Multi-Objective Task Assignment and Multiagent Planning with Hybrid GPU-CPU Acceleration

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Abstract. Allocation and planning of a collection of tasks to a group of agents under multiple-objective constraints is a challenging problem. Existing models which are used for model checking of these problems are prohibitively expensive. We show the convexity of our formal problem (MORAP), which can be decentralised, avoiding the exponential model size growth with agent-task numbers. Our main algorithm is a new point-oriented Pareto computation. Our algorithm checks whether a point corresponding to given cost and probability thresholds for our formal problem is feasible or not. If the given point is non-feasible, it also finds a new point which is closest to the given point and locates on the Pareto curve of the problem. We provide the first multi-objective model checking framework that simultaneously uses GPU and multi-core acceleration. Our framework manages CPU and GPU devices as a load balancing problem for parallel computation. Our experiments demonstrate that parallelisation achieves significant run time speed-up over serial methods.

Keywords: Multiagent Systems \cdot Task Allocation and Planning \cdot Model Checking \cdot GPU.

1 Introduction

Multiagent task allocation and planning is concerned with enabling a group of agents to divide up tasks amongst themselves and carry out their planning and execution, in possibly stochastic environments. For modern problems, coordination of agents usually involves conflicting solutions to the multiple objectives that such a multiagent system (MAS) is required to satisfy, for example, agents may need to balance execution time with energy consumption. Existing models for such systems typically increase exponentially in size with a linear increase in the number of agents in the system [5]. The major challenges in computation of models describing these systems include: (i) efficient computation and storage of such models; (ii) optimal task allocation to satisfy multi-objective constraints is an NP-hard problem; (iii) safety critical systems require rigorous guarantees that agents can carry out the set of tasks successfully.

Probabilistic model checking (PMC) is a verification technique to establish rigorous guarantees about the correctness of real-life stochastic systems [2]. PMC provides methods to incorporate LTL properties and query whether or not a system is capable of satisfying the property. We can substitute a property for a task specification by restricting the LTL specification to co-safe LTL[18] or LTL_f[8] which completes in finite time. In this work, we use co-safe LTL which has the benefit of a corresponding deterministic finite automaton (DFA) representation [2]. Task execution in finite time is important because we typically want to reuse the MAS to execute further tasks. In settings where an agent may carry out multiple actions in a given state, and the outcome of actions are uncertain, Markov Decision Processes (MDP) [25] is a fundamental model in PMC and is often used for multiagent planning [1]. When verification multi-objectives are concerned, we require the Multi-objective MDP (MOMDP) extension of the standard MDP [27] whose reward structure specifies cost vectors (rather than scalars). The solution space of the MOMDP is a convex polyhedron and we can construct a randomised scheduler in polynomial time by approximating a Pareto curve along its convex hull [12, 9].

The classical assignment problem finds an assignment, namely a one-to-one mapping from tasks to agents, such that the assignment reward for some given reward function is maximised. The multi-objective assignment problem is to determine an assignment such that the vectorised assignment reward is Pareto optimal. The classical assignment problem can be solved efficiently (e.g., using the Hungarian algorithm [17]), but the multi-objective assignment problem is much harder [30]. The multi-objective random assignment (MORA) problem pursues a randomised distribution over assignments such that the expected assignment reward is Pareto optimal.

Multi-objective MDP model checking is a technique concerned with a tradeoff between different objectives, and is implemented in Prism and Storm. Currently this technique includes three kinds of queries [12]: The achievability query is the most basic query, which asks whether there exists a scheduler to fulfil all objective thresholds. The numerical query is a numerical variant of the first query, which computes the optimal value of one objective while fulfiling all other objective thresholds. The Pareto query is the most expensive query, which computes approximately the Pareto curve of all objectives.

In this paper, we extend multi-objective MDP model checking to a setting of multi-objective random assignment and planning (MORAP) in MAS and present a novel implementation with hybrid GPU-CPU acceleration. Our main contributions are as follows:

- We show the convexity of our formal problem (MORAP), and that a practical approach to solve this problem can rely on a decentralised model, which avoids the exponential model size growth with agent-task numbers.
- Our main algorithm is a new point-oriented Pareto computation complementing the existing achievability and Pareto queries [12]. For a given point corresponding to cost and probability thresholds, our algorithm finds a point which is feasible for the MORAP problem and closest to the given point under a general vector norm.

To the best of our knowledge, we provide the first multi-objective model checking framework that utilises simultaneous GPU and multi-core acceleration. Our framework manages CPU and GPU devices as a load balancing problem for parallel computation.

Formal proofs of theorems are included in the appendix of the long version of this paper [26]. The remainder of this paper is organised as follows: Section 2 provides the preliminaries for the problem; Section 3 gives the approach to the problem, model construction and algorithms; Section 4 provides details on the hybrid implementation and parallel architecture; Section 5 analyses the performance of our approach; Section 6 provides related work; and finally Section 7 summarises and proposes future work.

2 Preliminaries

Deterministic finite automata. A deterministic finite automaton (DFA) \mathcal{A} is given by the tuple $(Q, q_0, Q_F, \Sigma, \delta)$ where (i) Q is a set of locations, (ii) $q_0 \in Q$ is an initial location, (iii) $Q_F \subseteq Q$ is a set of accepting locations, (iv) $\Sigma = 2^{AP}$ (where AP is a non-empty set of atomic propositions) is the alphabet, and (v) $\delta: S \times \Sigma \to S$ is the transition function. Let $\mathcal{A}[q]$ be a DFA by changing the initial location of \mathcal{A} to q. Let $Q_R = \{q \in Q \mid acc(\mathcal{A}[q]) \cap acc(\mathcal{A}) = \emptyset\}$. If $\delta(q, W) = q'$ for some $W \subseteq AP$, we call q a predecessor of q' and q' a successor of q. Let pre(q) and predecessor of q and predecessor of q and q' a successor of q and q' and q' and q' and q' a successor of q. Let pre(q) and predecessor of q and q' and q' a successor of q and q' and q' a successor of q and q' and q' and q' a successor of q and q' and q' and q' a successor of q and q' and q'

Co-Safe LTL. Automata are often considered too low-level in practice. LTL is a compact representation of linear time properties. The syntax of LTL is $\varphi := \top \mid \mathbf{a} \mid \neg \varphi \mid \varphi \land \varphi \mid \mathsf{X} \varphi \mid \varphi \, \mathsf{U} \varphi$ where $\mathbf{a} \in AP$. The operators X and U stand for "next" and "until", respectively. Let $\mathsf{F} \varphi := \top \, \mathsf{U} \varphi$, and $\mathsf{G} \varphi := \neg \, \mathsf{F} \, \neg \varphi$. The semantic relationship $\sigma \models \varphi$ where $\sigma \in \varSigma^\omega$ is standard. We are interested in the *co-safe* fragment of LTL formulas. Informally, φ is co-safe if any σ such that $\sigma \models \varphi$ includes some *good prefix* (which is accepting in some DFA). Syntactically, any LTL formula containing only the temporal operators X (next), U (until), and F (eventually) in positive normal form (PNF) is co-safe. A formal characterisation in the semantic level is included the appendix of [26].

