

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 Y-chart 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 widely-used 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 BitBucket¹.

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 4.5 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,

¹<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 is composed of three parts: software applications, an execution platform, and an allocation scheme. In this section, we show models of the different parts and describe them in detail. For the sake of clarity, first we list the main mathematical notations that are used to define the system model and the software allocation problem.

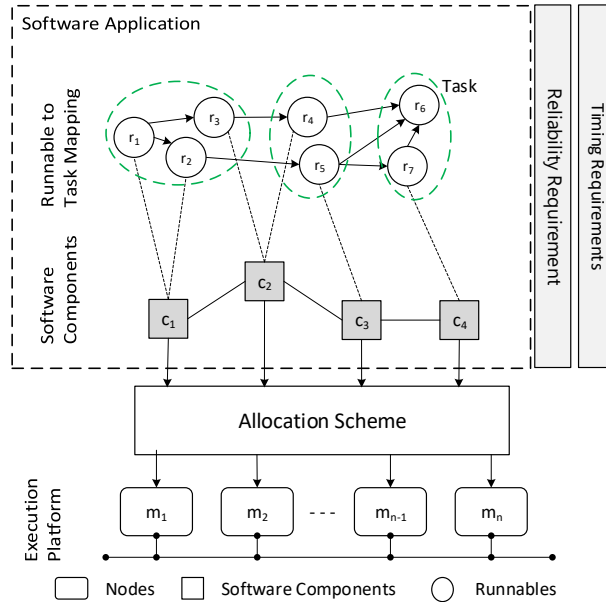


Figure 1: System Model.

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 [18][19]. A software application A_k is associated with user-defined requirements represented by the tuple $(\bigcup_i e2ereq_i, relreq, cl) : \mathbb{R}^+ \times [0, 1] \times \mathbb{I}^+$, where the elements of the tuple denote the end-to-end timing requirements, reliability requirements, and requirements on the criticality level of the application respectively. The

| Notation | Description |
|---|---|
| /* Application related */ | |
| • $A = \{A_k : k = 1, \dots, n_A\}$ | software applications* |
| • $C^{(k)} = \{c_i^{(k)} : i = 1, \dots, n_C\}$ | software component types of A_k ** |
| • $Q_i^{(k)} = \{q_{i,j}^{(k)} : j = 1, \dots, n_{Q_i}\}$ | component replicas of type $c_i^{(k)}$ |
| • $R_i^{(k)} = \{r_{i,j}^{(k)} : j = 1, \dots, n_{R_i}\}$ | runnables that implement $c_i^{(k)}$ |
| • $H_{i,j}^{(k)} = \{r_{i,j,h}^{(k)} : h = 1, \dots, n_{H_{i,j}}\}$ | runnables that implement $q_{i,j}^{(k)}$ |
| • $g_r^{(k)}$ | directed acyclic graph of $\bigcup Q_i^{(k)}$ |
| • $V(g)$ | node/vertices of graph g |
| • $E(g)$ | links/edges of graph g |
| /* 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 |
| • $\Gamma^{(k)} = \{\Gamma_i^{(k)} : i = 1, \dots, n_\Gamma\}$ | end-to-end chains |
| • $\Gamma_i^{(k)} = (e_i)_{i=1}^Z$ | a chain of tasks or messages e |
| • $\pi_i(\Gamma_i^{(k)})$ | the i^{th} element in $\Gamma_i^{(k)}$ |
| • τ, c, m, γ denote iterator variables, respectively for task, component, chain and node, e.g., $\forall \tau \in T^{(k)}$.*** | |
| /* Mapping related */ | |
| • $\mathbf{x}^{(k)} = \{\mathbf{x}_k^{(k)} : k = 1, \dots, n_{\mathbf{x}}\} : \bigcup Q_i^{(k)} \mapsto M$ | a mapping vector from $Q^{(k)}$ to M |
| • k, i, j denote iterator index-variables, respectively for the mapping vector \mathbf{x} , and rows and columns of the matrix $\mathbf{x}^{(k)}$, e.g., $x_{ij}^{(k)}$.*** | |
| • $g_\tau^{(k)}(\mathbf{x})$ | directed acyclic graph of tasks |
| /* Functions related */ | |
| • $Power(\mathbf{x})$ | 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 $\tau \in V(g_\tau^{(k)})(\mathbf{x})$ |
| • $Delay_\gamma(\mathbf{x})$ | age delay of $\gamma \in \Gamma^{(k)}$ in \mathbf{x} |

*Note: the total elements in a set S is denoted by n_S , e.g., n_A denotes the number of applications in the set S , essentially it refers to its cardinality.

** For readability, we prefer to use $S_i^{(k)}$ in place of $S_i^{(A_k)}$.

*** If for other uses of the iterators, they are defined in the context.

elements in the requirements tuple of A_k are accessed through the projection symbol π as $\pi_{e2ereq_i}(A_k)$, e.g., $\pi_{e2ereq_1}(A_2)$ means the first end-to-end timing requirement of application A_2 . The end-to-end timing requirements define 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 defines the expected reliability goal of an application which is discussed further in Subsection 4.1. Finally, 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 [?].

The software applications are run in parallel and therefore can potentially be distributed on different computation nodes $M = \{m_i : i = 1, \dots, n_M\}$, and we assume the nodes are heterogeneous with respect to processor speed, failure-rate and power consumption as indicated by the tuple (hz, λ, p) , respectively.

Definition 1 (Software Application Model). A software application 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

task and node need to be clarified...”graph of nodes, where each represents a periodic task”

V_τ , where $a_{ij} \in L_\tau$ refers to the data-flow link from node τ_i to node τ_j and $i \neq j$. The computation cost $w(\tau) = \langle \bigcup e_{m_i}, D, P, m \rangle$ refers to the worst-case execution time WCET of the task on node M , deadline and period, and mapping of the task to the node, where $m \in M$.

The last sentence also needs clarification...mapping of what to what?

Multiple applications can be executed on the same computation node(s) and can share the on-board network, e.g., the CAN bus. Since the applications can have different criticality requirements, the execution platforms should provide a 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 [20], 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-critical

this needs to be changed to mixed-criticality throughout the paper

applications on *uniprocessor* systems [20]. In our problem, though distributed applications, each task is mapped to a single node

confusion of the node with the node in the acyclic graph. Needs clarity

, and the mapping is static. In this case, the schedulability of tasks can be performed per node, that is using the approach applied to uniprocessor nodes. Therefore, in the context of this work, the distributed applications are schedulable if the tasks, messages, and the cause-effect chains distributed over multiple nodes in the system meet their respective timing requirements in the midst of power consumption and reliability constraints.

