques) to estimate the power consumption of a computing node. In this work, we use a technique based on processor load (or utilization) to estimate the average power consumption of a computinh node. Specifically, we use the linear polynomial model proposed by Fan et al. [31], which is shown in (9). The model states that the power consumption of

a node is directly proportional to its load, and is inductively formulated from experimental results:

$$\mathcal{P}(u) = P_{idle} + (P_{busy} - P_{idle}) * u, \tag{9}$$

where u is the utilization of a computing unit,  $P_{idle}$  and  $P_{busy}$ , respectively refer to the power consumption measured at minimum and maximum processor loads. The measurements can be obtained by running performance benchmark suits, e.g., MiBench [32], AutoBench [33], etc. The utilization of the computing units for a partition  $\mathbf{x}$  is computed as a sum of the utilization of their respective constituent tasks as shown in Equation (11). Finally, the total power-consumption of the applications is the sum of the power-consumption of the units as shown in Equation (12).

$$\mathcal{U}(\tau) = \frac{\text{WCET}_{\tau}}{P_{\tau}} \tag{10}$$

$$(u_1, ..., u_{n_N}) \leftarrow \sum_{k=1}^{n_A} \sum_{\tau \in V(q_{\tau}^{(A_k)}(\mathbf{x}))} (\mathcal{U}(\tau)|Node_{\tau} = n_h, h = 1, ..., n_N)$$
(11)

$$\mathcal{P}_{total}(\mathbf{x}) = \sum_{h=1}^{n_N} u_h(\mathbf{x}) \tag{12}$$

where  $\mathcal{U}$ , in Equation (10), computes utilization of task  $\tau$  as a ration of its worst-case execution time and period, and  $u_h$  is utilization of node  $n_h$ .

### 4.3. Software-Applications Reliability Constraints

The applications reliability constraints ensure the mapping  $\mathbf{x}$  satisfies the user-defined reliability requirements, that is  $\forall k \; Reliability_{A_k}(\mathbf{x}) \leq RelReq_{A_k}$ . The reliability of each application is computed over t period of time from the computing units  $N^{(A_k)}$  and the shared network bus B, where  $N^{(A_k)}$  hosts  $A_k$ . The reliability is computed assuming exponentially distributed and constant failure rates of the units  $\lambda_{n_h}$  as well as the network bus  $\lambda_B$ . Thus, the reliability of an application is computed as a product of the reliability of the units and the network bus as shown using Equation (13). Note: if application does not use the shared bus  $Reliability_B = 1$ . Equation (14) finds the units  $N^{(A_k)}$  that the application  $A_k$  uses by traversing the partition  $\mathbf{x}$  in linear time.

$$Reliability_{A_k}(\mathbf{x}) = Reliability_{N(A_k)}(\mathbf{x}) * Reliability_B$$
 (13)

$$N^{(A_k)} = \{ e \in N | \forall ij \ e = m_h \}, \text{ where } h = x_{ij}^{(k)}$$
 (14)

Note: we assume applications are mutually exclusive, that is no shared components exist between any two applications, therefore, we can safety calculate the reliability of applications independently. Consequently, to increase readability, we remove the superscript  $(A_k)$  in the rest of this subsection.

The reliability of the units is  $Reliability_N(\mathbf{x}) = e^{-\lambda_N(\mathbf{x})t}$ , where is  $\lambda_N(\mathbf{x})$  is the failure rate of an N-unit system over the partition  $\mathbf{x}$ . The system failure-rate is computed using the state enumeration as shown in [34], which is an exact technique to calculate reliability, as opposed to using series-parallel technique motivated in Subsection 3.4. By applying the state enumeration technique, the system failure-rate can be defined as the probability a software application fails in the probability space  $\langle \Omega, \xi, p, f \rangle$ .

- $\Omega = \{0,1\}$  are the possible outcomes (or states) of a computing unit. Assume the Boolean variable  $s_h \to \Omega$ , which indicates the state of  $n_h$ , then  $s_h = 0$  indicates  $n_h$  fails and  $s_h = 0$  indicates  $n_h$  operates. Thus, for computing units  $N = \{n_1, ..., n_{n_N}\}$ , the states of the units (or configuration) is indicated by the N-cardinality set  $S = \{s_1, ..., s_{n_N}\}$ .
- $\xi = \Omega^S$  are elementary events that correspond to the possible configurations of the units N, therefore, the events are mutually exclusive. Consider  $N = \{n_1, n_2, n_3\}$ , Table (3) shows the their possible configurations  $\xi$ . Assume the configuration  $s \in \xi = \{0, 1, 0\}$ , it shows  $n_1$  and  $n_3$  fail as indicated by  $s_1 = 0$ ,  $s_3 = 0$ , respectively, and  $n_2$  operates as indicated by  $s_2 = 1$ .
- $p:\xi \to [0,1]$  assings the configurations probabilities using

$$\forall s \in \xi \ p_s = \prod_{h=1}^{n_N} \lambda_{n_h} * (1 - s_h) + (1 - \lambda_{n_h}) * s_h$$

where  $\lambda_{n_h}$  is the failure-rate of  $n_h$ . The probability  $p_s$  is the product of the probability of having the state  $s_h$ , which is  $\lambda_{n_h}$  if  $n_h$  fails, otherwise,  $(1 - \lambda_{n_h})$  if  $n_h$  operates.

•  $f: \xi \to \{0,1\}$  determines the status of the application in each state  $s \in \xi$ , that is  $f_s = 0$  means the application fails, otherwise,  $f_s = 1$  means the application operates, at the sate s.