Markov Decision Process. A (labelled) MDP is given by the tuple $\mathcal{M} = (S, s_0, A, P, L)$ where (i) S is a finite nonempty state space, (ii) $s_0 \in S$ is an initial state, (iii) A is a set of actions, (iv) $P: S \times A \times S \to [0, 1]$ is a transition probability function such that $\sum_{s' \in S} P(s, a, s') \in \{0, 1\}$, and (v) $L: S \to \Sigma$ is a labelling function. A reward function or structure for \mathcal{M} is a function $\rho: \{(s, a) \in S \times A \mid a \in A(s)\} \to \mathbb{R}$. We write $\mathcal{M}[\rho]$ to explicitly indicate the reward structure ρ for \mathcal{M} . Let $A(s) = \{a \in A \mid \sum_{s' \in S} P(s, a, s') = 1\}$, i.e., A(s) is the set of enabled actions at s. An (infinite) $path \pi$ is a sequence $s_1a_1s_2a_2\ldots$ such that $P(s_i, a_i, s_{i+1}) > 0$ for all $i \geq 1$. Let $L(\pi)$ denote the word $L(s_1)L(s_2)\ldots \in$

 Σ^{ω} . Let IPath be the set of paths in \mathcal{M} and IPath(s) be the subset of IPath containing the paths originating from s. The set of probability distributions over A is denoted by Dist(A). A scheduler (or memoryless scheduler) for \mathcal{M} is a mapping $\mu: s \mapsto Dist(A(s))$ for all $s \in S$. If μ is a simple (or pure) if $\mu(s)(a) = 1$ for each $s \in S$ and some $a \in A(s)$. The set of schedulers (resp., simple schedulers) is denoted by $Sch(\mathcal{M})$ (resp., $Sch_S(\mathcal{M})$).

Reachability Reward. Given any LTL formula FB with B being a Boolean formula, let $\rho(\pi|\mathsf{F}B) = \sum_{i=1}^n \rho(s_i, a_i)$ where $\pi = s_1 a_1 s_2 a_2 \ldots \in \mathrm{IPath}(s)$ and n is the smallest number such that $L(s_n) \models B$ and $L(s_i) \not\models B$ for all i < n; if such n does not exist, let $\rho(\pi|\mathsf{F}B) = \infty$. Let $\mathbf{Pr}^{\mathcal{M},\mu}$ be the probability measure on $\mathrm{IPath}(s_0)$ for \mathcal{M} under μ [2]. The expectation $\mathbf{E}^{\mathcal{M}[\rho],\mu}(\mathsf{F}B) \doteq \int_{\pi} \rho(\pi|\mathsf{F}B) \mathrm{d}\mathbf{Pr}^{\mathcal{M},\mu}$, a.k.a. reachability reward [20], is the expected reward accumulated in a path of \mathcal{M} under μ until reaching states satisfying B. We say $\mathcal{M}[\rho]$ is reward-finite w.r.t. $\mathsf{F}B$ if $\sup_{\mu \in Sch(\mathcal{M})} \mathbf{E}^{\mathcal{M}[\rho],\mu}(\mathsf{F}B) < \infty$.

Product MDP. Given $\mathcal{M} = (S, s_0, A, P, L)$ and $\mathcal{A} = (Q, q_0, Q_F, \Sigma, \delta)$, a product MDP is a tuple $\mathcal{M} \otimes \mathcal{A} = (S \times Q, (s_0, q_0), A, P', L')$ where (i) $P' : S \times Q \times A \times S \times Q \rightarrow [0, 1]$ is a transition probability function such that

$$P'(s,q,a,s',q') = \begin{cases} P(s,a,s') & \text{if } q' = \delta(q,L(s')) \\ 0 & \text{otherwise} \end{cases}$$

and (ii) $L': S \times Q \to 2^{\Sigma}$ is a labelling function s.t. L'(s,q) = L(s). Let $\mathcal{M}[\rho] \otimes \mathcal{A}$ refer to $(\mathcal{M} \otimes \mathcal{A})[\rho]$ where $\rho(s,q,a) = \rho(s,a)$ for all $(s,q) \in S \times Q, a \in A(s)$.

Geometry. For a vector $\boldsymbol{v} \in \mathbb{R}^n$ for some n, let v_i denote the i^{th} element of \boldsymbol{v} . A weight vector \boldsymbol{w} is a vector such that $w_i \geq 0$ and $\sum_{i=1}^n w_i = 1$. The dot product of \boldsymbol{v} and \boldsymbol{u} , denoted $\boldsymbol{v} \cdot \boldsymbol{u}$, is the sum $\sum_{i=1}^n v_i u_i$. For a set $\boldsymbol{\Phi} = \{\boldsymbol{v}_1, \ldots, \boldsymbol{v}_m\} \subseteq \mathbb{R}^n$, a convex combination in $\boldsymbol{\Phi}$ is $\sum_{i=1}^m w_i \boldsymbol{v}_i$ for some weight vector $\boldsymbol{w} \in \mathbb{R}^m$. The downward closure of the convex hull of $\boldsymbol{\Phi}$, denoted $down(\boldsymbol{\Phi})$, is the set of vectors such that for any $\boldsymbol{u} \in down(\boldsymbol{\Phi})$ there is a convex combination $\boldsymbol{v} = w_1 \boldsymbol{v}_1 + \ldots + w_m \boldsymbol{v}_m$ such that $\boldsymbol{u} \leq \boldsymbol{v}$. Let $\boldsymbol{\Psi} \subseteq \mathbb{R}^n$ be any downward closure of points. A vector $\boldsymbol{u} \in \boldsymbol{\Psi}$ is Pareto optimal if $\boldsymbol{u}' \geq \boldsymbol{u}$ implies $\boldsymbol{u}' = \boldsymbol{u}$ for any $\boldsymbol{u}' \in \boldsymbol{\Psi}$. A Pareto curve in $\boldsymbol{\Psi}$ is the set of Pareto optimal vectors in $\boldsymbol{\Psi}$. The follow lemma is from the separating hyperplane and supporting hyperplane theorems.