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 [21, 22, 23], though less efficient. It is interesting to note that 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 policy* (FPPS) [?]. Initially, applications are prioritized based on their criticality levels following the PA technique, and within each application the tasks are prioritized with the *deadline monotonic* (DM) priorities assignment.

$$cri(A_h) > cri(A_l) \implies \forall \tau_1 \in \bigcup T_i^{(h)} \tau_2 \in \bigcup T_i^{(l)} Pri(\tau_1) > Pri(\tau_2)$$

where cri, pri are predicates which determine the criticality and priority of tasks τ_1, τ_2 , respectively; $\bigcup T_i^{(h)}, \bigcup T_i^{(l)}$ are the set of tasks which implement the applications A_i, A_j , respectively.

The schedulability of tasks assigned to a node is performed using the classical response-time analysis shown in Equation (1) [24, 24], which computes the worst-case response time of each task, denoted by R_τ . 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.

$$R_\tau = c_\tau + \sum_{\gamma \in HP(\tau)} \left\lceil \frac{R_\tau}{P_{\tau_{hi}}} * c_\gamma \right\rceil, \quad (1)$$

where $\gamma \in HP(\tau)$ is element of the higher-priority tasks returned by HP .

In this work, we assume heterogeneous computation nodes, therefore the schedule that delivers lower power-consumption of a node is considered the effective and efficient. The power-consumption of a node is computed linearly

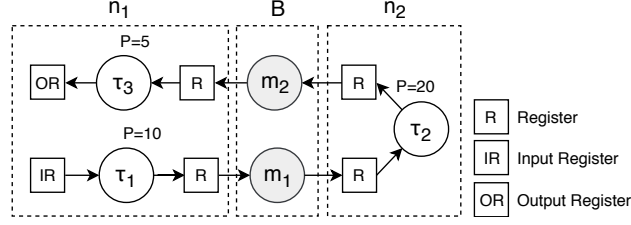


Figure 2: A Cause-effect Chain, mapped on nodes n_1 and n_2 .

from the utilization of a taskset mapped to a specific node as well as from its power-specification parameters, and is discussed in detail in Subsection 4.

Unlike the tasks, the messages in the CAN bus B are scheduled using a non-preemptive and fixed scheduling policy. Similar to the mixed-criticality of tasks, the messages in the can should be separated as well for different criticality applications, which is achieved by applying the PA technique. In this case though, the priorities of messages are inherited from the send tasks, $pri(m) = pri(\tau) | \tau = pre(m)$, where $pre(\tau)$ is a predicate that computes the predecessor of task τ from the tasks graph g_τ . The schedulability of messages is checked using the classical response-time analysis of messages in CAN network [?], as shown in Equation ()

Better to use the equation from the analysis by Rob Davis et. al [?]”

$$R_\tau = c_\tau + \sum_{\gamma \in HP(\tau)} \left\lceil \frac{R_\tau}{P_{\tau_{hi}}} * c_\gamma \right\rceil, \quad (2)$$

3.2.2. Scheduling Cause-effect Chains

The software application can be considered as a set of *cause-effect chains* $\Gamma^{(k)} = \{\Gamma_i^{(k)} : i = 1, \dots, n_\Gamma\}$, which are directed paths in the graph, annotated by end-to-end timing requirements $End2end_i^{(k)}$. 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 τ_1, τ_2, τ_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 End-to-End Timing Analysis of Cause-Effect Chains in Automotive Embedded Systems

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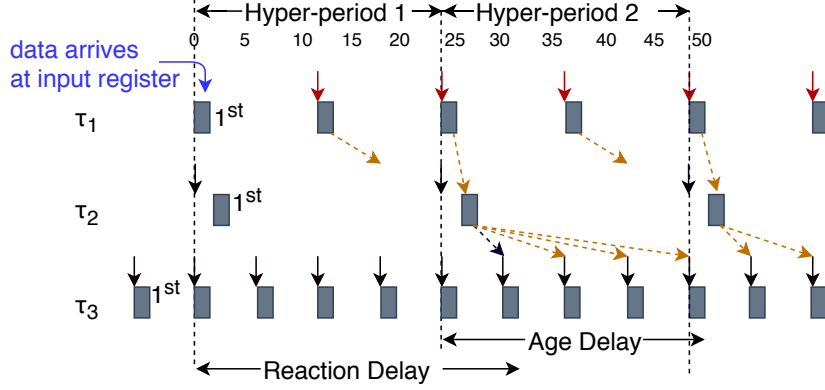


Figure 3: Reaction and Age Delays of the Cause-effect Chain, Shown in Figure 2.

The end-to-end delay in a chain is the duration between the reading of data from the input register by the source task $Source(\Gamma_i^{(k)})$ to the writing of same data to the output register by the last (or sink) task $Sink(\Gamma_i^{(k)})$. 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 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 τ_1 and τ_2 execute on node n_1 , whereas task τ_3 executes on node n_2 . Note: τ_2 communicates with τ_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 3rd instance of τ_1 and the 10th instance of τ_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 1st instance of τ_1 execution. The data corresponding to this event is not read by the current instance of τ_1 . In fact, the data will be read by the 2nd instance of τ_1 . The earliest effect of this data at the output of the chain will appear at the 7th instance of τ_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 analytically calculated using Equation 3, and is explained as

follows.

$$AgeDelay(\Gamma) = \begin{cases} \alpha(Sink(\Gamma)) - \alpha(Source(\Gamma)) + \delta(Sink(\Gamma)) \\ \text{(if the chain is mapped on a single node)} \\ \sum_{a \in Part(\Gamma)} AgeDelay(\Gamma) + \sum_{m \in M'} \delta(m) \\ \text{(if the chain mapped to multiple nodes)} \end{cases} \quad (3)$$

Assume $\Gamma \in \Gamma_i^{(k)}$ is a chain, if the chain is mapped on a single node, the age delay is a mere difference between the activation of the sink task $\alpha(Sink(\Gamma))$ and the activation of the source task $\alpha(Source(\Gamma))$ plus the worst-case response time of the sink task $\delta(Sink(\Gamma))$ 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 delay can be compositionally computed [?] as follows: the chain is partitioned into a set of sub-chains per node, indicated by the predicate $Part(\Gamma)$ and for each partitioned chain $a \in Part(\Gamma)$, the age delay is computed recursively using the method to compute the age delay in a single node, and the result is added to the response-times of the messages involved in the chain M' .

3.3. AUTOSAR System

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 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 specifications. Therefore, the software application model 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 $g_r = \langle V_r, L_r, w, v \rangle$ 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 at the node refers to the timing model of the runnable, where the tuple elements, respectively denote execution time on node m , deadline and period. Example: Figure 4(a).

