Deep Learning of Heuristics for Domain-independent Planning

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Abstract: Automated planning deals with the problem of finding a sequence of actions leading from a given state to a desired state. The state-of-the-art automated planning techniques exploit informed forward search guided by a heuristic, where the heuristic (under)estimates a distance from a state to a goal state. In this paper, we present a technique to automatically construct an efficient heuristic for a given domain. The proposed approach is based on training a deep neural network using a set of solved planning problems from the domain. We use a novel way of generating features for states which doesn’t depend on usage of existing heuristics. The trained network can be used as a heuristic on any problem from the domain of interest without any limitation on the problem size. Our experiments show that the technique is competitive with popular domain-independent heuristic.

1 INTRODUCTION

Heuristic learning is a relatively new field which studies how machine learning (ML) can be used to construct a heuristic used in informed forward-search algorithms such as $A^*$ and IDA*. The objective is to train a regression-based ML model to be able to estimate goal-distances of states and then to use the trained model as a heuristic function during search.

Automated planning, which deals with finding a sequence of actions leading to a goal state, exploits heuristics heavily. Domain-specific heuristics are hand-tailored for a specific domain and hence they are very efficient. However, developing them requires tremendous human effort and access to expert knowledge. Domain-independent heuristics are more popular in automated planners as they are designed once and then they work across all domains. Nevertheless, none of them works well on all domains. Again, a significant human effort would be required to select the best performing heuristic for the problem at hand, otherwise the search might be inefficient. This issue can partially be solved by using a portfolio-planner.

In this paper, we present a ML technique that automates the process of developing domain-specific heuristics for planning. We work with the standard STRIPS planning and we use supervised learning, with a multi-layered feed-forward neural network as the ML model. Expert knowledge about the domain is extracted automatically from a set of training samples. Using ML techniques for this task can be advantageous because in this particular case, the training samples can be obtained without human assistance, unlike in typical applications of ML where samples need to be human-labeled.

Our technique falls into category of zero-learning as the heuristic is constructed from scratch, without any human-knowledge initially added. This approach allows us to combine wide usability of domain-independent heuristics with accuracy of domain-specific ones without any assistance of a human expert.

Our main contribution is twofold. We propose a novel way of assigning features to states based on counting subgraphs of a graph-based representation of the state. The method produces fixed-size feature vectors without depending on existing heuristics. Second, we study the effect of the choice of loss function used during the training, on performance of the learned heuristic during the search. We show that Mean Squared Error (MSE) alone might not be the best loss function for heuristic learning task, and also that low error on training and test sets doesn’t automatically imply good performance of the learned heuristic.
2 BACKGROUND AND RELATED WORKS

2.1 STRIPS Planning

We work with classical planning problems, that is, with finding a sequence of actions transferring the world from a given initial state to a state satisfying certain goal condition (Nau et al., 2004). World states are represented as sets of predicates that are true in the state (all other predicates are false in the state). For example the predicate \( \text{at}(s, l, 1) \) represents information that some object \( r1 \) is at location \( l \). Actions describe how the world state can be changed. Each action \( a \) is defined by a set of predicates \( \text{prec}(a) \) as its precondition and two disjoint sets of predicates \( \text{eff}^+(a) \) and \( \text{eff}^-(a) \) as its positive and negative effects. Action \( a \) is applicable to state \( s \) if \( \text{prec}(a) \subseteq s \) holds. If action \( a \) is applicable to state \( s \) then a new state \( \gamma(a, s) \) defines the state after application of \( a \) as \( \gamma(a, s) = (s \cup \text{eff}^+(a)) \setminus \text{eff}^-(a) \) Otherwise, the state \( \gamma(a, s) \) is undefined. The goal \( g \) is defined as a set of predicates that must be true in the goal state. Hence the state \( s \) is a goal state if and only if \( g \subseteq s \).

The satisficing planning task is formulated as follows: given a description of the initial state \( s_0 \), a set \( A \) of available actions, and a goal condition \( g \), there is a sequence of actions \( (a_1, \ldots, a_n) \), called a solution plan, such that \( a_i \in A \), \( a_i \) is applicable to state \( s_0 \), each \( a_i \) s.t. \( i > 1 \) is applicable to state \( \gamma(a_{i-1}, \ldots, a_1, s_0) \), and \( g \subseteq \gamma(a_n, a_{n-1}, \ldots, a_1, s_0) \). Assume that each action \( a \) has some cost \( c(a) \). An optimal planning task is about finding a solution plan such that the sum of costs of actions in the plan is minimized.