Lemma 1 ([6]). Let $\Psi \subseteq \mathbb{R}^n$ be any downward closure of points. For any $\mathbf{v} \notin \Psi$, there is a weight vector \mathbf{w} such that $\mathbf{w} \cdot \mathbf{v} > \mathbf{w} \cdot \mathbf{x}$ for all $\mathbf{x} \in \Psi$. We say that \mathbf{w} separates \mathbf{v} from Ψ . Also, for any \mathbf{u} on the Pareto curve of Ψ , there is a weight vector \mathbf{w}' such that $\mathbf{w}' \cdot \mathbf{u} \geq \mathbf{w}' \cdot \mathbf{x}$ for all $\mathbf{x} \in \Psi$. We say that $\{\mathbf{x} \in \mathbb{R}^n \mid \mathbf{w}' \cdot \mathbf{x} = \mathbf{w}' \cdot \mathbf{u}\}$ is a supporting hyperplane of Ψ .

Bistochastic Matrix. For a matrix $U \in \mathbb{R}^{n \times n}$ for some n, let $u_{i,j}$ denote the element of U in the i^{th} row and j^{th} column. U is bistochastic if $u_{i,j} \geq 0$ and $\sum_{i'=1}^n u_{i',j} = \sum_{j'=1}^n u_{i,j'} = 1$ for all $1 \leq i,j \leq n$. A bistochastic matrix U is a permutation matrix if U has exactly one element with value 1 in each row and each column. We recall the following Birkhoff-von Neumann Theorem:

Lemma 2 ([3]). A bistochastic matrix U equals to a convex combination of permutation matrices U_1, \ldots, U_k for some $k \le n^2 - 2n + 2$.

Random Assignment. Given a set I (resp., J) of agents (resp., tasks) with |I| = |J|, a (balanced) assignment is a bijective function $f: J \to I$. Denote the set of assignments of J to I by \mathcal{F} . A random assignment ν is a randomised distribution over \mathcal{F} (or, equivalently, a convex combination of assignments in \mathcal{F}). For convenience, let $I = J = \{1, \ldots, n\}$. Let $\nu_{j \to i} = \nu(\{f \in \mathcal{F} \mid f(j) = i\})$, i.e., the marginal probability of assigning task j to agent i according to ν . Clearly, any assignment is equivalent to a permutation matrix. By Lemma 2, a bistochastic matrix U is equivalent to a random assignment ν such that $u_{i,j} = \nu_{j \to i}$.

3 Problem and Approach

3.1 Problem Statement

In our MAS setting, each agent is an MDP (with a reward structure) and each task is a DFA (or a co-safe LTL formula), and the rewards are the probabilities (as probabilistic rewards) of accomplishing the tasks and the costs (as negative rewards) that agents execute the tasks. Therefore, our problem is the computation of a random assignment and a scheduler for each agent and task, which must address multiple probability and cost requirements. Intuitively, we consider the task assignment and agent planning scenario which satisfies the following two conditions [29]:

- **C1.** The tasks are mutually independent.
- **C2.** The behaviours of agents do not impact each other.

For each $(i, j) \in I \times J$, we define an agent-task (product) MDP $\mathcal{M}_{i \otimes j}[\rho_i] \doteq \mathcal{M}_i[\rho_i] \otimes \mathcal{A}_j$ and include an atomic proposition done, such that

$$L_{i,j}(s,q) \models \mathtt{done}_j \text{ iff } q \in Q_{j,F} \cup Q_{j,R}$$

which indicates "task j is ended (either accomplished or failed)." For each $j \in J$ we define a designated reward function $\rho_{j+|I|}: \bigcup_{i \in I} (S_i \times Q_j \times A_i) \to \{0,1\}$ such that $\rho_{j+|I|}(s,q,a)=1$ iff $q \notin Q_{j,F}$ and $\mathrm{suc}(q)\subseteq Q_{j,F}$. If such a presink q does not exist, we can modify \mathcal{A}_j to include q without altering $acc(\mathcal{A}_j)$. In words, $\rho_{j+|I|}$ provides a one-off unit reward whenever an accepting location will be traversed for the first time. Informally, $\rho_{j+|I|}$ expresses "the probability of accomplishing task j." As the atomic proposition done_j is fixed for each $\mathcal{M}_{i\otimes j}$, we abbreviate $\mathbf{E}^{\mathcal{M}_{i\otimes j}[\rho_k],\mu_{i,j}}(\mathsf{Fdone}_j)$ as $\mathbf{E}^{\mathcal{M}_{i\otimes j}[\rho_k],\mu_{i,j}}$ where k=j or k=j+|I|. Similar to the multi-objective verification literature [12,14], we require that $\mathcal{M}_{i\otimes j}[\rho_i]$ is reward-finite (w.r.t. Fdone_j) for all $(i,j)\in I\times J$.

¹ Throughout the paper we assume $I = J = \{1, ..., n\}$ for some n unless explicitly stated otherwise, however we still use symbols I, J to indicate the agent or task references of indexing integers.

$$\begin{cases} \text{Maximise} \\ \begin{cases} \sum_{j \in J} \sum_{(s,q) \in S_i \times Q_j} \sum_{a \in A_i(s)} \rho_i(s,q,a) x_{s,q,a} & \forall i \in I \\ \sum_{i \in I} \sum_{(s,q) \in S_i \times Q_j} \sum_{a \in A_i(s)} \rho_{j+|I|}(s,q,a) x_{s,q,a} & \forall j \in J \end{cases} \\ \text{Subject to } \forall i \in I, j \in J, (s,q) \in S_i \times Q_j : \\ \begin{cases} \sum_{a \in A_i(s)} x_{s,q,a} - \mathbf{I}_{(s,q) = (s_{i,0},q_{j,0})} x_{i,j} \\ = \sum_{(s',q') \in S_i \times Q_j} \sum_{a' \in A_i(s')} P_{i,j}(s',q',a',s,q) x_{s',q',a'} \\ x_{s,q,a} \geq 0; \ x_{i,j} \geq 0; \ \sum_{i' \in I} x_{i',j} = 1; \ \sum_{j' \in I} x_{i,j'} = 1 \end{cases} \end{cases}$$

Fig. 1: The multi-objective linear program for MORAP

Definition 1 (MORAP). A multi-objective random assignment and planning (MORAP) problem is finding a bistochastic matrix $(x_{i,j})_{i\in I,j\in J}$ and a set of schedulers $\{\mu_{i,j}\in Sch(\mathcal{M}_{i\otimes j})\mid i\in I,j\in J\}$ such that the following two groups of requirements are satisfied:

(Probability)
$$\sum_{i \in I} x_{i,j} \mathbf{E}^{\mathcal{M}_{i \otimes j}[\rho_{j+|I|}], \mu_{i,j}} \geq p_j \text{ for all } j \in J,$$

(Cost) $\sum_{j \in J} x_{i,j} \mathbf{E}^{\mathcal{M}_{i \otimes j}[\rho_i], \mu_{i,j}} \geq c_i \text{ for all } i \in I,$

where the probability thresholds $(p_j)_{j\in J} \in [0,1]^{|J|}$ and the cost thresholds $(c_i)_{i\in I} \in \mathbb{R}^{|I|}$ are given. If the above requirements are satisfied, we say that the MORAP problem is feasible with given thresholds or just that the thresholds are feasible.

