# Minimizing State Space Partitionings Using Decision Trees

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Abstract—Lorem ipsum dolor sit amet, consectetuer adipiscing elit. Ut purus elit, vestibulum ut, placerat ac, adipiscing vitae, felis. Curabitur dictum gravida mauris. Nam arcu libero, nonummy eget, consectetuer id, vulputate a, magna. Donec vehicula augue eu neque. Pellentesque habitant morbi tristique senectus et netus et malesuada fames ac turpis egestas. Mauris ut leo. Cras viverra metus rhoncus sem. Nulla et lectus vestibulum urna fringilla ultrices. Phasellus eu tellus sit amet tortor gravida placerat. Integer sapien est, iaculis in, pretium quis, viverra ac, nunc. Praesent eget sem vel leo ultrices bibendum. Aenean faucibus. Morbi dolor nulla, malesuada eu, pulvinar at, mollis ac, nulla. Curabitur auctor semper nulla. Donec varius orci eget risus. Duis nibh mi, congue eu, accumsan eleifend, sagittis quis, diam. Duis eget orci sit amet orci dignissim rutrum.

### I. INTRODUCTION

Motivate for discrete reinforcement learning ... (list tools that produce policies that are similar)

However, discrete reinforcement learning suffers from some issues with the policies. Even if it succeeds, the constructed policies might be too large for humans to interpret. This challenges their explainability (even if we can verify them) ...

In this paper we look at the problem of decreasing the size of the policies expressed as decision trees ...

Our contributions include: ...

The experiments show that: ...

The paper proceeds as follows. ...

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### II. PRELIMINARIES

Definition 1: A partitioning  $\mathcal{A}$  of the state space  $\mathcal{S} \subseteq \mathbb{R}^K$  is a set of non-overlapping subsets  $\mathcal{A} \subseteq \mathcal{P}(\mathcal{S})$  covering  $\mathcal{S}$ , so  $\bigcup_{\nu \in \mathcal{A}} \nu = \mathcal{S}$  and whenever  $\nu, \nu' \in \mathcal{A}, \nu \neq \nu'$  then  $\nu \cap \nu' = \emptyset$ . do we assume that the state space  $\mathcal{S}$  is rectangular?

For an axis aligned partitioning, each region  $\nu$  can be expressed in terms of two corner points,  $s^{\min}, s^{\max} \in \mathcal{S}$ , so that for each  $s = (s_1, \ldots, s_K) \in \nu$  it holds that  $s_i^{\min} < s_i \leq s_i^{\max}$  for  $i = 1, \ldots, K$ . In this work we exclusively consider axis aligned partitionings and we define all regions as a tuples  $\nu = (s^{\min}, s^{\max})$ . Note that the entire state space  $\mathcal{S} \in \mathbb{R}^K$  can be described as a region: if  $\mathcal{S}$  is unbounded in all dimensions (meaning its limits are positive and negative infinity) then  $s_i^{\min}$  and  $s_i^{\max}$  for the entire state space is  $-\infty$  and  $\infty$  respectively for  $i = 1, \ldots, K$ .

Definition 2 (Decision tree): A binary decision tree over the domain  $\mathcal{S} \in \mathbb{R}^K$  is a tuple  $\mathcal{T} = (\eta_0, \mathcal{N}, \mathcal{L}, \mathcal{D})$  where  $\eta_0 \in \mathcal{N}$  is the root node of the tree,  $\mathcal{N}$  is a set of branching nodes and  $\mathcal{L}$  is a set of leaf nodes, each of which is assigned a decision  $\delta$  from the set of decisions  $\mathcal{D}$ . Each branch node  $\eta \in \mathcal{N}$  consists of two child nodes and a predicate function of the form  $\rho_{\eta}(s) = s_i \leq c$  with  $s \in \mathcal{S}$  and c being a constant

Given a state  $s \in \mathcal{S}$  and a decision tree  $\mathcal{T}$ , we can evaluate  $\mathcal{T}(s)$  to obtain a decision  $\delta$  by following the path from the root node to a leaf node given by the repeated evaluation of the predicate function  $\rho_{\eta}(s)$  at each node  $\eta$ , starting with the root node and continuing with the left child if  $\rho_{\eta}(s)$  evaluates to true and with the right child if it evaluates to false. When we encounter a leaf node  $\ell$ , we return the decision assigned to  $\ell$ . Further, we also allow evaluating a region of  $\mathcal{S}$ . Given a region  $\nu = (s^{\min}, s^{\max}), \ [\delta]_{\nu} = \mathcal{T}(\nu)$  is the set of all decisions that can be obtained evaluating configurations of  $\nu$ , ie.  $\mathcal{T}(\nu) = \{\mathcal{T}(s) \mid s \in \nu\}$ .

