Learning Probabilistic Action Models from Interpretation Transitions

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Abstract

There have been great advances in the probabilistic planning community during recent years, and planners can now provide solutions for very complex probabilistic tasks. However, planners require to have a model that represents the dynamics of the system, and in general these models are built by hand. In this paper, we present a framework to automatically infer probabilistic models from observations of the state transitions of a dynamic system. We propose an extension of previous works that perform learning from interpretation transitions. These works consider as input a set of state transitions and build a logic program that realizes the given transition relations. Here we extend this method to learn a compact set of probabilistic planning operators that capture probabilistic dynamics. Finally, we provide experimental validation of the quality of the learned models.

<u>KEYWORDS</u>:probabilistic planning operators, action model learning, probabilistic planning, inductive logic programming, learning from interpretation transitions

1 Introduction

Lately the performance of probabilistic task planners has improved to the point where they can find solutions in many complex scenarios (Kolobov et al. 2012; Keller and Eyerich 2012). Planners have been applied successfully in several fields, such as robotics (Chanel et al. 2014; Martínez et al. 2015) and aerospace (Surovik and Scheeres 2015). However, in most cases an expert is required to define the action model to be used by the planner. Creating a model is an error-prone task that requires in-depth knowledge about the problem and the planner, as well as extensive testing to assure the correctness of the model. We propose a new method to automatically learn action models based on a set of given state transitions.

Most previous approaches have only tackled the problem of learning deterministic action models (Zhuo et al. 2010), including uncertainty in the perceptions (Mourao et al.

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Fig. 1. Overview of the learning framework. The modules that interact with planning data and operators are in the blue rectangle on the left, while the modules related to logic programming are included inside the red rectangle on the right. The input of the method are grounded transitions, which are then converted to symbolic transitions to generalize between different objects. To obtain rule candidates, *LFIT* requires that the symbolic transitions are represented with propositional atoms. Finally, after rule candidates are obtained, they are transformed to planning operators to select the best set of operators.

2012). Regarding logic programming, abductive action learning has been studied based on abductive event calculus (Eshghi 1988), an abductive extension of event calculus, and has been extended for applications to planning, e.g., (Shanahan 2000). (Moyle 2003) uses an ILP technique to learn a causal theory based on event calculus (Kowalski and Sergot 1989), given examples of input-output relations. Probabilistic logic programs to maximize the probabilities of observations are learned by (Corapi et al. 2011; Sykes et al. 2013). They use parameter estimation to find the probabilities associated with each atom and rule, but it requires to code manually the restrictions for the initial set of rule candidates. Action models with uncertain effects have also been learned by (Rodrigues et al. 2012; Pasula et al. 2007) in STRIPS-like languages. Their optimization looks for planning operators that define as many effects as possible at once, which makes it a better choice for preconditions-effects based PPDDL domains (Younes and Littman 2004), such as the domains in past planning competitions (IPPC 2004-2008). In contrast our approach finds the best planning operators for each single action effect (i.e. rule head), which makes it a better choice to learn variable-based RDDL domains (Sanner 2010) used in newer planning competitions (IPPC 2011-2014).

Our approach learns an action model encoded as a set of planning operators. Each operator describes how the value of a predicate changes based on a set of preconditions given that an action is executed. We present a novel method to learn in two levels as shown in Fig. 1. On the right, the Learning From Interpretation Transitions (LFIT) framework (Inoue et al. 2014; Ribeiro and Inoue 2014) is used to generate rules. Given a set of state transitions of a system, it can learn a normal logic program that captures the system dynamics. The resulting rules are all candidates that have to be considered to select the best planning operators. On the left, the data is generalized between different objects by using a relational representation, and an optimization method is used to select the best subsets of planning operators that explain input transitions while maintaining generality. The dependency relations between the planning operators are used to efficiently select the best candidates. Our method is designed to learn RDDL-like (Younes and Littman 2004) operators, where each variable is updated separately based on a set of preconditions. To that end, in this paper we present the following novel improvements:

- The integration of logic programming to efficiently limit the number of rule candidates with a relational representation that provides better generalization, and an optimization method to select the best subsets of planning operators.
- The *LUST* algorithm: an extension of the *LFIT* framework to learn probabilistic dynamics from uncertain state transitions. The new algorithm that we propose can construct a model of a non-deterministic system by learning probabilistic rules. LUST also integrates multi-valued variables, which allows us to represent actions more efficiently since every transition has only one action of the many possible actions.
- An optimization method that makes use of the dependency relation between rules to select subsets of planning operators efficiently. It uses a score function that balances the likelihood of the operators and their generality. Moreover, the confidence is also used so that generality is strongly preferred for uncertain estimations.

The paper is organized as follows. First we will introduce some background required to understand the logic rule learning, and the algorithms proposed to that end. Afterwards, the action model and the translations needed between logic programs and planning domains are explained. Finally we show experimental results and conclusions of the presented approach.

2 Preliminaries

In this section we recall some preliminaries of logic programming. We also explain the basis of learning from interpretation transitions in order to make easier the understanding of its extension to multi-valued variable and probabilistic rule learning.

2.1 Logic Programming

We consider a first-order language and denote the Herbrand base (the set of all ground atoms) as \mathcal{B} . A (*normal*) *logic program* (NLP) is a set of *rules* of the form

$$A \leftarrow A_1 \wedge \dots \wedge A_m \wedge \neg A_{m+1} \wedge \dots \wedge \neg A_n \tag{1}$$

where A and A_i are atoms $(n \ge m \ge 0)$. For any rule R of the form (1), the atom A is called the *head* of R and is denoted as h(R), and the conjunction to the right of \leftarrow is called the *body* of R. An (*Herbrand*) *interpretation* I is a subset of \mathcal{B} . For a logic program P and an Herbrand interpretation I, the *immediate consequence operator* (or T_P operator) (Apt et al. 1988) is the mapping $T_P : 2^{\mathcal{B}} \to 2^{\mathcal{B}}$:

$$T_P(I) = \{ h(R) \mid R \in ground(P), b^+(R) \subseteq I, b^-(R) \cap I = \emptyset \}.$$
 (2)

2.2 Learning from Interpretation Transitions

LF1T (Inoue et al. 2014) is an *any time algorithm* that takes a set of one-step state transitions E as input. These one-step state transitions can be considered as positive examples. From these transitions, the algorithm learns a logic program P that represents the dynamics of E. To perform this learning process, we can iteratively consider one-step transitions. In *LF1T*, the set of all atoms \mathcal{B} is assumed to be finite. In the input *E*, a state transition is represented by a pair of interpretations. The output of *LF1T* is a logic program that realizes all state transitions of *E*.

Learning from 1-Step Transitions (LF1T)

Input: *E* a set of state transitions (I, J) of a system *S* and *B* the set of all possible atoms that can appear in *I* and *J*.

Output: A logic program P such that J = next(I) holds for any $(I, J) \in E$.

To build a logic program with *LF1T*, in (Ribeiro and Inoue 2014) we used a bottom-up method that generates hypotheses by *specialization* from the most general rules, that are fact rules, until the logic program is consistent with all input state transitions. Learning by specialization ensures to output the most general valid hypothesis.

2.3 Learning Multi-valued logic programs

Research in multi-valued logic programming has proceed along three different directions (Kifer and Subrahmanian 1992): bilattice-based logics (Fitting 1991; Ginsberg 1988), quantitative rule sets (Van Emden 1986) and annotated logics (Blair and Subrahmanian 1989; Blair and Subrahmanian 1988). The multi-valued logic representation used in our new algorithm is based on annotated logics. Here, to each atom corresponds a given set of values. In a rule, a literal is an atom annotated with one of these values. It allows us to represent annotated atoms simply as classical atoms and thus to remain in the normal logic program semantics.