Definition 1 is an adequate formulation in the presence of conditions C1 and C2. First, since tasks are mutually independent (C1), the probability requirements only need to address the successful probability of each task. Second, since the execution of any task by each agent does not impact other agents (C2), the cost requirements only need to consider the cost of each agent. In practice, we can relax the condition |I| = |J| to $|I| \ge |J|$ (e.g., adding dummy tasks whose probability threshold is 0).

3.2 Convex Characterisation and Centralised Model

An essential characteristic of the MORAP problem is the *convexity*. More specifically, the downward closure of probability and cost thresholds such that the MORAP problem is feasible is a convex polytope (i.e., the downward convex hull of some finite set of points). This follows from the fact that the MORAP problem can be expressed as a multi-objective linear program (LP) by using a similar technique which underpins multi-objective verification of MDPs [12, 23, 9]. Fig. 1 includes the multi-objective LP for MORAP. Intuitively, for each $(i,j) \in I \times J$, $x_{i,j}$ represents the probability of assigning j to i (c.f., Lemma 2), and for each $(s,q) \in S_i \times Q_i$, $x_{s,q,a}$ is the expected frequency of visiting (s,q) and taking action a. A memoryless scheduler can be defined as follows: $\mu_{i,j}(s,q)(a) = x_{s,q,a}/x_{s,q}$ where $x_{s,q} = \sum_{a \in A_i(s)} x_{s,q,a}$. Therefore, the MORAP problem has the following complexity:

Theorem 1. The MORAP problem is solvable in polynomial time.

LP is not efficient for large problems, and value- and policy-iteration methods are more scalable methods in practice. For this purpose, we define a centralised MDP model which combines all agent-task MDPs and includes an additional variable indicates which agents have been assigned with tasks.

Definition 2 (Centralised MDP). A centralised MDP is $\mathcal{M}^{\mathsf{ct}} = (S^{\mathsf{ct}}, s_0^{\mathsf{ct}}, A^{\mathsf{ct}}, P^{\mathsf{ct}}, L^{\mathsf{ct}})$ where (i) $S^{\mathsf{ct}} = \bigcup_{i \in I} \bigcup_{j \in J} S_i \times Q_j \times 2^I$, (ii) $s_0^{\mathsf{ct}} = (s_{1,0}, q_{1,0}, \emptyset)$, (iii) $A^{\mathsf{ct}} = \bigcup_{i \in I} A_i \cup \{b_1, b_2, b_3\}$, (iv) $P^{\mathsf{ct}} = S^{\mathsf{ct}} \times A^{\mathsf{ct}} \times S^{\mathsf{ct}} \to [0, 1]$ such that:

- $-P^{\mathsf{ct}}(s,q,\sharp,a,s',q',\sharp) = P_{i,j}(s,q,a,s',q') \text{ if } s,s' \in S_i, \ q,q' \in Q_j, \ a \in A_i(s)$ and $i \in \sharp$ for some i,j.
- $-\ P^{\mathsf{ct}}(s,q,\sharp,b_1,s,q,\sharp \cup \{i\}) = 1 \ \textit{if} \ s = s_{i,0}, \ q = q_{j,0} \ \textit{and} \ i \notin \sharp \ \textit{for some} \ i,j,$
- $P^{\text{ct}}(s, q, \sharp, b_2, s', q, \sharp) = 1$ if $s = s_{i,0}$, $q = q_{j,0}$, $\sharp \subseteq I$, and $s' = s_{i',0}$ with $i' = \arg\min\{i'' \in I \mid i'' > i, i'' \notin \sharp\}$ for some i, j,
- $P^{\text{ct}}(s, q, \sharp, b_3, s', q', \sharp) = 1$ if $s \in S_i$, $q \in Q_{j,F} \cup Q_{j,R}$, $i \in \sharp$, $s' = s_{i',0}$ with $i' = \arg\min\{i'' \in I \mid i'' \notin \sharp\}$, and $q' = q_{j+1,0}$ for some i, j < |J|.

(v)
$$L^{\operatorname{ct}}: S^{\operatorname{ct}} \to 2^{\{\operatorname{done}\}}$$
 such that $L^{\operatorname{ct}}(s,q) \models \operatorname{done} iff q \in Q_{j,F} \cup Q_{j,R}$

Intuitively, \sharp contains agents who have worked on some tasks; b_1 indicates "a task is assigned to the current agent"; b_2 indicates "a task is forwarded to the next agent"; and b_3 indicates "the next task is considered". The model behaves as an individual product MDP when working on the assigned tasks.

Given any reward structure ρ for $\mathcal{M}_{i\otimes j}$, we view ρ as a reward structure for $\mathcal{M}^{\mathsf{ct}}$ by letting $\rho(s,q,\sharp,a) = \rho(s,q,a)$ a reward structure and $\rho(s,q,\sharp,a) = 0$ otherwise for all (s,q,\sharp,a) . Similarly, given any reward structure ρ for $\mathcal{M}^{\mathsf{ct}}$, a restriction of ρ on $S_i \times Q_j \times A_i$ is a reward structure for $\mathcal{M}_{i\otimes j}$. Similar to agent-task MDPs, we abbreviate $\mathbf{E}^{\mathcal{M}^{\mathsf{ct}}[\rho],\mu}(\mathsf{Fdone})$ as $\mathbf{E}^{\mathcal{M}^{\mathsf{ct}}[\rho],\mu}$ for a given ρ .

Theorem 2. The MORAP problem in Definition 1 is feasible with respect to $(p_j)_{j\in J}$ and $(c_i)_{i\in I}$ if and only if there is $\mu \in Sch(\mathcal{M}^{\mathsf{ct}})$ such that $\mathbf{E}^{\mathcal{M}^{\mathsf{ct}}[\rho_{j+|J|}],\mu} \geq p_j$ and $\mathbf{E}^{\mathcal{M}^{\mathsf{ct}}[\rho_i],\mu} \geq c_i$ for all $i \in I, j \in J$.