According to AUTOSAR specification [26], 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. Therefore, through the mappings, eventually runnables graphs are refined by tasks graphs as shown in Figure4 (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 $v \in V(g_r)$, if the

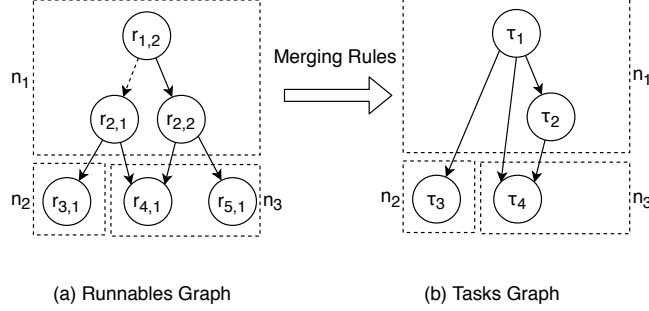


Figure 4: Example of a Software Application, Modeled as Directed Acyclic Graph, where $--\rightarrow$, $--\rightarrow$ denote triggering link, data-flow link, respectively.

| $r_{i,j}^{(k)}$ | 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) | |

Table 1: Runnables Timing Specifications.

| $\tau_i^{(k)}$ | $\bigcup r_{i,j}^{(k)}$ | (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.

following rules satisfy': (i) the runnables are co-hosted in the same computation node, i.e., $a \mapsto m \wedge b \mapsto m$ (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 on the tasks.

3.4. Execution Platform

The execution platform provides computation and communication resources to the user applications, and is modeled as a *complete* graph $\langle M, L^m \rangle$ of computation nodes, where $(m_i, m_j) \in L^m \wedge i \neq j$ refer to the communication links of the nodes, which are realized by a network bus, e.g., CAN. The computation nodes are heterogeneous with respect to parameters defined as a tuple $\langle hz, \lambda, p \rangle$, respectively denote processor speed, failure-rate and power consumption specifications. The allocation scheme is a mapping table $f : C \mapsto M$ from software components to computation nodes, where $C = \bigcup_{i=1}^{|A|} V(A_i)$ is the *infinitary* union of the user applications' vertices, which denote nodes o software components.

3.5. Fault-tolerant Software Application Model

Redundancy is the most common way to increase the reliability of an application. It can be implemented according to different schemes, such as hot stand-by, cold stand-by, etc [27]. 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.

We denote the k^{th} replica of a software component c as c^k , with $1 \leq k \leq K$; where K is the maximum number of replicas allowed for each application component.

4. Software Allocation Problem

The software allocation problem is a type of job shop scheduling with constraints, as such it is a discrete optimization problem []. The solution to the allocation problem is represented by a vectormatrix $\mathbf{x} = \{\mathbf{x}^{(k)} : k = 1, \dots, N_a\}$, where $\mathbf{x}^{(k)}$ is a matrix of size $N_c \times K$, and $x_{ij}^{(k)} = k \in \{1, \dots, N_m\}$ represents the mapping of the software component replica $c_{ij}^{(k)}$ to the computation node m_k .

$$\mathbf{x}^{(k)} = \begin{bmatrix} x_{11}^{(k)} & x_{12}^{(k)} & \dots & x_{1K}^{(k)} \\ x_{21}^{(k)} & x_{22}^{(k)} & \dots & x_{2K}^{(k)} \\ \vdots & \vdots & \ddots & \vdots \\ x_{N_c 1}^{(k)} & x_{N_c 2}^{(k)} & \dots & x_{N_c K}^{(k)} \end{bmatrix} \quad (4)$$

In this work, the main objective of the allocation problem is to satisfy the user-defined requirements, namely reliability requirements, end-to-end timing requirements, and criticality of the software applications A_i by effectively mapping the software components to the computation nodes, $C^{(k)} \mapsto M$. Furthermore, the components are allocated efficiently to minimize the total power consumption $Power(\mathbf{x}) = \sum_{m \in M'} P_m(\mathbf{x})$ of the applications by selecting lower-power consuming nodes $M' \subseteq M$, provided the requirements are met, where $P_m(\mathbf{x})$ is the power consumption of node m on the mapping \mathbf{x} . Power consumption, in this context, refers to the energy usage of electronic components in the integrated circuits of the node, e.g., processor, memory, I/O devices, etc., per time unit.

There are several power consumption models and different techniques to estimate the power consumption of a computing node. In this work, we employ a technique based on processor load (or *Processor Utilization*) to estimate the

average power consumption of a computation node. Specifically, we use the linear polynomial model proposed by Fan et al. [28], which is shown in (5). The model states that the power consumption of a node is directly proportional to its load, and is inductively formulated from experimental results:

$$f_p(u) = P_{idle} + (P_{busy} - P_{idle}) * u, \quad (5)$$

where u is the utilization (or load) of a computation node, p_{idle} and p_{busy} , respectively, refer to the power consumption of a node measured at minimum and maximum processor loads. Such measurements can be obtained by running performance benchmark suits, e.g., MiBench [29], AutoBench [30], etc., which is computed based on the utilization of the node via the linear power consumption model shown in Equation (5).

Consequently, the power consumption of a node m for a given mapping \mathbf{x} is computed using Equations (6-8), by calculating first the node's utilization $U_m(\mathbf{x})$ using Equation (7). The node's utilization is computed from the set of components allocated to it (which are $\forall_{ij} x_{ij} = m$) using Equation (7). And the utilization of a component on the node m is computed from its constituent tasks T_c using Equation (8).

$$nodPow(\mathbf{x}) = f_p(nodUtil(\mathbf{x}, m)) \quad (6)$$

$$nodUtil(\mathbf{x}, m) = \sum_k \sum_i \sum_j comUtil(m, c) | x_{ij}^{(k)} = m \quad , \text{ where } c = C_i^{(k)} \quad (7)$$

$$comUtil(m, c) = \sum_{r \in c.R} \frac{r.e_m}{r.\tau.P}, \quad (8)$$

where $c.R$ is set of runnables in component c , $r.e_m$ and $r.\tau.P$ are the execution of runnable r on node m and its period, respectively.

The applications requirements are modeled as constraints that need to be satisfied in the allocation problem. The constraints formulations are shown in the following subsections, respectively for reliability, timing and other design constraints such as related to runnables-to-tasks merging and replication.

4.1. Software Application Reliability Constraints

The applications reliability constraints ensure the mapping \mathbf{x} satisfies the user-defined reliability requirements, that is $\forall k \in [1, n_A] \text{ Reliability}_{A_k}(\mathbf{x}^{(k)}) \leq RelReq_{A_k}$. The reliability is computed from the execution framework that is provided to run the application, which consists of computation nodes $M^{(k)}$ and the shared CAN bus B, as shown in Equation (9). The nodes $M^{(k)}$ host the components $C^{(k)}$ and are determined by searching the mapping $\mathbf{x}^{(k)}$ in polynomial time using Equation (10).

$$Reliability_{A_i}(\mathbf{x}) = Reliability_{A_i}(M^{(k)}) * Reliability(B) \quad (9)$$

$$M^{(k)} = \{e | e \in M \wedge \forall ij (e = m_h)\}, \text{ where } h = x_{ij}^{(k)} \quad (10)$$

