Combining Optimal Path Search With Task-Dependent Learning in a Neural Network

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Abstract—Finding optimal paths in connected graphs requires determining the smallest total cost for traveling along the graph’s edges. This problem can be solved by several classical algorithms, where, usually, costs are predefined for all edges. Conventional planning methods can, thus, normally not be used when wanting to change costs in an adaptive way following the requirements of some task. Here, we show that one can define a neural network representation of path-finding problems by transforming cost values into synaptic weights, which allows for online weight adaptation using network learning mechanisms. When starting with an initial activity value of one, activity propagation in this network will lead to solutions, which are identical to those found by the Bellman–Ford (BF) algorithm. The neural network has the same algorithmic complexity as BF, and, in addition, we can show that network learning mechanisms (such as Hebbian learning) can adapt the weights in the network augmenting the resulting paths according to some task at hand. We demonstrate this by learning to navigate in an environment with obstacles as well as by learning to follow certain sequences of path nodes. Hence, the here-presented novel algorithm may open up a different regime of applications where path augmentation (by learning) is directly coupled with path finding in a natural way.

Index Terms—Graph neural networks (GNNs), navigation, reinforcement learning (RL), sequences, shortest path problem.

I. INTRODUCTION

This study addresses the so-called single-source shortest-path (SSSP) problem. Possibly, the most prominent goal for any path-finding problem is to determine a path between two vertices (usually called source and destination) in a graph, such that the total summed costs for traveling along this path are minimized. Usually, (numerical) costs are associated with the edges that connect the vertices, and in this way, costs are accumulated when traveling along a set of them. In the SSSP problem, the aim is to find the shortest paths (SPs) from one vertex (source) to all other remaining vertices in the graph.

The SSSP problem has a wide range of applications, e.g., in computer networks (SP between two computers), social networks (SP between two persons or linking between two persons), trading and finance (e.g., currency exchange), multiagent systems, such as games, task, and path planning in robotics, and so on, just to name a few.

The most general way to solve the SSSP problem is the Bellman–Ford (BF) algorithm [1], [2], [3], which can also deal with graphs that have some negative cost values. In this work, we present a neural implementation, which is mathematically equivalent to the BF algorithm with an algorithmic complexity, which is the same as for BF.

The neural implementation relies on the multiplication of activities (instead of adding of costs). As a consequence, it is directly compatible with (Hebbian) network learning, which is not the case for any additively operating path-planning algorithm. To demonstrate this, we are using Hebbian-type three-factor learning [4], [5], [6] to address SSSP tasks under some additional constraints, solved by the three-factor learning.

Our paper is structured as follows. First, we will provide an overview of the state-of-the-art methods and state our contribution with respect to that. Next, we will describe the details of the BF algorithm and the proposed neural network (NN-BF). This will be followed by a comparison between the BF algorithm and NN-BF and by examples of combining NN-BF-based planning with three-factor learning. Finally, we will conclude our study with a summary and provide an outlook for future work.

II. RELATED WORK

A. State of the Art

There are mainly two types of approaches to solve the SSSP problem: classical algorithms and approaches based on artificial neural networks. Classical algorithms exist with different complexity and properties. The simplest and fastest algorithm is breadth-first search (BFS) [7], [8]. However, it can only solve the SSSP problem for graphs with uniform costs, i.e., all costs equal to 1. Dijkstra’s algorithm [9] as well as the BF algorithm [1], [2] can deal with graphs that have arbitrary costs. From an algorithmic point of view, Dijkstra’s algorithm is faster than the BF algorithm; however, Dijkstra only works on graphs with positive costs, whereas the
BF algorithm can also solve graphs where some of the costs are negative. Furthermore, Dijkstra is a greedy algorithm and requires a priority queue, which makes it not so well suited for parallel implementation as compared with BF.

It is worth noting that there are two versions of the BF algorithm (see [3]). Both versions have the same basic algorithmic complexity but version 2 operates on graph nodes and not on graph edges (as version 1), which allows implementing version 2 in a totally asynchronous way where computations at each node are independent of other nodes [3]. In spite of this, version 2 is largely ignored in the literature, and users will usually be directed to version 1 when doing a (web-)search.