With the above theorem, one can work on the centralised MDP $\mathcal{M}^{\mathsf{ct}}$ (e.g., by using value-iteration) to solve a MORAP problem. Therefore, existing probabilistic model checking tools for multi-objective MDP verification (e.g., Prism [19] and Storm [15]) can be employed. However, the state space of $\mathcal{M}^{\mathsf{ct}}$ is exponential with respect to the agent team size |I|. Therefore, this approach is hard to scale to a relatively large |I| (which equals to |J|).

3.3 Point-Oriented Pareto Computation by Decentralised Model

We present a decentralised method solve a given MORAP problem, especially when the agent number (i.e., task number) is large. Besides deciding whether the problem is feasible or not, for a non-feasible problem our method also computes

Algorithm 1: Point-oriented Pareto computation

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Input: \{\mathcal{M}_{i\otimes j}\}_{(i,j)\in I\times J}, \boldsymbol{\rho}=\{\rho_k\}_{k=1}^{|I|+|J|}, \boldsymbol{t} (a concatenation of \boldsymbol{c} and \boldsymbol{p}), \varepsilon\geq 0
 1 t_{\uparrow} := -\infty; t_{\downarrow} := t; \Phi := \emptyset; \Lambda := \emptyset; w := (1, 0, \dots, 0);
      while \| \boldsymbol{t}_{\downarrow} - \boldsymbol{t}_{\uparrow} \| > \varepsilon do
 3
             if \Phi \neq \emptyset then
                     Find x \in down(\Phi) minimising ||t - x||;
  4
                     t_{\uparrow} := x;
  5
                     oldsymbol{w} := oldsymbol{M}(oldsymbol{t} - oldsymbol{t}_{\uparrow}) / \|oldsymbol{M}(oldsymbol{t} - oldsymbol{t}_{\uparrow})\|_1;
  6
              Find r s.t. \{y \mid w \cdot y = w \cdot r\} is a supporting hyperplane of \mathscr C;
              \Phi := \Phi \cup \{r\}; \ \Lambda := \Lambda \cup \{(\boldsymbol{w}, \boldsymbol{r})\};
 8
              if w \cdot r < w \cdot t_{\downarrow} then
 9
                     Find z minimising ||t - z|| s.t. w' \cdot r' \ge w' \cdot z for all (w', r') \in \Lambda;
10
11
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a new feasible threshold vector on the Pareto curve of the problem, and nearest the original threshold vector up to some numerical tolerance.

Let $\mathscr{C}_0 = \{(\mathbf{E}^{\mathcal{M}^{\operatorname{ct}}[\rho_k],\mu})_{1 \leq k \leq |I|+|J|} \mid \mu \in Sch(\mathcal{M}^{\operatorname{ct}})\}$. The reward-finiteness implies that \mathscr{C}_0 is non-empty and bounded. Let \mathscr{C} be the downward closure of \mathscr{C}_0 , i.e., namely, \mathscr{C} is the set of feasible threshold vectors in Definition 1. The main algorithm for our method is presented in Algorithm 1 with the supporting hyperplane computation (i.e., Line 7) detailed in Algorithm 2. Algorithm 1 works by iteratively refining a lower approximation, encoded as Φ , and an upper approximation, encoded as Λ , for \mathscr{C} . It computes a vector \mathbf{t}_{\uparrow} (resp., \mathbf{t}_{\downarrow}) which is the closest point from the origin threshold vector \mathbf{t} to the lower (resp., upper) approximation such that \mathbf{t}_{\uparrow} and \mathbf{t}_{\downarrow} converge eventually.

The algorithm uses a general norm $\|\cdot\|$ to measure the distance between vectors, because in practice one may prefer to differentiate the importance of probability and cost thresholds. An inner product of $\boldsymbol{v}, \boldsymbol{u} \in \mathbb{R}^m$ (m a positive integer), denoted $\langle \boldsymbol{v}, \boldsymbol{u} \rangle$, is the matrix-vector multiplication $\boldsymbol{v}^T \boldsymbol{M} \boldsymbol{u}$, where \boldsymbol{M} is a symmetric positive-definite matrix. Note that if \boldsymbol{M} is the identity matrix then $\langle \boldsymbol{v}, \boldsymbol{u} \rangle$ is $\boldsymbol{v} \cdot \boldsymbol{u}$. $\|\cdot\|_1$ refers to vector 1-norm. The weight vector \boldsymbol{w} computed in Line 6 provides the orthogonal orientation between the point \boldsymbol{t} and the convex set $down(\Phi)$. $\boldsymbol{w} \cdot \boldsymbol{\rho}$ refers to a weighted combination of reward functions in $\boldsymbol{\rho}$.

Theorem 3. Algorithm 1 terminates. Throughout the execution of Algorithm 1, the following properties hold: (i) $\mathbf{t}_{\uparrow} \in \mathscr{C}$. (ii) If $\mathbf{t} \in \mathscr{C}$ then $\mathbf{t}_{\downarrow} = \mathbf{t}$. (iii) $\|\mathbf{t} - \mathbf{t}_{\downarrow}\| \leq \min_{\mathbf{u} \in \mathscr{C}} \|\mathbf{t} - \mathbf{u}\| \leq \|\mathbf{t} - \mathbf{t}_{\uparrow}\|$.

Corollary 1. Let $\varepsilon = 0$. Then, after Algorithm 1 terminates, (i) $\mathbf{t}_{\uparrow} = \mathbf{t}_{\downarrow}$ and (ii) $\mathbf{t} \in \mathscr{C}$ if and only if $\mathbf{t}_{\downarrow} = \mathbf{t}$.

Algorithm 2 finds a supporting hyperplane of \mathscr{C} for a given orientation \boldsymbol{w} . As probabilistic model checking are employed in the two inner loops, it is usually very expensive in computation. To see the significance of Algorithm 2, we point out that \mathscr{C} is a convex set defined on the centralised model $\mathcal{M}^{\mathsf{ct}}$ whose size is

Algorithm 2: Supporting hyperplane computation in Line 7 of Alg. 1

```
Input: \{\mathcal{M}_{i\otimes j}\}_{(i,j)\in I\times J}, \, \rho=\{\rho_k\}_{k=1}^{|I|+|J|}, \, \mathbf{w}

1 foreach (i,j)\in I\times J do

\begin{array}{c} /^*\text{ Line 2 is computed by policy iteration.} & */\\ \mathbf{2} & c_{i,j}:=\mathbf{E}^{\mathcal{M}_{i\otimes j}[\mathbf{w}\cdot\rho],\mu_{i,j}} \text{ with } \mu_{i,j}:=\arg\max_{\mu}\mathbf{E}^{\mathcal{M}_{i\otimes j}[\mathbf{w}\cdot\rho],\mu}; \\ \mathbf{3} \text{ Find an assignment } f\in\mathcal{F} \text{ maximising } \sum_{j\in J} c_{f(j),j}; \\ \mathbf{4} \text{ foreach } j\in J \text{ do} & \\ & /^*\text{ Lines 5-6 are computed by value iteration.} & */\\ \mathbf{5} & r_{j+|I|}:=\mathbf{E}^{\mathcal{M}_{f(j)\otimes j}[\rho_{j+|I|}],\mu_{f(j),j}}; \\ \mathbf{6} & \mathbf{r}_{f(j)}:=\mathbf{E}^{\mathcal{M}_{f(j)\otimes j}[\rho_{f(j)}],\mu_{f(j),j}}; \\ \mathbf{7} \text{ return } (r_k)_{k=1}^{|I|+|J|}; \end{array}
```