**Definition 3 (Software Application Failure).** A software application fails in the configuration  $s \in \xi$  if there exists a component type  $c_i$  where all of its replicas  $Q_i$  fail, otherwise, it functions, as shown using Equation (15). The component replica  $q_i, j \in Q_i$  of type  $c_i$  fails if  $n_h$  fails, that is  $s_h = 0$ .

$$f_s(\mathbf{x}) = \begin{cases} 0 & \text{if } \exists i \ c_i | \forall j \ s_h = 0 \\ 1 & \text{otherwise} \end{cases}$$
 where  $h = x_{ij}$  (15)

Thus, the failure rate of the N-unit system  $\lambda_N(\mathbf{x})$  is the sum of the probabilities in which the application fails, that is

$$\lambda_N(\mathbf{x}) = \sum_{s \in \xi \mid f_s(\mathbf{x}) = 0} p_s(\mathbf{x})$$

$s \in \xi$	$p_s$	$\forall i \ s_{c_i}$	$f_s$
$\{0,0,0\}$	0.0000000000	$\{0, 0, 0\}$	0
$\{0,0,1\}$	0.0000000099	$\{0, 0, 1\}$	0
$\{0,1,0\}$	0.0000000099	$\{1, 0, 0\}$	0
$\{0,1,1\}$	0.0000999800	$\{1, 1, 1\}$	1
$\{1,0,0\}$	0.0000000099	$\{1, 0, 1\}$	0
$\{1,0,1\}$	0.0000999800	$\{1, 1, 1\}$	1
$\{1,1,0\}$	0.0000999800	$\{1, 1, 1\}$	1
$\{1,1,1\}$	0.9997000299	$\{1, 1, 1\}$	1

Table 3: Example of Application Reliability Calculation using State Enumeration Over 10-years Operational Lifetime: an Application with Component Types  $C = \{c_1, c_2, c_3\}$ , Replicas  $Q = \{c_{1,1}, c_{1,2}; c_{2,1}, c_{2,2}; c_{3,1}, c_{3,2}\}$  Partitioned on  $N = \{n_1, n_2, n_3\}$  according to Figure (5), the Variable  $s_{c_i} \in \{0, 1\}$  Indicates if the Replicas of Type  $c_i$  Fails or Functions, Respectively.

**Example 1 (Reliability Calculation).** Let us assume we want to calculate the reliability of the application in Table (3) over a 10-year (or 87600h) operational lifetime. The reliability of the units is Reliability<sub>N</sub> =  $e^{-\lambda_N t}$  = 0.99736671, where  $\lambda_N = p_1 + p_2 + p_3 + p_5 = 0.0000000301$ . Assume  $\lambda_B = 0.00000001$ , hence Reliability<sub>B</sub> =  $e^{-\lambda_B t}$  = 0.99912438. Then, the reliability of the application is Reliability<sub>N</sub> \* Reliability<sub>B</sub> = 0.99649339932.

### 4.4. Timing constraints

The timing constraints ensure that the schedulability of each applications on the execution platform, that is, the tasks and the cause-effect chains of each application should meet their timing specifications. The schedulability of each task is checked using the worst-case response-time anlaysis presented in Subsection 3.2.1, and the cause-effect chains using the age-delay analysis shown in Subsection 3.2.2.

### 4.5. Tasks Timing constraints

The timing constraints over the tasks ensure that the worst-case response time of each task in the system meets its respective deadline, in the partition  $\mathbf{x}$ , that is  $\forall k \forall \tau \in V(g_r(\mathbf{x}))$  Response $Time_{\tau} \leq Deadline_{\tau}$ , where  $V(g_r(\mathbf{x})_i^{(A_k)})$  is the nodes in the tasks graphs. To compute the worst-case response time of the tasks, first we arrange the tasks per node as represented by the  $T_{n_h}$ , by traversing the tasks graphs using Equation (16). The complexity of this equation, considering an adjucency matrix implementation, is linear time  $O(N_a*n^{(A_k)})$ , where  $n^{(A_k)}$  is the sum of nodes and the links of graph  $g_{\tau}^{(A_k)}(\mathbf{x})$ .

$$T_{n_h} = \{ e \in V(g_{\tau}^{(A_k)}(\mathbf{x})) | Node_e^{(A_k)} = n_h \}$$
 for all  $h = 1, ..., n_N$ , (16)

where where  $h = x_{ij}^{(k)}$ .

Then, we calculate the response time of each task  $\tau \in T_m$  by invoking the response-time analysis formula, and construct the tasks timing constraints as shown in Equation (17).

$$\forall \tau \in T_m \ ResponseTime_{\tau}(\mathbf{x}) \le Deadline_{\tau} \tag{17}$$

### 4.6. Cause-effect Chains Timing constraints

For a mapping x, the age delays of cause-effect chains should meet thier respective end-to-end requirements, that is  $\forall k \forall ij \; AgeDelay_{\Gamma}(\mathbf{x}) \leq E2eReq_{\Gamma}^{(A_k)}$ , where  $\Gamma \in \Gamma_i^{(A_k)}$ . To calculate the age delays, first we identify the messages scheduled by the CAN bus for the mapping x using Equation (18).

$$M = \{e | \forall (a, b) \in g_{\tau}(\mathbf{x}) \forall n \in N \ (a \mapsto n \land b \mapsto n = false) \implies createMsg(e)\},$$
(18)

where Period(e) = Period(a), that is the message inherits the period of its predecessor (or sender) task. Accordingly, we update only the chains that communicate over the shared CAN bus to incorporate the messages, that is  $\Gamma_i^{(A_k)} = \{\tau_1, e^*\}$ , where  $\tau_1, \tau_2$  are  $Source(\Gamma_i^{(A_k)}) and Sink(\Gamma_i^{(A_k)})$ , respectively, and  $e \in V(g_{\tau}(\mathbf{x})) \cup M$ . Then, the cause-effect timing constrains are formulated over the updated list of chains  $\Gamma_i^{(A_k)}$  using Equation (19).

$$\forall \gamma \in \Gamma_i^{(A_k)} \ Delay_{\gamma}(\mathbf{x}) \le E2eReq_{\gamma} \tag{19}$$

# 4.7. Software Allocation Optimization

The software allocation is defined as a single-objective optimization problem. The objective function  $Power(\mathbf{x})$  is a cost function which minimizes the total power consumption of the software applications as deployed in the heterogenuous computing units, where  $\mathbf{x}$  is the decision variable (or solution) of the optimization. The cost function is formulated in Equation 20, with inequality constraints shown by Equation (21, 22,23). The constraints ensure the solution meet the reliability requirements, the tasks deadlines, and the chains end-to-end requirements. Furthermore, the overlapping constrain shown in Equation (24) ensure that replicas are not allocated to the same computing units.