In practice, the planning problem is specified in two components: a planning domain file and a planning problem file. The domain file specifies the names of predicates describing properties of world states and actions that can be used to change world states. The problem file then specifies a particular goal condition and an initial state and hence it also gives the names of used objects (constants) and their types.

In the rest of the text we use the following notation. \( P \) denotes a planning problem, \( s^p_0 \) denotes the initial state of \( P \), \( S^p \) the set of all states of \( P \), \( \text{Dom}(P) \) a set of all planning problems from the same domain as \( P \) and \( S^{\text{dom}(P)} \) a set of all states of all problems from the same domain as \( P \). For a state \( s^p \in S^p \), \( h^p(s^p) \) denotes the goal-distance of \( s^p \) in \( P \), i.e. the cost of the optimal plan from \( s^p \), or \( \infty \) if there is no path from \( s^p \) to a goal state.

2.2 Heuristic Learning

A heuristic learning (HL) system works with a set of training samples, where each training sample is a pair \( (s_i, h^*(s_i)) \), \( s_i \) is a state of some planning problem and \( h^*(s_i) \) is its goal-distance. The system involves a ML model \( M \) that is trained to predict \( h^*(s_i) \) from \( s_i \). Most of ML models work with fixed-size real valued vectors as their inputs, hence another component is required that transforms states into this form. We call this component a features generator and denote it by \( F \). We call \( F(s) \) the features of \( s \), and \( h^*(s) \) the target of \( s \). There are several variants of HL based on their usage scenarios:

**Type I:** several problems \( P_1, P_2, \ldots, P_j \) and \( P \) from the same domain are given as input and the task is solving the problem \( P \) as quickly as possible. Time required for generating the training data and training the model is considered a part of the solving process. The ML model is trained specifically for \( P \), it is not intended to work on different problems.

**Type II:** several problems \( P_1, P_2, \ldots, P_j \) from the same domain \( \text{Dom} \) are given and the task is solving new problems from \( \text{Dom} \). Time required for generating the training data and training the model is NOT considered a part of the solving process. It is considered a pre-processing, or a domain analysis phase. The ML model captures knowledge about the whole domain, i.e., it can generalize to other, previously unseen problems from the same domain. The training phase can in this case take several hours or even days. After the model is trained, new problems from the same domain can be solved quickly. The Type II HL system requires a much more flexible features generator, as features of states of different problems must be comparable. Those different problems might contain different numbers of objects and might have different goal conditions.

**Type III:** Training samples from several different domains are given and the ML model serves as a multi-domain heuristic, or a domain-independent heuristic. To our best knowledge, no serious attempt has been made in this area.

In this paper, we work with the Type II HL and our approach is domain-independent in a sense that the domain from which the training problems come might be arbitrary. We conducted experiments on domains without action costs but the technique is applicable to domains with costs as well.
2.3 Related Works

Many attempts have been made to utilize ML in planning or in general search (Jiménez et al., 2012). ML has been used to learn reactive policies (Grosheva et al., 2017; Martín and Geffner, 2004; Grosheva et al., 2018), control knowledge (Yoon et al., 2008), for plan recognition (Bisson et al., 2015) and for other planning-related tasks (Konidaris et al., 2018). ML tools are also often used to combine several heuristics (Samadi et al., 2008; Fink, 2007) and in particular to help portfolio-based planners to efficiently combine multiple search algorithms (Cenamor et al., 2013). A lot of papers exist that utilize ML in neoclassical planning paradigm (partial observability, non-deterministic actions, extended goals etc.), like robotics (Takahashi et al., 2019). These techniques are not directly related to our work.

Heuristic learning was investigated by (Arfaee et al., 2010) where the authors use a bootstrapping procedure with a NN to successively learn stronger heuristics using a set of small planning problems for training. The paper proposed an efficient way of generating training data based on switching between learning and search phases. The technique became popular and was successfully used by other authors (Chen and Wei, 2011; Thayer et al., 2011). We use a modified version of this technique as well. A domain-independent generalization of this approach was published later (Geissmann, 2015).