 $O(2^{|I|})$. But instead of dealing with \mathcal{M}^{ct} , Algorithm 2 works on a decentralised model consisting of $\{\mathcal{M}_{i\otimes j}\}_{(i,j)\in I\times J}$. The first inner loop includes $|I|\times |J|$ (i.e., $|I|^2$) policy-iteration processes to compute optimal schedulers and reachability rewards. The second inner loop uses 2|I| value-iteration processes under a fixed scheduler.² The model selection is computed by using the Hungarian algorithm [17] (Line 3) whose run time is $O(|I|^3)$. Another important implication of using a decentralised model is the parallel execution of the two inner loops, which we elabrate on in Section 4. Also notice that if $\mathcal{M}_{i\otimes j} = \mathcal{M}_{i'\otimes j'}$ for some $(i,j)\neq (i',j')$, then the two inner loops can skip the computation for some models.

Note that if $\varepsilon=0$, then after the algorithm terminates, $t_{\uparrow}=t_{\downarrow}$ and they are on the Pareto curve of \mathscr{C} . But in the implementation we should choose some positive ε for the following three reasons: First, the policy and value iterations for computing the two inner loops are approximate. Second, small numerical inaccuracy (e.g., rounding) usually occur in the solving optimisation problems in the algorithm. Third, while the number of iterations in Algorithm 1 may be very large in the worst case (i.e., exponential on the model size and agent number [12]), a suitable ε can terminate the algorithm earlier with an approximate threshold vector whose precision is acceptable in practice.

For synthesis purposes, we can extract a random assignment and a collection of schedulers. Assume that the while loop iterates ℓ times in total. Let $\{\mu_{i,j}^t \mid i \in I, j \in J, \}$ and f_{ι} be generated in Lines 2-3 in Algorithm 1, respectively, in the ι^{th} iteration. Let $v_1 r_1 + \ldots + v_{\ell} r_{\ell} \geq t_{\uparrow}$ for some weight vector \boldsymbol{v} (this \boldsymbol{v} exists since $\boldsymbol{t}_{\uparrow} \in down(\Phi)$). The convex combination of assignments $v_1 f_1 + \ldots + v_{\ell} f_{\ell}$ defines a random assignment (i.e., bistochastic matrix). After an assignment f_{ι} is chosen randomly according to probability v_{ι} , the schedulers for planning are those from $\{\mu_{i,j}^{\iota} \mid j \in J, f_{\iota}(j) = i\}$.

Example. An example demonstrating execution of Algorithm 1. The agent MDP is shown in Fig. 2a. Construct a task $\varphi := \neg x \cup y$, and $\varepsilon = 0.0001$. Let an achievable target threshold be $t_a = (-2.5, 0.7)$ shown in Fig. 2b. Initially,

² For completeness, the detailed methods are included in the appendix of [26].

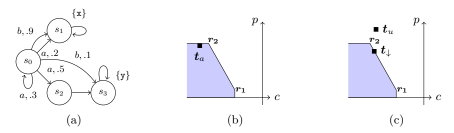


Fig. 2: Example MOMDP agent and corresponding execution of Algorithm 1.

choose $\boldsymbol{w}=(1,0)$ and corresponds to $\boldsymbol{r}_1=(-1.1,0.1)$ $|\boldsymbol{t}_{\downarrow}-\boldsymbol{t}_{\uparrow}||=.6$ so we do another iteration. Find $\boldsymbol{w}=(0.4,0.6)$ which corresponds to $\boldsymbol{r}_2=(-2.1,.71)$, and \boldsymbol{t}_a is contained in $down(\{\boldsymbol{r}_1,\boldsymbol{r}_2\})$ and the algorithm terminates. Fig. 2c shows the unachievable case. Let $\boldsymbol{t}_u=(1.8,.9)$. Similar to before we find \boldsymbol{r}_1 with $\boldsymbol{w}=(1,0)$. On finding \boldsymbol{r}_2 where $\boldsymbol{w}=(0.4,0.6)$, $\boldsymbol{w}\cdot\boldsymbol{r}_2<\boldsymbol{w}\cdot\boldsymbol{t}_u$. We find a new threshold $\boldsymbol{t}_{\downarrow}=(-1.97,0.61)$ satisfying the problem in Line 10 shown in Fig. 2c. The distance $|\boldsymbol{t}_{\downarrow}-\boldsymbol{t}_{\uparrow}||<\varepsilon$ and the algorithm terminates.

4 Hybrid GPU-CPU Implementation

In modern systems, GPU and multi-core CPU hardware is readily available. We developed an implementation for our MORAP framework, which utilises heterogeneous GPU and multi-core CPU resources to accelerate the computation. The acceleration is based on non-shared data within the two probabilistic model checking loops in Algorithm 2, which takes up the majority of run time for Algorithm 1 in practice. Parallel execution on GPUs and CPUs is by allocating models to each available GPU device and CPU core. For GPUs, further (massive) parallelisation can be achieved on the low-level matrix operations for probabilistic model checking.

Goal. The main goal of the implementation is to maximise throughput and parallelism. Combination of multiple devices is a load balancing problem in which we can effectively schedule model checking problems to keep all devices optimally busy, and reduce run time. We say that computations run on GPUs are called device operations. A multi-core processor can leverage shared memory with negligible latency before computing. The main concern with parallelism on when using a multi-core processor is thread-blocking and context switching overhead which should be avoided. Moreover, because low level computations are sequential a processor's execution run time will correspond to the size of a models state space. Conversely, the major issue with computing on GPUs is data transfer between the host and the device.

Design. Fig. 3 shows the parallel architecture of our framework, and Table 1 explains the roles of thread types. There are k FIFO queues controlled by k threads for each GPU device, and a single FIFO queue and corresponding thread controlling the processor, designated manager threads. Queues are data structures responsible for holding a subset of models. The processor thread is

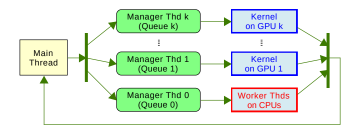


Fig. 3: Parallel architecture of MORAP framework.