The reliability of the nodes, $Reliability_{A_i}(M^{(k)})$ with respect to the application A_k is calculated using the *state-enumeration* technique [31] as shown in Equations (11). According to the technique, the reliability is basically the total probability that the application A_k functions under mutually exclusive and uniformly distributed failure-events of the nodes, represented by $\mathcal{F}^{(k)}(M^{(k)}) = \{\mathbf{0}, \mathbf{1}\}^{M^{(k)}}$, where $\mathbf{0}$ means node fails, $\mathbf{1}$ means node is operational. An event $\omega \in \mathcal{F}_{A_k}(M^{(k)})$ represents the states of nodes by a sequence of 0-1 variables (b_1, \dots, b_n) , where $\omega.b_i$ denote the state of node $m_i \in M^{(k)}$.

$$Reliability_{A_i}(M^{(k)}) = \sum_{s \in \mathcal{F}^{(k)} | g(A_i, s) = 1} probability(s) \quad (11)$$

The fact that an application functions $g(A_k, s)$ is defined via its inverse, which is *software application failure*, deductively as follows:

Definition 3 (Software Application Failure). The application A_k fails in the event $\omega \in \mathcal{F}^{(k)}(M^{(k)})$ if there exists a component type $c_i^{(k)}$ where all of its replica $Q_i^{(k)}$ fail, otherwise, the application functions, as shown in Equation (12). A component replica $q_{ij}^{(k)} \in Q_i^{(k)}$ fails if the node that host it m_h fails, where $h = x_{ij}^{(k)}$.

$$g(a, \omega) = \begin{cases} \mathbf{0} & \text{if } \exists i \forall j (m_h \in M_\omega^{(k)}) \\ \mathbf{1} & \text{otherwise} \end{cases} \quad (12)$$

$$M_\omega^{(A_k)} = \{e | e \in M^{(k)} \wedge (\omega.b_e = 1)\}, \quad (13)$$

where $M_\omega^{(k)}$ denote the nodes that function at the event ω , and can be found by searching the nodes $M^{(k)}$ with state $\omega.b_e = 1$ in constant time, as show in Equation (14).

The probability that a nodes-failure event ω occurs is computed as a product of the probabilities of its constituent elements $\omega.b$ as shown in Equation (??).

$$\prod_{(m,b) \in (M, \omega)} \lambda_m * (1 - b) + (1 - \lambda_m) * b \quad (14)$$

where λ_m is the failure-rate of node m .

4.2. Timing constraints

The timing constraints ensure that the applications are schedulable on the execution platform, that is the tasks and the cause-effect chains in each application meet their timing requirements. The schedulability of each task is checked using the worst-case response-time analysis presented in Subsection 3.2.1, and for the cause-effect chains using the end-to-end data-propagation delay analysis shown in Subsection 3.2.2. Before we formulate the timing constraints, first we derive the task graph for a mapping \mathbf{x} from the runnables graph $g_r^{(k)}$, that is

using Equation (16). Using Equation (15), we update the runnables with nodes information, to which they are mapped by traversing elements of the mapping \mathbf{x} , $x_{ij}^{(k)}$, in linear-time complexity $O(n)$, where $n = |\mathbf{x}| * N_r$, and N_r is the total number of runnables in the system. Consequently, we traverse the runnables graphs, and apply the merging rules stated in the Subsection 3.3 to derive the tasks graphs. In the case that only runnables from the same component are mapped to at least one task, the derivation of tasks graphs is independent of the mapping \mathbf{x} , and thus can be performed before the mapping activity. However, the tasks graphs have to be updated with the nodes information after the mapping \mathbf{x} is identified.

$$\forall k, ij \forall r \in H_{i,j}^{(k)} r.node = n_h, \text{ where } h = x_{ij}^{(k)} \quad (15)$$

$$\forall k \ g_r^{(k)}(\mathbf{x}) \xrightarrow{\text{Eqn. (13); Merging Rules}} g_\tau^{(k)}(\mathbf{x}) \quad (16)$$

where $H_{i,j}^{(k)}$ is the set of runnables that implement component $q_{i,j}^{(k)}$.

4.3. Tasks Timing constraints

The tasks timing constraints states that the worst-case response time of each task in the system meets its respective deadlines for a mapping \mathbf{x} , that is $\forall k \forall \tau \in V(g_\tau(\mathbf{x})) \text{ ResponseTime}(\tau) \leq \text{Deadline}(\tau)$, where $V(g_\tau(\mathbf{x}))^{(k)}$ is the nodes in the tasks graphs. To compute the worst-case response time of the tasks, first we partition the tasks per node, that is tasks mapped to the same node grouped, represented by the T_{n_h} , by traversing the tasks graphs using Equation (17). The complexity of this equation, considering an adjacency matrix tasks graphs representation is linear-time $O(N_a * n^{(k)})$, where $n^{(k)}$ is the sum of the order and the size of graph $g_\tau^{(k)}(\mathbf{x})$.

$$T_{m_h} = \{e \in V(g_\tau^{(k)}(\mathbf{x})) | e.node == m_h, \} \quad \text{where } h = x_{ij}^{(k)} \text{ for all } h = 1, \dots, n_N \quad (17)$$

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 (18).

$$\forall \tau \in T_m \text{ ResponseTime}((\mathbf{x}, \tau) \leq \text{Deadline}(\tau) \quad (18)$$

4.4. Cause-effect Chains Timing constraints

For a mapping \mathbf{x} , the age delays of cause-effect chains should meet their respective end-to-end requirements, that is $\forall k \forall ij \text{ AgeDelay}_\Gamma(\mathbf{x}) \leq E2eReq_\Gamma^{(k)}$, where $\Gamma \in \Gamma_i^{(k)}$. To calculate the age delays, first we identify the messages scheduled by the CAN bus for the mapping \mathbf{x} using Equation (19).

$$M = \{e | \forall (a, b) \in g_\tau(\mathbf{x}) \forall n \in N (a \mapsto n \wedge b \mapsto n = \text{false}) \implies \text{createMsg}(e)\}, \quad (19)$$

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 $\Gamma_i^{(k)} = \{\tau_1, e^*\}$, where τ_1, τ_2 are $Source(\Gamma_i^{(k)})$ and $Sink(\Gamma_i^{(k)})$, respectively, and $e \in V(g_\tau(\mathbf{x})) \cup M$. Then, the cause-effect timing constraints are formulated over the updated list of chains $\Gamma_i^{(k)}$ using Equation (20).

$$\forall \Gamma \in \Gamma_i^{(k)} \quad AgeDelay(\mathbf{x}, \Gamma) \leq E2eReq(\Gamma) \quad (20)$$

4.5. Software Allocation Optimization Problem

In this subsection, we define the allocation problem, that is the mapping of software components to heterogeneous computation nodes, which are networked over a shared CAN bus, with constraints. The allocation problem objective is to minimize the total power-consumption of the distributed software applications as formulated in Equation 21. The inequality constraints shown by Equation (22, 23, 24), respectively ensure the feasible solution meet the reliability requirements of the applications, the deadlines of the tasks in the system and the end-to-end requirement of the cause-effect chains, and the last inequality constraint ensures that replicas are not allocated to the same computation nodes.

