Discovering dispatching rules from data using imitation learning

Case study for the job-shop problem

Helga Ingimundardottir · Thomas Philip Runarsson

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Abstract Dispatching rules can be automatically generated from scheduling data. This paper will demonstrate that the key to learning an effective dispatching rule is through the careful construction of the training data, $\{\mathbf{x}_i(k), y_i(k)\}_{k=1}^K \in$ \mathcal{D} , where: i) features of partially constructed schedules \mathbf{x}_i should necessarily reflect the induced data distribution \mathscr{D} for when the rule is applied. This is achieved by updating the learned model in an active imitation learning fashion; ii) y_i is labelled optimally using a MIP solver, and iii) data needs to be balanced, as the set is unbalanced with respect to the dispatching step k. Using the guidelines set by our framework the design of custom dispatching rules, for a particular scheduling application, will become more effective. In the study presented three different distributions of the job-shop will be considered. The machine learning approach considered is based on preference learning, i.e., which dispatch (post-decision state) is preferable to another.

Keywords Scheduling · Composite dispatching rules · Performance Analysis · Imitation Learning · DAgger · Preference Learning

H. Ingimundardottir

Dunhaga 5, IS-107 Reykjavik, Iceland

Tel.: +354-525-4704 Fax: +354-525-4632 E-mail: hei2@hi.is

T.P. Runarsson

Hjardarhagi 2-6, IS-107 Reykjavik, Iceland

Tel.: +354-525-4733 Fax: +354-525-4632 E-mail: tpr@hi.is

1 Introduction

Hand crafting heuristics for scheduling is an ad hoc approach to finding approximate solutions to problems. The practice is time-consuming and its performance can even vary dramatically between different problem instances. The aim of this work is to increase our understanding of this process. In particular, the learning of new problem specific priority dispatching rules (DR) will be addressed for a subclass of scheduling problems known as the job-shop scheduling problem (JSP).

A recent editorial the state-of-the-art approaches [6] in advanced dispatching rules for large-scale manufacturing systems reminds us that: "... most traditional dispatching rules are based on historical data. With the emergence of data mining and on-line analytic processing, dispatching rules can now take predictive information into account." The importance of automated discovery of dispatching rules was also emphasised by [24]. Data for learning can also be generated using a known heuristic on a set of problem instances. Such an approach is taken in [20] for a single machine where a decision tree is learned from the data to have similar logic to the guiding dispatching rule. However, the learned method cannot outperform the original dispatching rule used for the data generation. This drawback is confronted in [22, 35,25] by using an optimal scheduler or policy, computed off-line. The resulting dispatching rules, as decision trees, gave significantly better schedules than using popular heuristics in that field, and a lower worst-case factor from optimality. Although using optimal policies for creating training data gives vital information on how to learn good scheduling rules an experimental study will show that this is not sufficient. Once these rules make a suboptimal dispatch then they are in uncharted territory and the effects are relatively unknown. This work will illustrate the sensitivity of learned dispatching rule's performance in the way the training data is sampled. For this purpose, JSP is used as a case study to illustrate a methodology for generating meaningful training data, which can be successfully learned using preference-based imitation learning (IL).

The competing alternative to learning dispatching rules from data would be to search the dispatching rule space directly. The prevalent approach in this case would be using an evolutionary algorithm, such as genetic programming (GP). The predominant approach in hyper-heuristics is a framework for creating new heuristics from a set of predefined heuristics via genetic algorithm optimisation [3]. Adopting a two-stage hyper-heuristic approach to generate a set of machine-specific DRs for dynamic job-shop, [27] used genetic programming (GP) to evolve CDRs from basic features, along with an evolutionary algorithm to assign a CDR to a specific machine. The problem space consists of jobshops in semiconductor manufacturing, with additional shop constraints, as machines are grouped into similar work centres, which can have different set-up times, workloads, etc. In fact, the GP emphasised efficient dispatch at the work centres with set-up requirements and batching capabilities, which are rules that are non-trivial to determine manually.

With meta-heuristics existing DRs can be used, and for example portfolio-based algorithm selection [29,9,36], either based on a single instance or class of instances, to determine which DR to choose from. Implementing ant colony optimisation to select the best DR from a selection of nine DRs for JSP, experiments from [19] showed that the choice of DR does affect the results and for all performance measures considered. They showed that it was better to have all the DRs to choose from rather than just a single DR at a time. A simpler and more straightforward way to automate selection of composite priority dispatching rules (CDR), [13], translated dispatching rules into measurable features which describe the partial schedule and optimise directly what their contribution should be via evolutionary search.

Using case based reasoning for timetable scheduling, training data in [2] is guided by the two best heuristics in the literature. They point out that in order for their framework to be successful, problem features need to be sufficiently explanatory and training data needs to be selected carefully so they can suggest the appropriate solution for a specific range of new cases. When learning new dispatching rules there are several important factors to consider. First and foremost the context in which the training data is constructed will influence the quality of the learned dispatching rule [2]. Since the training data consists of a collection of features, the quality of training data is interchangeable with the predictability of features. The training data is necessarily also problem instance specific. In addition to addressing these aspects, the paper will show that during the scheduling process, the most critical moment to make the 'right' dispatch will vary. Furthermore, depending on the distribution of problem instances, these critical moments can vary greatly. Moreover, a supervised learning algorithm will optimize classification accuracy, while it is the actual end-performance of the dispatching rule learned that will determine the success of the learning method.

The outline of the paper is as follows, with Section 2 giving the mathematical formalities of the scheduling problem and Section 3 describing the main features for job-shop and illustrating how schedules are created with dispatching rules. Section 4 sets up the framework for learning from optimal schedules, in particular, the probability of choosing optimal decisions and the effects of making a suboptimal decision. Furthermore, the optimality of common single priority dispatching rules is investigated. With these guidelines presented, Section 5 goes into detail on how to create meaningful composite priority dispatching rules using preference learning, focusing on how to compare operations and collect training data with the importance of the sampling strategy applied. Sections 6 and 7 explain the trajectories for sampling meaningful schedules used in preference learning, either using passive or active imitation learning. Experimental results are jointly presented in Section 8 with comparison for a randomly generated problem space. Furthermore, some general adjustments for performance boost are also considered. Following a discussion in Section 9 the main conclusion and future work are presented in Section 10.

2 Job-shop Scheduling

JSP involves the scheduling of jobs for a set of machines. Each job consists of a number of operations which are then processed on the machines in a predetermined order. An optimal solution to the problem will depend on the specific objective.

This study will consider the $n \times m$ JSP, where n jobs, $\mathscr{J} = \{J_j\}_{j=1}^n$, are scheduled on a finite set, $\mathscr{M} = \{M_a\}_{a=1}^m$, of m machines. The index j refers to a job $J_j \in \mathscr{J}$ while the index a refers to a machine $M_a \in \mathscr{M}$. Each job requires a number of processing steps or operations, and the pair (j,a) refers to the operation, i.e., processing the task of job J_j on machine M_a .

Each job J_j has an indivisible operation time (or cost) on machine M_a , p_{ja} , which is assumed to be integral and finite. Starting time of job J_j on machine M_a is denoted $x_s(j,a)$ and its end time is denoted $x_e(j,a)$. Each job J_j has a specified processing order through the machines. It is a permutation vector, $\boldsymbol{\sigma}_j$, of $\{1,\ldots,m\}$, representing a job J_j which can be processed on $M_{\boldsymbol{\sigma}_j(a)}$ only after it has been completely processed on $M_{\boldsymbol{\sigma}_j(a-1)}$, namely:

$$x_s(j, \boldsymbol{\sigma}_j(a)) \ge x_e(j, \boldsymbol{\sigma}_j(a-1)) \tag{1}$$

for all $J_j \in \mathcal{J}$ and $a \in \{2,..,m\}$. Note, that each job can have its own distinctive flow pattern through the machines

which is independent of the other jobs. However, in the case that all jobs share the same fixed permutation route, it is referred to as flow-shop (FSP).

The objective function is to minimise the schedule's maximum completion times for all tasks, commonly referred to as the makespan, $C_{\rm max}$. This family of scheduling problems is denoted by $J||C_{\rm max}|$ [28]. Additional constraints commonly considered are job release-dates and due-dates or sequence dependent set-up times; however, these will not be considered here.