$$\min_{\mathbf{x} \in X} Power(\mathbf{x})$$
 subjected to: (20)

$$Reliability_{A_k}(\mathbf{x}) \le RelReq_{A_k}$$
 for all  $k = 1, ..., n_{A_k}$  (21)

$$Reliability_{A_k}(\mathbf{x}) \leq RelReq_{A_k}$$
 for all  $k = 1, ..., n_{A_k}$  (21)  
  $\forall \tau \in T_{m_h} \ ResponseTime_{\tau}(\mathbf{x}) \leq Deadline_{\tau}$  for all  $h = 1, ..., n_M$  (22)

$$\forall \gamma \in \Gamma^{(A_k)} \ Delay_{\gamma}(\mathbf{x}) \le E2eReq_{\gamma} \qquad \text{forall } k = 1, ..., n_A \quad (23)$$

$$\forall k \forall ij \ x_{ij}^{(k)} \neq x_{ij}^{(k')},$$
 where  $k \neq k' = 1, ..., n_{rep}$  (24)

where X is the search space of the problem,  $\mathbf{x} \in X$  is a feasible solution, and  $x_{ij}^{(k)} \in \mathbf{x}$  is a mapping of a component  $q_{i,j}^{(A_k)}$  to the node  $m_h$ , where  $h = x_{ij}^{(k)}$ 

In the next section, we discuss our proposed method to address the considered optimization problem.

#### 5. Solution using Hybrid Particle Swarm Optimization (PSO)

In our previous work, we provided an ILP model for the same optimization problem, then the CPLEX solver returned optimal solutions to problems in the range *small* and *medim*, where the small problem refers to a software application with software components less than 10, chains less than 30, and similarly the medium problem refers to applications with components less than 15, and chains less than 40. The problems specifications are stipulated from the real automotive benchmark proposed by Kramel et al. [14]. However, the ILP approach, as also shown for similar problems, suffered from the scalability problem of large software allocation problems. In this section, we propose a metaheuristic approach based on the particle-swarm optimization (PSO), evolutionary, differential evolution (DE), hybrid PSO with DE, hill-climbing and stochastic hill-climbing.

Metaheuristics does not guarantee optimal solutions, nevertheless, the solutions can be good enough (or acceptable) in practice. Thus, although the power consumption of the applications may not be optimal, the solution can be deemed acceptable. PSO has been applied to solve a wide range of problems, including a task allocation problem [35], and DE is shown to scale well for problems with high dimensions. In fact, PSO and DE are used together for improved performance in several optimization problems [?], likewise, PSO is used with local search techniques such as Hill climbing to intensify the search [?]. Finally, we evaluate the different meta-heuristic methods based on solution quality and computation time for different software allocation problems.

A meta-heuristic algorithm comprises two major parts: solution representation and an objective function. The solution representation shows the data structure that is used to represent each point in the problem space, and it has a significant impact on the performance of the meta-heuristic algorithm. The fitness function is used to evaluate the quality of candidate solutions based on their fitness to meet the problem objectives. A solution that delivers lower power consumption and violates less constraints is indicated by a lower fitness value. In the following, we describe the solution representation and the objective function proposed in our solution framework to run the meta-heuristic algorithms.

# 5.1. Solution Representation

The software allocation is a type of Assignment optimization problem, as such, the solutions are discrete values. There are two commonly used solution encodings (or representations) are binary (0-1) and integer. The binary variable indicates if a component is allocated to a computing node or not. In the integer representation, the variable indicate the computing-node identifier to which the component is allocated. The two representations are demonstrated in Figure 9 using the example provided in Figure 5.

Г1	Ω	٦٥	Γο	1	٦٦	[1	2
0	1	0		0	1	$\begin{bmatrix} 1 \\ 2 \\ 1 \end{bmatrix}$	3
$\frac{1}{1}$	1	0	$\begin{bmatrix} 0 \\ 0 \\ 0 \end{bmatrix}$	0	1	[1	3_

Figure 7: Binary (0-1) Representation.

Figure 8: Integer Representation.

Figure 9: Solution Representations for Components  $\{c_1, c_2, c_3\}$  Mapped to Computiong Nodes  $\{n_1, n_2, n_3\}$  based on Table 5b.

In this work, consider the integer representation due to efficient encoding (much fewer variables), and it is computationally more efficient, considering the various operations in the optimization problem, e.g., the binary representation shown in Figure 9 uses a pair of metrices to represent the primary and secondary replicas using 18 binary variables versus only 6 variables used in the case of integer representation shown in Figure 8.

#### 5.2. Fitness Function

The fitness function  $f: \mathbf{x} \to \mathbb{R}$  is a type of objective function that summerizes the contriutions of the decision variables via real numbers. The fitness value is used to compare feasible solutions, that is the higher the fitness, the better. In the context of metaheuristics, it is highly desirable to integrate the goal function and all constraints into one function that can be used as a fitness function. [cite to talbi2009metaheuristics and faragardi2018efficient]. Thus, it combines the objective function, which is the power-consumption minimization, with the constraints such as the reliability and timing constraints into a single function by using a penality function.

The benefit of using a single function, including all penalty functions, is to provide a metric to distinguish between two unfeasible solutions. For example, let us assume that A and B are two different solutions for the allocation problem while both violate some constraints of the problem. Let us also assume that solution A slightly violates only one constraint, whereas solution B significantly violates multiple constraints. If the heuristic algorithm can perceive the difference between A and B in terms of being far away from a feasible solution, it guide the search toward a feasible solution more efficiently, compared to the case that the heuristic algorithm only knows that they are both infeasible. The integration of the goal function with all the penalty functions is a promising solution to provide knowledge about how far an unfeasible solution is from a feasible solution.

Consequently, the original constrained optimization problem is transformed into unconstrained optimization problem, by extending the objective function  $P(\mathbf{x})$  with the constrains, represented by a set of penalty functions  $\{\phi_{reliability}(\mathbf{x}), \phi_{deadline}(\mathbf{x}), \phi_{e2e}(\mathbf{x}), \phi_{rep}(\mathbf{x})\}$ . The first penalty function corresponds to the reliability constraint which returns 0 if the reliability constrain is not violated, otherwise returns a positive number denoting how far the reliability constraint is violated. The further violation, the higher value of the penalty function.