$$\min_{\mathbf{x} \in X} Power(\mathbf{x}) \quad \text{subjected to:} \quad (21)$$

$$Reliability_{A_k}(\mathbf{x}) \leq RelReq_{A_k} \quad \text{for all } k = 1, \dots, n_{A_k} \quad (22)$$

$$\forall \tau \in T_{m_h} \quad ResponseTime_\tau(\mathbf{x}) \leq Deadline_\tau \quad \text{for all } h = 1, \dots, n_M \quad (23)$$

$$\forall \gamma \in \Gamma^{(k)} \quad Delay_\gamma(\mathbf{x}) \leq E2eReq_\gamma \quad \text{for all } k = 1, \dots, n_A \quad (24)$$

$$\forall k \forall j x_{ij}^{(k)} \neq x_{ij}^{(k')} \quad \text{where } k \neq k' = 1, \dots, n_A \quad (25)$$

$$(26)$$

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 $q_{i,j}^{(k)}$ to the node m_h , where $h = x_{ij}^{(k)}$

In the next section, we propose a hybrid Particle Swarm Optimization (PSO) to solve the allocation problem.

5. Solution Using Hybrid Particle Swarm Optimization

Although the proposed ILP approach provides exact solutions, that is with an AUTOSAR software allocation with minimum power consumption, the approach does not scale well for large applications. Thus, in this section, we propose an approximation approach based on several metaheuristic techniques to address the scalability challenge. Metaheuristic techniques assumes little of the problem in question, and therefore are ideal to solve high dimensional and complex optimization problems, that is, problems that are difficult or practically impossible to solve by exact methods, e.g., linear programming, or heuristic techniques, e.g., local search algorithms. They have improved over the last

decades with respect to effectiveness, efficiency, and ease of use by providing fewer user-configurable parameters. Metaheuristic techniques do not guarantee optimal solutions, nevertheless, the returned solutions can be good enough (or acceptable) in the eyes of the system designer, for this particular problem, it means although the power consumption of the system is not optimal, the solution can be deemed acceptable, that is as long as the constraints are fulfilled. In the opposite case, where the constraints could not be fulfilled, the algorithms can be rerun several times until the desired results are found, or the design can be relaxed by weakening the timing and reliability constraints of the system.

Metaheuristic techniques perform differently for different problem types, size, and complexity. In this section, we show application of different metaheuristic techniques that employ swarm intelligence and evolutionary approaches in finding the (near) optimal solutions. Specifically, we apply the Particle Swarm Intelligent (PSO), Differential Evolution techniques primarily, and further hybrid PSO with DE and Hill-climbing for improved performance. PSO has been applied to solve a wide range of problems, including a task allocation problem [32], 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 metaheuristic methods based on solution quality and allocation (or computation) time for different software allocation problems.

5.1. Solution Representation

In contrast to the $0-1$ representation, the integer-linear representation uses much lower number of variables, that is $N * K(L - 1)$, e.g., for a software allocation problem with $N = 10, L = 8, K = 2$, 140 variables are saved. Of course the possible values in the former representation is two whereas in the latter representation, it is L , which is usually higher and results larger solution space. Nevertheless, the integer-linear representation is compact and computationally more efficient.

5.2. Fitness Function Definition

Since a metaheuristic method functions over the meta of a problem, the quality of candidate solutions is evaluated based on their fitness to meet the problem's objective. A solution that delivers lower power consumption and violates less constraints is indicated by a lower fitness value. The fitness function $f(\mathbf{x})$ combines the original objective function $P(\mathbf{x})$ and the constraints $Timing(\mathbf{x}), Reliability(\mathbf{x})$ in order to compute real-valued numbers that indicate quality of the candidate solutions.

Consequently, the original constrained optimization problem is transformed into unconstrained optimization problem, by extending the objective function $P(\mathbf{x})$ with the constraints, represented by a *penalty function* $\Phi(\mathbf{x})$. The function returns 0 if no constraints are violated otherwise returns a positive number,

essentially to penalize the candidate solution by increasing its fitness (for our minimization problem), thus discriminating the solution. The function is a combination of $\beta \sum g_i(\mathbf{x})$ and $\gamma h(\mathbf{x})$ functions, respectively computes the timing violations and a software application reliability violation, and each function is weighted to indicate the size of the penalty separately. Moreover, the penalty function $\Phi(\mathbf{x})$ is weighted to indicate the size of penalty that imposed on the combined violations of timing and reliability. **How to compute the parameters [Hamid]**

$$\min_{\mathbf{x} \in X} f(\mathbf{x}) = P(\mathbf{x}) + \alpha \Phi(\mathbf{x}) \quad (27)$$

$$\Phi(\mathbf{x}) = \sum \phi_i(\mathbf{x}) \quad (28)$$

5.3. Penalty Function

5.4. Metaheuristic Algorithms

5.4.1. 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 [33]. 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 real-world problems, e.g., diagnosis and classification of diseases, efficient engineering designs, tuning control design parameters, scheduling problems, etc [34].

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 [35]. 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 29. 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 (30).

$$\mathbf{v} \leftarrow \omega \mathbf{v} + c_1 \text{Rand}() \circ (\mathbf{y} - \mathbf{x}) + c_2 \text{Rand}() \circ (\mathbf{z} - \mathbf{x}) \quad (29)$$

$$\mathbf{x} \leftarrow \mathbf{x} + \mathbf{v} \quad (30)$$

where ω 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 [36] or integer-linear by approximation to the nearest integer values [37], 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 29 and 30, the solution is discretized by rounding off the its elements to the nearest integer values, that is $\mathbf{x} \leftarrow \lfloor \mathbf{x} \rfloor$.

5.4.2. 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 1995 [38], 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 [39]. 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 (32), and (33), respectively. A mutant vector v is created from randomly selected elements $\{a, b, c\} \in PN$ according the mutation operation shown in Equation (31), 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.

$$\mathbf{v} \leftarrow \mathbf{a} + F \circ (\mathbf{b} - \mathbf{c}) \quad (31)$$

$$u_{ik} \leftarrow \begin{cases} v_{ik} & \text{if } U(0,1) \leq CF \text{ and } h = (i * K + k) \\ x_{ik} & \text{if } U(0,1) > CF \text{ and } h \neq (i * K + j) \end{cases} \quad (32)$$

$$\mathbf{x} \leftarrow \begin{cases} \mathbf{u} & \text{if } f(\mathbf{u}) < f(\mathbf{x}) \text{ functions} \\ \mathbf{x} & \text{otherwise} \end{cases} \quad (33)$$

5.4.3. 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.

```

input : PSO parameters, DE parameters
output: Software allocation solution sBest.x

Particles  $P \leftarrow \text{initPSO}()$ ;
while termination criteria do
     $P \leftarrow P$ ;
     $\text{sBest} \leftarrow P$ ;
    foreach  $p \in P$  do
        computeParticleVelocity( $p$ ) according to Equation (29);
        computeParticlePosition( $p$ ) according to Equation (30);
    end
    if interval criteria then
         $P \leftarrow \text{optimizeUsingDE}(P)$ ;
        //  $P \leftarrow \text{optimizeUsingHC}(P)$ 
        //  $P \leftarrow \text{optimizeUsingSHC}(P)$ 
    end
end

```

Algorithm 1: Hybrid PSO Algorithms.