In order to represent multi-valued variables, we now restrict all atoms of a logic program to the form var^{val} . The intuition behind this form is that var represents some variable of the system and val represents the value of this variable. In annotated logics, the atom var is said to be annotated by the constant val. We consider a *multi-valued logic program* as a set of *rules* of the form

$$var^{val} \leftarrow var_1^{val_1} \land \dots \land var_n^{val_n}$$
 (3)

where var^{val} and $var^{val_i}_i$'s are atoms $(n \ge 1)$. Like before, for any rule R of the form (3), left part of \leftarrow is called the *head* of R and is denoted as h(R), and the conjunction to the right of \leftarrow is called the *body* of R. We represent the set of literals in the body of R of the form (3) as $b(R) = \{var_1^{val_1}, \ldots, var_n^{val_n}\}$. A rule R of the form (3) is interpreted as follows: the variable var takes the value val in the next state if all variables var_i have the value val_i in the current state. An interpretation of a multi-valued program provides the value of each variable of the system and is defined as follows.

Definition 1 (Multi-valued Interpretation)

Let \mathcal{B} be a set of atoms where each element has the form var^{val} . An *interpretation* I of a set of atoms \mathcal{B} is a subset of \mathcal{B} where $\forall var^{val} \in \mathcal{B}, \exists var^{val'} \in I$ and $\forall var^{val'} \in I, \exists var^{val''} \in I, val' \neq val''$.

For a system S represented by a multi-valued logic program P and a state s_1 represented by an interpretation I, the successor of s_1 is represented by the interpretation:

$$next(I) = \{h(R) \mid R \in P, b(R) \subseteq I\}$$

The state transitions of a logic program P are represented by a set of pairs of interpretations (I, next(I)).

Definition 2 (Multi-valued Consistency)

Let R be a rule and (I, J) be a state transition. R is *consistent* with (I, J) iff $b(R) \subseteq I$ implies $h(R) \in J$. Let E be a set of state transitions, R is consistent with E if R is consistent with all state transitions of E. A logic program P is *consistent* with E if all rules of P are *consistent* with E.

Definition 3 (Subsumption)

Let R_1 and R_2 be two rules. If $h(R_1) = h(R_2)$ and $b(R_1) \subseteq b(R_2)$ then R_1 subsumes R_2 . Let P be a logic program and R be a rule. P subsumes R if there exists a rule $R' \in P$ that subsumes R.

We say that a rule R_1 is more general than another rule R_2 if R_1 subsumes R_2 . In particular, a rule R is most general if there is no rule $R' \neq R$ that subsumes $R(b(r) = \emptyset)$.

Example 1

Let R_1 and R_2 be the two following rules: $R_1 = (a^1 \leftarrow b^1)$, $R_2 = (a^1 \leftarrow a^0 \land b^1)$, R_1 subsumes R_2 because $(b(R_1) = \{b^1\}) \subset (b(R_2) = \{a^0, b^1\})$. When R_1 appears in a logic program P, R_2 is useless for P, because whenever R_2 can be applied, R_1 can be applied.

To learn multi-valued logic programs with **LF1T** we need to adapt the ground resolution of (Inoue et al. 2014) and the least specialization of (Ribeiro and Inoue 2014) to handle non-boolean variables.

Definition 4 (complement)

Let R_1 and R_2 be two rules, R_2 is a complement of R_1 on var^{val} if $var^{val} \in b(R_1)$, $var^{val'} \in b(R_2)$, $val \neq val'$ and $(b(R_2) \setminus \{var^{val'}\}) \subseteq (b(R_1) \setminus \{var^{val}\})$.