Similarly, the other penalty functions correspond to the deadline, the end-toend timing requirement, and the replication constraints, respectively. Indeed, the penalty functions penalize the candidate solution by increasing its fitness (for our minimization problem), thus discriminating the solution. Section 5.3 explains how our solution framework formulates the penalty functions.

The fitness function f(x) is computed as follows.

$$\min_{\mathbf{x} \in X} f(\mathbf{x}) = P(\mathbf{x}) + \beta_1 \phi_{reliability}(\mathbf{x}) + \beta_2 \phi_{deadline}(\mathbf{x}) + \beta_3 \phi_{e2e}(\mathbf{x}) + \beta_4 \phi_{rep}(\mathbf{x})$$
(25)

where  $\beta_1$  to  $\beta_4$  are penalty coefficients used to tune the weight of the penalty functions with regard to the range of the objective function. In Section 5.4, the proper value of the penalty coefficients is discussed in more details.

#### 5.3. Penalty Function

$$\phi_{reliability}(\mathbf{x}) = \sum_{k=1}^{n_{A_k}} Max\{0, Reliability_{A_k}(\mathbf{x}) - RelReq_{A_k}\}$$
 (26)

$$\phi_{deadline}(\mathbf{x}) = \sum_{\forall \tau \in T_{m_h}} Max\{0, ResponseTime_{\tau}(\mathbf{x}) - Deadline_{\tau}\}$$
 (27)

$$\phi_{e2e}(\mathbf{x}) = \sum_{\forall \gamma \in \Gamma^{(A_k)}} Max\{0, Delay_{\gamma}(\mathbf{x}) - E2eReq_{\gamma}\}$$
 (28)

$$\phi_{rep}(\mathbf{x}) = \sum_{\forall c_i} \sum_{\forall n_j} \sum_{k=1}^{n_{rep}-1} \sum_{k'=k+1}^{n_{rep}} x_{ij}^{(k)} x_{ij}^{(k')}$$
(29)

#### 5.4. Penalty Coefficients

# 5.5. Metaheuristic Algorithms

# 5.5.1. Particle Swarm Optimization

PSO is a population-based technique proposed by Eberhart and Kennedy in 1995 to study social behavior, as inspired by natural swarm intelligence observed from the flocking of birds and schooling of fishes [36]. Since then, it is extended in order to address various metaheuristic optimization challenges, such as intensification, diversification, convergence analysis, local optima, parameter tuning, computation time, etc. It is successfully applied on several complex realworld problems, e.g., diagnosis and classification of diseases, efficient engineering designs, tuning control design parameters, scheduling problems, etc [37].

In PSO, the population (or swarm)  $PN = \{p_1, p_2, \dots, p_N\}$  is a collection of particles  $p_i \in PN$ , organized according to a certain population topology [38]. A particle has a position  $\mathbf{x}$  and a velocity  $\mathbf{v}$ , which denote current location and

direction of the particle's motion, and current momentum, respectively. It is a memory-based technique, that is, it remembers the best performance of every particle as well as the best performance of the swarm  $\mathbf{z}$  in order to plan for the next move of the particles, where  $\mathbf{y}, \mathbf{z}$  are position vectors and have the same dimensions as  $\mathbf{x}$ . The velocity of a particle is the resultant vector of its current velocity and the particles attraction vectors  $(\mathbf{y}-\mathbf{x}), (\mathbf{z}-\mathbf{x})$ , respectively, known as *cognitive* and *social* components of the particle's velocity formula, as shown in Equation 30. The attraction vectors impose force of attraction on the particle to move closer to their respective components. Thus, the next position of a particle is the resultant of its current position and its next velocity as shown in Equation (31).

$$\mathbf{v} \leftarrow \omega \mathbf{v} + c_1 Rand() \circ (\mathbf{z} - \mathbf{x}) + c_2 Rand() \circ (\mathbf{z} - \mathbf{x})$$
 (30)

$$\mathbf{x} \leftarrow \mathbf{x} + \mathbf{v}$$
 (31)

where  $\omega$  is the weight of the velocity, also known as *inertia coefficient*, and controls the convergence of the algorithm,  $c_1, c_2$  are acceleration coefficients and controls the weight of attraction towards the cognitive and social components, respectively,  $Rand() \in U(0,1)$  is a random function, along the acceleration coefficients, is element-wise multiplied with the components to improve diversity of the search by introducing stochastic behavior.

Although PSO was originally proposed for continuous problem, it is applied to discrete problems successfully as well. In the latter case, the solutions are represented by 0-1 integer variables [39] or integer-linear by approximation to the nearest integer values [40], which is the representation employed adopted in our problem as it is compact, hence fewer decision variables. Accordingly, after the new position (or candidate solution) is determined, following Equations 30 and 31, the solution is discretized by rounding off the its elements to the nearest integer values, that is  $\mathbf{x} \leftarrow [\mathbf{x}]$ .

## 5.5.2. Differential Evolution

Similar to PSO, Differential Evolution (DE) is a population-based metaheuristic technique for the global optimization which includes non-linear and non-differentiable problems. It was initially proposed by Storn and Price in 19995 [41], since then it has improved with regard to the different operators of DE such as mutation and crossover, and variants over population topology and hybridization [42]. It is a parallel search technique, therefore, is ideal for computationally intensive problems, and employs mutation and crossover operators that allow the search to skip local minima as opposed to PSO.