5.4.4. 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.

5.4.5. 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 $\exists i, j | x_{ij} \neq x'_{ij}$, 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 the proposed approach using synthesized automotive applications that conform to the automotive benchmark proposed by Kramel et al. [14]. The number of runnables, timing specification and activation patterns within the cause-effect chains are also selected according to the benchmark. In order to show the scalability of our approach and to assess the scope of its applicability in practice, in some cases, we use higher specifications standard than what is indicated in the benchmark, e.g., the maximum number of activation patterns is extended from three to four.

In the rest of the section, we describe the setup and method of evaluation, followed by discussion of the evaluation results.

6.1. Setup

The evaluation setup consists of three hardware platforms with different computing capacities, i.e., processing speed and memory size as shown in Table 3. The evaluation on different platforms can be used as performance indicator and also to identify performance bottlenecks in the model. HP EliteBook and Lenovo 20378 are personal computers with core-i5 and core-i7 processors, respectively, whereas PowerEdge is a workstation with much higher processing and memory specification than the personal computers.

| Hardware Model | Pro. Model | #Pro. | #Core | Cache | RAM |
|---------------------------|------------------------------|-------|-------|-------|------|
| ¹ HP EliteBook | ³ Core i5, 2.2GHz | 1 | 2 | 3M | 8G |
| Lenovo 20378 | ⁴ Core i7, 2.6GHz | 1 | 4 | 6M | 16G |
| ² PowerEdge | ⁵ Xeon(R), 2.4GHz | 24 | 6 | 15M | 256G |

Table 3: Summary of the Hardware Specifications.

¹HP EliteBook 840 G2, ²PowerEdge R730 Rack Server

³Intel® Core™ i7-4720HQ @ 2.60GHz, ⁴Intel® Core™ i7-4720HQ @ 2.60GHz ⁵Intel(R) Xeon(R) CPU E5-2620 v3 @ 2.40GHz

6.1.1. Software Application Benchmark

In order to evaluate the proposed approach on different ranges of applications, we automatically generate synthetic examples that comply with the automotive benchmark [14]. The examples denote simple to complex automotive functions such as reverse-parking assistance system and engine control system. For simplicity, we identify three classes of applications with different sizes and complexity, shown as tuple (c, r, t, g) , respectively for the number of software components, runnables, tasks, and cause-effect chains. Table 4 shows the range of values used in the applications for evaluation.

| Parameter | Spec.-I | spec.-II | spec.-III |
|------------------------------|-------------------|------------|-----------|
| components, c | ≤ 10 | ≤ 15 | > 15 |
| runnables, r | ≤ 50 | ≤ 100 | > 100 |
| tasks, t | ≤ 30 | ≤ 60 | > 60 |
| cause-effect chains, g | ≤ 30 | ≤ 40 | > 60 |
| activation-pattern | $[2, 3, 4]$ | | |
| share of activation-patterns | $[0.7, 0.2, 0.1]$ | | |

Table 4: Specification of the Applications for Evaluation.

6.1.2. Platform Benchmark

The specification of nodes can be obtained from simulation, vendor product specification and experience. Figure 5 shows the range of values that are used in the nodes' specification. In the experiment, the values are randomly generated while respecting the benchmark.

| Parameter | Range |
|--------------------------------|------------------|
| nodes | $4 - 10$ |
| power consumption (Watt), p | $10 - 200$ |
| failure-rate (/Mhr), λ | $10^4 - 10^{-2}$ |
| speed factor, hz | $0.0 - 1.0$ |

Table 5: Range of Values for the Specification of Nodes.

6.2. Result

We conduct three experiments that assess the proposed approach for scalability in terms of *allocation time*. The allocation time is defined as the time required to prepare and solve the allocation problem using the proposed ILP method. Furthermore, we show resource efficiency in terms of saving nodes (i.e., using smaller number of nodes). The experiments consist of: i) varying the size of applications in order to observe the effect of increasing components, runnables and tasks in the system, ii) varying the complexity of applications in order

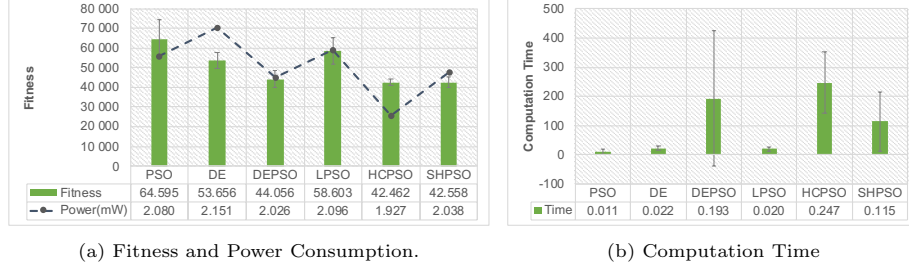


Figure 5: Allocation of Software Application ($N_c = 20, N_r = 200, N_T = 30$) on N_m number of Heterogeneous Nodes using Different Optimization Methods.

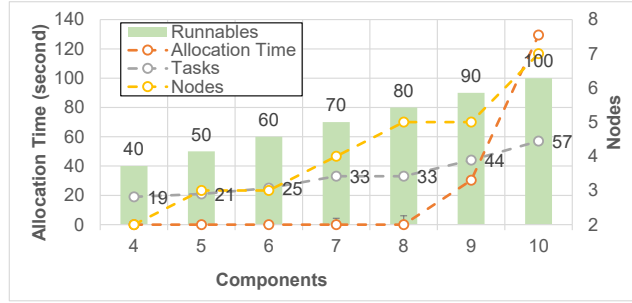


Figure 6: Effect of Varying the Application Size on the Allocation Time and Number of Utilized Nodes.

to observe the effect of cause-effect chains in the system, and iii) varying the replications. The experiments are discussed in detail in the next subsection.

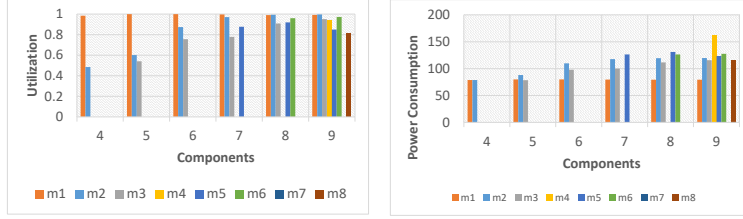
A preliminary analysis of the evaluation indicates a successful termination of the allocation for Spec-I&II of the applications. Whereas for Spec-III, the allocation problem is intractable. In fact, it took days on the PowerEdge machine and many times it did not terminate successfully. Therefore, the experimental results that are shown in this paper are conducted on the Spec-I&II classes.