Definition 5 (multi-valued ground resolution)

Let R be a rule, P be a logic program and \mathcal{B} be a set of atoms, R can be generalized on var^{val} if $\forall var^{val'} \in \mathcal{B}, val \neq val', \exists R' \in P$ such that R' is a complement of R on var^{val} :

$$generalise(R, P) = h(R) \leftarrow b(R) \setminus var^{val}$$

Definition 6 (Multi-valued least specialization)

Let R_1 and R_2 be two rules such that $h(R_1) = h(R_2)$ and R_1 subsumes R_2 . Let \mathcal{B} be a set of atoms. The least specialization $ls(R_1, R_2, \mathcal{B})$ of R_1 over R_2 w.r.t \mathcal{B} is

$$ls(R_1, R_2, \mathcal{B}) = \{h(R_1) \leftarrow b(R_1) \land var^{val'} | var^{val} \in b(R_2) \backslash b(R_1), var^{val'} \in \mathcal{B}, val' \neq val\}$$

Least specialization can be used on a rule R to avoid the subsumption of another rule with a minimal reduction of the generality of R. By extension, least specialization can be used on the rules of a logic program P to avoid the subsumption of a rule with a minimal reduction of the generality of P. Let P be a logic program, \mathcal{B} be a set of atoms, R be a rule and S be the set of all rules of P that subsume R. The least specialization $ls(P, R, \mathcal{B})$ of P by R w.r.t \mathcal{B} is as follows:

$$ls(P, R, \mathcal{B}) = (P \setminus S) \cup (\bigcup_{R_P \in S} ls(R_P, R, \mathcal{B}))$$

LF1T starts with an initial logic program $P = \{var^{val} \leftarrow | var^{val} \in \mathcal{B}\}$. Then LF1T

iteratively analyzes each transition $(I, J) \in E$. For each labeled atom A that **does not** appear in J, LF1T infers an anti-rule R_A^I :

$$R_A^I = A \leftarrow \bigwedge_{B_i \in I} B_i$$

A rule of P that subsumes such an anti-rule is not consistent with the transitions of E and must be revised. The idea is to use least specialization to make P consistent with the new transitions (I, J) by avoiding the subsumption of all anti-rules R_A^I inferred from (I, J). After least specialization, P becomes consistent with the new transition while remaining consistent with all previously analyzed transitions (theorem 3 of (Ribeiro and Inoue 2014)). When all transitions of E have been analyzed, LF1T outputs the rules of the system that realize E.

3 Learning from Uncertain State Transitions

In this section we extend the *LFIT* framework to learn probabilistic dynamics by proposing an extension of *LFIT* for learning from uncertain state transitions. Where other work like (Gutmann et al. 2011) perform inferences from a probabilistic logic program, what we do is inferring the rules of such logic program. The programs infered by our new algorithm are similar to paraconsistent logic program of (Blair and Subrahmanian 1988). The use of annotated atoms allows the learned programs to induce multiple values for the same represented variable. It allows us to represent multi-valued models and capture nondeterministic state transitions. Our semantics differs from previous work like (Fierens et al. 2013). There, the authors consider probabilistic logic programs as logic programs in which some of the facts are annotated with probabilities. But in our method, its the rules that have probabilities and they are independent.

3.1 Formalization

An non-deterministic system can be represented by a set of logic programs where the rules have the following form:

$$R = value(var, val, i, j) \leftarrow var_1^{val_1} \land \dots \land var_n^{val_n}$$

where var^{val} and var^{val}_i are atoms $(n \ge 1)$, value(var, val, i, j) is the head of R again denoted h(R) and i, j are natural numbers, $i \le j$. Let I be a multi-valued interpretation. R means that i times over j, var takes the value val in the successor of I if $b(R) \subset I$.