In every generation, the population undergoes mutation, crossover, and selection according to the formulas shown in Equation , (33), and (34), respectively. A mutant vector v is created from randomly selected elements  $\{a, b, c\} \in PN$  according the mutation operation shown in Equation (32), that is by adding the base matrix to the weighted difference matrix  $F \circ (b - c)$ , where F controls the

amplification of the  $(\mathbf{b} - \mathbf{c})$  variation.

$$\mathbf{v} \leftarrow \mathbf{a} + F \circ (\mathbf{b} - \mathbf{c}) \tag{32}$$

$$u_{ik} \leftarrow \begin{cases} v_{ik} & \text{if } U(0,1) \le CF \text{ and } h = (i*K+k) \\ x_{ik} & \text{if } U(0,1) > CF \text{ and } h \ne (i*K+j) \end{cases}$$
 (33)

$$\mathbf{x} \leftarrow \begin{cases} \mathbf{u} & \text{if } f(\mathbf{u}) < f(\mathbf{x}) \text{ functions} \\ \mathbf{x} & \text{otherwise} \end{cases}$$
 (34)

# 5.5.3. Hybrid Particle Swarm Optimization

The canonical PSO technique uses the constriction factors to balance exploitation and exploration of the search space, that is to deliver better quality solutions. Nevertheless, it still suffers from local minima especially for complex and large problems that exhibit especially multimodal behavior. Hybridization of PSO is one the most widely studied approach in the improvement of the the PSO technique. Basically, it combines other optimization techniques, for instance to intensify local search, and improve diversification by introducing stochastic search. However, hybridization of PSO usually incurs additional computation time. Therefore, the benefit of hybridization has to be studied carefully in conjunction to computation time. Moreover, it should not complicate the user-configurable parameters, to be inline with the philosophy of PSO for ease-of-use.

PSO is hybridized with several optimization techniques, such as Genetic Algorithm (GA), DE, local searches (e.g., Hill-climbing, gradient decent, etc.), ant colony, simulated annealing, etc. Of which, it is shown to perform better when hybridized with DE on constrained, discrete, large benchmarks. Furthermore, it is shown to perform better when hybridized with Hill-climbing (specifically Steepest-descent variant) for software allocation problem [] in particular. In this paper, we hybridize PSO with DE (DEPSO) and Hill-climbing (HCPSO) to the solve the software allocation problem as formulated in Equation (x). In the latter case, we also apply the stochastic variant of Hill-climbing (SHPSO) in order to offset stagnation of the steepest Hill-climbing when applied on large software allocation problems.

### 5.5.4. Differential Evolution PSO (DEPSO)

DE complements the classical PSO by introducing stochastic behavior via the evolutionary operators such as mutation, cross-over and selection. In this specific hybridization approach, we allow the DE algorithm to run intermittently for some number of generations before the next PSO generation starts.

# 5.5.5. Hill-climbing PSO

Hill-climbing is a popular local search based on the notion of *neighborhood*, that is, the candidate solution (or neighbor) that performs better is selected iteratively until no improvements can be made. The software allocation solution  $\mathbf{x}$  is neighbor to  $\mathbf{x}'$  if  $\mathbf{x} = \mathbf{x}'$  except  $\exists i, j \mid x_{ij} \neq x'_{ij}$ , that is, a single mapping is different. In every iteration, the best neighbor is selected, and subsequently

replaces the current candidate solution if it performs better, and continues until maximum iteration, this variant is known as Steepest-descent Hill-climbing (SHC).

Since SHC exhaustively checks all neighbors before moving to the next iteration, the computation time is high especially for high-dimensional problems. To offset this problem, we also apply the stochastic version of Hill-climbing. In the later case, the neighbor is selected randomly, first by selecting the dimension, that is the component  $c_{ij}$ , where i = U(1, I) and j = U(1, K), second, selecting the value, that is the node  $n_j$ , where j = U(1, J). If the neighbor improves the current candidate solution sufficiently, the search moves to the next iteration, which is until no more improvements can be made.

#### 6. Evaluation

In this section, we evaluate our proposed hybrid PSO algorithms for the allocation of software applications on heterogenous computing units, which conform to the system model presented in Section 3. The algorithms are evaluated against different specifications of automotive software applications and execution platforms with regard to effectiveness, stabilitity and scalability. The software-application specifications consist of the number of software components c, runnables r, tasks t and cause-effect chains g. The specifications are synthesized from the automotive benchmark proposed by Kramel et al. [14]. The benchmark indicates a strong correlation between runnables and cause-effect chains in terms of timing and activation patterns. It shows the timing specifications of runnables and their shares in an engine management system. Moreover, it shows the activation patterns of cause-effect chains, the runnables per activation and their shares in the system. The engine management system is one of the most complex automotive systems in the vehicular electical/electronic execution platform.

Software Applications Benchmark. Based on our experience in the automotive industry, the benchmark results are extrapolated to characterize different classes of automotive software applications specifications, that is by varying the parameters related to the software components, runnables, and cause-effect chains. The different classes of specifications range from Spec-I to Spec-V as shown in Table ??. The specification classes are useful to evaluate and discusse the effectiveness and scalability of the different optimization algorithms. The first specification class Spec-I encompases small software applications with number of components less than 10, runnables less than 50, tasks 30, cause-effect chains less than 30. The Spec-I and Spec-II classes represent medium and large software applications, and the last specification class introduced to strech the performance analysis. Note: the classification is superficial and maynot be realistic and could also be different in different domains.

Execution Platform Specifications. Likewise, the specifications for an execution platform consist of the processor speed, power specifications and failure rates of computing units, and we assume the range of values to these parameters as shown in Table 6.

Parameter	SpecI	SpecII	SpecIII	SpecIV
Components $c$	≤ 10	$\leq 15$	≤ 20	<u>≤</u> 80
Runnables $r$	$\leq 50$	$\leq 100$	$\leq 500$	$\leq 1000$
Tasks $t$	$\leq 30$	$\leq 60$	$\leq 80$	$\leq 100$
Cause-effect chains $g$	$\leq 30$	$\leq 40$	$\leq 60$	$\leq 100$
Activation-pattern		{2,	3, 4}	
share of activation-patterns		{0.7,0	0.2, 0.1}	

Table 4: Specification of the Applications for Evaluation.

Parameter	Range
EE	$100n_{\Gamma}$
RL	0.99999999
CL	$\{A,B,C,D\}$

Parameter	Range
Nodes $n_N$ $P_{min}, P_{max}$ (Watt) $\lambda_n \ (h^{-1})$ $\lambda_B \ (h^{-1})$	$ 4 - 10  1 - 10  10^{-4} - 10^{-2}  10^{-4} - 10^{-2} $
Hz (MHz)	80 - 800

Table 5: Ranges of Values for Applications Requirements.

Table 6: Ranges of Values for Execution Platforms.