6.2.1. ILP Evaluation

Varying the Size of Application. This refers to increasing the number of software components, as well as runnables, tasks, and cause-effect chains in the system. Figure 6 shows the effect of increasing the size of an application from ($c4, r40, t19, g30$) to ($c10, r100, t57, g60$) on the allocation time and the number of nodes utilized. The applications are allocated to a pool of 8 heterogeneous nodes sharing a single network and with specifications shown in Table ???. The specifications are generated randomly, with uniform distribution, within the scope of Table 5.

The CPLEX solver returns an optimal solution for the application ($c8, r80, t44, g30$) within 6.06 sec. The allocation time increased sharply to 30.3 sec and 129.4 sec respectively for 9 and 10 components. Even if not indicated in this chart, the

| Application | Measure | PSO | LPSO | DE | DEPSO | HCPSO | SHPSO |
|-------------|----------|-----|------|----|-------|-------|-------|
| App1 | Mean | | | | | | |
| | Std.Dev. | | | | | | |
| App2 | | | | | | | |



(a) Utilization of Nodes. (b) Power Consumption of Nodes.

Figure 7: Allocation of Applications on Heterogeneous Nodes.

solver, the solver returns an optimal solution within 45 min for components reaching 15 on the PowerEdge machine and OutOfMemory error on the Lenovo and HP machines. Figure 7a and Figure 7b show the utilization and power consumption on each node for the different application sizes. The optimal allocation, in the general case, favor nodes with higher processor speed and lower power consumption specifications.

Varying the Number of Cause-effect Chains. In order to observe the effect of increasing the cause-effect chains on the allocation time, we vary the number of chains in the application from 10 to 60, which is consistent with the benchmark [14]. The share of activation patterns also increases proportionally with the ratio $1 : [0.7, 0.2, 0.1]$, respectively, for two, three, and four activation patterns. For instance, out of 10 cause-effect chains, there are 7 chains (with two activation patterns), 3 chains (with three activation patterns), and 1 chain (with four activation patterns). The experiment is conducted on two cases of schedulability analysis, namely response time analysis (RTA) and utilization bound (UB), and their result is shown in Figure 8 for increasing number of chains.

The figures in the data table show an exponential growth of allocation time whenever the cause-effect chains are increased linearly for a specific application, in both cases of schedulability analysis. In the case of RTA, the overhead in the pre-optimization is higher than the overhead in the optimization for 50 or less chains. Whereas in the UB case, the computational time of pre-optimization is almost always less than the computational time of the optimization. The results are consistent with our expectation that the RTA computation, in the preparation of the timing assertions, is expensive, albeit provides schedulable tasks allocation based on the fixed-priority scheduling policy. In contrast, the UB computation time is relatively low; however, the search space gets larger due to more and more feasible tasks partitioning and chains fulfilling the timing constraints. As a result, the optimization time in the case of UB is usually higher

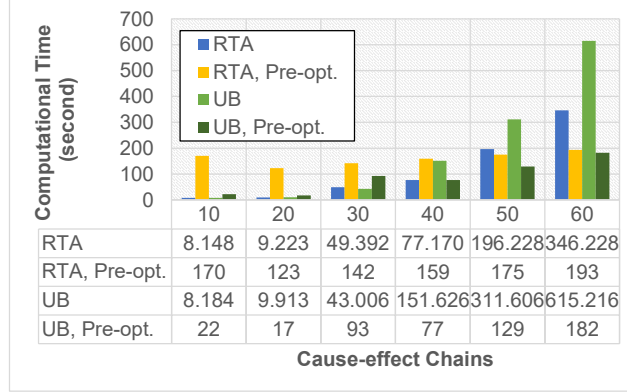
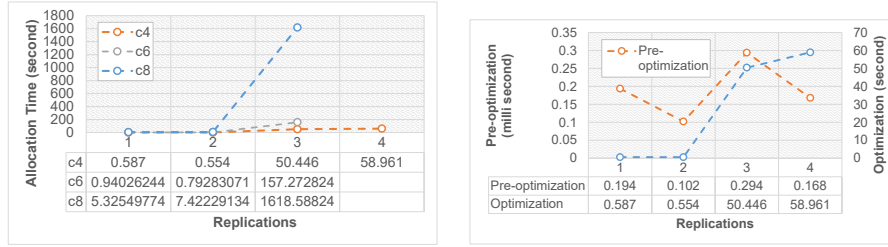


Figure 8: Effect of Increasing Cause-effect Chains (under RTA and UB) on the Allocation Time.

as compared to the case of RTA. Therefore, for applications with chains not more than 40 and naive scheduling assumption using UB, the experiments favor the UB assumption. Whereas, the allocation with the RTA assumption should be selected for applications with exact scheduling requirements. Note, the scalability of this experiment should be seen in conjunction with the experiment discussed in the previous subsection.

Varying Replications. In this experiment, we evaluate the allocation time of the applications with the increase in the number of replications. For the applications specification shown in Figure 9, we vary the replications from 1 to 4. The allocation in all applications took not more than 10 sec for replications 1 and 2. For Spec-I with replication 3 and 4, the allocation time went up close to 1 min. For Spec-III with replication 3, the allocation time went up rapidly to 30 min, and took extremely large time for replication 4 which is also the case for Application-II.



(a) Effect on Allocation Time.

(b) Effect on Pre-optimization.

Figure 9: Effect of Varying the Component Replications on the Allocation Time.

Figure 9b shows the effect of shared constraints on the overhead of replications during the pre-optimization and optimization phases of the allocation. In the pre-optimization phase, the allocation time was stable for the increased size of applications. This is due to the fixed constraints regardless of the replications.

In contrast, the allocation time increases during optimization as the constraints are applied for the various combination of task partitions, cause-effect chains, and reliability states generated as the result of replications.

6.3. Metaheuristic Evaluation

Clearly, the ILP evaluation shows that the ILP approach does not scale to large software allocation problems, that is, that allocation time increases exponentially as the size of the software application increases. In fact, the allocation does not return results for applications consisting more than 15 software components. In this subsection, we evaluate ILP in conjunction with the PSO, DE, and hybrid PSO techniques for different problem sizes.

Table 6 shows the fitness values and allocation time of the different algorithms for the increasing sizes of the software allocation problems. For no constraints violations, the fitness values denote the total power consumption, which is the case for the first three problems whereas it is not the case for the last three problems, thus the fitness values extremely increased as more violations are made. Furthermore, the algorithms highlighted in bold shows the benchmark for the corresponding specific problems.

The result shows that the ILP approach returned solutions for the first three problems, in which case it becomes the benchmark for the quality assessment of the metaheuristic algorithms. Whereas, for the last three problems, ILP does not return solution, which conforms the previous ILP evaluation results. The metaheuristic results are further discussed as follows, that is with regard to solution quality and computation time.