In order to find an optimal (or near optimal) solution for scheduling problems it is possible to use either exact methods or heuristics methods. Exact methods guarantee an optimal solution. However, job-shop scheduling is strongly NP-hard [8]. Any exact algorithm generally suffers from the curse of dimensionality, which impedes the application in finding the global optimum in a reasonable amount of time. Using state-of-the-art software for solving scheduling problems, such as LiSA (A Library of Scheduling Algorithms) [1], which includes a specialised version of branch and bound that manages to find optimums for job-shop problems of up to 14×14 [34]. However, problems that are of greater size become intractable. Heuristics are generally more time efficient but do not necessarily attain the global optimum. Therefore, job-shop has the reputation of being notoriously difficult to solve. As a result, it's been widely studied in deterministic scheduling theory and its class of problems has been tested on a plethora of different solution methodologies from various research fields [23], all from simple and straight forward dispatching rules to highly sophisticated frameworks.

3 Priority Dispatching Rules

Priority dispatching rules determine, from a list of incomplete jobs, \mathcal{L} , which job should be dispatched next. This process, where an example of a temporal partial schedule of six jobs scheduled on five machines, is illustrated in Fig. 1. The numbers in the boxes represent the job identification j. The width of the box illustrates the processing times for a given job for a particular machine M_a (on the vertical axis). The dashed boxes represent the resulting partial schedule for when a particular job is scheduled next. Moreover, the current C_{max} is denoted by a dotted vertical line. The object is to keep this value as small as possible once all operations are complete. As shown in the example there are 15 operations already scheduled. The *sequence*, χ , of dispatches used to create this partial schedule is:

$$\mathbf{\chi} = (J_3, J_3, J_3, J_4, J_4, J_5, J_1, J_1, J_2, J_4, J_6, J_4, J_5, J_3) \quad (2)$$

This refers to the sequential ordering of job dispatches to machines, i.e., (j,a); the collective set of allocated jobs to

machines is interpreted by its sequence, which is referred to as a schedule. A scheduling policy will pertain to the manner in which the sequence is determined from the available jobs to be scheduled. In our example, the available jobs are given by the job-list $\mathcal{L}^{(k)} = \{J_1, J_2, J_4, J_5, J_6\}$ with the five potential jobs to be dispatched at step k = 16 (note that J_3 is completed).

However, deciding which job to dispatch is not sufficient as one must also know where to place it. In order to build tight schedules, it is sensible to place a job as soon as it becomes available and such that the machine idle time is minimal, i.e., schedules are non-delay. There may also be a number of different options for such a placement. In Fig. 1 one observes that J_2 , to be scheduled on M_3 , could be placed immediately in a slot between J_3 and J_4 , or after J_4 on this machine. If J_6 had been placed earlier, a slot would have been created between it and J_4 , thus creating a third alternative, namely scheduling J_2 after J_6 . The time in which machine M_a is idle between consecutive jobs J_j and $J_{j'}$ is called idle time or slack:

$$s(a,j) := x_s(j,a) - x_e(j',a)$$
 (3)

where J_i is the immediate successor of $J_{i'}$ on M_a .

Construction heuristics are designed in such a way that it limits the search space in a logical manner while not excluding the optimum. Here, the construction heuristic, Υ , is to schedule the dispatches as closely together as possible, i.e., minimise the schedule's idle time. More specifically, once an operation (j,a) has been chosen from the job-list $\mathscr L$ by some dispatching rule, it can then be placed immediately after (but not prior) to $x_e(j,\sigma_j(a-1))$ on machine M_a due to constraint Eq. (1), this could be considered the release time for J_j . However, to guarantee the disjunctive condition (i.e. no overlapping of jobs on machines) is not violated, idle times M_a from Eq. (3) are inspected as they create a slot which J_j can occupy, but only considering those that occur after J_j has been released from its previous machine, namely:

$$\tilde{s}(a,j') := x_s(j'',a) - \max\{x_e(j',a), x_e(j,\sigma_i(a-1))\}$$
 (4)

for all already dispatched jobs, $J_{j'}, J_{j''} \in \mathscr{J}_a$ where $J_{j''}$ is $J_{j'}$ successor on M_a . Moreover, since preemption is not allowed, the only applicable slots for a new dispatch, \tilde{S}_{ja} , are those idle times that can process the entire operation, namely:

$$\tilde{S}_{ja} := \{ J_{i'} \in \mathscr{J}_a : \tilde{s}(a, j') \ge p_{ja} \}.$$
 (5)

The placement rule applied to decide which slot from Eq. (5) is used to place J_j is intrinsic to the construction heuristic, which is chosen independently of the priority dispatching rule that is applied. Different placement rules could be considered for selecting a slot, e.g., if the main concern were to

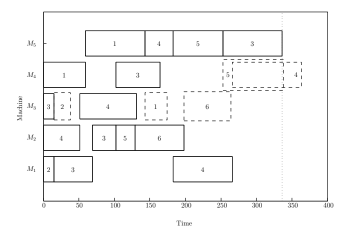


Fig. 1: Gantt chart of a partial JSP schedule after 15 dispatches: Solid and dashed boxes represent χ and $\mathcal{L}^{(16)}$, respectively. Current C_{max} denoted as dotted line.

utilise the slot space, then choosing the slot with the smallest idle time would yield a closer-fitted schedule and leave greater idle times undiminished for subsequent dispatches on M_a . In our experiments, cases were discovered where such a placement could rule out the possibility of constructing optimal solutions. However, this problem did not occur when jobs are simply placed as early as possible, which is beneficial for subsequent dispatches for J_j . For this reason, it will be the placement rule applied here.

Priority dispatching rules will use features of operations, such as processing time, in order to determine the job with the highest priority. Consider again Fig. 1; if the job with the shortest processing time (SPT) were to be scheduled next, then J_2 would be dispatched. Similarly, for the longest processing time (LPT) heuristic, J_5 would have the highest priority. Dispatching can also be based on features related to the partial schedule. Examples of these are dispatching the job with the most work remaining (MWR) or alternatively the least work remaining (LWR). A survey of more than 100 of such rules are presented in [26]. However, the reader is referred to an in-depth survey for simple or *single priority dispatching rule* (SDR) by [12]. The SDRs assign an index to each job in the job-list and is generally only based on a few features and simple mathematical operations.

Designing priority dispatching rules requires recognising the important features of the partial schedules needed to create a reasonable scheduling rule. These features attempt to grasp key attributes of the schedule being constructed. Which features are most important will necessarily depend on the objectives of the scheduling problem. The features used in this study applied for each possible operation encountered are given in Table 1, where the set of machines already dispatched for J_j is $\mathcal{M}_j \subset \mathcal{M}$, and similarly, M_a has already had the jobs $\mathcal{J}_a \subset \mathcal{J}$ previously dispatched. The features of particular interest were obtained by inspect-

Table 1: Feature space \mathscr{F} for JSP where job J_j on machine M_a given the resulting temporal schedule after operation (j,a).

φ	Feature description	Mathematical formulation								
	job-related									
ϕ_1	job processing time	p_{ja}								
ϕ_2	job start time	$x_s(j,a)$								
ϕ_3	job end time	$x_e(j,a)$								
ϕ_4	job arrival time	$x_e(j,a-1)$								
ϕ_5	time job had to wait	$x_s(j,a) - x_e(j,a-1)$								
ϕ_6	total processing time for job	$\sum_{a \in \mathcal{M}} p_{ja}$								
ϕ_7	total work remaining for job	$\sum_{a' \in \mathcal{M} \setminus \mathcal{M}_i} p_{ja'}$								
ϕ_8	number of assigned operations for job	$ \mathscr{M}_j $								
	machine-related									
ϕ_9	when machine is next free	$\max_{j' \in \mathscr{J}_a} \{x_e(j', a)\}$								
ϕ_{10}	total processing time for machine	$\sum_{i \in \mathscr{I}} p_{ja}$								
ϕ_{11}	total work remaining for machine	$\sum_{j' \in \mathscr{J} \setminus \mathscr{J}_a} p_{j'a}$								
ϕ_{12}	number of assigned operations for machine	$ \mathcal{J}_a $								
ϕ_{13}	change in idle time by assignment	$\Delta s(a,j)$								
ϕ_{14}	total idle time for machine	$\sum_{j' \in \mathscr{J}_a} s(a, j')$								
ϕ_{15}	total idle time for all machines	$\sum_{a' \in \mathscr{M}} \sum_{j' \in \mathscr{J}_{a'}} s(a', j')$								
ϕ_{16}	current makespan	$\max_{(j',a')\in\mathscr{J}\times\mathscr{M}_{j'}}\{x_f(j',a')\}$								

ing the aforementioned SDRs. Features ϕ_1 - ϕ_8 and ϕ_9 - ϕ_{16} are job-related and machine-related, respectively. In fact, [27] note that in the current literature, there is a lack of global perspective in the feature space, as omitting them won't address the possible negative impact an operation (j,a) might have on other machines at a later time. It is for this reason features such as ϕ_{13} - ϕ_{15} are considered, since they are slack related and are a means of indicating the current quality of the schedule.