Definition 7 (Non-deterministic successors)

Let I be the multi-valued interpretation of a state of a non-deterministic system S represented by a set of logic programs P. Let P' be a logic program, one of the successors of I according to P' is

$$next(I, P') = \{var^{val} | R \in P', b(R) \subseteq I, h(R) = value(var, val, i, j), 0 < i \leq j\}$$

The set of *successors* of I in S according to P is

 $successor(I, P) = \{J | J \in next(I, P_i), P_i \in P\}$

Example 2

Let R_1 and R_2 be two rules such that $R_1 = (value(a, 1, 10, 100) \leftarrow a^1)$ and $R_2 = (value(a, 2, 90, 100) \leftarrow a^1)$. Let S be a probabilistic system represented by a set of logic programs P such that $P = \{\{R_1\}, \{R_2\}\}$. The possible next states of I in S are $successor(I, P) = \{\{a^1\}, \{a^2\}\}$. The likelihood of having a^1 in the next state of I is 10% and the one of having a^2 is 90%.

3.2 Algorithm

We now present **LUST**, an extension of *LFIT* for Learning from Uncertain State Transition. **LUST** learns a set of deterministic logic programs. The main idea is that when two transitions are not consistent we need two different programs to realize them. The first program will realize the first transition and the second one will realize the second transition. The algorithm will output a set of logic programs such that every transition given as input is realized by at least one of those programs. The rules learned also provide the probability of the variable values in the next state. The probability of each rule $R = value(var, val, i, j) \leftarrow b(R)$ is simply obtained by counting how many transitions (I, J) it realizes (when $b(R) \subseteq I$ and $var^{val} \in J$), represented by *i*, over how many transitions it matches (when $b(R) \subseteq I$), represented by *j*.

LUST algorithm:

- Input: a set of pairs of interpretations E and a set of atoms \mathcal{B} .
- Step 1: Initialize a set of logic programs P with one program P_1 with fact rules for each atom of \mathcal{B} .
- Step 2: Pick (I, J) in E, check consistency of (I, J) with all programs of P:
- if there is no logic program in P that realizes (I, J) then

— copy one of the logic programs P_i into a P'_i and add rules in P'_i to realize (I, J). — Use full ground resolution to generalize P'_i .

- Step 3: Revise all logic programs that realize (I, J) by using least specialization.
- Step 4: If there is a remaining transition in *E*, go to step 2.
- Step 5: Compute the probability of each rule of all programs P_i according to E.
- Output: P a set of multi-valued logic programs that realizes E.

4 Integration of Logic Programming and Planning Domains

In this section we describe the formalization used to represent planning operators, as well as the data conversions needed to learn symbolic operators from grounded transitions while using propositional logic programming.

4.1 Planning Model

Although *LUST* uses a propositional representation, the planning model uses a relational representation to provide a better generalization between different states. Relational domains represent the state structure and objects explicitly. These domains are described by using a vocabulary of predicates \mathcal{P} and actions \mathcal{A} , and a set of objects C_{π} . Predicates and actions take objects as arguments to define their ground counterparts.

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Example 3

Let on(X,Y) be a symbolic predicate, and { box1,box2,box3} be a set of objects. Three possible groundings of the symbolic predicate on(X,Y) with the given atoms are on(box1, box2), on(box1, box3) and on(box3, box1).

A ground predicate or action is equivalent to an atom. In a planning domain, a planning state s is a set of ground predicates $s = \{p_1, p_2, ..., p_n\}$ that is equivalent to an Herbrand interpretation.

Our planner operators represent a subset of RDDL domains (Sanner 2010). For each variable, we define the probability that it will become true in the next state based on a set of preconditions. In contrast to the full RDDL specification, these preconditions can only consist of an action and a set of predicates. Thus we define a planning operator as a tuple $o = \langle o_{p^{val}}, o_{act}, o_{prec}, o_{prob} \rangle$ where:

- $o_{p^{val}}$ is a predicate p whose value can change to val by applying the operator. It is equivalent to the head of a logic rule.
- o_{act} is the action that has to be executed for the operator to be applicable.
- *o*_{prec} is a set of predicates that have to be satisfied in the current state so that the planning operator is applicable. It is equivalent to the body of a logic rule.
- o_{prob} is the probability that p^{val} will be true.