Solution Quality. For the first problem, which models the smallest software allocation over heterogeneous nodes, the algorithms DE, LPSO, HCPSO, SHPSO returned the optimal results besides ILP, with zero deviations. However, DEPSO performed relatively worse but better than PSO. The same results are repeated out of the second allocation problem for the algorithms HCPSO, and SHPSO, which are optimal. In contrast, DE and LPSO performed worse followed by DEPSO, and PSO performed relatively the worst. For the third problem, no metaheuristic algorithms returned the optimal solutions on average. Furthermore, the stability of the solutions for the algorithms DE and PSO, as indicated by the standard deviations of the fitness values, are quite higher than the rest metaheuristic algorithms.

As the size of the problem further increases, that is *c20m10g30* and *c20m10g30*, the hybrid PSO performed the far better, with HCPSO the best followed by SHPSO and DEPSO. However, as the size of the problem further increases, that is with *c80m20g60*, HCPSO did not return solutions at reasonable time. In contrast, its stochastic variant HCPSO performed the best followed by DEPSO, LPSO, and PSO, in order of their solutions quality, and DE performed the last.

Allocation Time. In the case of metaheuristic algorithms, the allocation time refers to the amount of time needed to return the near (optimal) result. It is calculated over a maximum of 5000 iterations (or generations) and maximum

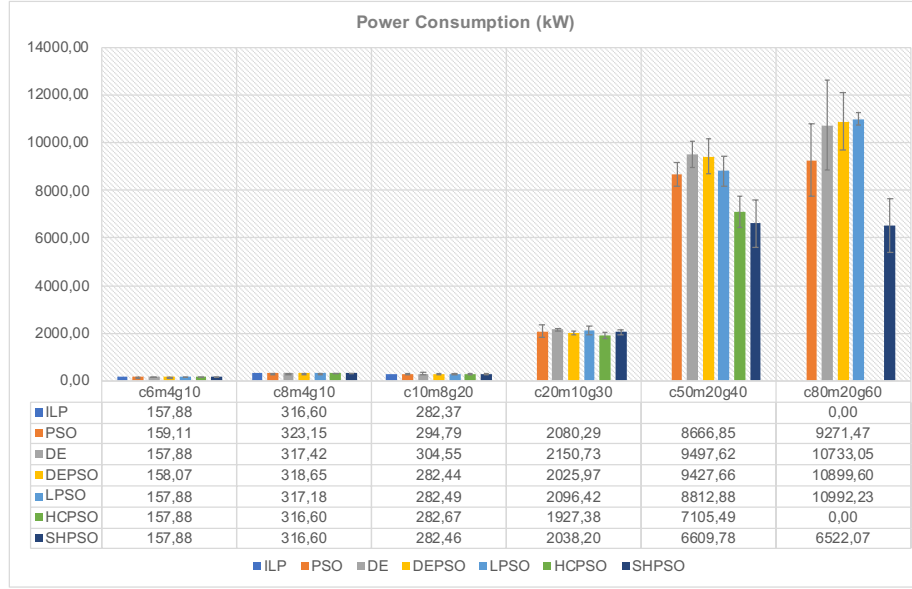


Figure 10: (Near) Optimal Power Consumption of the Different Software Allocation Problems.

steady time of 5mins, that is, the period in which no changes are observed in the fitness values. Figure 11 shows the allocation time of the algorithms for the problems listed in Table 6. ILP rapidly increased to 14sec for the problem *c10m8g20* which became intractable for larger problems. Whereas HCPSO rapidly increased to 17sec for the problem *c50m20g40* and became intractable for larger problems. However, for the rest of the algorithms, the allocation time did not exceed 6secs for all of the allocation problems.

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 [40][41][32][42]. Interleaving real-time constraints into the problem adds more complexity to reliability-aware task allocation in distributed systems [43]. 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

| Problem | Algorithm | Fitness | | Time (ms) | | Quality |
|-----------|-----------|------------|----------|-----------|----------|---------|
| | | Mean | SD | Mean | SD | |
| c6m4g10 | 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 |
| c8m6g20 | 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 |
| c10m8g20 | ILP | 442.37 | 0 | 14049.1 | 150.84 | 100.00 |
| | 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 |
| c20m10g30 | 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 |
| c50m20g60 | 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 |
| c80m20g60 | ILP | NA | NA | NA | NA | NA |
| | 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 |
| | SHPSO | 2466535.41 | 89380.36 | 2147.79 | 357.58 | 100.00 |

Table 6: Fitness and Allocation Time of the ILP and the Metaheuristic Techniques, for the Increasing Sizes of the Software Allocation Problem.

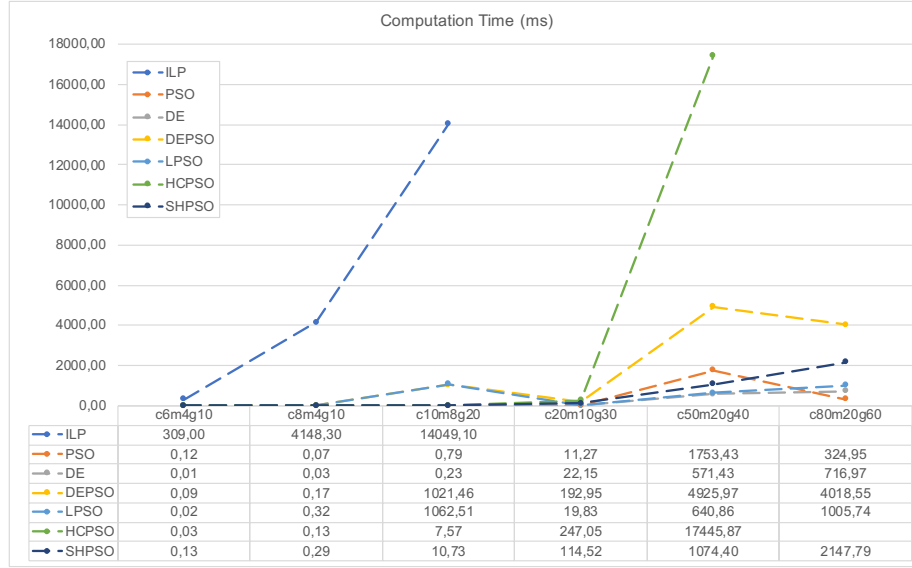


Figure 11: Computation Time of the Various Algorithms for Solving Different Instances of the Software Allocation Problem.

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, [44] 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 [45] 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) [46][47], and ii) task consolidation to minimize the number of used computing and communication units [48], 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 [49][50]. 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) [51] 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 heterogeneous 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.

Acknowledgement

This work is supported by the Swedish Governmental Agency for Innovation Systems (Vinnova) through the VeriSpec project, and the Swedish Knowledge

Foundation (KKS) through the projects HERO and DPAC.

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