Most papers deal with the Type I HL scenario and almost all of them use a set of simple heuristics as features (Arfaee et al., 2011; Brunetto and Trunda, 2017). A serious attempt to use other kind of features was made in (Yoon et al., 2008).

Authors mostly use simple ML models like linear regression or a shallow NN. ML models are often just used as a tool and ML-related issues like generalization capabilities, number and distribution of training samples, choice of loss function, etc. are not analyzed at all. Few exceptions exist: in (Thayer et al., 2011) the authors proposed a modification to the loss function used during the training to bias the model towards under-estimation which increased quality of solutions found during the subsequent search.

3 THE FRAMEWORK

We will use heuristic learning in automated planning using the following framework. First, during the training phase, the deep NN will learn the heuristic from example plans. Then, during the deployment phase, the obtained NN will be used to calculate heuristic values that will be exploited by A* search to find plans. The focus of this paper is on the training phase, in particular, on novel approach to generating features for training and on selecting appropriate loss function.

3.1 Obtaining the Training Data

Ideally, the training data should be given as inputs. That might be possible in some specific situations when historical data are available, but in general the training data need to be generated. This involves two tasks: generating states $s_i$ and computing $h'(s_i)$ for those states. As computing $h'(s_i)$ is very time-consuming, majority of existing works use states for which it is easy to compute $h'(s_i)$, such as the states close to goal.

From the ML perspective, it is important that training data come from the same probability distribution as the data encountered during the deployment. This would require to predict what kind of states will $A^*$ expand during the search. Making such predictions is tricky as the set of expanded nodes depends on the heuristic used which depends on how well the model is trained and that depends back on the choice of training data.

We adopt a popular technique (Arfaee et al., 2010), which solves this issue by an iterative procedure that combines training and search steps. The first set of training samples is generated by random walks from the goal state. Then, in each iteration, the model is trained on current set of samples and a time-limited search is performed on the training problems using the trained model as the heuristic. States that the algorithm expanded are collected and used as training samples in the next iteration. This process continues until sufficient amount of samples is generated or all training problems can be solved within the time limit.

To speed up the training phase, we use ad-hoc solvers to calculate goal-distances of samples. This allows us to work with larger training problems in reasonable time and not having to rely on backward search to calculate $h'$. Pseudocode for the training phase is presented as Algorithm 1.

4 GENERATING FEATURES

The important step in ML is selecting features that will be used in learning. Given a set of planning problems $\{P_1, P_2, \ldots, P_j\} \subset Dom$, the feature generator $F$ realizes a mapping $F: S^{Dom} \rightarrow \mathbb{R}^k$, i.e., assigns a real valued vector to any state of any problem from the domain of interest.
Algorithm 1: Training phase.

**Input:** Set of planning problems \( \{P_j\} \subset \text{Dom} \) used for training features generator \( F \)

**Output:** Trained model \( M \) that realizes mapping from \( F[S^{\text{dom}}] \rightarrow \mathbb{R} \)

1. \( L := 1 \);
2. \( \text{trainingStates} := \) initial states of all \( P \in \{P_j\} \);
3. repeat
   4. compute \( h^*(s_i) \) for each state \( s_i \in \text{trainingStates} \);
   5. assign features \( f_i = F(s_i) \) to all states \( s_i \in \text{trainingStates} \);
   6. \( M := \) train neural net on data \( \{(f_i, h^*(s_i))\} \);
   7. foreach problem \( P \in \{P_j\} \) do
      8. run IDA* on \( P \) with time limit \( L \) using \( M \) as heuristic;
      9. \( T := \) states of \( P \) that were expanded during the search;
   10. \( \text{trainingStates} := \text{trainingStates} \cup T \);
   11. end
   12. \( L := L + 1 \);
4. until termination criterion is met;
5. return \( M \); 

Length of the feature vector needs to be fixed and independent of the specific planning problem. Features should also be informative in a sense that states with different goal-distances should have different features, and comparable among problems from the whole domain so that knowledge is transferable to previously unseen problems.

When assigning features to state \( s \in S^P \), properties of \( P \) that affect \( h^*(s) \) have to be taken into account, namely the set of available actions and the goal condition. The model is trained on problems from a single domain and for such problems the set of actions is always the same so it is not necessary to encode it into features of states. Goal conditions, however, must be encoded so that the learned knowledge is transferable to problems with different goal states.