Table 1: Component functionalities in MORAP implementation

Component	Functionality							
Main Thread	Running all computation except the two (inner) loops in							
	Alg. 2; generating and allocating models to queues for worker							
	threads and kernels							
Manager Thread	Managing the bounded FIFO queues; (one thread) spawning							
	CPU Worker Threads; (the other threads) calling GPU ker							
	nels, incl. copying data between the host memory and GPU							
	device memory; communicating with each other via a mes-							
	saging channel for load balancing							
Worker Thread	Computing the loops in Alg. 2; each thread bounded on one							
	CPU core and handling one model							
Kernel on GPU	Computing the loops in Alg. 2; each kernel running on one							
	GPU device and handling one model							

responsible for spawning worker threads whose count is the minimum of the available CPU-cores, or $\mathcal{M}_{i\otimes j}$ remaining in its queue. The worker threads are bound to CPUs while management threads are not. For a given worker, we require its bound CPU-core to be dedicated to computation, resulting in minimal response time and context switching. Management threads are not required to be bound to CPUs as program management is not demanding. We manage the computation workload scheduling through a work stealing [4] approach and bounded FIFO queues shown in Fig. 3. In the work stealing model, if the processor or a devices are idle and its queue is empty, then its thread can request (steal) a model $\mathcal{M}_{i\otimes j}$. In this way, the fastest processor gets the majority of the work and threads operate asynchronously of each other

Programming. The framework uses multiple languages including Rust (framework API), CUDA C (GPU device control), and a Python user interface. The API uses a foreign function interface (FFI) for Rust calls to C, and Python. We use the *affine* property [24] of the Rust [21] type system to ensure that *owned* variables can be used at most once in the application with move-only types. This is particularly useful in the parallel architecture, as $M_{i\otimes j}$ can be owned by at most one thread at a time, ensuring mutability of destruction of a model does not affect other computations. Isolating data access to each $\mathcal{M}_{i\otimes j}$ mitigates the requirement of shared memory access, freeing the framework from data races, or

data starving. Consequently the problem is *embarrassingly* parallel. Constructing the architecture in this way ensures that we approach the upper-bound of parallelism.

We use explicit data structures $P_{i,j}[(s,q,a),(s',q')] > 0$ for each $a \in A(s)$ and $R_{i,j}[(s,q,a),k]$, where k is an objective, for MOMDP transition functions and reward structures respectively. Low level matrix operations are managed on $device_k$ using the CUDA cuSPARSE API which is thread-safe. The action-value comparison requirement of finding an optimal policy in Line 2 can also be managed in parallel on $device_k$ with a single kernel launch involving a reduce operation. Parallelism on the device is based on the state space $|S \times Q|$ of $\mathcal{M}_{i \otimes j}$. Optimal occupancy for a GPU kernel is managed through a kernel launcher and a call to CUDA cudaOccupancyMaxPotentialBlockSize.

5 Experiments

One realistic example for our MORAP problem is a smart-warehousing or robotic mobile fulfilment system (RMFS), which usually controls tasks centrally with limited communication between robots [31]. The environment is a $W \times H$ two-dimensional grid typically consisting of movable racks (shelves), storage locations, and workstations where order picking and replenishment takes place [22]. Robots maneuver in the warehouse to carry out order picking, replenishment and other tasks. The state of robots is described by the robot position, the internal robot state (e.g., carrying a rack or not) and the environment parameters (e.g., the rack locations) and is discrete. Robots can perform such actions as Rotate Left/Right, Go Forward, Load/Unload Rack. The MORAP problem in this example is (random) assignment n tasks to n robots, and task planning for robots, under the multi-objective requirements of running costs and task fulfilment probabilities.

A replenishment task was modelled as co-safe LTL using φ_R below.

$$\varphi_R := \neg \mathtt{carry} \, \mathsf{U} \, \mathtt{rack}_{X_1,Y_1} \wedge \, \mathsf{X}(\mathtt{carry} \, \mathsf{U} \, \mathtt{feed} \, \wedge \, \mathsf{X}(\mathtt{carry} \, \mathsf{U} \, \mathtt{rack}_{X_2,Y_2} \, \wedge \, \mathsf{X} \, \neg \mathtt{carry}))$$

Picking tasks can be similarly modelled.

We conducted two experiments to evaluated our MORAP implementation in our smart warehousing example. **Experiment 1** included a scalability comparison of the decentralised model with the centralised model, and a run time comparison of hybrid GPU-CPU computation with (pure) GPU accelerated, multi-core CPU accelerated and non-accelerated computation. To benchmark our implementation, **Experiment 2** restricted our GPU accelerated and non-accelerated computation to a verification-only mode for the centralised model, and compared the performance with the Prism and Storm model checkers (which are not designed for task assignment in MAS)

All experiments were conducted on Debian with an AMD 2970WX 24 Core 3.0GHz Processor PCIe 3.0 32Gbps bandwidth, 3070Ti 1.77GHz 8Gb 6144 CUDA Cores GPU, and 32Gb of RAM. An artefact to reproduce the experiments is

available online³. A single GPU was used and therefore k=1 for the number of GPU management threads. Prism configuration included using explicit data structures, the Java heap size and hybrid maxmem were set to 32Gb to avoid memory exceptions. The default configuration was sufficient for Storm. The value of epsilon for value iteration was $\varepsilon_{VI}=10^{-6}$. Running time cut-off was set to 180 seconds, if the run time exceeds the cut-off time a timeout error was recorded. If the GPU device runs out of memory, a memory was recorded.

Experiment 1. The results for Experiment 1 are demonstrated in Table 2. When comparing the centralised and decentralised models, it can be observed that, in general, the run time performance of the decentralised model is improved over the centralised model. As expected, the centralised model run time scales exponentially for increasing state-spaces, while the scaling for the decentralised model is linear.

Table 2: This table evaluates the average run time for Lines 3-11 of Algorithm 1, for centralised and decentralised models, and an increasing number of agents, and an unachievable query. The number of tasks assigned are equal to the number of agents. Run times are in seconds. The decentralised model size parameter is measured using the reachable $|S| = \sum |S_i \times Q_j|$, note the number reachable states may be smaller than the total sum, and $|P| = \sum |\{P(s_i, q_j, a, s_i', q_j') > 0\}|$, for each $i \in I, j \in J$.