Some heuristic search algorithms, such as A* [10] and its variants (see [11], [12], [13], [14]), exist, which are faster than Dijkstra or BF, but they can only solve the single-source single-target SP problem. The most general, but slowest, algorithm to solve the SSSP problem is the Floyd–Warshall algorithm [15], [16], which finds all-pairs SPs (APSPs), i.e., all SPs from each vertex to all other vertices. Another way to solve the APSP problem is by using Johnson’s algorithm [17], which utilizes BF and Dijkstra’s algorithms and is—under some conditions—faster than the Floyd–Warshall algorithm (essentially, this is the case for sparse graphs).

Different algorithms exist, which utilize artificial neural networks to solve SP problems. Some early approaches, which were dealing with relatively small graphs (below 100 nodes), were based on Hopfield networks [18], [19], [20] or Potts neurons and a mean field approach [21]. These approaches, however, may not always give optimal solutions or may fail to find solutions at all, especially for larger graphs.

Some other bioinspired neural networks were proposed [22], [23], [24], [25], [26], [27], [28] for solving path-planning problems. These approaches work on grid structures, where activity in the network is propagated from the source neuron to the neighboring neurons until activity propagation within the whole network is finished. SPs can then be reconstructed by following the activity gradients. The drawback of these approaches is, however, that they are specifically designed for grid structures and cannot be applied at general graphs with arbitrary costs.

Several deep learning approaches were proposed to solve SP problems, for example, using a deep multilayer perceptron (DMLP; [29]), fully convolutional networks [30], [31], [32], or a long short-term memory (LSTM) network [33]. Inspired by classical reinforcement learning (RL) algorithms, e.g., Q learning, SARSA [34], some deep RL algorithms had been employed [35], [36], [37] too. These approaches are designed to solve path-planning problems in 2-D or 3-D spaces and cannot deal with graphs with arbitrary costs.

In addition to this, deep learning approaches based on graph neural networks (GNNs) have been employed to solve path problems, too [38], [39], [40], [41], [42], [43], mostly in the context of relation and link predictions. While deep learning approaches may lead to a better run-time performance as compared with classical approaches due to fast inference, all deep learning-based approaches need to learn their (many) synaptic weights before they are functional. This usually requires a large amount of training data. Another disadvantage of these approaches is that optimal solution is not guaranteed, since networks intrinsically perform function approximation based on the training data. Moreover, in some of the cases, networks may even fail to find a solution at all, especially, in cases where training data are quite different from new cases (generalization issue for out-of-distribution data).

Recently, graph-based neural networks have been proposed too, such as spatial–temporal graph convolutional networks (ST-GCN) for skeleton-based action recognition [44], spatial–temporal graph network for video supported machine translation [45], and graph transformer network for zero-shot temporal activity detection (TN-ZSTAD) from videos [46]. Other deep networks are based on scene graph generation (SGG), e.g., conditional random field (CRF)-based SGG [47], TransE-based SGG [48], convolutional neural network (CNN)-based SGG [49], recurrent neural network (RNN)/LSTM-based SGG [50], [51], [52], and GNN-based SGG [53], [54]. While these approaches have been mostly applied for text and image/video analysis, such as text-to-image generation, image/video captioning, image retrieval, scene understanding, and human–object interaction (for a comprehensive survey, see [55]), none of them have been exploited to specifically address SP/SSSP problems.

Furthermore, some approaches exist for an automated search (tuning) of generative adversarial network (GAN) architectures, such as neural architecture search (NAS) [56], [57], AutoGAN [58], and automated generative adversarial network (AGAN) [59], and for the simultaneous learning of network architecture parameters and weights (ZeroNAS, [60]). Although these approaches are related to our proposed method, there are two principle differences. First, the aforementioned approaches are used to optimize network architectures or architecture and weights to perform discriminating tasks, e.g., image classification or object detection, whereas in our approach, we are dealing with the SSSP problem, where we build the network architecture and update connection weights based on local learning rules. Hence, we do not specifically optimize network architecture and weights to map input to outputs. Second, in contrast to NAS approaches, weights in our network correspond to costs, which allow using activity propagation within the network to find optimal solutions for the SSSP tasks.