Priority dispatching rules are attractive since they are relatively easy to implement, perform fast, and find reasonable schedules. In addition, they are relatively easy to interpret, which makes them desirable for the end-user. However, they can also fail unpredictably. A careful combination of dispatching rules has been shown to perform significantly better [16]. These are referred to as *composite priority dispatching rules* (CDR), where the priority ranking is an expression of several dispatching rules. CDRs deal with a greater number of more complicated functions and are constructed from the schedule features. In short, a CDR is a combination of several DRs. For instance, let π be a CDR comprised of d DRs, then the index I for $J_j \in \mathcal{L}^{(k)}$ using π is:

$$I_j^{\pi} = \sum_{i=1}^d w_i \pi_i(\boldsymbol{\chi}^j) \tag{6}$$

where $w_i > 0$ and $\sum_{i=0}^{d} w_i = 1$ with w_i giving the weight of the influence of π_i (which could be an SDR or another CDR) to π . Note: each π_i is a function of J_j 's features from the current sequence χ , where χ^j implies that J_j was the latest dispatch, i.e., the partial schedule given $\chi_k = J_j$.

At each step k, an operation is dispatched which has the highest priority. If there is a tie, some other priority measure

is used. Generally the dispatching rules are static during the entire scheduling process. However, ties could also be broken randomly (RND).

While investigating 11 SDRs for JSP, [21] a pool of 33 CDRs was created. This pool strongly outperformed the original CDRs by using multi-contextual functions based on either job waiting time or machine idle time (similar to ϕ_5 and ϕ_{14} in Table 1), i.e., the CDRs are a combination of either one or both of these key features and then the SDRs. However, no combinations of the basic SDRs were explored, only those two features. Similarly, using priority rules to combine 12 existing DRs from the literature, [37] had 48 CDR combinations which yielded 48 different models to implement and test. It is intuitive to get a boost in performance by introducing new CDRs, since where one DR might be failing, another could be excelling, so combining them together should yield a better CDR. However, these approaches introduce fairly ad hoc solutions and there is no guarantee the optimal combination of dispatching rules have been found.

The composite priority dispatching rule presented in Eq. (6) can be considered as a special case of the following general linear value function:

$$\pi(\boldsymbol{\chi}^j) = \sum_{i=1}^d w_i \phi_i(\boldsymbol{\chi}^j). \tag{7}$$

when $\pi_i = \phi_i$, i.e., a composite function of the features from Table 1. Finally, the job to be dispatched, J_{j^*} , corresponds to the one with the highest value, namely:

$$J_{j^*} = \underset{J_i \in \mathcal{L}}{\operatorname{argmax}} \ \pi(\boldsymbol{\chi}^j) \tag{8}$$

Similarly, single priority dispatching rules may be described by this linear model. For instance, let all $w_i = 0$, but with following exceptions: $w_1 = -1$ for SPT, $w_1 = +1$ for LPT, $w_7 = -1$ for LWR and $w_7 = +1$ for MWR. Generally, the weights \mathbf{w} are chosen by the designer or the rule apriori. A more attractive approach would be to learn these weights from problem examples directly. The following section will investigate how this may be accomplished.

4 Performance Analysis of Priority Dispatching Rules

In order to create successful dispatching rules, a good starting point is to investigate the properties of optimal solutions and hopefully be able to learn how to mimic the construction of such solutions. For this, optimal solutions (obtained by using a commercial software package [10]) are followed and the probability of SDRs being optimal is inspected. This serves as an indicator of how hard it is to put our objective up as a machine learning problem. However, the end goal, which is minimising deviation from optimality, ρ , must also

be taken into consideration because its relationship to stepwise optimality is not fully understood.

In this section the concerns of learning new priority dispatching rules will be addressed. At the same the time experimental set-up used in the study is described.

4.1 Problem Instances

The class of problem instances used in our studies was the job-shop scheduling problem described in Section 2. Each instance will have different processing times and machine ordering. Each instance will therefore create different challenges for a priority dispatching rule. Dispatching rules learned will be customised for the problems used for their training. For real world application using historical data would be most appropriate. The aim would be to learn a dispatching rule that works well on average for a given distribution of problem instances. To illustrate the performance difference of priority dispatching rules on different problem distributions within the same class of problems, consider the following three cases. Problem instances for JSP are generated stochastically by fixing the number of jobs and machines to ten. A discrete processing time is sampled independently from a discrete uniform distribution from the interval $I = [u_1, u_2]$, i.e., $\mathbf{p} \sim \mathcal{U}(u_1, u_2)$. The machine order was a random permutation of all of the machines in the job-shop. Two different processing times distributions were explored, namely $\mathscr{P}_{j.rnd}^{n \times m}$ where I = [1,99] and $\mathscr{P}_{j.rndn}^{n \times m}$ where I = [45,55]. These instances are referred to as random and random-narrow, respectively. In addition, the case where the machine order is fixed and the same for all jobs, i.e. $\sigma_j(a) = a$ for all $J_j \in \mathscr{J}$ and where $\mathbf{p} \sim \mathscr{U}(1,99)$, was also considered. These jobs are denoted by $\mathscr{P}_{f.rnd}^{n\times m}$ and are analogous to $\mathcal{P}_{j,rnd}^{n \times m}$. The problem spaces are summarised in

The goal is to minimise the makespan, C_{\max} . The optimum makespan is denoted $C_{\max}^{\pi_{\star}}$ (using the expert policy π_{\star}), and the makespan obtained from the scheduling policy π under inspection by C_{\max}^{π} . Since the optimal makespan varies between problem instances the performance measure is the following:

$$\rho = \frac{C_{\text{max}}^{\pi} - C_{\text{max}}^{\pi_{\star}}}{C_{\text{max}}^{\pi_{\star}}} \cdot 100\% \tag{9}$$

which indicates the percentage relative deviation from optimality. Note: Eq. (9) measures the discrepancy between predicted value and true outcome, and is commonly referred to as a loss function, which should be minimised for policy π .

Figure 2 depicts the box plot for Eq. (9) when using the SDRs from Section 3 for all of the problem spaces from Table 2. These box plots show the difference in performance of the various SDRs. The rule MWR performs on average the

Table 2: Problem space distributions used in experimental studies. Note, problem instances are synthetic and each problem space is i.i.d.

name	size $(n \times m)$	$N_{\rm train}$	N _{test}	note		
$\mathscr{P}_{j.rnd}^{10 \times 10}$	10×10	300	200	random		
$\mathscr{P}_{j.rndn}^{10 imes 10}$	10×10	300	200	random-narrow		
$\mathscr{P}_{f.rnd}^{10 imes 10}$	10×10	300	200	random		

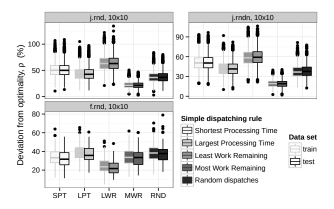


Fig. 2: Box plot for deviation from optimality, ρ , (%) for SDRs

best on the $\mathscr{P}_{j.rnd}^{n\times m}$ and $\mathscr{P}_{j.rndn}^{n\times m}$ problem instances, whereas for $\mathscr{P}_{f.rnd}^{n\times m}$ it is LWR that performs best. It is also interesting to observe that all but MWR perform statistically worse than a random job dispatching on the $\mathscr{P}_{j.rnd}^{n\times m}$ and $\mathscr{P}_{j.rndn}^{n\times m}$ problem instances.