We denote the path  $\lambda(\ell)$  and define it as an ordered list of tuples of the form  $(\eta_j,b)$  where  $\eta_j$  is the jth node on the path  $(\eta_0$  will always be the root node) and b is a binary value indicating wether the path continues with the left child (b=1) or right child (b=0).[inline]Not sure if it makes sense or is necessary to state which value of b means what. The reasoning for my choice is that b=1 indicates 'true' and it is when  $\rho_{\eta_j}(s)$  is true, that we choose the left path. The

path can then be said to define a region where the corner points  $s^{\min}$  and  $s^{\max}$  are given by compiling the bounds on each dimension  $i=1,\ldots,K$  given by the predicate function  $\rho_{\eta_j}=i_{\eta_j}\leq c_{\eta_j}$  for each  $\eta_j\in\lambda(\ell)$  into points. The coordinates for each point is given by

$$\begin{split} s_i^{\min} &= \max( \{ \, c_\eta \mid (\eta, b) \in \lambda(\ell), \ i_\eta = i, \ b = 1 \, \} ) \\ s_i^{\max} &= \min( \{ \, c_\eta \mid (\eta, b) \in \lambda(\ell), \ i_\eta = i, \ b = 0 \, \} ) \end{split}$$

[inline]I struggled a lot with coming up with a good way of writing this definition. Do let me know if it works (and if it is even necessary to describe how a region is constructed from a leaf node).

for all  $i=1,\ldots,K$ . We write  $\nu_{\ell}$  to denote the region associated with the leaf node  $\ell$ .

The set of regions obtained from all the leaf nodes of a decision tree constitutes a complete partitioning of a state space  $\mathcal S$  in accordance with Definition1. We thus say that  $\mathcal T$  induces a partitioning  $\mathcal A_{\mathcal T}=\{\nu_\ell\mid \ell\in\mathcal L\}$ . For any region  $\nu$  and a decision tree  $\mathcal T$  we say, that  $\nu$  has singular mapping in  $\mathcal T$  if for all  $p\in \nu$ ,  $\mathcal T(p)=\delta$  for some  $\delta\in\mathcal D$ . Naturally, all regions in  $\mathcal A_{\mathcal T}$  has singular mapping in  $\mathcal T$ . For any partitioning  $\mathcal B$  of the same state space, we say  $\mathcal B$  respects  $\mathcal T$  if and only if every region  $\nu\in\mathcal B$  has singular mapping in  $\mathcal T$ .

#### III. MAXPARTITIONS ALGORITHM

Since state space discretization for Reinforcement Learning is usually done *before* any learning takes place, it tends to be conservative. For this reason, discretization is likely to create adjacent discrete states that are mapped to the same optimal action. The question we would then like to answer is this: if  $\mathcal{T}$  is a decision tree representing a trained strategy and  $\mathcal{A}_{\mathcal{T}}$  is its induced partitioning, can we find another partitioning  $\mathcal{B}$  which is smaller than  $\mathcal{A}_{\mathcal{T}}$  but still respects  $\mathcal{T}$ ?

As an example, consider a case where we have a state space  $\mathcal{S} \in \mathbb{R}^2$  over variables x and y, both of which are defined in the interval [0,9], and a set of actions  $Act = \{action1, action2\}$ . Before learning, we might decide to discretize both x and y into 3 distinct bins, giving us a partitioning of  $\mathcal{S}$  with  $3\times 3=9$  regions. After training, we end up with a Q-table that maps states to action in a way that is shown in Figure III. This same mapping can also be represented by a decision tree, as shown in Figure III.

However, we can (in this small toy example) easily see that we do not need 9 regions to represent this exact state-action mapping. For example, the two regions given by ((3,0),(6,3)) and ((6,0),(9,3)), respectively, both assign action2 as the optimal action, but this mapping would still be preserved if we replaced those two regions with a larger one ((3,0),(9,3)). After a little bit of inspection, we can actually see here that we could represent the same state-action mapping with a partitioning consisting of only 5 regions.