Note that planning operators use a relational representation (i.e. symbolic predicates), while rules learned by *LUST* are propositional (i.e. annotated atoms, which are equivalent to ground predicates). Moreover, a symbolic planning operator can be grounded by replacing each variable by an object.

4.2 Data Representation

To have a more general and compact model, we are using a relational representation at the planning level. The input of our method consists of state transitions that are tuples $t = \langle s, act, s' \rangle$ where s and s' are the states before and after executing the action *act*. The states consist of sets of ground predicates, and *act* is a grounded action. On the other hand, the output is a set of symbolic (i.e. non-grounded) planning operators. Therefore, our method transforms initial grounded data to symbolic planning operators. Moreover, *LUST* works with propositional atoms, so a transformation from symbolic predicates to atoms and back to symbolic predicates is also needed. Figure 1 shows the needed data conversions, which are explained in more detail below.

Transform grounded transitions to symbolic transitions:

- Input: a set of grounded transitions $T = [t_1, t_2, ..., t_n]$.
- For each transition $t = \langle s, act, s' \rangle$:
 - Take every argument χ of the action *act*.
 - Substitute χ for a default symbolic parameter in s, act, and s'.
 - Create a new transition t' with the symbolic predicates in s and s' and the symbolic action act.
 - Add the new transition t' to T'.
- **Output**: set of symbolic transitions T'.

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Transform a symbolic transition to an interpretation transition :

- **Input**: a symbolic transition $t = \langle s, act, s' \rangle$.
- Assign an atom to each predicate in s, act and s'.
- *I* = atoms that correspond to *s*.
- Add *act* as an atom in *I* that represents the action.
- J = atoms that correspond to s'.
- **Output**: interpretation transition (*I*, *J*).

To transform a planning symbolic transition to an interpretation transition, a labeled atom that encodes the action is added to the body of the interpretation transition. As each transition has exactly one action, a multi-valued variable represents it more efficiently than boolean, otherwise every action would have to be represented as different variables with only one being true. Moreover, each symbolic predicate value is represented by one labeled atom. After the logic programs are generated, the labeled atoms are translated back to symbolic predicates by using the same conversion.

Transform a logic program to a set of planning operators:

- **Input**: a logic program *P*.
- For every rule R ∈ P such that h(R) = value(var, val, i, j), a planning operator o is created so that:
 - $o_{p^{val}} = var^{val}.$
 - o_{act} = the action in the atoms of b(R).
 - o_{prec} = the set of atoms in b(R) that represent predicates.
 - $o_{prob} = i/j.$
 - Add o to O
- **Output**: set of planning operators \mathcal{O} .

5 Selecting the Set of Planning Operators

In this section we present the method to select the best subset of probabilistic planning operators by using the set of logic programs generated by *LF1T*. First, the requirements that the planning operators have to satisfy are presented. Afterwards, we explain the preferences to decide which are the best planning operators. Finally, the method to obtain the desired planning operators is described.

5.1 Planning Operators Requirements

Probabilistic planners require that only one planning operator can be applied in each stateaction pair. Therefore the planning model has to be defined with a set of non-conflicting planning operators, so the planner can always decide which operator to apply for each state-action pair.

Definition 8 (Conflicting planning operators)

Let o_1 and o_2 be two planning operators that represent the same action $o_{1,act} = o_{2,act}$ and change the same predicate $o_{1,p^{val}} = o_{2,p^{val}}$ with different probabilities $o_{1,prob} \neq o_{2,prob}$. A conflict exists between both planning operators if $\exists s \mid o_{1,prec} \subset s, o_{2,prec} \subset s$.