Applications Requirements Specifications . Table ?? shows the range of values used in our experiment to specify the requirements of software applications, that include the end-to-end timing requirements EE of chains, the reliability requirement RL and the criticality level CL. The end-to-end requiremens are assumed as a function of length of the chain  $n_{\Gamma}$ , that is the longer the chain, the higher the number. The reliability range of safety-critical automotive application is usually given in higher degree of 9, for operation of over a long period of time, which implies almost no failure during the specified duration.

Evaluation Setup. The evaluation is conducted on a MacBook Pro laptop comptuer, with hardware specifications as follows: Intel Core i7 processor type, 2.6.GHz processor speed, 6 Cores, 9 MB L3 cache, and 16 GB memory.

# 6.1. Result

We conduted two experiments: i) the first experiment is designed to compare the performance such as converge time, computation time, optimality (or solutions quality), and stability of solutions of the meta-heuristic algorithms used in this paper, ii) the second experiment is designed to evaluate the overhead of increasing replication on the optimization especially on the computation of cause-effect chains, and also to evaluate the effect of the approximation algorithms proposed in Subsection x to reduce the overheadd and maybe trade-off with optimality of solutions.

Experiment 1. According to the specifications of the range discussed, we synthesized six optimization problems as shown in Table 10. The problems emulate the

Algorithm	Parameters Settings
PSO	Particle Swarm Optimization: learning factors $c_1 = c_2 = 1.49445 \in$
	[0, 4], number of particles 40, iterations 5000
DE	Differential Evolution: crossover $CR=0.5\in[0,1],$ scale factor $F=$
	$0.7 \in [0, 2]$
PF	Penality Function: $\beta_1 = \beta_2 = \beta_3 = \beta_3 = \beta_3$

Table 7: Parameters Settings of the Metaheuristic Optimization. .

Problem id	Components $c$	Runnables $r$	Chains $g$	Units $n$
$c_6 g_{10} n_4$	6	60	10	4
$c_8 g_{20} n_6$	8	80	20	6
$c_{10}g_{20}n_8$	10	100	20	8
$c_{20}g_{30}n_{10}$	20	200	30	10
$c_{50}g_{40}n_{20}$	50	500	60	20
$c_{80}g_{60}n_{20}$	80	800	60	20

Table 8: Specifications of the Optimization Problems, Sampled from Table 5 and Table ??, and Used in Experiment 1 and 2.

software allocation safety-critical distributed automotive applications on a CAN network of heterogenous computing nodes. The problems are identified by handlers of type  $\langle c_i g_i n_i \rangle$  to improve readability, where the c, g, n variables indicate respectively the number of components, cause-effect chains and computing nodes. The  $c_6g_{10}n_4$  and  $c_8g_{20}n_6$  problems conform to Spec-I and denote a small (or light) optimization problem, the  $c_{10}g_{20}n_8$  problems is based on Spec-II and denote a medium size problem, the  $c_{20}g_{30}n_{10}$ ,  $c_{50}g_{40}n_{20}$  and  $c_{80}g_{60}n_{20}$ are based on Spec-III and denote large size problem. The optimization problems are executed each 30× using our ILP method proposed in [43] and the metaheuristic algorithms presented in Section 5. The optimization parameters such as the penality function coefficient and the meta-heuristic parameters control the metaheuristics optimization, and their settings are shown in Table 7. The settings are obtained from literature as best practices of using the algorithms, as well as from our experimentaion of the algoriths with the problems at hand. Subsequently, we recorded the computation time, fitness values, power-consumption delivered by each algorithm.

Table 9 shows a summary of the evaluation results from executing Experiment 1 such as the average and standard deviation of the computation times and fitness values, as well as the quality of solutions. The latter is determined by comparing the power-consumption outcomes delivered from each algorithms agains the optimal or best solutions found (or benchmarks), which are indicated by the **boldface** type. It simply indicates how optimal or good the solution is as compared to the benchmark. In the first three optimization problems, the ILP is the benchmark since it returned optimal solutions. Similarly, the SHPSO is the benchmark in the problems  $c_{20}g_{30}n_{10}$  and  $c_{50}g_{40}n_{20}$ , and SHPSO is the

benchmark in the last probem  $c_{80}g_{60}n_{20}$ .

Experiment 2. Usually the replication exerts heavy computation over the calculation of the cause-effect delays due its combinatorial nature. The approximation technique, which is presented in Subsection ??, optimizes the calculation of the cause-effec chain delays in the presence of replication. We executed the optimization problems  $c_{50}g_{40}n_{20}$  and  $c_{80}g_{60}n_{20}$  with 2 and 3 degrees of replication, and also with and without the approximation technique applied according to the specification in Table 10. The degree of replication indicates the multiplicity of each component in the software applications.

### 6.2. Analysis

In this subsection, we analyze the results from experiement 1 and 2, respectively.

#### 6.2.1. Analysis of Experiment 1

We analyze the results over three metrices: computation and convergence time, solution quality and stability, respectively.

Solution Quality. In the  $1^{st}$  sample  $c_6g_{10}n_4$ , the ILP, DE, LPSO, HCPSO, SHPSO returned the optimal power consumption, which is 227KW, but DEPSO and PSO returned near optimal solutions with > 99% quality measures. In the  $2^{nd}$  sample  $c_8g_{20}n_6$ , similar results are obtained from ILP, HCPSO and SHPSO, which are optimal, but this time, in contrast, DE, LPSO and DEPSO performed worse by less than 1% but better than PSO by 2%. In the  $3^{rd}$  sample  $c_{12}g_{20}n_8$ , only ILP returned the optimal solution, followed by DEPSO, LPSO, HCPSO, SHPSO with near optimal solutions with > 99% quality measures, and rest performed worse. In the last three samples  $c_{20}g_{30}n_{10}$ ,  $c_{50}g_{40}n_{20}$  and  $c_{80}g_{60}n_{20}$ , ILP did not return solutions due to extremly large computation time, hence terminated manually. However, the hybrid algorithms based on hill-climbing such as HCPSO and SHCPSO performed well, followed by DEPSO in the samples  $c_{20}g_{30}n_{10}$  and  $c_{80}g_{60}n_{20}$ . However, HCPSO failed to returned solutions in the largest sample  $c_{80}g_{60}n_{20}$  but its stochastic verion SHCPSO did.