W.H.	agent- task num.	iter.	Decentralised						Centralised			
			Dec.		Time per iter.				Cent.		Time per iter.	
size			Model Size		Hybrid Mult.		GPU CPU		Model Size		CPU	GPU
Size			states	trans.	CPU			states trans.				
6×6	2	4	16k	104k	0.01	0.016	0.017	0.025	21.2K	136K	0.059	0.017
	5	4	106k	652k	0.036	0.035	0.2	0.36	3.5M	22.5M	6.1	2.35
	6	6	152K	940K	0.3	0.038	0.96	0.38	24.9M	162M	timeout	15.2
	50	7	10.6M	$65.3\mathrm{M}$	1.36	1.0	27.2	11.1	memerr	memerr	-	-
	100	9	42.4M	261M	4.8	3.9	timeout	31.9	-	-	-	-
	2	4	254k	1.5M	0.18	0.16	0.12	0.5	190k	1.2M	1.08	1.5
12×12	4	4	1.0M	6.1M	0.36	0.55	0.42	1.78	635k	4.2M	9.8	2.1
	6	7	2.2M	$13.8\mathrm{M}$	0.6	1.4	0.9	4.0	memerr	memerr	-	-
	8	6	4.1M	24.5M	1.1	2.5	1.6	7.2	-	-	-	-
	10	15	6.4M	38.3M	1.8	4.23	2.46	11.7		-	-	-
	20	7	25.4M	153M	6.5	17.3	9.8	45.7	-	-	-	-
	30	8	57.2M	$345\mathrm{M}$	15.3	38.8	22.1	timeout	-	-	-	-

Table 3: Evaluates the run time comparison for a centralised model

W.H Grid Size	Mode	l Size	Time per iter.				
	states	${\rm trans}$	CPU	GPU	Prism	Storm	
3×3	334	2.17k	1e-4	0.03	0.005	0.038	
6×6	4.2k	18.8k	0.004	0.038	0.025	0.058	
8×8	12.9k	78.5k	0.017	0.041	0.081	0.114	
10×10	30.9k	187k	0.048	0.046	0.17	0.33	
12×12	63.5k	383k	0.12	0.056	-	0.66	

³ https://github.com/tmrob2/hybrid-motap

Overall, Table 2 demonstrates that parallelism of some form achieves improved run time performance. For smaller warehouse sizes, i.e. 6×6 , multiple CPUs achieve an almost 10 times improvement over single-CPU. For larger models, warehouse size of 12×12, the hybrid GPU-CPU achieved a similar performance increase. This confirms that the best resource deployment depends on the size of the product model $\mathcal{M}_{i\otimes j}$ being verified. For example, the multi-CPU model checking significantly outperformed the GPU performance. To explain this hardware disparity, the assignment problem involves copying 10,000 small models to the GPU and exposes the PCI bus data transfer as a bottleneck. In other words, when the model size is small and there are many models, parallelism from the GPU does not outweigh the data transfer cost resulting in the GPU being less effective. Conversely, taking the case of the randomised task assignments involving 30 agents. Larger model verification approximately results in a 65% reduction in run times when using hybrid implementation over multiple CPUs. The major reason for this is that run time improvement on the GPU is at a maximum when the GPU occupancy is highest, meaning the GPU can consume more work in the device queues faster. Moreover, the model checking component of the run time as a ratio to model copying time is larger, meaning it is more effective to use the GPU.

The hybrid implementation is observed to achieve better scaling performance than either pure GPU or CPU data allocations. However, the hybrid allocation mechanism results in a 23-50% increase in run time for a large number of smaller models over purely using multiple CPUs. Therefore, while the hybrid implementation has better scaling, there is a clear threshold for large MAS with smaller model sizes.

Experiment 2. Prism and Storm comparisons were conducted using a centralised model regarding one agent and task. We excluded the model build times from run time comparisons. For all platforms, we examined the achievability query that agent cost did not exceed c_1 , and the probability of reaching **done** was greater than p_1 .

The comparison results of our Algorithm 1 against Prism and Storm multiobjective routines are presented in Table 3. Because of model translation incompatibility, we were unable to generate the 12×12 model for Prism in a reasonable time. These results demonstrate, for a standard problem, our algorithm has competitive multi-objective run time performance across a number of increasing model sizes. While initially the GPU only run time for Algorithm 1 suffers an overhead penalty, it scales better than the CPU only implementation.

6 Related Work

Multi-objective optimisation considers the domain of planning where objectives are conflicting, and we are often interested in global maximisation of all objectives. These problems are often the focus of multi-objective model checking [27, 11, 12, 14, 9, 16, 13]. Pareto optimal solutions using value iteration, and the synthesis of schedulers, are included in [12–14]. Section 3, in particular, elaborates

on the difference between our approach and existing multi-objective queries of [12]. Additionally, we reduce the number of computations required by finding an optimal task assignment using the Hungarian algorithm before progressing to verification of objectives.

GPU acceleration for MDP is studied in [7] and implements GPU acceleration for MOMDP. When compared with our implementation, [7] is limited to a specific problem, scalarises the multi-objective problem, and does not implement model checking for MAS. Moreover, there is no comparison against modern model checking frameworks for performance benchmarks, and does not handle co-safe LTL inputs. A parallel GPU accelerated sparse value iteration is presented in [28] which is similar to our implementation of value iteration within Line 2 including the reduce kernel operation for action comparison. However, key two differences are (1) we combine the sparse transition matrix dense value vector multiplication and rewards summation in one operation, reducing kernel overhead; and (2) the explicit data structure that we use organises the sparse matrix to exploit consecutive memory addresses for actions. Also, the work in [28] does not consider multi-objective value iteration, model checking properties, or MAS task assignment.

With similar reasoning to this paper, the approach of [10] reduces the redundant complexity in the multi-agent MDP [5] model for problems in which agents do not share task segments, only that an agent may optimally complete its allocated tasks concurrently. In contrast, we consider the classical randomised assignment problem for which agents may only work independently on a single task. Therefore, the model generated using [10] is not suitable for solving this problem as it has no way of keeping track of which agents have already been assigned a task. In contrast, our work presents a way of solving the randomised task assignment and constructs a decentralised model removing all transition redundancy and scales linearly.

7 Conclusion

In this paper we present an approach addressing the classical assignment problem maximising the multi-objective assignment reward for a set of tasks to a multi-agent system. We demonstrate that our problem is convex and can be solved in polynomial time, and we can synthesise optimal schedulers in a decentralised way. We provide a hybrid CPU-GPU multi-objective model checking framework which optimally manages the computational load on GPU devices and multiple CPU-cores. We conduct two experiments to show that decentralising the problem results in a parallel implementation which achieves significant run time improvements and linear scaling, and our multi-objective model checking performance is competitive with Prism and Storm model checkers. Future work consists of further optimisation of the implementation utilising CUDA streams to alleviate the PCI bottleneck for smaller models. We are also interested in ω -regular LTL long-running task assignments to a MAS where we have some way of injecting interrupts into the system.

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