5.2 Score Function

LUST provides the minimal set of rules that describe all possible transitions. However, the best subset of non-conflicting planning operators has to be selected to create the model. To decide which are the best set of operators the algorithm uses a score function. The algorithm prefers operators with a high likelihood (i.e. that can successfully explain the state transitions), but also has a regularization term to avoid overfitting to the training data. The regularization is based on the number of planning operators and preconditions in those operators, so that general operators are preferred when the likelihood of both is similar. This regularization penalty is bigger when there are few training transitions to estimate each operator, as general operators are preferred to poorly estimated specific ones. On the other hand, the regularization penalty decreases as our estimate improves with more transitions because the method can be more confident about the operators. The following functions are used to define the score function.

- The likelihood P(t|O) = P(s'|s, act, o) | (o ∈ O, o_{prec} ∈ s, o_{act} = act) is the probability that the transition t is covered by the set of planning operators O.
- The penalty term $Pen(\mathcal{O}) = |\mathcal{O}| + \frac{1}{|\mathcal{O}|} \sum_{o \in \mathcal{O}} |o_{prec}|$ is the number of planning operators plus the average number of preconditions that they have.
- The confidence in a planning operator $Conf(\mathcal{O}, T)$ is bounded by using the Hoeffding's inequality. The probability that our estimate $\widehat{o_{prob}}$ is accurate enough $|\widehat{o_{prob}} - o_{prob}| \leq \epsilon$ with a number of samples |T| is bounded by $Conf(\mathcal{O}, T) \leq 1 - 2e^{-2\epsilon^2|T|}$.

Finally, the proposed score function is defined as

$$S(\mathcal{O},T) = \frac{1}{|T|} \left(\sum_{t \in T} P(t|\mathcal{O}) \right) + \alpha \frac{Pen(\mathcal{O})}{Conf(\mathcal{O},T)}$$
(4)

where α is a scaling parameter for the penalty term. Also note that $Conf(\mathcal{O}, T) \simeq 1$ when the number of input transitions is large, so the penalty term will always be present to ensure general operators are preferred.

5.3 Selecting the Best Planning Operator Subset

In this section we present the algorithm used to select the subset of non-conflicting planning operators that maximizes the score function. To do it efficiently we make use of the dependency relations between operators by using the parent relation graph defined below.

Definition 9 (Parent relation)

Let o_1 and o_2 be two planning operators that represent the same action $o_{1,act} = o_{2,act}$ and the same effect $o_{1,p^{val}} = o_{2,p^{val}}$. o_1 is a parent of the operator o_2 if $o_{1,prec} \subset o_{2,prec}$.

Definition 10 (Parent graph)

The parent graph $G_{\mathcal{O}}$ of a set of planning operators $\mathcal{O} = \{o_1, ..., o_n\}$ is a directed graph with arcs (o_i, o_j) when o_i is parent of o_j and $|o_{j,prec}| - |o_{i,prec}| = 1$. Figure 2 shows an example of a parent graph. We will call *leaves* $L(G_{\mathcal{O}})$ of the graph all nodes that do not have a child.

The parent graph orders the rules in levels that represent the generality of the operator: the less predicates in the preconditions the more general the operator is. This graph provides two advantages: generalizing operators is easier as we only have to follow the arcs,



Fig. 2. Example of a parent graph. In each node, the planning operator preconditions are shown. Leaves are the nodes painted in blue.

and it reduces the number of conflicts to check because general operators won't be checked if other more specific operators that represent the same dynamics exist. An optimal solution can be obtained through backtracking, but as it is computationally intensive, we present the greedy approach that provides similar experimental results and can be applied efficiently in difficult domains with many input transitions.

The first step to select the set of planning operators is to create the parent graph. Afterwards, two algorithms are applied iteratively until no better sets of planning operators can be found. The first algorithm prunes all conflicting leaf operators, leaving only the subset of leaves with the highest score. The second one tries to generalize leaf operators, and removes them from the tree if it can find a better and more general subset.