This example showcases how discretization techniques can easily end up with redundancy in the partitioning. This can

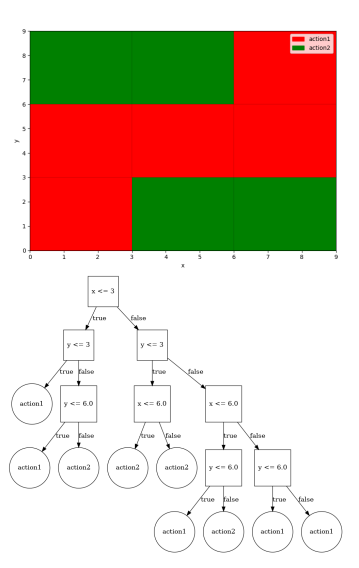


Fig. 1. Two representations of a strategy mapping states to actions. In section III the strategy is represented as a partitioning with colors representing the assigned action. In section III the strategy is given as a decision tree, which also induces the partitioning. For this small example, it is obvious to see that an equivalent state-action mapping could be achieved with fever regions/leaves.

be very difficult to anticipate before learning, especially since a very fine-grained discretization is typically needed for the learning to capture essential information and details for the strategy. Furthermore, for other learning techniques, such as the online partitioning refinement scheme of UPPAAL STRATEGO [1], regions can be created on-the-fly, not be arranged in a straight grid and/or vary in size, which can enhance the problem of redundancy.

We propose MAXPARTITIONS, an algorithm that postprocesses a decision tree inducing a partitioning of a state space in order to minimize the partitioning by *maximizing* the size of the individual regions (or partitions). The output of MAXPARTITIONS is a new partitioning, ie. a list of regions with associated actions, which can then be arranged into a new decision tree.

### A. Details of the algorithm

We write  $\mathcal{T}_i$  for the (ascendingly) sorted list of bounds on dimension i in the policy given by the tree  $\mathcal{T}$ . The first bound in the list is defined to be negative  $\infty$  and the last is positive  $\infty$ . By  $\mathcal{T}_{i,j}$  we write the jth smallest bound on dimension i for each  $j=1,2,\ldots,|\mathcal{T}_i|$ . This can be precomputed as a matrix in log-linear time by collecting and sorting the bounds on all branch nodes in  $\mathcal{T}$  and allows accessing  $\mathcal{T}_{i,j}$  in constant time. Further, in a slight abuse of notation, we define  $\mathcal{T}_{i,|\mathcal{T}_i|+1}$  to be some *sentinel* value representing that we are outside the boundaries of dimension i. Correspondingly, we define a sentinel action  $\alpha$ , and we say that  $\mathcal{T}(s_{\mathcal{T}}^p) = \alpha$  if and only if  $\exists p_i \in p, \ p_i = |\mathcal{T}_i| + 1$ .

Exploiting this notation, let p be a K-dimensional vector of integers, such that  $p_i \leq |\mathcal{T}_i| + 1$  for all  $i = 1, \ldots, K$ , then we can define a point at an intersection of bounds in  $\mathcal{T}$  as  $s_{\mathcal{T}}^p = (\mathcal{T}_{1,p_1}, \mathcal{T}_{2,p_2}, \ldots, \mathcal{T}_{K,p_K})$ . To avoid cluttering the notation, will for the most part omit the subscript  $\mathcal{T}$  on  $s_{\mathcal{T}}^p$ . To make things clear, we will write s to refer to an actual point in the state space of  $\mathcal{T}$ , ie.  $s \in \mathcal{S}$ , and we will write s to refer to a vector of integers representing indicies of bounds in  $\mathcal{T}$ .

The algorithm (given in pseudo-code in Algorithm 1) works by maintaining a pair  $(p^{\min}, p^{\max})$ , and iteratively incrementing  $p^{\max}$  in one dimension at a time until a region  $\nu = (s^{p^{\min}}, s^{p^{\max}})$  cannot be expanded further. When this happens, the region is added to a tree  $\mathcal{T}_{track}$ , which is used to track which areas of the state space have been covered and to provide new a starting point for each iteration of the algorithm. Expansion in dimension i is disallowed if one of the following three *expansion rules* are violated by the expanded region  $\nu'$ :

Definition 3 (Expansion rules): Let  $\nu'$  be a candidate region for a new partitioning derived from  $A_T$ . Then  $\nu'$  is valid if it adheres to the following rules:

- 1)  $\nu'$  does not have singular mapping in  $\mathcal{T}$
- 2)  $\nu'$  intersects with one or more regions already in  $\mathcal{T}_{track}$
- 3)  $\nu'$  intersects with a region  $\nu_o$  in the original partitioning, such that the disjunction  $\nu' \cap \nu_o$  cannot be described by a single region of the form  $(s^{\min}, s^{\min})$

The first two cases are directly related to the definition of the problem, ie. the produced partitioning should respect  $\mathcal{T}$  and only have non-overlapping regions. The third case is required in order to guarantee that in each iteration the algorithm on average will add *at least* one region from the original partitioning to the new partitioning. To see this, consider ... [inline] Make an example to show.

How do we determine this expansion? Let  $(p^{\min}, p^{\max})$  define region in the original partitioning (or potentially the remains of a region previously cut in two). We then want to find a  $\Delta_p \in \mathbb{Z}^K$  such that  $(p^{\min}, p^{\min} + \Delta_p)$  defines a region that follows the three expansion rules and such that incrementing in any one dimension would result in a violation. By definition, a valid value for  $\Delta_p$  is when  $\Delta_p = p^{\max} - p^{\min}$ , since this would just produce the original region. We are therefore guaranteed to at least find this

region. This gives rise to the following definition.

Definition 4 (The expansion vector  $\Delta_p$ ): Given  $p^{\min} \in \mathbb{Z}^K$ , a decision tree  $\mathcal{T}$  over a K-dimensional space and a decision tree  $\mathcal{T}_{track}$  of already found regions,  $\Delta_p \in \mathbb{Z}^K$  is a vector such that for  $p^{\max} = p^{\min} + \Delta_p$  the region  $\nu = (s_{\mathcal{T}}^{p^{\min}}, s_{\mathcal{T}}^{p^{\max}})$  does not violate any of the expansion rules in Definition 3 and where for any other  $\Delta_p' = (\Delta_{p_1}, \dots, \Delta_{p_i} + 1, \dots, \Delta_{p_K})$  this would not hold.

A greedy approach to finding  $\Delta_p$  starts with  $\Delta_p = p^{\max} - p^{\min}$ , for some (remainder of a) region  $\nu = (s_{\mathcal{T}}^{p^{\min}}, s_{\mathcal{T}}^{p^{\max}})$ . We then iteratively increment a single dimension chosen non-deterministically untill the invariants are violated. Let  $\hat{\mathbf{e}}_i$  denote the unit vector parallel to axis i, such that  $\Delta_p + \hat{\mathbf{e}}_i = (\Delta_{p_1}, \ldots, \Delta_{p_i} + 1, \ldots, \Delta_{p_K})$ . At each increment, we define a candidate region  $\nu'$  from  $p^{\min}$  and  $p^{\max} = p^{\min} + \Delta_p$  and check for singular mapping (Rule 1) and no overlap with regions in  $\mathcal{R}$  (Rule 2). If any of these two do not hold, we mark dimension i as exhausted, roll back the increment and continue with a new dimension not marked as exhausted.

If Rule 3 is violated, the algorithm wil initiate an attempt at *healing* the candidate expansion, by continuing the expansion to the largest bound in the expansion dimension of any of the broken regions. This way we try to see if the violation can be overcome by simply expanding more aggressively. However, care is required to ensure, that we can roll back this extra expansion if it did not work (or if we inadverdently broke any of the other rules in the process).

When all dimensions have been exhausted,  $\Delta_p$  adheres to Defintion III-A.