4.2 Reconstructing optimal solutions

When building a complete schedule, $K = n \cdot m$ dispatches must be made sequentially. A job is placed at the earliest available time slot for its next machine, whilst still responding to the fact that each machine can handle at most one job at each time, and jobs need to have been finished by their previous machines according to their machine order. Unfinished jobs from the job list \mathcal{L} are dispatched one at a time according to a deterministic scheduling policy (or heuristic). This process is given as a pseudo-code in Algorithm 1. After each dispatch the schedule's current features are updated based on the half-finished schedule, χ . For each possible post-decision state, $J_j \in \mathcal{L}^{(k)}$, the temporal features, ϕ_j , are collected (cf. Line 5) forming the feature set, Φ , based on

Algorithm 1 Pseudo code for constructing a JSP sequence using a deterministic scheduling policy rule, π , for a fixed construction heuristic, Υ .

```
1: procedure ScheduleJSP(\pi, \Upsilon)
                                                    ▷ initial current dispatching sequence
2:
3:
            for k \leftarrow 1 to K = n \cdot m do

    b at each dispatch iteration

                  for all J_j \in \mathcal{L}^{(k)} \subset \mathscr{J} do
 4:
                                                                                      5:
                        \boldsymbol{\phi}^{j} \leftarrow \boldsymbol{\phi} \circ \Upsilon \left( \boldsymbol{\chi}^{j} \right)
                                                                       \triangleright temporal features for J_i
 6:
                                                                                       \triangleright priority for J_i
 7:
 8:
                       \leftarrow \operatorname{argmax}_{j \in \mathcal{L}^{(k)}} \{ I_j^{\pi} \}
                                                                         9:
                                                                                            \triangleright dispatch j^*
10:
            \mathbf{return}\; C^{\pi}_{\max} \leftarrow \Upsilon(\pmb{\chi})
                                                               ⊳ makespan and final schedule
12: end procedure
```

the union of all N_{train} problem instances available, namely:

$$\Phi := \bigcup_{\substack{\{\mathbf{x}_i\}_{i=1}^{N_{\text{train}}}}} \left\{ \phi^j : J_j \in \mathcal{L}^{(k)} \right\}_{k=1}^K \subset \mathscr{F}$$
 (10)

where the feature space \mathscr{F} is described in Table 1, and are based on job and machine features which are widespread in practice.

It is easy to see that the sequence of task assignments is by no means unique. Inspecting a partial schedule further along in the dispatching process such as in Fig. 1, then let's say J_1 would be dispatched next, and in the next iteration J_2 . Now this sequence would yield the same schedule as if J_2 had been dispatched first and then J_1 in the next iteration, i.e., these are jobs with non-conflicting machines. In this particular scenario, one cannot infer that choosing J_1 is better and J_2 is worse (or vice versa) since they can both yield the same solution. Furthermore, there may be multiple optimal solutions to the same problem instance. Hence not only is the sequence representation 'flawed' in the sense that slight permutations on the sequence are in fact equivalent in terms of the end-result, but very varying permutations on the dispatching sequence (although given the same partial initial sequence) can result in very different complete schedules yet can still achieve the same makespan.

The redundancy in building optimal solutions using dispatching rules means that many different dispatches may yield an optimal solution to the problem instance. Let's formalise the probability of optimality (or stepwise classification accuracy) for a given policy π , yet following the expert policy π_* , as:

$$\xi_{\pi}^{\star} := \mathbb{E}_{\pi_{\star}} \{ \pi_{\star} = \pi \} \tag{11}$$

that is to say, the mean likelihood of our policy π being equivalent to the expert policy π_* . The probability that a job chosen by a SDR yields an optimal makespan on a step-by-step basis, i.e., $\xi_{\langle \text{SDR} \rangle}^*$, is depicted in Fig. 3. These probabilities vary quite a bit between the different problem instance

¹ Dispatch and time step are used interchangeably.

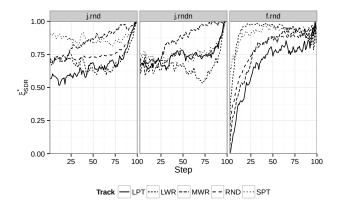


Fig. 3: Probability of SDR being optimal, $\xi_{\langle \text{SDR} \rangle}^{\star}$

distributions studied. From Fig. 3 it is observed that ξ_{MWR}^{\star} has a higher probability than random guessing, in choosing a dispatch which may result in an optimal schedule. This is especially true towards the end of the schedule building process. Similarly, ξ_{LWR}^{\star} selects dispatches resulting in optimal schedules with a higher probability. This would appear to support the idea that the higher the probability of dispatching job that may lead to an optimal schedule, the better the SDRs performance, as illustrated by Fig. 2. However, there is a counter example; ξ_{SPT}^{\star} has a higher probability than random dispatching of selecting a jobs that may lead to an optimal solution. Nevertheless, the random dispatching performs better than SPT on problem instances $\mathcal{P}_{j,rnd}^{10\times 10}$ and $\mathcal{P}_{j,rnd}^{10\times 10}$.

As shown in Fig. 3, $\mathcal{P}_{j.rnd}^{10\times10}$ has a relatively high probability (70% and above) of selecting an optimal job at random. However, it is imperative to keep making optimal decisions, because the consequences of making suboptimal dispatches are unknown. To demonstrate this Fig. 4 depicts mean worst and best case scenario of the resulting deviation from optimality, ρ , once off the optimal track, defined as follows:

$$\zeta_{\min}^{\star}(k) := \mathbb{E}_{\pi_{\star}} \left\{ \min_{J_{j} \in \mathscr{L}^{(k)}} (\rho) : \forall C_{\max}^{\chi^{j}} \geqslant C_{\max}^{\pi_{\star}} \right\}$$
 (12a)

$$\zeta_{\max}^{\star}(k) := \mathbb{E}_{\pi_{\star}} \left\{ \max_{J_{j} \in \mathscr{L}^{(k)}} (\rho) : \forall C_{\max}^{\mathbf{z}^{j}} \geqslant C_{\max}^{\pi_{\star}} \right\}$$
 (12b)

Note, that it is a given that there is only one non-optimal dispatch. Generally, there will be more, and then the compound effects of making suboptimal decisions are cumulative.

It is interesting to observe that for $\mathscr{P}_{j.rnd}^{10\times 10}$ and $\mathscr{P}_{j.rndn}^{10\times 10}$ making suboptimal decisions later impacts on the resulting makespan more than making a mistake early. The opposite seems to be the case for $\mathscr{P}_{f.rnd}^{10\times 10}$. In this case it is imperative to make good decisions right from the start. This is due to the major structural differences between JSP and FSP,

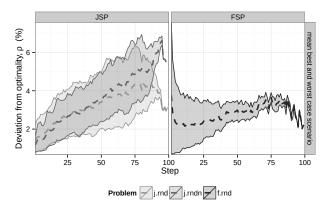


Fig. 4: Mean deviation from optimality, ρ , (%), for best and worst case scenarios of making one suboptimal dispatch (i.e. ζ_{\min}^{\star} and ζ_{\max}^{\star}), depicted as lower and upper bounds, respectively, for $\mathcal{P}_{j,rnd}^{10\times10}$, $\mathcal{P}_{j,rndn}^{10\times10}$ and $\mathcal{P}_{f,rnd}^{10\times10}$. The mean suboptimal move is given as a dashed line.

namely the latter having a homogeneous machine ordering and therefore constricting the solution immensely.

4.3 Blended dispatching rules

A naive approach to creating a simple blended dispatching rule (BDR) would be to switch between SDRs at a predetermined time. Observing again Fig. 3, a presumably good BDR for $\mathscr{P}_{j,rnd}^{10\times 10}$ would be to start with ξ_{SPT}^{\star} and then switch over to ξ_{MWR}^{\star} at around time step k = 40, where the SDRs change places in outperforming one another. A box plot for ρ for the BDR compared with MWR and SPT is depicted in Fig. 5 and its main statistics are reported in Table 3. This simple swap between SDRs does outperform the SPT heuristic, yet doesn't manage to gain the performance edge of MWR. Using SPT downgrades the performance of MWR. A reason for this lack of performance of our proposed BDR is perhaps that by starting out with SPT in the beginning, it sets up the schedules in such a way that it's quite greedy and only takes into consideration jobs with the shortest immediate processing times. Now, even though it is possible to find optimal schedules from this scenario, as Fig. 3 shows, the inherent structure that's already taking place might make it hard to come across by simple methods. Therefore, it is by no means guaranteed that by simply swapping over to MWR will handle the situation that applying SPT has already created. Figure 5 does show, however, that by applying MWR instead of SPT in the latter stages does help the schedule to be more compact in terms of SPT. However, the fact remains that the schedules have diverged too far from what MWR would have been able to achieve on its own.

SDR #1	SDR #2	k	Set	Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
SPT	_	K	train	20.38	41.15	50.70	51.31	59.18	94.20
SPT	_	K	test	22.75	41.39	49.53	50.52	58.60	93.03
MWR	_	K	train	4.42	17.84	21.74	22.13	26.00	47.78
MWR	_	K	test	3.37	17.07	21.39	21.65	25.98	41.80
SPT	MWR	10	train	5.54	17.98	21.75	21.99	25.43	44.02
SPT	MWR	10	test	5.87	17.29	20.78	21.28	24.67	44.47
SPT	MWR	15	train	4.76	18.24	22.04	22.49	26.65	49.86
SPT	MWR	15	test	7.42	17.60	21.38	21.83	25.45	45.98
SPT	MWR	20	train	5.76	18.98	22.46	23.01	26.97	41.59
SPT	MWR	20	test	8.31	18.64	22.92	23.29	27.10	49.93
SPT	MWR	30	train	9.77	20.89	25.60	25.76	30.01	50.94
SPT	MWR	30	test	4.39	21.20	26.08	26.25	30.58	49.88
SPT	MWR	40	train	13.04	23.42	28.12	28.94	33.67	54.98
SPT	MWR	40	test	8 55	24 20	28 16	28 98	33.20	57.21

Table 3: Main statistics for $\mathscr{P}_{j,rnd}^{10\times 10}$ deviation from optimality, ρ , using BDR that changes from SDR at a fixed time step k.