Convergence Time. In the case of metaheuristics, the convergence time refers to the amount of time taken by the algorithm to return solutions before the steady state where new fitness values are observed. In this evalution, it is calculated over a maximum of 5000 iterations (or generations) only for the duration before steady period, which is bounded by 5 minutes. Note: the steady time, where no fitness values change within the maximum iterations, is not considered in the converge time. Figure 11 summarizes the computation time of the algorithms for the samples listed in Table 9. For the samples the ILP method returned solutions, the computation times are usually larger than the rest, which are in milliseconds for the  $1^{st}$  sample and in seconds for the  $2^{st}$  and  $3^{rd}$ . The the meta-heuristic algorithms, the convergence time is in milliseconds for the first four samples, and is in seconds for the rest. However, the computation times

Sample	Algorithm	$\operatorname{Fitn}$	ess	Time	e (ms)	Quality
		Mean	SD	Mean	SD	
$c_6g_{10}n_4$	ILP	227.88	0	309	57.74	100.00
	PSO	229.11	2.38	0.12	0.34	99.46
	DE	227.88	0	0.01	0	100.00
	DEPSO	228.07	0.31	0.09	0.01	99.92
	LPSO	227.88	0	0.02	0.02	100.00
	HCPSO	227.88	0	0.03	0	100.00
	SHPSO	227.88	0	0.13	0.03	100.00
$c_8 g_{20} n_6$	$\mathbf{ILP}$	406.6	0	4148.3	95.77	100.00
	PSO	415.15	12.4	0.07	0.15	97.94
	DE	407.42	1.05	0.03	0.02	99.80
	DEPSO	409.65	8.8	0.17	0.01	99.26
	LPSO	407.18	0.53	0.32	0.73	99.86
	HCPSO	406.6	0	0.13	0.06	100.00
	SHPSO	406.6	0	0.29	0.14	100.00
$c_{10}g_{20}n_8$	ILP	442.37	0	14049.1	150.84	100.00
Ü	PSO	448.79	12.61	0.79	1.37	98.57
	DE	451.55	17.72	0.23	0.41	97.97
	DEPSO	442.44	0.19	1021.46	2263.76	99.98
	LPSO	442.49	0.17	1062.51	2338.73	99.97
	HCPSO	442.67	0.21	7.57	22.68	99.93
	SHPSO	442.46	0.19	10.73	61.31	99.98
$c_{20}g_{30}n_{10}$	ILP	NA	NA	NA	NA	NA
	PSO	64595.28	9544.82	11.27	9.73	65.74
	DE	53655.73	4134.84	22.15	7.95	79.14
	DEPSO	44055.97	4237.81	192.95	230.83	96.38
	LPSO	58603.42	6617.49	19.83	6.98	72.46
	HCPSO	42462.38	1643.71	247.05	104.36	100.00
	SHPSO	42558.2	2770.52	114.52	102.41	99.77
$c_{50}g_{60}n_{20}$	ILP	NA	NA	NA	NA	NA
	PSO	1298680.85	38557.68	1753.43	776.16	98.26
	DE	1460553.62	34599.66	571.43	248.46	87.37
	DEPSO	1384474.66	32550.41	4925.97	4809.57	92.17
	LPSO	1430847.88	32045.32	640.86	320.33	89.18
	HCPSO	1276036.05	65320.02	17445.87	15796.87	100.00
	SHPSO	1336679.78	98051.36	1074.4	339.83	95.46
$c_{80}g_{60}n_{20}$	ILP	NA	NA	NA	NA	NA
23300 20	PSO	2692638.14	46015.42	324.95	103.66	91.60
	DE	2737416.39	23780.06	716.97	207.19	90.10
	DEPSO	2604249.6	46945.89	4018.55	12.37	94.71
	LPSO	2650992.23	35813.35	1005.74	375.25	93.04
	HCPSO	NA	NA	NA	NA	NA
		2466535.41	89380.36	2147.79	357.58	100.00

Table 9: Fitness and Allocation Time of the ILP and the Metaheuristic Techniques, for the Increasing Sizes of the Software Allocation Problem.

Identifier	Chain $g$	Replication $d$	Problem id.
$g_{30}d_{2}$	30	2	$c_{50}g_{40}n_{20}$
$g_{30}d_{2}$	30	3	$c_{50}g_{40}n_{20}$
$g_{30}d_{2}$	60	2	$c_{80}g_{60}n_{20}$
$g_{30}d_{2}$	60	3	$c_{80}g_{60}n_{20}$

Table 10: Specifications of Samples, tobe Used in Experiment 2.

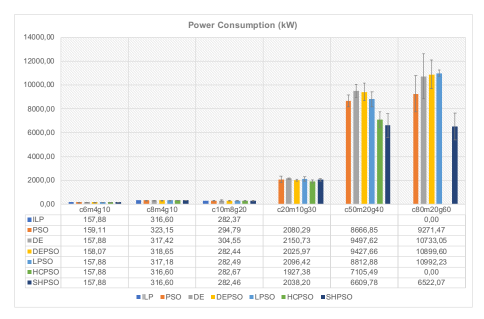


Figure 10: (Near) Optimal Power Consumption of the Different Software Allocation Problems.

of the meta-heuristic algorithms, which are not shown in the table usually took less than 50 minutes for the larest sample.

Solutions Stability. the PSO, DE are characterized by random search which enables exploration of higher dimenstional problems possible. However, somestime this creates instability in the solutions, that is for the same problem, it is possible to observe different performance, e.g., fitness values, computation time, etc. The stability of the solutions depend on the nature of the algorithms as well as the problems at hand. Therefore, it is crucial to evaluate the stability of the meta-heuristic algorithms used in this work. One way measuring the stability is using standar deviation, and Figure x and Figure y show the deviation of each algorithms for the (near) optimal power consumption of the different samples.