# Algorithm 1 MaxPartitions

18: return  $\mathcal{R}$ 

```
Require: \mathcal{T}: A binary decision tree over the domain \mathbb{R}^K
       inducing the partitioning A_T
  1: \mathcal{T}_{track} \leftarrow \text{empty tree}
  2: \mathcal{R} \leftarrow \{\}
  3: while \mathcal{T}_{track} has unexplored regions do
             (p^{\min}, p^{\max}) \leftarrow \text{select region bounds of unexplored}
       region
             \Delta_p \leftarrow p^{\text{max}} - p^{\text{min}}
  5:
             while not all dimensions have been exhausted do
  6:
                   d \leftarrow randomly select unexhausted dimension
  7:
                   \Delta_{p_d} \leftarrow \Delta_{p_d} + 1 \\ \nu' \leftarrow (s_{\mathcal{T}}^{p^{\min}}, s_{\mathcal{T}}^{p^{\min} + \Delta_p})
  8:
                   if \nu' violates Rule 1 or 2 (Definition 3) then
 10:
11:
                         \Delta_{p_d} \leftarrow \Delta_{p_d} - 1
12:
                          mark d as exhausted
                   else if \nu' violates Rule 3 (Definition 3) then
13:
                          attempt healing in dimension d
14:
            \begin{array}{l} \nu' \leftarrow (s_{\mathcal{T}}^{p^{\min}}, s_{\mathcal{T}}^{p^{\min} + \Delta_p}) \\ \text{PUT}(\mathcal{T}_{track}, \, \nu') \end{array}
15:
16:
             \mathcal{R} \leftarrow \mathcal{R} \cup \{\nu'\}
```

### B. Analyzing the algorithm

In the following we provide an upper bound of the running time of MAXPARTITIONS and a proof of correctness.

- 1) Running time:
- 2) Proof of correctness:

## C. From regions to decision tree

The output of the MAXPARTITIONS algorithm is a list of regions with associated actions. For this to be of any use, we need to construct a new decision tree to represent these stateaction pairs. To this goal, we face the issue that it is not given (and in fact, very unlikely) that the suggested partitioning can be perfectly represented by a decision tree, as this would require the existence of enough 'clean splits' (ie. predicates on some variable that perfectly divides the regions into two sets with an empty intersection) to arrange the entire set of regions.

Therefore, we suggest a brute-force algorithm that tries to separate the regions as cleanly as possible. Let  $\mathbf{R}$  be a list of regions and let  $a_{\nu}$  be the action associated with the region  $\nu=(s^{\min},s^{\min})$ . In the following, we refer to  $s^{\min}$  and  $s^{\max}$  of a region  $\nu$  by  $\nu_{\min}$  and  $\nu_{\max}$  respectively, and to the value of a specific dimension i in one such boundary point as  $\nu_{\min,i}$  or  $\nu_{\max,i}$ .

We iteratively create a branch node that splits  $\mathbf{R}$  into two,  $\mathbf{R}_{low}$  and  $\mathbf{R}_{high}$ , based on a predicate function  $\rho(x) = x_i \le c$  with  $c \in \mathbb{R}$  so that  $\mathbf{R}_{low} = \{ \nu \in \mathbf{R} \mid \rho(\nu_{\min}) \text{ is True} \}$  and  $\mathbf{R}_{high} = \{ \nu \in \mathbf{R} \mid \rho(\nu_{\max}) \text{ is False} \}$ . When the list only contains a single element  $\nu$ , we create a leaf node with action  $a_{\nu}$  and return.

The question is how to determine  $\rho(x)$ , more specifically which dimension i to predicate on and at which value c. Ideally, we want to split  $\mathbf{R}$  in two equally sized subsets and in a way that no single region would have to occur in both, ie. we would like  $\mathbf{R}_{low} \cap \mathbf{R}_{high} = \emptyset$ . For this we define an impurity measure  $I(\mathbf{R}_{low}, \mathbf{R}_{high})$  that penalises the difference in size between  $\mathbf{R}_{low}$  and  $\mathbf{R}_{high}$  and the size of the intersection between the two. Let abs(a) be the absolute value of a and let |b| denote the size of a set b, then

$$I(\mathbf{R}_{low}, \mathbf{R}_{high}) = abs(|\mathbf{R}_{low}| - |\mathbf{R}_{high}|) + |\mathbf{R}_{low} \cap \mathbf{R}_{high}|$$

Our brute-force way of finding the predicate that minimizes I is to iterate over the dimensions in  $\mathcal S$  and for each dimension i we sort the regions according to their upper bound. Let  $\mathbf R_i = \{\nu^1, \nu^2, \dots, \nu^n\}$  be the list sorted according to the i th dimension so that for all  $\nu^j, \nu^{j+1} \in \mathbf R_i$  it holds that  $\nu^j_{\max,i} \leq \nu^{j+1}_{\max,i}$ . If we then let  $\rho(x) = x_i \leq c$  with  $c = \nu^j_{\max,i}$  we have  $|\mathbf R_{low}| = j$  and  $|\mathbf R_{high}| = n - j$ . For determining the size of  $\mathbf R_{low} \cap \mathbf R_{high}$  we simply need to count the number of regions  $\nu^{j+m}$  for  $m=1,2,\dots,n-j$  whose lower bound is less than our predicate bound c, since these regions will appear both in  $\mathbf R_{low}$  (because then, by definition,  $\rho(x) = x_i \leq c$  will be true for  $x_i = \nu^{j+m}_{\min,i}$  and  $c = \nu^j_{\max,i}$ ) and in  $\mathbf R_{high}$  (because our sorting ensures that for all  $\nu^j, \nu^{j+m}$  it holds that  $\nu^j_{\max,i} \leq \nu^{j+m}_{\max,i}$ ).