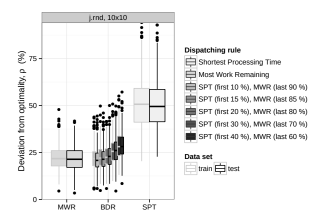


Fig. 5: Box plot for deviation from optimality, ρ , (%) for BDR where SPT is applied for the first 10%, 15%, 20%, 30% or 40% of the dispatches, followed by MWR

In Fig. 3 the stepwise optimality was inspected, given that all committed dispatches were based on the optimal trajectory. As mistakes are bound to be made at some points, it is interesting to see how the stepwise optimality evolves for its intended trajectory, π , thereby updating Eq. (11) to:

$$\xi_{\pi} := \mathbb{E}_{\pi} \left\{ \pi_{\star} = \pi \right\} \tag{13}$$

Figure 6 shows the log likelihood for $\xi_{\langle SDR \rangle}$ using $\mathscr{P}_{j.rnd}^{10\times 10}$. There one can see that even though ξ_{SPT} is generally more likely to find optimal dispatches in the initial steps, then shortly after k=15, ξ_{MWR} becomes a contender again. This could explain why our BDR switch at k=40 from Fig. 5 was unsuccessful. However, changing to MWR at $k\leq 20$ is not statistically significant from MWR (the boost in mean ρ is at most -0.5%). But as pointed out for Fig. 4, it's not so fatal to make bad moves in the very first dispatches for $\mathscr{P}_{j.rnd}^{10\times 10}$, hence there is little gain with improved classification accuracy in that region. However, after k>20 then the BDR performance starts diverging from that of MWR.

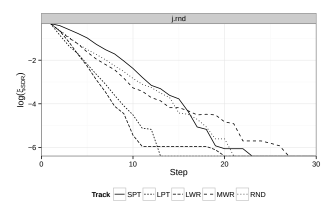


Fig. 6: Log likelihood of SDR being optimal for $\mathscr{D}_{j.rnd}^{10\times 10}$ when following its corresponding SDR trajectory, i.e., $\log\left(\xi_{\langle \mathrm{SDR}\rangle}\right)$

5 Preference Learning

Section 4.3 demonstrated there is something to be gained by trying out different combinations of DRs; however, it is non-trivial. This section presents one approach to learning how such combinations can work. Learning models considered in this study are based on ordinal regression in which the learning task is formulated as learning preferences, and in the case of scheduling, learning which operations are preferred to others. Ordinal regression has been previously presented in [33] and in [14] for JSP, and given here for completeness.

The optimum makespan is known for each problem instance. At each time step k, a number of feature pairs are created. Let $\phi^o \in \mathscr{F}$ denote the post-decision state when dispatching $J_o \in \mathscr{O}^{(k)}$ corresponds to an optimal schedule being built. All post-decision states corresponding to suboptimal dispatches, $J_s \in \mathscr{S}^{(k)}$, are denoted by $\phi^s \in \mathscr{F}$.

The approach taken here is to verify analytically, at each time step, by fixing the current temporal schedule as an initial state, whether it is possible to somehow yield an optimal schedule by manipulating the remainder of the sequence. This also takes care of the scenario that having dispatched a job resulting in a different temporal makespan would have resulted in the same final makespan if another optimal dispatching sequence had been chosen. That is to say, the training data generation takes into consideration when there are multiple optimal solutions² to the same problem instance.

Let's compare features from Eq. (10), and define the difference of optimal from suboptimal ones as, $\psi^o = \phi^o - \phi^s$, and vice versa, $\psi^s = \phi^s - \phi^o$, and label the differences by $y_o = +1$ and $y_s = -1$ respectively. Then, the preference learning problem is specified by a set of preference pairs:

$$\Psi = \left\{ (\boldsymbol{\psi}^{o}, y_{o}), (\boldsymbol{\psi}^{s}, y_{s}) : \forall (J_{o}, J_{s}) \in \mathcal{O}^{(k)} \times \mathcal{S}^{(k)} \right\}_{k=1}^{K}$$

$$\subset \Phi \times Y$$
(14)

where $\Phi \subset \mathbb{R}^d$ is the training set of d = 16 features (cf. Table 1), $Y = \{+1, -1\}$ is the outcome space from job pairs $J_o \in \mathcal{O}^{(k)}$ and $J_s \in \mathcal{S}^{(k)}$, for all dispatch steps k.

To summarise, each job is compared against another job of the job list, $\mathcal{L}^{(k)}$, and if the makespan differs, i.e., $C_{\max}^{\pi_{\star}(\boldsymbol{\mathcal{X}}^{s)}} \geq C_{\max}^{\pi_{\star}(\boldsymbol{\mathcal{X}}^{o})}$ an optimal/suboptimal pair is created. However, if the makespans are identical the pair is omitted since they give the same optimal makespan. In this way, only features from a dispatch resulting in a suboptimal solution is labelled undesirable.

Now let's consider mappings from solutions to ranks, such function π induces an ordering on the solutions by the following rule:

$$\boldsymbol{\chi}^i \succ \boldsymbol{\chi}^j \iff \pi(\boldsymbol{\chi}^i) > \pi(\boldsymbol{\chi}^j)$$
 (15)

where the symbol \succ denotes *is preferred to* with respect to the solutions' ranks. Referring to the ranks in a discrete manner as follows:

$$r_1 \succ r_2 \succ \cdots \succ r_{n'} \quad (n' \le n)$$
 (16)

implies r_1 is preferable to r_2 , and r_2 is preferable to r_3 , etc.

The function used to bring about preference is a linear function in the feature space, previously defined in Eq. (7). Logistic regression learns the optimal parameters $\mathbf{w}^* \in \mathbb{R}^d$. For this study, L2-regularised logistic regression from the LIBLINEAR package [7] without bias is used to learn the preference set Ψ , defined by Eq. (14). Hence the job chosen to be dispatched, J_{j^*} , is the one corresponding to the highest

preference estimate, i.e., Eq. (8) where π is the classification model obtained by the preference set.

Preliminary experiments for creating step-by-step model was done in [14] resulting in a local linear model for each dispatch for a total of K linear models for solving $n \times m$ JSP. However, the experiments showed that by fixing the weights to each mean value throughout the dispatching sequence the results remained satisfactory. A more sophisticated way would be to create a new linear model, where the preference set, Ψ , is the union of the preference pairs across the K dispatches, as described in Eq. (14). This would amount to a substantial preference set, and for Ψ to be computationally feasible to learn, Ψ has to be reduced. For this several ranking strategies were explored in [15], and the results showed that it is sufficient to use partial subsequent rankings with combinations of r_i and r_{i+1} for $i \in \{1, ..., n'\}$, added to the preference set in such a manner that there is more than one operation with the same ranking and only one from that rank is needed to be compared to the subsequent rank. Moreover, for this study, which dealt with 10×10 problem instances instead of 6×5 , the partial subsequent ranking became necessary, as full ranking was computationally infeasible due to its size. Defining the size of the preference set as $l = |\Psi|$, then if l is too large, re-sampling to size l_{max} may be needed in order for the ordinal regression to be computationally feasible.

The training data from [14] was created from optimal solutions of randomly generated problem instances, i.e., traditional passive imitation learning (PIL). As JSP is a sequential decision making process, errors are bound to emerge. Due to the compound effect of making suboptimal dispatches, the model leads the schedule astray from learned feature-space, resulting in the new input being foreign to the learned model. Alternatively, training data could be generated using suboptimal solution trajectories as well, as was done in [15], where the training data was also incorporated following the trajectories obtained by applying successful SDRs from the literature. The reasoning behind it was that they would be beneficial for learning, as they might help the model to escape from local minima once off the coveted optimal path. Simply aggregating training data obtained by following the trajectories of well-known SDRs yielded better models with lower deviation from optimality, ρ .

Inspired by the work of [30,31], the methodology of generating training data will now be such that it will iteratively improve upon the model, such that the feature-space learned will be representative of the feature-space the eventual model would likely encounter, known as DAgger for active imitation learning (AIL) and thereby, eliminating the ad hoc nature of choosing trajectories to learn, by instead letting the model lead its own way in a self-perpetuating manner until it converges.