### 4.1 Heuristics as Features

Vast majority of papers on HL use a fixed set of simple heuristics as features. Given a sequence of heuristics \( H = (h_1, h_2, \ldots, h_k) \), we can define \( F^H(s) = (h_1(s), h_2(s), \ldots, h_k(s)) \). Most papers use a set of pattern database heuristics (PDBs). This approach is popular as the feature vector has fixed length, is quite informative and its computation is fast.

This approach is advantageous in the Type I HL scenario but not so in Type II that we deal with. PDB is based on a pattern: a set of objects from the planning problem. It is possible to us it if all the problems are given in advance. In the Type II scenario, however, the model needs to be applicable to new, unseen problems from the domain. If we use a fixed set of PDBs, new problems might not contain the same objects, and even if they do, meaning of those objects might be different so the features would not be comparable.

Also we want to avoid dependency on existing human-designed heuristics as such dependence might prevent the model from achieving super-human performance. Development in the fields of sound, image, and language processing during the last ten years showed that ML systems with little to none expert knowledge encoded might achieve better results than sophisticated human-designed tools.

### 4.2 Graph-based Features

We use a direct encoding of the state to an integer-valued vector. We transform the PDDL representation of both the current state and the goal condition to a labeled graph and use this graph to generate features. We select a set of small connected graphs and then use the number of occurrences of these graphs in the original graph as features. The idea is inspired by Bag-Of-Words model (Goldberg, 2017) that is used to assign fixed-length feature vectors to variable-length texts by counting number of occurrences of selected words or phrases.

#### 4.2.1 Object Graph

An object graph for a state \( s \) of a planning problem \( A \) (denoted \( G(s^A) \)) is a vertex-labeled graph \( G(s) = (V, E) \) defined as follows. The set of vertices is composed by four disjoint sets. There is a vertex \( v_c \) for every constant \( c \) in the problem, a vertex \( v_{P} \) for every predicate symbol used in the definition of the problem, a vertex \( v_q \) for every instantiated predicate \( q \) that is true in \( s \), and a vertex \( v_{\neg q} \) for every predicate \( q \) in goal conditions of \( A \) (we don’t support negative goal conditions as they can be compiled-away). The set \( E \) contains an edge \( e_{dp} \) from \( v_q \) or \( v_{\neg q} \) to \( v_P \) if instantiated predicate \( q \) uses the predicate symbol \( P \), and an edge \( e_{cq} \) from \( v_c \) to \( v_q \) or \( v_{\neg q} \) if instantiated predicate \( q \) contains constant \( c \).

The labelling function \( w : V \mapsto \mathbb{N}^0 \) looks as follows. Every \( v_c \) is assigned the same number 0. Every \( v_P \) is assigned a unique number from \( [1, 2, \ldots, \#P] \) where \( \#P \) is the number of predicate symbols. Every \( v_q \) is assigned the same number \( \#P + 1 \) and every \( v_{\neg q} \) is assigned the same number \( \#P + 2 \). Types are treated
as unary predicate symbols. Labels were intentionally chosen such that the number of labels does not depend on the size of the problem.

In figure 1 there is an example of object graph for the initial state of the problem *pfile2* of the *zenotravel* domain. Constants are colored red, types blue, predicate symbols pink, initial predicates green and goal predicates gold.

The object graph is an equivalent representation of the current state and goal condition. The original PDDL representation of the state and the goal can be reconstructed back from the graph.

### 4.2.2 Generating Features

The size of the object graph depends on the number of constants and the number of valid predicates in the state and in the goal. Hence we need a method to encode information from \(G(s)\) in a feature vector whose length is independent of state and goal. This is where we will use the node labels and the subgraph-counting method.

Let \(B_k^q\) be a sequence of all connected non-isomorphic vertex-labeled graphs containing at most \(q\) vertices where the labels are from \(\{1,2,\ldots,k\}\). See figure 2 for an example of graphs \(B_k^3\) — colors represent labels.

**Occurrence** of a graph \(G_1\) in graph \(G_2\) is a set of vertices \(T\) of \(G_2\) such that induced subgraph of \(G_2\) on \(T\) is isomorphic to \(G_1\).

Given a state \(s\) and \(q \in \mathbb{N}\), the feature vector of \(s\) (denoted \(F_q(s)\)) is an integer-valued vector of size \(|B_k^q|\) whose \(i\)-th component is the number of occurrences of the \(i\)-th graph from \(B_k^q\) in \(G(s)\).