B. Contribution

The comparison of the above discussed approaches for solving SP/SSSP problems is graphically summarized in Fig. 1, and details are provided in Table I. Here, we compare our approach with respect to the following criteria: 1) ability to deal with positive and negative costs; 2) solution optimality; and 3) algorithmic complexity. Given these criteria, one can see that our approach most closely relates to the neural approaches by Xia and Wang [61] and Zhang and Li [62]. The approach by Xia and Wang [61] utilizes a relatively complex RNN architecture. It can deal with optimal and negative costs and find optimal solutions, but it has a higher complexity as compared with the BF algorithm. Zhang and Li [62],...
as in our approach, proposed a neural implementation of the BF algorithm with the same complexity; however, it uses additive activity propagation (akin to additive cost propagation in the BF algorithm) as compared with multiplicative activity propagation as proposed by our algorithm. Moreover, neither Xia and Wang [61] nor Zhang and Li [62] show a combination of addressing the SP problem together with neural learning.

In summary, we present a neural implementation of the BF algorithm, which finds optimal solutions for graphs with arbitrary positive and (some) negative costs with the same algorithmic complexity as BF. The advantage of this is that, different from BF and other neural implementations of the BF algorithm [61], [62], we show that we can also directly apply neuronal learning rules that can dynamically change the activity in the network and lead to rerouting of paths according to some requirements. We show this by two cases where neuronal learning is used to structure the routing of the paths in a graph in different ways.

### III. Methods

#### A. Definitions

We denote a weighted graph as $G(V, E, C)$ with vertices $V$, edges $E$, and corresponding edge costs $C$. In the first version of the BF algorithm, a so-called relaxation procedure (cost minimization) is performed for all edges, whereas in the second version of the BF algorithm, relaxation is performed for all nodes. The relaxation procedure is performed for a number of iterations until convergence, i.e., approximate distances are replaced with shorter distances until all distances finally converge.

The BF algorithm is guaranteed to converge if relaxation of all edges is performed $|V| - 1$ times.\(^1\) After convergence, SPs (sequences of nodes) can be found from a list of predecesor nodes backward from target nodes to the source node.

\(^1\)Note that this corresponds to the worst case. See also algorithmic complexity analysis below.

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**Table I**

| Algorithm | Problem: SP/SSSP | Numerical calculation basis | Cost type | Algorithmic procedure | Training | Optimality | Complexity | Online learning |
|-----------|------------------|-----------------------------|-----------|-----------------------|----------|------------|------------|----------------|
| A* [10]   | SP               | Cost values                 | Only positive | Additive cost summation | NO       | Optimal    | $O(|E|)$   | Not directly possible |
| Dijkstra [9] | SSSP             | Cost values                 | Only positive | Additive cost summation | NO       | Optimal    | $O(|E| + |V|^2)$ or $O(|E| + |V| \log |V|)$ | Not directly possible |
| BF [11], [2] | SSSP             | Cost values                 | Positive and negative | Additive cost summation | NO       | Optimal    | $O(|V|^2)$ | Not directly possible |
| Recurrent | Neural network   | SSSP                        | Weight values = cost values | Additive activity propagation | NO       | Optimal    | Not stated | Not shown |
| Xiao, 2000 [61] |                  |                             | Positive and negative | Additive activity propagation |          |            |            |                |
| DMLP [29] | SP               | Weight values*              | n/a        | Additive/Multiplicative activity propagation | YES      | Not guaranteed | Not stated | n/a |
| PCN [30]–[32] | SP               | Weight values*              | n/a        | Additive/Multiplicative activity propagation | YES      | Not guaranteed | Not stated | n/a |
| LSTM [33] | SP               | Weight values*              | n/a        | Additive/Multiplicative activity propagation | YES      | Not guaranteed | Not stated | n/a |
| RL [34]   | SSSP             | Value tables                | Reward or punishment | Reward propagation | NO       | Not guaranteed | $O(n^2)$ | Reinforcement learning |
| Deep-LR [35]–[37] | SSSP           | Weight values*              | n/a        | Additive/Multiplicative activity propagation | YES      | Not guaranteed | Not stated | Reinforcement learning |
| GNN [38]–[43] | SSSP             | Weight values*              | n/a        | Additive/Multiplicative activity propagation | YES      | Not guaranteed | Not stated | n/a |
| T-POOL [28] | SSSP             | Weight values = cost values | Only positive (uniform) | Additive/Multiplicative activity propagation | NO       | Optimal    | $O(|V| \cdot |E|)$ | n/a |
| NN-BF (this study) | SSSP             | Weight values = inverse of cost values | Positive and negative | Multiplicative activity propagation | NO       | Optimal    | $O(|V| \cdot |E|)$ | Feedback learning, Reinforcement learning |