 $^{^2}$ There can be several optimal solutions available for each problem instance. However, it is deemed sufficient to inspect only one optimal trajectory per problem instance as there are $N_{\rm train}=300$ independent instances, which gives the training data variety.

Furthermore, in order to boost training accuracy, two strategies were explored:

Boost.1 increasing number of preferences used in training (i.e. varying $l_{\max} \leq |\Psi|$),

Boost.2 introducing more problem instances (denoted EXT in experimental setting).

Note, the following experimental studies will address Boost.2, whereas preliminary experiments for Boost.1 showed no statistical significance in boost of performance. Hence, the default set-up will be $l_{\rm max} = 5 \cdot 10^5$ which is roughly the amount of features encountered from one pass of sampling a K-stepped trajectory using a fixed policy π for the default $N_{\rm train} = 300$.

Another way to adjust training accuracy is to give different weights to various time steps. To address this problem, two different stepwise sampling biases (or data balancing techniques) are here considered:

Bias.1 (equal) where each time step has equal probability. This was used in [13,15] and serves as a baseline.

Bias.2 (adjdbl2nd) where each time step is adjusted to the number of preference pairs for that particular step (i.e. each step now has equal probability irrespective of quantity of encountered features). This is done with re-sampling. In addition, there is superimposed twice as much likelihood of choosing pairs from the latter half of the dispatching process.

Remark: as the following sections require repeated collection of training data and since labelling is a very time intensive task, the remainder of the paper will solely focus on $\mathcal{P}_{j.rnd}^{10\times10}$. The experiments in [15] generally took several hours to collect for $N_{\rm train}^{6\times5}=500$. Going to higher dimension and using $N_{\rm train}^{10\times10}=300$ computational intensity becomes an issue, as $\mathcal{P}_{j.rnd}^{10\times10}$ needs a few days and $\mathcal{P}_{j.rndn}^{10\times10}$ or $\mathcal{P}_{f.rnd}^{10\times10}$ require several weeks using a personal computer.

6 Passive Imitation Learning

Using the terms from game theory used in [4], then our problem is a basic version of the sequential prediction problem where the predictor (or forecaster), π , observes each element of a sequence χ of jobs, where at each time step $k \in \{1,...,K\}$, before the k-th job of the sequence is revealed, the predictor guesses its value χ_k on the basis of the previous k-1 observations.

6.1 Prediction with Expert Advice

Let us assume the expert policy π_* is known, which can query what is the optimal choice of $\chi_k = j^*$ at any given

Algorithm 2 Pseudo code for choosing job J_{j^*} following a perturbed leader.

```
Require: Ranking r_1 \succ r_2 \succ \cdots > r_{n'} (n' \le n) of \mathcal{L}
                                                                                          \triangleright query \pi_{\star}
 1: procedure PERTURBEDLEADER(\mathcal{L}, \pi_{\star})
                                                                               ▷ likelihood factor
           \varepsilon \leftarrow 0.1
 3:
            p \leftarrow \mathcal{U}(0,1) \in [0,1]

    b uniform probability

 4:
            \mathscr{O} \leftarrow \left\{ j \in \mathscr{L} : r_j = r_1 \right\}
                                                                                ⊳ optimal job-list
 5:
            \mathscr{S} \leftarrow \left\{ j \in \mathscr{L} : r_j > r_1 \right\}
                                                                          ⊳ sub-optimal job-list
 6:
           if p < \varepsilon and n' > 1 then
 7:
                 return j^* \in \{j \in \mathscr{S} : r_j = r_2\}
                                                                          ⊳ any second best job
 8:
 9:
                 return j^* \in \mathcal{O}
                                                                                ⊳ any optimal job
10:
            end if
11: end procedure
```

time step k. Now let's use Eq. (8) to back-propagate the relationship between post-decision states and $\hat{\pi}$ with preference learning via our collected feature set, denoted Φ^{OPT} , i.e., collecting the feature set corresponding to the following optimal tasks J_{j^*} from π_{\star} in Algorithm 1. This baseline sampling trajectory originally introduced in [14] for adding features to the feature set is a pure strategy where at each dispatch an optimal task is dispatched.

By querying the expert policy, π_{\star} , the ranking of the joblist, \mathcal{L} , satisfies Eq. (16). In this study, then it's known that the best rank $r_1 \propto C_{\max}^{\pi_{\star}}$, hence the optimal job-list is the following:

$$\mathscr{O} = \left\{ J_j : r_1 \propto \min_{J_j \in \mathscr{L}} C_{\max}^{\pi_{\star}(\boldsymbol{\chi}^j)} \right\}$$
 (17)

found by solving the current partial schedule to optimality using a MIP solver.

When $|\mathcal{O}^{(k)}| > 1$, there can be several trajectories worth exploring. However, only one is chosen at random. This is deemed sufficient as the number of problem instances, N_{train} , is relatively large.

6.2 Follow the Perturbed Leader

By allowing a predictor to randomise it's possible to achieve improved performance [4,11]. This is the inspiration for our next strategy called Follow the Perturbed Leader, denoted OPT ε . Its pseudo code is given in Algorithm 2 and describes how the expert policy (i.e. optimal trajectory) from Section 6.1 is subtly "perturbed" with $\varepsilon = 10\%$ likelihood, by choosing a job corresponding to the second best C_{max} instead of an optimal one with some small probability.

6.3 Experimental study

Results for Sections 6.1 and 6.2 using the $\mathscr{P}_{j,rnd}^{10\times10}$ box plot of deviation from optimality, ρ , is given in Fig. 7 and the main statistics are reported in Table 4. To address Boost.2,

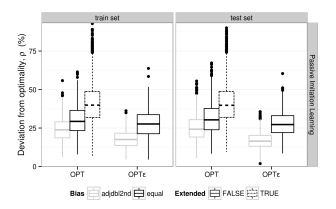


Fig. 7: Box plot for $\mathscr{P}_{j.rnd}^{10\times10}$ deviation from optimality, ρ , using either expert policy and following perturbed leader.

the extended training set was simply obtained by iterating over more examples, namely $N_{\text{train, EXT}}^{\text{OPT}} = 1000$. However, one can see that the increased number of varied features dissuades the preference models from achieving a good performance in terms of ρ . It's preferable to use the default $N_{\text{train}}^{\text{OPT}} = 300$ allowing slight perturbations of the optimal tra- $T_{\text{train}} = S_{\text{train}} = S_{\text$ head has not managed to surpass MWR in performance, except for $\Phi^{\text{OPT}\varepsilon}$ using Bias.2 with a $\Delta \rho \approx -4.24\%$ boost in mean performance. Otherwise, for Bias.1, there is a loss of $\Delta \rho \approx +6.23\%$ in mean performance. This is likely due to the fact that if equal probability is used for stepwise sampling, then there is hardly any emphasis given to the final dispatches as there are relatively few (compared to previous steps) preference pairs belonging to those final stages. Revisiting Fig. 4, then the band for $\{\zeta_{\min}^{\star}, \zeta_{\max}^{\star}\}$ is quite tight as the problem is immensely constricted and few operations to choose from. However, the empirical evidence from using Bias.2 shows that it is imperative to make the right decisions at the very end.

Based on the results from [14] the expert policy is a promising starting point. However, that was for 6×5 dimensionality (i.e. K = 30), which is a much simpler problem space. Notice that in Fig. 6 there was virtually no chance for $\xi_{\pi}(k)$ of choosing a job resulting in optimal makespan after step k = 28. Since job-shop is a sequential prediction problem, all future observations are dependent on previous operations. Therefore, learning sampled features that correspond only to optimal or near-optimal schedules isn't of much use when the preference model has diverged too far. Section 4.3 showed that good classification accuracy based on ξ_{π}^{\star} does not necessarily mean a low mean deviation from optimality, ρ . This is due to the learner's predictions affects future input observations during its execution, which violates the crucial i.i.d. assumptions of the learning approach, and ignoring this interaction leads to poor performance. In

fact, [30] proves that assuming the preference model has a training error of ε , then the total compound error (for all K dispatches), the classifier induces itself and grows quadratically, $O(\varepsilon K^2)$, for the entire schedule, rather than having linear loss, $O(\varepsilon K)$, as if it were i.i.d.

7 Active Imitation Learning

To amend performance from Φ^{OPT} -based models, suboptimal partial schedules were explored in [15] by inspecting the features from successful SDRs, $\Phi^{\langle \mathrm{SDR} \rangle}$, by passively observing a full execution of following the task chosen by the corresponding SDR. This required some trial-and-error as the experiments showed that features obtained by SDR trajectories were not equally useful for learning.