### 4.2.3 Example

Consider the graph \(G\) in the left-hand side of figure 2 with two different labels represented by colors. We use \(q = 2\), i.e. we count occurrences of connected subgraph of size up to 2. On the right-hand side of figure 2 there are graphs from \(B_2^2\) that we will use to assign features to \(G\). The resulting vector is \(F_2(G) = (5,3,5,2,1)\) which corresponds to number of occurrences of the individual graphs in \(G\). For example: occurrences of the second graph are \(\{2\}\), \(\{7\}\), \(\{8\}\), occurrences of the fourth one are \(\{3,6\}\) and \(\{5,6\}\), etc.

The length of \(F_q(.)\) can be controlled by adjusting the parameter \(q\). With low \(q\), the vector will be short and will contain less information about the state but its computation will be faster, and vice versa. Given a graph with \(n\) vertices, it is not known whether or not \(F_q(G)\) uniquely determines the graph for some \(q < n\).

It is an open problem in graph theory known as the **Reconstruction conjecture**.

Length of the vector is at most \(\sum_{i=1}^{q} 2^{\frac{(i-1)}{i(i-1)}} C_i(k)\), where \(k\) is the number of labels, \(q\) is the maximum size of subgraphs considered, \(2^{\frac{(i-1)}{i(i-1)}}\) is the number of graphs on \(i\) vertices and \(C_i(k) = \frac{(k+i-1)!}{2!(k-1)!i!}\) is the number of combinations with repetition of size \(i\) from \(k\) elements. In practice, \(|F_q(.)|\) is much lower since we only use subgraphs that occurred at least once in the training data and most graphs do not occur due to the way how the object graph is defined. E.g. every predicate symbol has its own label \(l_i\) and every object graph contains exactly one vertex with such label so subgraphs that contain more than one vertex labeled \(l_i\) can never occur.

From the planning perspective, features capture relations between objects in the given state. E.g. in blocksworld, occurrence of a certain subgraph of size 4 can capture the fact, that there are 2 blocks \(A\) on top of \(B\) and \(B\) is not correctly placed (so both of them have to be moved), etc.

### 4.2.4 Computing features from Scratch

Given the object graph \(G(s)\), feature vector \(F_q(s)\) can be computed by a recursive procedure that iterates through all connected subgraphs with size up to \(q\) in the graph and its time complexity is proportional to the number of such subgraphs. It is difficult to estimate the number in general as it strongly depends on the structure of the graph. E.g. a cycle with \(n\) vertices contains just \(n\) connected induced subgraphs of size \(q < n\), while a clique on \(n\) vertices contains \(\binom{n}{q}\) such subgraphs. The number depends on the edge-connectivity of the graph as well as on degrees of vertices. Experiments show that the time complexity grows exponentially in both \(n\) (size of the graph) and \(q\).

### 4.2.5 Computing Features Incrementally

\(A^*\) expands nodes in a forward manner and two successive states differ only locally. It is therefore possible and useful to calculate the features incrementally. Given a state \(s\), its feature vector \(F(s)\) and an action \(a\), we can calculate \(F(\gamma(a,s))\) of the successor state without having to enumerate all its subgraphs again.

Unfortunately, \(F(s)\) and \(a\) alone are not sufficient to determine features of the successor. For any fixed \(q\) there exist states \(s_1 \neq s_2\) and action \(a_1\) such that \(F_q(s_1) = F_q(s_2)\) but \(F_q(\gamma(a_1,s_1)) \neq F_q(\gamma(a_1,s_2))\) so a more sophisticated approach is needed. Applying action to a state \(s\) can be viewed as performing some local changes in \(G(s)\). These changes can be
decomposed into a sequence of several atomic operations of 3 types: AddVertex, RemoveVertex and AddEdge. For example, in Zenotravel, there is an action $a = \text{load}(\text{person1, city1, plane1})$. Given $G(s)$, we can construct $G'(a, s)$ by first removing vertex that represents predicate $\text{at}(\text{person1, city1})$, then adding vertex for predicate $\text{in}(\text{person1, plane1})$ and then successively adding edges between the new vertex and vertices for $\text{in}$, $\text{person1}$ and $\text{plane1}$.