Now we can write our impurity measure in terms of these quantities:

$$I(\mathbf{R}_{low},\mathbf{R}_{high}) = abs(j-(n-j)) + \sum_{m=1}^{n} \mathbb{1}(\rho(\nu_{\min}^{j+m})), \quad \text{for all } \nu^{j} \in \mathbf{R}_{i}$$

where  $\mathbb{1}$  is the indicator function,  $\mathbf{R}_i$  is the list of regions sorted according to upper bounds in dimension i and  $\mathbf{R}_{low}$  and  $\mathbf{R}_{high}$  are the subsets resulting from splitting on the predicate function  $\rho(x) = x_i \leq c$  with  $c = \nu_{\max,i}^j$  so that  $\mathbf{R}_{low} \subsetneq \mathbf{R}$ ,  $\mathbf{R}_{high} \subsetneq \mathbf{R}$  and  $\mathbf{R} \supseteq \mathbf{R}_{low} \cup \mathbf{R}_{high}$ .

Finding the best split, ie. the one that minimizes the impurity, is a  $O(Kn^2)$  operation, as it requires a nested loop through all the regions for each of the Kdimensions (the nested loop being the final summation term over  $m=1,2,\ldots,n-j$  for all  $j=1,2,\ldots,n-1$ ). In this work, we have not attempted to find a faster implementation as we found that the size of  ${\bf R}$  obtained by using our MAXPARTITIONS algorithm did not cause performance issues.

#### IV. EXPERIMENTS

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TABLE I

COMPARING MAXPARTITIONS AND VIPER FOR PURE SHIELDS AND SHIELDED ORACLES.

| Model              | Input | Pure shield MAXPARTITIONS | VIPER | Input | Shielded oracle MAXPARTITIONS | VIPER |
|--------------------|-------|---------------------------|-------|-------|-------------------------------|-------|
| Bouncing ball      | XXX   | XXX                       | XXX   | XXX   | XXX                           | XXX   |
| Random walk        | XXX   | XXX                       | XXX   | XXX   | XXX                           | XXX   |
| Cartpole           | XXX   | XXX                       | XXX   | XXX   | XXX                           | XXX   |
| Cruise             | XXX   | XXX                       | XXX   | XXX   | XXX                           | XXX   |
| DCDC boost control | XXX   | XXX                       | XXX   | XXX   | XXX                           | XXX   |

### REFERENCES

[1] M. Jaeger, P. G. Jensen, K. Guldstrand Larsen, A. Legay, S. Sedwards, and J. H. Taankvist, "Teaching stratego to play ball: Optimal synthesis for continuous space MDPs," in *Automated Technology for Verification and Analysis* (Y.-F. Chen, C.-H. Cheng, and J. Esparza, eds.), (Cham), pp. 81–97, Springer International Publishing, 2019.

#### APPENDIX

The goal is to take all the leafs of a Q-forest and from these build a single binary decision tree, that through evaluation of a state arrives at a leaf node indicating the best action to take. The general idea of the algorithm is to repeatedly take the next leaf  $L_i$  in  $\mathcal{L}$ , which we (because of our sorting) know to have the best Q-value, and insert its action into the decision tree we are building so that it respects  $S_i$ . This means, that the internal nodes of the tree will still be checks on variables in  $\mathcal{V}$ .

### A. Creating the root

We initialise the tree from the lowest valued leaf in  $\mathcal{L}$  which we will denote  $L_0$ . The operation of inserting  $L_0$  into the at this point empty tree requires special attention, since we have to create split nodes for all the variable bounds  $(l_{0,i},u_{0,i})$  in  $S_0$ . We denote this operation MakeRoot and its pseudocode is given in Algorithm 2.