To automate this process, inspiration from AIL presented in [31] was sought, called Dataset Aggregation (DAgger) method, which addresses a no-regret algorithm in an on-line learning setting. The novel meta-algorithm for IL learns a deterministic policy guaranteed to perform well under its induced distribution of states. The method is closely related to Follow-the-leader (cf. Section 6), only with a more sophisticated leverage to the expert policy. In short, it entails the model π_i that queries an expert policy as in Section 6.1, π_{\star} , that it's trying to mimic, but also ensuring the learned model updates itself in an iterative fashion, until it converges. The benefit of this approach is that the featurestates that are likely to occur in practice are also investigated and as such used to dissuade the model from making poor choices. In fact, the method queries the expert about the desired action at individual post-decision states which are both based on past queries, and the learner's interaction with the current environment.

DAgger has been proven successful in a variety of benchmarks [31,32], such as the video games Super Tux Kart and Super Mario Bros., handwriting recognition and autonomous navigation for large unmanned aerial vehicles. In all cases it greatly improved on traditional supervised IL approaches.

7.1 DAgger

The policy of AIL at iteration i > 0 is a mixed strategy given as follows:

$$\pi_i = \beta_i \pi_\star + (1 - \beta_i) \hat{\pi}_{i-1} \tag{18}$$

where π_{\star} is the expert policy and $\hat{\pi}_{i-1}$ is the learned model from the previous iteration. Note, for the initial iteration, i = 0, a pure strategy of π_{\star} is followed. Hence, $\hat{\pi}_0$ corresponds to the preference model from Section 6.1 (i.e. $\Phi^{\text{IL}0} = \Phi^{\text{OPT}}$).

Algorithm 3 Pseudo code for choosing job J_{j^*} using imitation learning (dependent on iteration i) to collect training set $\Phi^{\text{IL}i}$; either by following optimal trajectory, π_{\star} , or preference model from previous iterations, $\hat{\pi}_{i-1}$.

```
Require: i \ge 0
Require: Ranking r_1 \succ r_2 \succ \cdots > r_{n'} \ (n' \le n) of \mathscr{L}
                                                                                     \triangleright query \pi_{\perp}
 1: procedure ACTIVEIL(i, \hat{\pi}_{i-1}, \pi_{\star})
           p \leftarrow \mathcal{U}(0,1) \in [0,1]

    □ uniform probability

 3:
           if i > 0 then (unsupervised)
 4:
                \beta_i \leftarrow 0
                                                                 5:
           else (fixed supervision)
 6:
                                      ▷ always follow expert policy (i.e. optimal)
 7:
           end if
 8:
           if p > \beta_i then
                return j^* \leftarrow \operatorname{argmax}_{i \in \mathcal{L}} \{I_i^{\hat{\pi}_{i-1}}\} \triangleright best job based on \hat{\pi}_{i-1}
 9:
10:
                 \mathscr{O} \leftarrow \{j \in \mathscr{L} : r_j = r_1\}
11:
                                                                           ⊳ optimal job-list
                 return j^* \in \mathscr{O}
                                                                           ⊳ any optimal job
12:
13:
           end if
14: end procedure
```

Algorithm 4 DAgger: Dataset Aggregation for JSP

```
Require: T \ge 1
 1: procedure DAGGER(\pi_{\star}, \Phi^{OPT}, T)
 2:
             \Phi^{\text{IL}0} \leftarrow \Phi^{\text{OPT}}

    initialize dataset

             \hat{\pi}_0 \leftarrow \text{Train}(\Phi^{\text{IL}0}) \quad \triangleright \text{ initial model, equivalent to Section 6.1}
 3:
 4:
             for i \leftarrow 1 to T do

    b at each imitation learning iteration

  5:
                   Let \pi_i = \beta_i \pi_* + (1 - \beta_i) \hat{\pi}_{i-1}
 6:
                   Sample K-step tracks using \pi_i \triangleright \text{cf. ACTIVEIL}(i, \hat{\pi}_{i-1}, \pi_{\star})
 7:
                            = \{(s, \pi_{\star}(s))\} \triangleright \text{ visited states for } \pi_i \text{ and actions by } \pi_{\star}
                    \Phi^{\mathrm{DA}i} \leftarrow \Phi^{\mathrm{DA}i-1} \cup \Phi^{\mathrm{IL}i} \triangleright \text{aggregate datasets from Eq. (10)}
 8:
                    \hat{\pi}_{i+1} \leftarrow \text{TRAIN}(\Phi^{\text{DA}i}) \triangleright preference model from Eq. (7)
 9:
10:
             end for
11:
             return best \hat{\pi}_i on validation
                                                                             ⊳ best preference model
12: end procedure
```

Equation (18) shows that β_i controls the probability distribution of querying the expert policy π_{\star} instead of the previous imitation model, $\hat{\pi}_{i-1}$. The only requirement for $\{\beta_i\}_i^{\infty}$ according to [31] is that $\lim_{T\to\infty}\frac{1}{T}\sum_{i=0}^T\beta_i=0$ to guarantee finding a policy $\hat{\pi}_i$ that achieves ε surrogate loss under its own state distribution limit.

Algorithm 3 explains the pseudo code for how to collect partial training set, $\Phi^{\text{IL}i}$ for i-th iteration of AIL. Subsequently, the resulting preference model, $\hat{\pi}_i$, learns on the aggregated datasets from all previous iterations, its update procedure is detailed in Algorithm 4.

7.2 Results

Due to time constraints, only T=3 iterations will be inspected here. In addition, preliminary experiments using DAgger for JSP favoured a simple parameter-free version of β_i in Eq. (18). Namely, the mixed strategy for $\{\beta_i\}_{i=0}^T$

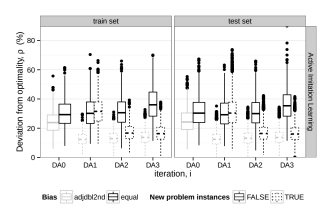


Fig. 8: Box plot for $\mathscr{P}_{j.rnd}^{10\times 10}$ deviation from optimality, ρ , using DAgger for JSP

was unsupervised with $\beta_i = I(i = 0)$, where *I* is the indicator function.³

Regarding Boost.2 strategy, Section 6 showed that adding new problem instances did not boost performance for the expert policy (which is equivalent for the initial iteration of DAgger). Hence, for active IL, the extended set now consists of each iteration encountering N_{train} new problem instances. This way, the extended training data is used sparingly, as labelling for each problem instance is computationally intensive. As a result, the computational budget for DAgger is the same regardless of whether or not new problem instances are used, i.e., $|\Phi^{\text{DA}i}| \approx |\Phi^{\text{DA}i}_{\text{EXT}}|$.

problem instances are used, i.e., $|\Phi^{\mathrm{DA}i}| \approx |\Phi^{\mathrm{DA}i}_{\mathrm{EXT}}|$. The results for $\mathscr{P}^{10\times 10}_{j.rnd}$ box plot of deviation from optimality, ρ , are given in Fig. 8 and the main statistics are reported in Table 4. As can be seen, DAgger is not fruitful when the same problem instances are continually used. This is due to the fact that there is not enough variance between $\Phi^{\text{IL}i}$ and $\Phi^{\text{IL}(i-1)}$, hence the aggregated feature set $\Phi^{\mathrm{DA}i}$ is only slightly perturbed with each iteration. Section 6.3 has showed this was not a very successful modification for the expert policy; even though it's noted that by introducing suboptimal feature-space the preference model is not as drastically bad as the extended optimal policy, even though $|\Phi^{\mathrm{DA}i}| \approx |\Phi^{\mathrm{OPT}}_{\mathrm{EXT}}|.$ However, when using new problem instances at each iteration, the feature set becomes varied enough that situations arise that can be learned to achieve a better represented classification problem which yields a lower mean deviation from optimality, ρ .

8 Summary of Imitation Learning

A summary of $\mathscr{P}_{j.rnd}^{10\times 10}$ best PIL and AIL models in terms of deviation from optimality, ρ , from Sections 6.3 and 7.2, respectively, are illustrated in Fig. 9, and the main statistics are

 $^{^{3} \}beta_{0} = 1 \text{ and } \beta_{i} = 0, \forall i > 0.$

given in Table 4. To summarise, the following trajectories were used: i) expert policy, trained on Φ^{OPT} ; ii) perturbed leader, trained on $\Phi^{\mathrm{OPT}\varepsilon}$, and iii) imitation learning, trained on $\Phi^{\mathrm{DA}i}_{\mathrm{EXT}}$ for iterations $i=\{1,\ldots,3\}$ using an extended training set. As a reference, the single priority dispatching rule MWR is shown at the edges of Fig. 9.