Given graph $G$, its vertex $v$ and a set of graphs $B_q$, we define contribution of a vertex $v$ to $F(G)$ denoted by $C(v)$ as a set of all occurrences of graphs from $B_q$ in $G$ which intersect with $v$. We will now show how each of the three atomic operations can be performed incremen tally given $G(s), F(s)$ and $C(v)$ for every vertex of $G(s)$.

**Remove Vertex:** For each $c \in C(v)$, remove $c$ from every $C(v')$ that contains it, decrease values in $F(s)$ accordingly. Remove $v$ from $G(s)$.

**Add Vertex:** Add $v$ to $G(s)$, add one new occurrence of a subgraph containing a single vertex with the given label to $C(v)$, increment the the corresponding element of $F(s)$.

**Add Edge:**
1. replace every occurrence $c \in C(v_1) \cap C(v_2)$ by occurrence of a graph with the same vertex set and one more edge added at the corresponding location.
2. for every occurrence $c \in C(v_1) \setminus C(v_2)$ such that $|c| \leq q - 1$ create a new occurrence on vertices $c \cup \{v_2\}$. If $c$ contained some vertex adjacent to $v_2$, edges between these vertices and $v_2$ will be taken into account when determining which graph from $B_q$ occurred on $c \cup \{v_2\}$.
3. repeat the previous step symmetrically for $v_2$, then add the edge to $G(s)$.

Using a carefully designed data structure, we can perform AddVertex, RemoveVertex as well as the step 1 of AddEdge in time $O(1)$, steps 2 and 3 can be performed in time $O(|C(v_1)|)$ and $O(|C(v_2)|)$ respectively.

## 5 ERROR FUNCTION FOR THE TRAINING

We train the network by a standard gradient-descent optimizer which iteratively updates parameters of the network in order to minimize the given loss function. We are solving a regression task - predicting a real number for each state. We experimented with two loss functions: standard MSE defined as

$$\text{MSE} = \frac{1}{n} \sum_{i=1}^{n} (Y_i - \hat{Y}_i)^2$$

and a MSE transformed by a logarithm (denoted LogMSE), defined as

$$\text{LogMSE} = \sum \log(Y_i) - \log(\hat{Y}_i)$$

where $Y_i$ is target of the i-th sample (i.e., the real goal distance of the i-th state), $\hat{Y}_i$ is output of the model on the i-th sample (i.e. the value that would be used as a heuristic estimate for the i-th state) and $n$ is the number of samples. The sum goes over all samples.

Figure 3 presents a histogram of fit of the trained model when MSE is used. On the x-axis there is difference between target value and output, the height of the column represents the number of samples that fall into each category. Yellow columns show training data, purple ones show test data. In Figure 4 we can see accuracy of the model on samples with different targets (blue columns). On the x-axis there is target and the height of the column represents average of absolute values of absolute error among all samples with
corresponding target value. The model is well trained having small error on both training and test data. The use of MSE loss function leads to an even distribution of the error among all samples regardless of their target (up to goal distance about 50). For example, the model makes error of ±1.5 on states 50 steps from goal, but makes about the same error (±1.5) also on states that are 2 steps from goal.

During the deployment phase, however, the model perform poorly. The heuristic is very accurate on states far from goal, but our analysis showed that making relatively large mistakes on states close to goal hurts the performance a lot.

In order to improve the search efficiency, the accuracy of the heuristic on states close to goal (i.e. with target 0-10) needs to be much higher. This can be achieved by using LogMSE as the loss function. LogMSE minimizes the average of relative error, i.e., the ratio between target and output which enforces low absolute error on samples close to goal and tolerates larger absolute error on samples further from goal.

Figure 4 illustrates this behavior as it compares absolute and relative error on training and test samples for both loss functions. MSE (in blue) makes large relative error on states close to goal while LogMSE (orange) performs much better on these states but has slightly larger error on states further from goal. The total sum of error is similar for both functions but its distribution with respect to the target is quite different. Experiments show that using LogMSE leads to better performance during the search.

6 EXPERIMENTS

We conducted experiments on two standard benchmark domains: zonotravel and blocks because it is easy to obtain training data for these domains. For the purpose of the experiments, we implemented ad-hoc solvers for the two domains and used them to generate training data. Our solvers are based on a genetic algorithm combined with a greedy search, they are capable of solving most problems within a few seconds and provide optimal or close-to-optimal solutions. We test the method on 20 problems available for zonotravel1 and the first 27 problems from blocks2.