**Fig. 1**. Comparison of different methods for solving SP and SSSP problems.

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Algorithm 1 Pseudocode of Neural Network Algorithm

Input: list of neuron connections \((i, j)\) with weights \((w_{i,j})\), source neuron \((s)\), and costs \((c_{i,j})\)

Output: list of neuron activations (outputs), list of input nodes with maximal activity

```
for each neuron \(i\) do
    \(w[i, j] \leftarrow 1 - c[i, j]/K\)
    activations\([i]\) \(\leftarrow 0\)
    max_inputs\([i]\) \(\leftarrow\) null
end for

activations\([s]\) \(\leftarrow 1\)

for \(k \leftarrow 1\) to \(N - 1\) do
    for each neuron \(i\) do
        for each input \(j\) of neuron \(i\) with weight \(w[i, j]\) do
            weighted_input \(\leftarrow\) activations\([j]\) \(\times\) \(w[i, j]\)
            if weighted_input > activations\([i]\) then
                activations\([i]\) \(\leftarrow\) weighted_input
                max_inputs\([i]\) \(\leftarrow j\)
            end if
        end for
    end for
    activations\([s]\) \(\leftarrow 1\)
end for
```

Return activations, max_inputs

B. Neural Implementation of the BF Algorithm

In this section, we will describe our neural implementation of BF algorithm. Similar to BF, the proposed neural network finds SPs in a weighted graph from a source node (vertex) to all other nodes; however, SPs here are defined in terms of maximal activations from the source neuron to all other neurons.

The neural network has \(N\) neurons, where \(N\) corresponds to the number of vertices \(|V|\) in a graph, connections between neurons \(i\) and \(j\) \((i,j = 1, \ldots, N)\) correspond to the edges, and connection weights correspond to the (inverse) costs of edges, which are computed by

\[
 w_{i,j} = 1 - \frac{c_{i,j}}{K} \quad (1)
\]

where \(K > 0\). We can prove that the solutions obtained with the neural network are identical to the solutions obtained with BF under certain constraints (see Section IV-A1).

We present the pseudocode of the neural network algorithm (NN-BF) in Algorithm 1. Similar to BF version 2 (see [3]), we run NN-BF for a number of iterations until convergence operating on the graph nodes (neurons). In every iteration, we update the activation (output) of each neuron by

\[
 a_i = \max_j \{a_j w_{i,j}\} \quad (2)
\]

where \(a_j\) corresponds to the inputs of neuron \(i\) and \(w_{i,j}\) to the connection weights between input neurons \(j\) and output neuron \(i\). To start this process, we set the activation of the source neuron to 1. Thus, in this way, activity from the source neuron is propagated to all other neurons until activation has converged everywhere. After convergence, SPs (sequences of neurons) can be found from a list of maximal input activations backward from target neurons to the source neuron (similar to BF algorithm).\(^2\)

The NN-BF algorithm is guaranteed to converge if activity of the network is updated \(N - 1\) \((N = |V|)\) times. This can be shown by the worst-case scenario where neurons would be connected in a sequence, i.e., \(1 \rightarrow 2 \rightarrow 3 \rightarrow \cdots \rightarrow N\). Thus, to propagate activity from the first neuron to the last neuron in the sequence, one would need to perform \(N - 1\) updates.

A comparison of solving a simple graph using BF (version 1) and NN-BF is presented in Fig. 2. Here, we used a simple directed weighted graph with four nodes and six edges with one negative cost and five positive costs. We show the resulting distances in Fig. 2(a) and network activations in Fig. 2(b) after each iteration (numbers inside the nodes/neurons of the graph/neural network). Red edges/connections denote distance/activation propagation from the source node/neuron to all other nodes/neurons. For more details, please refer to the figure caption. In this case, the algorithm converged after two iterations when using BF and after one iteration when using NN-BF.