The operation iterates through all tuples  $(u_{0,i}, l_{0,i})$  in  $S_0$  and whenever it encounters a bound that is different from the limit (which we here assume to be infinity) it has to insert a new node into the tree. This is done via the MakeNewNode procedure, which we define as a helper function to avoid repetitions in the algorithm. This function takes the variable  $V_i$  that the encountered bound relates to, the value of the bound and the action of the leaf, we are inserting. It then creates a new internal node, that represents the partition according to the inequality  $V_i \leq b$  where  $b \in (u_{0,i}, l_{0,i})$ .

Now, MakeNewNode also takes two additional arguments, one boolean to indicate if the bound is an upper bound or not and the current parent node (which might be None). If the bound is an upper bound, then we know that  $Q(a_0)=q_0$  only holds if  $V_i \leq u_{0,i}$ . This means, that  $L_0$  should be inserted somewhere in the left ('low') subtree of the newly created node, while the right ('high') subtree should for now just be set to some random leaf with a very poor Q-score (we will find better values for these subtrees when we process the remaining leaf nodes of  $\mathcal{L}$ ).

Lastly, if the current parent node is not None, then we know it has one subtree that is set (like above) and one that

# Algorithm 2 MakeRoot

```
MAKENEWN-
 1: procedure
    ODE(var, bound, action, isHigh, prevNode)
        node \leftarrow Node(var, bound)
 2:
        if isHigh then
 3:
            node.high \leftarrow \texttt{Leaf}(action, \infty, \texttt{State}(\cdot))
 4:
 5:
        else
            node.low \leftarrow \texttt{Leaf}(action, \infty, \texttt{State}(\cdot))
 6:
 7:
        if prevNode is not None then
            if prevNode.low is None then
 8:
                prevNode.low \leftarrow node
 9:
10:
            else
                prevNode.high \leftarrow node
11:
        return node
12:
 1: function MAKEROOT(Leaf(a_0, q_0, S_0))
        rootNode \leftarrow \texttt{None}
 2:
        prevNode \leftarrow None
 3:
 4:
        for (l_{0,i}, u_{0,i}) in S_0 do
 5:
            if u_{0,i} < \infty then
                 prevNode
 6:
    MAKENEWNODE(V_i, u_{0,i}, a_0, True, prevNode)
                if rootNode is None then
 7:
                     rootNode \leftarrow prevNode
 8:
 9:
            if l_{0,i} > -\infty then
                prevNode
10:
    MAKENEWNODE(V_i, l_{0.i}, a_0, \text{False}, prevNode)
                if rootNode is None then
11:
                     rootNode \leftarrow prevNode
12:
        if prevNode.low is None then
13:
            prevNode.low \leftarrow \text{Leaf}(a_0, q_0, S_0)
14:
15:
        else
            prevNode.high \leftarrow Leaf(a_0, q_0, S_0)
16:
        return \quad rootNode
17:
```

itself is None as it is reserved for our newly created node. We therefore set the undefined child of the parent node to our new node and returns this new node, which is then marked as being the new parent node. If this our first new, we also mark it as our root node.

When we have processed all the bounds in  $S_0$ , we insert  $L_0$  at the 'free' spot in the parent node and return the root node. With this approach, we end up with an initial tree that is very shallow and basically only has one interesting leaf at

the maximum depth of the tree.

### B. Inserting the leaf nodes

To insert the remaining leaf nodes we need a couple of extra helper functions. The task is to identify or construct all the paths in the tree under construction that leads to leaf nodes where the action of the leaf that are being inserted is to be preferred.

When we insert  $L_i = (a_i, q_i, S_i)$ , we will always either encounter an internal branch node that defines a split on a variable and has two subtrees, or we encounter a leaf node storing an action, a Q-value and a state partition. We therefore define a general Put function (Algorithm 3), that takes a root node and a leaf triplet to be inserted and decides what to do based on the type of the root node (either a branch node or a leaf node).