Prune conflicting leaves:

- Input: Parent graph $G_{\mathcal{O}}$ of the set of operator candidates \mathcal{O} that represent one action.
- While leaf nodes $L(G_{\mathcal{O}})$ are conflicting, repeat:
 - Create an undirected graph $G_{conflict}$ that represents the conflicts between $L(G_{\mathcal{O}})$.
 - Calculate the score for each non-conflicting subset in $G_{conflict}$.
 - Remove all leaves from $G_{\mathcal{O}}$ with the exception of the subset with the best score.
- **Output**: Parent graph $G'_{\mathcal{O}}$ with non-conflicting leaves.

Generalize operators:

- Input: Parent graph $G'_{\mathcal{O}}$ with non-conflicting leaves.
- For each leaf parent $p \mid \exists (p, l) \in G'_{\mathcal{O}}, l \in L(G'_{\mathcal{O}})$ do:
 - Create a new set of operators with the leaves $\mathcal{O}' = L(G'_{\mathcal{O}})$.
 - Add p to \mathcal{O}' and remove its children leaves $l \mid \exists (p, l) \in G'_{\mathcal{O}}, l \in L(G'_{\mathcal{O}}).$
 - Calculate score $S(\mathcal{O}', T)$.
- If the best parent p improved the score:

- Remove $l \mid \exists (p, l) \in G'_{\mathcal{O}}, l \in L(G'_{\mathcal{O}})$ from $G'_{\mathcal{O}}$.

• **Output**: Parent graph $G'_{\mathcal{O}}$ with better generalization.

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Fig. 3. Results of learning the Triangle Tireworld domain and the Elevators domain from IPPC 2014. The results shown are the means and standard deviations obtained from 10 runs. The number of training transitions per action are shown.

6 Evaluation

In this section we provide an experimental evaluation of our approach by learning two domains of the 2014 International Probabilistic Planning Competition (IPPC). The experiments use transitions $t = \langle s, act, s' \rangle$ generated by randomly constructing a state s, randomly picking the arguments of the action *act*, and then applying the action to generate the state s'. The distribution of samples is biased to guarantee that half of the samples have a chance to change the state. To measure the quality of the learned models, the errors shown represent the differences between the learned operators and the ground truth ones. For each incorrect predicate in an operator preconditions, the number was increased by 1.

The left plot in Fig. 3 shows the results obtained in the Triangle Tireworld domain. In this domain, a car has to move to its destination, but it has a probability of getting a flat tire while it moves. The domain has 3 actions. A "Change Tire" action that has only one precondition, a "Load Tire" action that has 2 preconditions to change one predicate, and a "MoveCar" action that changes 3 predicates based on 3 different preconditions. The results show that easy actions can be learned with just a few transitions, while very complex actions require 100 transitions until average errors below 1 are obtained.

The right plot in Fig. 3 shows the results obtained in the Elevators domain. In this domain, we are only learning the actions that interact with the elevators, and not the dynamic effects related to people using them. The "openDoorGoing*" and "closeDoor" actions are easy to learn, but the "moveCurrentDir" action has two effects with 3 different preconditions each, and another two effects with 4 preconditions. Therefore, to learn successfully the dynamics of "moveCurrentDir" a large number of input transitions is required.

7 Conclusions

We have presented a new approach to learn planning operators. In contrast to previous approaches, it learns the dynamics of every predicate independently, and thus can be applied to learn RDDL domains. The presented approach can find good sets of planning operators efficiently, as it combines logic programming to restrict the set of candidates, and then the parent relations between operators are used to quickly prune bad candidates until the desired subset of operators is found (i.e. a set that generalizes as much as possible while explaining well enough the input transitions). Finally, experimental validation is provided to show that the planning operators can be learned with a reduced number of transitions.

As future work, we would like to work on planning operators independent of actions. This would require to have a more intelligent method to generate symbolic transitions from grounded transitions so that changes not represented by actions could be learned.

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