C. Performance Evaluation of the Neural Planner

We evaluated the performance of the neural network and compared it against the performance of the BF algorithm (version 1) with respect to the following: 1) solution equality; 2) algorithmic complexity; and 3) number of iterations until convergence.

\(^2\)The source code is available at https://doi.org/10.5281/zenodo.10029797.
For the numerical evaluation, we generated random directed weighted graphs of different density, from sparse graphs (only few connections per node) to fully connected graphs (every node connects to all other nodes). To generate graphs of different densities, we used the probability \( p_e \) to define whether two nodes will be connected or not. For example, a graph with 100 nodes and \( p_e = 0.1 \) will obtain a 10% connectivity rate and, thus, \( \approx 10 \) edges per node and \( \approx 1000 \) edges in total.

Costs for edges were assigned randomly from specific ranges (e.g., \([-10, -1]\) and \([1, 100]\)), where negative weights were assigned with probability \( p_n \) (e.g., \( p_n = 0.1 \)) and positive weights with probability \( 1 - p_n \). The specific parameters for each evaluation case will be provided in Section IV.

D. Combining Neural Planning Network With Neural Plasticity

The neural implementation of the BF algorithm allows us to use neuronal learning rules in order to dynamically change connection weights between neurons in the network and, thus, the outcome of the neural planner. The methods applied to achieve this will be presented directly in front of the corresponding results to allow for easier reading.

IV. RESULTS

A. Comparison of BF and NN-BF

In the following, we first provide a comparison between BF and NN-BF, and then, we will show results for NN-BF also on network learning.

1) Solution Equality: To prove equivalence, one can pairwise compare two paths \( A \) and \( B \) between some source and a target, where \( A \) is the best path under the BF condition and \( B \) is any another path. Path \( A \) is given by nodes \( a_i \) and \( B \) by \( b_i \). As described above, for a BF cost of \( a_i \), we define the weight of the NN-BF connection as \( w_i = 1 - \frac{a_i}{K} \) and likewise for \( b_i \). We need to show that

\[
\prod_{i=1}^{n+m} \left( 1 - \frac{a_i}{K} \right) > \prod_{i=1}^{n+m} \left( 1 - \frac{b_i}{K} \right). \tag{3}
\]

We start from the second line in (4) and rewrite it as follows:

\[
\frac{1}{K^2} \left[ K^2 - K(a_1 + a_2) + a_1a_2 \right] > \frac{1}{K^2} \left[ K^2 - K(b_1 + b_2) + b_1b_2 \right]. \tag{5}
\]
We simplify and divide by $K$, where $K > 0$, to
\[-a_1 - a_2 + \frac{a_1 a_2}{K} > -b_1 - b_2 + \frac{b_1 b_2}{K}. \tag{6}\]
Now, we let $K \to \infty$ and get
\[a_1 + a_2 < b_1 + b_2 \tag{7}\]
which proves conjecture (4) for large enough $K$.

To generalize this, we need to show that (3) holds. For this, we analyze the second inequality given by
\[
\prod_{i=1}^{n} \left(1 - \frac{a_i}{K}\right) > \prod_{i=1}^{n} \left(1 - \frac{b_i}{K}\right). \tag{8}
\]
From the structure of (5), we can deduce that
\[
\prod_{i=1}^{n} \left(1 - \frac{a_i}{K}\right) = \frac{1}{K^n} \left(K^n - K^{n-1} \sum a_i + K^{n-2} \sum_{i\neq j} a_i a_j - K^{n-3} \sum_{i\neq j\neq k} a_i a_j a_k \pm \cdots \right) \tag{9}
\]
and likewise for $b_i$, which allows writing out both sides of inequality (3). When doing this (not shown to save space), we see that the fore factor $(1/K^n)$ can be eliminated. Then, the term $K^n$ also abstracts away from both sides. After this, we can divide the remaining inequality as in (6), here using $K^{n-1}$ as divisor. This renders
\[-\sum a_i + \frac{1}{K} \sum_{i\neq j} a_i a_j - \frac{1}{K^2} \sum_{i\neq j\neq k} a_i a_j a_k \pm \cdots > -\sum b_i + \frac{1}{K} \sum_{i\neq j} b_i b_j - \frac{1}{K^2} \sum_{i\neq j\neq k} b_i b_j b_k \pm \cdots \]
\[\iff \sum a_i + D_a < \sum b_i + D_b \tag{10}\]
using the correct individual signs for $D$. Now, for $K \to \infty$, the disturbance terms $D$ vanish on both sides, and we get
\[\sum a_i < \sum b_i \tag{11}\]
as needed.