Algorithm 3 Build decision tree from leaf nodes of Q-tree

```
1: function PUT(root, Leaf(a, q, S))
                                                                                                  return PUTATBRANCHNODE(root, Leaf(a, q, S):

| Putatbranch | Putatb
                                                               if root is Node then
2:
3:
4:
                                                               else
                                                                                                                                                                                                  PUTATLEAFNODE(root, Leaf(a, q, S))
5:
```

We will first deal with situation where we encounter an internal node,  $N_i$ , which splits on variable  $V_k$  at bound b. We now need to check on what side of this split  $S_i$  falls (it might be both). So we test on both  $u_{i,k} > b$  and  $l_{i,k} < b$ . If the first check is true, then we know that  $S_i$  defines an area for  $V_k$  that can be larger than b, why we have to visit the right ('high') subtree of  $N_i$ . Likewise for the latter test, only then we have to continue our insertion in the left ('low') subtree. Note that both tests can be true.

We do, however, need to keep track of the implicit limitations we put on  $S_i$  as we go along. When continuing our insertion of  $L_i$  in the subtree of  $N_j$  defined by  $V_k > b$ , then we should reflect in  $S_i$  that now  $V_k$  has a lower bound b, that is, we should set  $l_{i,k} = b$ . We do this in the algorithm through an implicit helper function SetLower(state, var, bound) (and likewise SetUpper for updating the upper bound). The pseudocode for this is given in Algorithm 4.

Algorithm 4 PUT Q-leaf into an internal node

```
PUTATBRANCHN-
1: function
   ODE(Node(V_k, b, low, high), Leaf(a_i, q_i, S_i))
       if l_{i,k} < b then
2:
            S_i' \leftarrow \text{SetUpper}(S_i, V_k, b)
3:
            low \leftarrow Put(low, Leaf(a_i, q_i, S'_i))
4:
       if u_{i,k} > b then
5:
            S_i' \leftarrow \text{SetLower}(S_i, V_k, b)
6:
            high \leftarrow Put(high, Leaf(a_i, q_i, S'_i))
7:
       return Node(V_k, b, low, high)
8:
```

The second case is when we encounter a leaf node during the insertion operation. We denote this node as  $L_t$ 

to indicate that it is a leaf already present in the tree under construction. First, we check if the Q-value of  $L_t$  is better than that of  $L_i$ , in which case we do nothing and abort the insert operation. If  $q_i$  on the other hand is the better option, then we need to insert  $L_i$  but in a way that respects  $S_i$ .

It is guaranteed at this stage, that  $S_t$  contains  $S_i$ , that is  $u_{t,j} \geq u_{i,j}$  and  $l_{t,j} \leq l_{i,j}$  for all  $j = 1, 2, \ldots, k$ . But this also means that in the cases where the bounds on  $S_t$  are strictly larger or smaller than those of  $S_i$  then we need to insert a new internal node to ensure this partition before we can insert  $L_i$ . In other words, if  $u_{t,j} > u_{i,j}$ , then we need to create a branch node that splits on  $V_j$  at bound  $b = u_{i,j}$  and whose right ('high') subtree is the original leaf  $L_t$  but with an updated state  $S_t$  where  $l_{t,j} = u_{i,j}$ . The left ('low') subtree should also, for a start, be set to  $L_t$  but then we continue the insert operation on this side, either creating more branch nodes or eventually inserting  $a_i$  and  $q_i$  in place of  $a_t$  and  $q_t$ .

The pseudocode for the function is given in Algorithm 5.

```
Algorithm 5 Put Q-leaf into a leaf node
```

```
highState \leftarrow SetLower(state, var, bound)
        lowState \leftarrow \texttt{SetUpper}(state, var, bound)
        high \leftarrow \text{Leaf}(action, q, highState)
 4:
        low \leftarrow \texttt{Leaf}(action, q, lowState)
 5:
        return Node(var, bound, low, high)
 1: function
                                                   PUTATLEAFN-
    ODE(Leaf(a_t, q_t, S_t), Leaf(a_i, q_i, S_i))
        if q_t \leq q_i then
 2:
            return Leaf(a_t, q_t, S_t)
 3:
        for (l_{t,j}, u_{t,j}) in S_t do
 4:
            if l_{t,j} < l_{i,j} then
 5:
 6:
                 newNode \leftarrow SPLIT(a_t, q_t, V_i, l_{i,j})
 7:
                 return PUT(newNode, Leaf(a_i, q_i, S_i))
 8:
            else if u_{t,j} > u_{i,j} then
                 newNode \leftarrow SPLIT(a_t, q_t, V_i, u_{i,i})
 9:
                 return PUT(newNode, Leaf(a_i, q_i, S_i))
10:
        return Leaf(a_i, q_i, S_i)
11:
```