At first one can see that the perturbed leader ever-soslightly improves the mean for ρ , rather than using the baseline expert policy. However, AIL is by far the best improvement. With each iteration of DAgger, the models improve upon the previous iteration: i) for Bias.1 with Boost.2 then i = 1 starts with increasing $\Delta \rho \approx +1.39\%$. However, after that first iteration there is a performance boost of $\Delta \rho \approx -15.11\%$ after i = 2 and $\Delta \rho \approx -0.19\%$ for the final iteration i = 3, and ii) on the other hand when using Bias.2 with Boost.2, only one iteration is needed, as $\Delta \rho \approx -11.68$ for i = 1, and after that it stagnates with $\Delta \rho \approx +0.55\%$ for i=2 and for i=3 it is significantly worse than the previous iteration by $\Delta \rho \approx +0.75\%$. In both cases, DAgger outperforms MWR: i) after i = 3 iterations by $\Delta \rho \approx -5.31\%$ for Bias.1 with Boost.2, and ii) after i = 1 iteration by $\Delta \rho \approx -9.31\%$ for Bias.2 with Boost.2. Note, for Bias.1 without Boost.2, DAgger is unsuccessful, and the aggregated data set downgrades the performance of the previous iterations, making it best to learn solely on the initial expert policy for that model configuration.

Regarding Boost.2, it is not then successful for the expert policy as ρ increased approximately 10%. This could most likely be counteracted by increasing $l_{\rm max}$ to reflect the 700 additional examples. What is interesting, though, is that Boost.2 is well suited for AIL, using the same $l_{\rm max}$ as before. Note, the number of problems used for $N_{\rm train,EXT}^{\rm OPT}$ is equivalent to $T=2\frac{1}{3}$ iterations of extended DAgger. The new varied data give the aggregated feature set more information on what is important to learn in subsequent iterations, as those new feature-states are more likely to be encountered 'in practice.' Not only does the AIL converge faster, it also consistently improves with each iteration.

9 Discussion

Experiments in Section 6.3 show that following the optimal policy is not without its faults. There are many obstacles to consider in order to improve model configurations. When training the learning model, there is a trade-off between making the over-all best decisions (in terms of highest mean validation accuracy) versus making the right decision on crucial time points in the scheduling process, as Fig. 4 clearly illustrates. Moreover, before training the learning model, the preference set Ψ needs to be re-sampled to size $l_{\rm max}$. As the effects of making suboptimal choices varies as a function of time, the stepwise bias should rather take

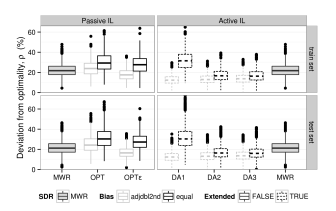


Fig. 9: Box plot for $\mathscr{P}_{j.rnd}^{10\times10}$ deviation from optimality, ρ , using either expert policy, DAgger or following perturbed leader strategies. MWR shown for reference.

into account the disproportional amount of features during the dispatching process. As the experimental studies in Sections 6.3, 7.2 and 8 showed, instead of equal probability (i.e. Bias.1) it was much more fruitful to adjust the set to its number of preference and doubling the emphasis on the second half (i.e. Bias.2). However, there are many other stepwise sampling strategies based on our analysis that could have been chosen instead, as here only a simplification of the trend from Fig. 4 was chosen. This also opens up the question of how validation accuracy should be measured, taking into account that the model is based on learning preferences, both based on optimal versus suboptimal, and then varying degrees of sub-optimality. Since ranks are only looked at in a black and white fashion, such that the makespans need to be strictly greater to belong to a higher rank, then it can be argued that some ranks should be grouped together if their makespans are sufficiently close. This would simplify the training set, making it (presumably) have fewer contradictions and be more appropriate for linear learning. Or simply the validation accuracy could be weighted in terms of the difference in makespan. During the dispatching process, there are some significant time points which need to be especially taken care of. Figure 4 shows how making suboptimal decisions is especially critical during the later stages for jobshop, whereas for flow-shop the earlier stages of dispatches are more critical.

Despite the information gathered by following an optimal trajectory, the knowledge obtained is not enough by itself. Since the learning model isn't perfect, it is bound to make a suboptimal dispatch eventually. When it does, the model is in uncharted territory as there is no certainty the samples already collected are able to explain the current situation. For this we propose investigating partial schedules from suboptimal trajectories as well, since future observations depend on previous predictions. A straight forward

Table 4: Main statistics for $\mathscr{P}_{j.rnd}^{10\times10}$ deviation from optimality, ρ , using either expert policy, imitation learning or following perturbed leader strategies.

π^a T^b	Bias	Set	$N_{\rm train}$	Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
OPT 0	adjdbl2nd	train	300	6.05	18.60	23.85	24.50	29.04	55.81
OPT 0	adjdbl2nd	test	300	5.56	19.16	24.24	25.19	30.42	55.52
OPT 0	equal	train	300	7.87	23.34	29.30	30.73	36.47	61.45
OPT 0	equal	test	300	8.31	23.88	30.32	31.46	37.70	67.24
DA1 1	adjdbl2nd	train	600	2.08	9.44	12.30	12.82	15.67	29.63
DA1 1	adjdbl2nd	test	300	0.00	9.22	12.39	12.73	15.85	35.17
DA1 1	equal	train	600	9.47	24.92	31.51	32.12	37.96	66.29
DA1 1	equal	test	300	4.77	23.77	30.34	31.40	37.81	73.73
DA2 2	adjdbl2nd	train	900	0.93	10.01	12.91	13.37	16.40	31.19
DA2 2	adjdbl2nd	test	300	0.39	9.84	13.13	13.44	16.62	34.57
DA2 2	equal	train	900	2.36	12.82	16.65	17.01	21.06	39.25
DA2 2	equal	test	300	1.72	12.57	16.38	16.89	20.66	42.44
DA3 3	adjdbl2nd	train	1200	0.93	10.45	13.71	14.12	17.15	32.91
DA3 3	adjdbl2nd	test	300	0.87	10.44	13.64	14.08	17.23	34.41
DA3 3	equal	train	1200	0.98	12.50	16.28	16.82	20.67	37.93
DA3 3	equal	test	300	0.26	12.32	16.01	16.52	20.22	41.62
ΟΡΤ ε 0	adjdbl2nd	train	300	4.64	13.63	17.56	18.07	21.66	36.25
OPT ε 0	adjdbl2nd	test	300	1.91	13.18	16.48	16.89	20.28	35.60
OPT ε 0	equal	train	300	4.52	21.31	27.63	28.04	33.69	63.74
ΟΡΤ ε 0	equal	test	300	8.54	22.03	27.26	27.94	33.02	60.38

^a For DAgger, then T = 0 is conventional expert policy (i.e. DA0 = OPT).

approach would be to inspect the trajectories of promising SDRs or CDRs. However, more information is gained when applying AIL inspired by the work of [30,31], such that the learned policy following an optimal trajectory is used to collect training data, and the learned model is iteratively updated. This can be done over several iterations, with the benefit being that the scheduling features that are likely to occur in practice are investigated and as such used to dissuade the model from making poor choices in the future.

The main drawback of DAgger is that it quite aggressively queries the expert, making it impractical for some problems, especially if it involves human experts. A way to confront that, [18,17] propose frameworks to minimise the expert's labelling effort. Or even circumvent the expert policy altogether by using a 'poorer' reference policy instead (i.e. π_{\star} in Eq. (18) is suboptimal) [5].

10 Conclusion

The single priority dispatching rules remain a popular approach to scheduling, as they are simple to implement and quite efficient. By inspecting optimal schedules, and investigating the probability that an optimal dispatch could be chosen by chance, and by looking at the impact of choosing suboptimal dispatches, some light is shed on how SDRs vary in performance. Furthermore, the problem instance space was varied, giving an even better understanding of the behaviour

of the SDRs. This analysis, however, also revealed that creating effective dispatching rules from data is by no means trivial.

Our analysis has shown that training data must be generated not only from optimal trajectories but also from the actual dispatching rule undergoing the training. The learning of effective dispatching rules is an iterative process with many calls to the expert (MIP solver). Techniques for reducing computationally expensive calls to the expert need to be addressed in the future. This study has been structured around the job-shop scheduling problem. However, it can easily be extended to other types of deterministic optimisation problems that involve sequential decision making. Applying the technique to other types of scheduling problems is of interest as future work.

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^b If T = 0 then passive imitation learning. Otherwise, for T > 0 it is considered active imitation learning.

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