We conclude that for $K \to \infty$, the neural network is equivalent to the BF algorithm.

In the following, we will show by an extensive statistical evaluation that the distribution of $K$ for different realistic graph configurations is well behaved, and we never found any case where $K$ had to be larger than $10^6$.

Evidently, $K$ can grow if $\sum a_i \approx \sum b_i$. This can be seen easily, as, in this case, we can express $\sum b_i = \sum a_i + \epsilon$, with $\epsilon$ a small positive number. When performing this setting, we get from (10) that $\epsilon > D_b - D_a$, which can be fulfilled with large values for $K$.

We have set the word “can” above in italics, because a broader analysis shows that—even for similar sums—$K$ gets somewhat larger only for very few cases only as can be seen from Fig. 3.
where one iteration. Thus, the worst algorithmic complexity of BF is be run for \(|V|\) or neuron activations were not changing anymore. For this analysis, we used 500 randomly generated graphs with 5000 nodes were analyzed for each case. The results are shown in Fig. 6, where we show the maximal number of iterations until convergence of BF version 1 (BF) or NN-BF versus number of edges |E|. The 500 randomly generated graphs with 5000 nodes were analyzed for each case.

Fig. 5. Success rate of optimal solutions (left) and mean cost ratio (NN-BF cost/BF cost) of suboptimal solution versus \(K_0\) value. The 1000 randomly generated graphs with costs in range [1, 100] were used with four different densities (1%, 10%, 50%, and 100%). Note that for the graphs with 1% density and with \(K_0 = 10\), solutions were not found, and for the graphs with 10% density and with \(K_0 = 10\), solutions were found for four graphs and were optimal, whereas for all other remaining graphs, solutions were not found.

Fig. 4. Analysis of \(K_0\) values for graphs of different sizes (500, 1000, and 2000), different densities (5%, 10%, 50%, and 100%), and different cost ranges ([1, 10], [1, 100], and [1, 1000]). For cases \(|V| = 500\) with graph densities 5% and 10% and cost range [1, 1000], and for \(|V| = 1000\) with graph density 5% and cost range [1, 1000], 10000 random graphs were used, whereas in all other cases, 1000 graphs were used.

2) Algorithmic Complexity of the Algorithms: In the following, we will show that all three algorithms have the same algorithmic complexity.

BF version 1 consists of two loops (see [3]), where the outer loop runs in \(O(|V|)\) time and the inner loop runs in \(O(|E|)\), where \(|V|\) and \(|E|\) define the number of graph vertices (nodes) and edges (links), respectively. In the worst case, BF needs to be run for \(|V| - 1\) iterations and in the best case only for one iteration. Thus, the worst algorithmic complexity of BF is \(O(|V| |E|)\), whereas the best is \(O(|E|)\).

The algorithmic structure of the second version of the BF algorithm (see [3]) is the same as that of our NN-BF algorithm (Algorithm 1). Hence, their complexity is the same, and we will provide the analysis for the NN-BF algorithm. NN-BF operates on neurons and their inputs and not on edges of the graph as in BF version 1. Hence, NN-BF is similar to BF version 2 that operates on the nodes. The outer loop, as in BF version 1, runs in \(O(N)\), where \(N\) is the number of neurons and corresponds to the number of vertices \(|V|\). The second loop iterates through all neurons, and the third loop iterates through all inputs of the respective neuron and runs in \(O(I_j)\), respectively. Here, \(I_j\) corresponds to the number of inputs of a particular neuron \(j\). Given the fact that \(N = |V|\) and \(\sum_j I_j = |E|\) (\(j = 1, \ldots, N\)), we can show that the worst and the best algorithmic complexity of the NN-BF algorithm is the same as for BF version 2, i.e., \(O(|V| |E|)\) and \(O(|E|)\), respectively. Hence, this is the same complexity as for BF version 1, too.