For each problem $P$, we train the model using the other problems from the domain (except $P$) as the training data, and then use the trained model as a heuristic with an $A^*$ algorithm to solve $P$. We compare the quality of the resulting heuristic with the Fast-Forward heuristic $h_{FF}$ (Hoffmann and Nebel, 2001). The heuristic has been around for quite some time now but it is still often used as a baseline in experiments, like in (Höller et al., 2019) for example.

The NN we used have 5 hidden layers with sizes of (256, 512, 128, 64, 32) neurons respectively, and two Dropout layers. We used ReLU activation function, Xavier weight initialization and Adam as the training algorithm (Goodfellow et al., 2016). The last layer contained a single neuron with a linear activation to compute the output. Architecture of the network was chosen according to best practices for this kind of scenario. The network is large enough to create efficient representation of the data and drop-out layers prevent overfitting. Similarly to other zero-learning scenarios, the amount of time required for training is quite high. For every problem, training the net took about 8 hours using over 1 million training samples for zonotravel and over 1.8 million for blocks.

We experimented with values of parameter $q$ (size of subgraphs) from 2 to 4. For values larger that 4, computing features of states is too costly and the heuristic is not competitive. We experimented with the two loss functions - MSE and LogMSE, and we also tried to include value of the $h_{FF}$ among the features of states. I.e., we first trained the network having

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1 api.planning.domains/json/classical/problems/17
2 api.planning.domains/json/classical/problems/112
Figure 5: Results of experiments.

only the graph-based features as its inputs and then another network that used both graph-based features and $h_{FF}$ value of the state as its inputs. We conducted experiments for all combinations of these parameters: $q \in \{2, 3, 4\}$, $\text{lossFunction} \in \{\text{MSE}, \text{LogMSE}\}$, $\text{FFasFeature} \in \{\text{true}, \text{false}\}$. This gives us 12 different neural net-based heuristics.

We used all 13 heuristics (12 NN-based + $h_{FF}$) to solve each of the problems. Search time was capped at 30 minutes per problem instance. None of the heuristic is admissible so they don’t guarantee finding optimal plans. We compared performance of heuristics using the IPC-Score. Given a search problem $P$, a minimization criterion $R$ (e.g. length of the plan) and algorithms $A_1, A_2, \ldots, A_k$, the IPC-Score of $A_i$ on problem $P$ is computed as follows: $\text{IPC}_R(A_i, P) = 0$ if $A_i$ didn’t solve $P$, or $\frac{R^*}{R_i}$ otherwise, where $R_i$ is value of the criterion for the i-th algorithm and $R^* = \min_i \{R_i\}$. For every problem $P$, $\text{IPC}_R(A_i, P) \in [0,1]$ and higher means better. We can then sum up the IPC-Score over several problem instances to get accumulated results. The IPC-Score takes into account both number of problems solved as well as quality of solutions found. We monitor four criteria: total number of problems solved, IPC-Score of time, IPC-Score of plan length and IPC-Score of number of expansions.

Figure 5 shows results for the two domains.

We can see that network trained by LogMSE is superior to the one trained by MSE in all criteria on both domains. As expected, higher values of $q$ lead to a more accurate heuristic: the number of expanded nodes as well as plan length are better. The difference is apparent especially for values 2 and 3. Using value $q = 4$ still helps but computing features in this case is slower and so $A^*$ expands less nodes per second and overall results are not that much better than for $q = 3$.

Adding $h_{FF}$ as feature has a mixed effect. It is very helpful on blocks domain when $q \in \{2, 3\}$, but not much helpful when $q = 4$. See figure 6. This indicates that subgraphs of size 2 and 3 cannot capture useful knowledge about a blocks problem hence the network rely on $h_{FF}$ as the source of information. Subgraphs of size 4 seem to be able to provide the required knowledge already and adding $h_{FF}$ doesn’t help anymore. This phenomenon is domain-dependent and should be analyzed further in the future. In general, adding $h_{FF}$ improves accuracy of the NN so the resulting heuristic is more informed which improves both number of expansions and plan length. Due to the slow-down, though, adding $h_{FF}$ doesn’t often improve number of problems solved.