In general, with regard to quality of the solutions, the hybrid PSO with hill-climbing are more stable in the first three samples, but also DE and LPSO in  $1^{st}$  sample, as compared to PSO and DEPSO. However, as the problem size increases to  $1^{st}$ ,  $2^{nd}$ ,  $3^{rd}$ , the hybrid PSO with hill-climbing performed worse and PSO and others improved. With regard to convergence time, the

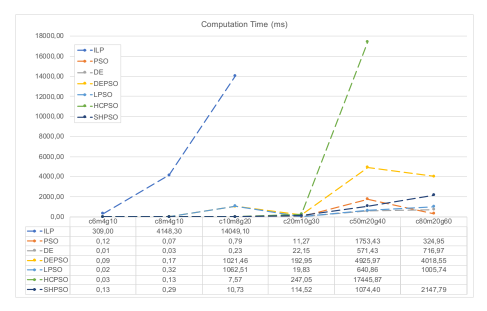


Figure 11: Computation Time of the Various Algorithms for Solving Different Instances of the Software Allocation Problem.

stability usually decreased, that is with uniformity for the PSO, DE, HCPSO and SHPSO, however, for the rest it is not uniform.

### 6.2.2. Analysis of Experiment 2

Table 12 shows results of executing experiment 2, which shows improvements of the computation time by applying the approximation algorithm in stead of the exact approach. In the case of the approximation, the delays are exaustively calculated in the presence of replication. However, the quality of the solutions are degraded as expected due to the approximation. Specifically, the result shown 61%-81% computation time improvement over the exac method while facing quality degradation only for samples  $g_{30}d_2$  and  $g_{30}d_2$ . The improvements are in seconds, which implies for a single usage (or run) of the meta-heuristic optimization algorithms, it is not significant. However, considering practical systems design process, which requires several iterations, the commulative effect of the algorithms can negatively impact the responsivess to engineers. Thus, the improvements can be in trade-off with optimality of the solutions.

#### 7. Discussion

#### 8. Related Work

In a heterogeneous distributed system where computing nodes and communications links could have various failure rates, a reliability-aware allocation of tasks to nodes, and using links with the lowest failure rates can noticeably improve the system reliability [44][45][35][46]. Interleaving real-time constraints into the

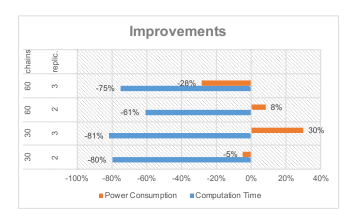


Figure 12: Effect of Approximate Algorithm over Delay Calculations with Replication.

problem adds more complexity to reliability-aware task allocation in distributed systems [47]. As opposed to [9][5], we assume that software applications are multirate, which increase the difficulty of software allocation due the complexity of their timing analysis, and increased search space as the result of increasing timed paths of cause-effect chains. Furthermore, we assume a fault-tolerant system model.

Although improving reliability of the system using a reliability-aware task allocation does not impose extra hardware/software cost, in reliability-based design approach, redundancy (or replication) of software or hardware components is frequently applied to improve reliability. In such systems not only optimal allocation of software components (or replicas) should be taken into account but also the cardinality of the replicas should be limited for improved efficiency while meeting the desired reliability requirement. The integration of these two approaches (i.e., reliability-aware task allocation and application redundancy) is a promising technique to deal with high criticality of the system to fulfill the required reliability. For example, [48] proposes a heuristic algorithm to maximize reliability of a distributed system using task replication while at the same time minimizing the makespan of the given task set. Furthermore, in systems with replication, it uses the Minimal Cut Sets method, which is an approximate algorithm, to calculate reliability of a system. In contrast, we apply an exact method based on state enumeration, which is applicable to the problem size assumed in this work.

In our problem, power consumption is the other criterion of the optimization problem. Several research work exist on improving power consumption in real-time distributed systems. The research work [49] shows a survey of different methods on energy-aware scheduling of real-time systems, which categorizes the study into two major groups: i) Dynamic Voltage Scaling (DVS) [50][51], and ii) task consolidation to minimize the number of used computing and communication units [52], which is the approach followed in our work.

In the context of automotive systems, there are few works considering the

reliability of a distributed system subject to real-time requirements of the automotive applications [53][54]. There are also other works discussing the allocation of software components onto nodes of a distributed real-time systems that consider other types of constraints other than reliability, for example, i) [55] which considers computation, communication and memory resources, and ii) [15] which proposes a genetic algorithm for a multi-criteria allocation of software components onto heterogeneous nodes including CPUs, GPUs, and FPGAs. Our approach also considers a hetrogeneous platform, i.e., nodes with different power consumption, failure-rate, and processor speed. In this work, we consider only the processor time; however, it can easily be extended to take into account different types of memory consumption that the software applications require.

#### 9. Conclusions and Future Work

Software to hardware allocation plays an important role in the development of distributed and safety-critical embedded systems. Effective software allocation ensures that high-level software requirements such as timing and reliability are satisfied, and design and hardware constraints are met after allocation. In fault-tolerant multirate systems, finding an optimal allocation of a distributed software application is challenging, mainly due to the complexity of cause-effect chains' timing analysis, as well as the calculation of software application reliability. The timing analysis is complex due to oversampling and undersampling effects, caused by the different sampling rates, and the complexity of the reliability calculation is caused by the interdependency of the computation nodes due to replicas. Consequently, the formulation of the problem, to find an optimal solution, becomes non trivial.

In this work, we propose an ILP model of the software allocation problem for fault-tolerant multirate systems. The objective function of the optimization problem is minimization of power consumption with the aim of satisfying timing and reliability requirements, and meeting design and hardware constraints. The optimization problem involves linearization of the reliability model with piecewise functions, formulating the timing model using logical constraints, and limiting the number of replicas that can be used in the allocation. Furthermore, the allocation consider two cases of timing analysis: response time analysis and utilization bound.

Our approach is evaluated on synthetic automotive applications that are developed using the AUTOSAR standard, based on a real-world automotive benchmark. Although we consider automotive applications for the evaluation, the proposed approach is equally applicable to resource-constrained embedded systems, especially with timing, power and reliability requirements, in any other domain that are developed using the principles of model-based development and component-based software development. Our approach effectively applies to medium-sized automotive applications, but does not scale for complex applications. Considering similar system models, we plan to extend the current work with heuristic methods, e.g., genetic algorithms, simulated annealing, particle swarm optimization, etc., to handle large systems.

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