Figure 6: Number of problems solved in blocks domain (sum of both MSE and LogMSE). On the X-axis there is $q$ value, blue columns correspond to networks trained without using $h_{FF}$ as feature, orange columns show networks trained with $h_{FF}$ included.

Among the neural-based heuristics, the setting with $q = 4$, $h_{FF}$ added and LogMSE performs best. If we compare it with the $h_{FF}$, we see that our method vastly outperforms the baseline on blocks where it solved 26 out of 27 problems while $h_{FF}$ can only solve 8 problems. Even on problems solved by both methods, the NN heuristic finds shorter plans and expands less nodes. On the zenotravel domain, our method outperforms $h_{FF}$ in all criteria except Time. As $h_{FF}$ can find suboptimal plans very quickly in zenotravel, it is difficult to achieve better score even though our method solved more problems within the time limit.
6.1 Performance Guarantees

The resulting heuristic is neither admissible nor $\varepsilon$-admissible for any $\varepsilon$ but we can still provide some theoretical guarantees of solution quality. We will only provide here a simple bound and sketch of a proof. Tighter bounds together with formal statements of theorems and full proofs will be published in a separate paper.

Due to the stochastic nature of the NN-training algorithm, output of the trained network on state $s$ (denoted $H(s)$) must be considered a random variable. Cost of the solution that $A^*$ finds with $NN$ as heuristic from state $s$ (denoted $A_H(s)$) is a random variable as well. Assuming that $\forall s_t \neq s_j, H(s_t)$ and $H(s_j)$ are independent, we can state the following theorem.

**Theorem 1.** If for all states $s$: $E[H(s)] = h^*(s)$, then

$$\forall c > 1 : \text{Prob}[A_H(s_0) \geq c \cdot (h^*(s_0))^2] < \frac{1}{c^2}$$

**Proof (sketch).** Let’s denote by $Opt(s_0)$ the optimal path from $s_0$ to goal (a sequence of states). The weighted $A^*$ efficiency theorem (Pearl, 1984) states that

$$\forall \varepsilon \geq 1 : \forall s : h(s) \leq \varepsilon \cdot h^*(s) \Rightarrow A_h(s_0) \leq \varepsilon \cdot h^*(s_0)$$

By a contraposition of the previous, we have

$$\forall \varepsilon \geq 1 : A_H(s_0) > \varepsilon \cdot h^*(s_0) \Rightarrow \exists s_i \in Opt(s_0) : H(s_i) > \varepsilon \cdot h^*(s_i)$$

hence

$$P[A_H(s_0) > \varepsilon \cdot h^*(s_0)] \leq \frac{1}{\varepsilon}$$

Now for given $c > 1$, we set $\varepsilon = c \cdot h^*(s_0)$ which gives us the required bound.

The proof works for planning without action costs, i.e. where cost of every action is 1 but the theorem still holds for planning with action costs. □

We don’t require that all $H(s_t)$ are identically distributed. Assumptions of independence and unbiasedness of $H(s_t)$ can be justified by analyzing the bias-variance tradeoff for NNs (Hastie et al., 2001). NNs in general have high variance and low bias hence for a large enough network, $H(s)$ should be unbiased and $\forall s_t, s_j : H(s_t)$ and $H(s_j)$ should be close to independent. Tighter bounds can be acquired if we take variance of $H(s_t)$ into account.

7 CONCLUSIONS & FUTURE WORK

We presented a technique to automatically construct a strong heuristic for a given planning domain. Our technique is domain-independent and can extract knowledge about any domain from a given set of solved training problems without any assistance from a human expert. The knowledge is represented by a trained neural network.

We analyzed how the choice of loss function used during the training affects performance of the learned heuristic. We showed that the Mean Squared Error – the most popular loss function – is not appropriate for heuristic learning task and we presented a better alternative.

We developed a novel technique for generating features for states where we encode the state description directly without using existing heuristics. The method allows to compute features incrementally which is very useful in planning application. Our approach falls into category of zero-learning as it works without any human-knowledge initially added. The presented technique significantly outperforms a popular domain-independent heuristic $h_{FF}$ in both number of problems solved and solution quality. We have also provided a simple theoretical bound on solution quality when using learned heuristic, similar to bounds for weighted $A^*$.

As a future work, we will provide better bounds on solution quality and conduct larger experiments on
more domains to properly back the claims made in this paper.

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