3) Convergence Analysis: The worst-case scenario would be to run the algorithms for \(|V| - 1\) iterations, where \(|V|\) is the number of graph nodes. However, in practice, this will be not needed, and significantly, fewer iterations will usually suffice. Thus, we tested how many iterations are needed until cost convergence of the BF algorithm (version 1) as compared with activation convergence of the neural network. In both cases, we stopped the respective algorithm, as soon as its node costs or neuron activations were not changing anymore.

For this analysis, we used 500 randomly generated graphs with 5000 nodes with positive costs \((p_n = 0)\) from the range [1, 10] and connectivity densities corresponding to approximately \(12.5 \times 10^3, 25 \times 10^3, 250 \times 10^3, 2.5 \times 10^6, 12.5 \times 10^6, \) and \(24.995 \times 10^6\) edges.

The results are shown in Fig. 6, where we show the maximal number of iterations obtained from 500 tested graphs until
The synaptic weight $w$ between pre- and postsynaptic neuron is then changed according to

$$\Delta w = \alpha O_{\text{pre}} O_{\text{post}} R$$

(12)

where $\alpha > 0$ is the learning rate, usually set to small values to prevent too fast learning. The driving force of the learning process is the reward signal $R$, which can be positive or negative and will lead to weight increase [long-term potentiation (LTP)] or decrease [long-term depression (LTD)], respectively. This learning rule represents, thus, a three-factor rule [5], where, here, the third-factor is the reward signal $R$.

A diagram of the learning scheme is presented in Fig. 7, where we show the components of the learning rule to change the synaptic weight between two neurons in the planning network.

Note that in our learning scheme, weights were bounded between 0 and 1, i.e., $0 \leq w < 1$.

C. Combining Learning With Path Planning—Two Examples

In the following, we will show the examples of network learning combined with path finding. Examples stem from two common problem sets (navigation learning and sequence learning), and we can demonstrate here the main advantage of NN-BF, which is that learning and path finding rely on the same numerical representation. To achieve the same with BF (or any other additively operating algorithm), one would have to transform costs (for path finding) to activations (for network learning) back and forth. Examples are kept small, but upscaling to large problems is straightforward.

1) Navigation Learning: In the first learning scenario, we considered a navigation task in 2-D space, where an agent had to explore an unknown environment and plan the SP from a source location to a destination.

We assume a rectangular arena; see Fig. 8(a) and (b), iteration 0, with some locations (blue dots) and obstacles (gray boxes). The locations are represented by neurons in the planning network, whereas connections between neurons correspond to possible transitions from one location to other neighboring locations. Note that some transitions between neighboring locations, and, thus, connections between neurons are not possible due to obstacles. The positions of locations and possible transitions between them were generated in the following way. The position of a location $k$ is defined by

$$(x_k, y_k) = (hi + \xi_i, h j + \xi_j)$$

(13)

where $i, j = 1, 2, \ldots, m$ and $\xi$ is noise from a uniform distribution $U(0, b)$. In this study, we used $m = 5$ ($k = m^2 = 25$), $h = 0.2$, $a = -0.05$, and $b = 0.05$.

Possible transitions between neighboring locations and, thus, possible connections between neurons were defined based on a distance threshold $\theta_d$ between locations, i.e., connections were only possible if the Euclidean distance between two locations $d_{ij} = ||(x_i, y_i) - (x_j, y_j)|| \leq \theta_d$. In one case [Fig. 8(a)], we used $\theta_d = 0.25$, and in another case [Fig. 8(b)], we used $\theta_d = 0.35$.

Synaptic weights between neurons were updated after each transition (called iteration) from the current location $k$ (presynaptic neuron) to the next neighboring location $l$. 

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Fig. 7. Schematic of the learning architecture. $I_e$—external input. $R$—reward signal. $O_{\text{pre}}$ and $O_{\text{post}}$—output of presynaptic and postsynaptic neuron, respectively. $w$—synaptic weight between pre- and postsynaptic neuron. The multiplication symbol followed by a red arrow indicates that the synaptic weight $w$ is changed by a three-factor multiplicative (correlation) process [see (12)].

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C++ CPU implementation on Intel Core i9-9900 CPU, 3.10 GHz, and 32.0-GB RAM.
(postsynaptic neuron) according to the learning rule as described above, where pre- and postsynaptic neurons obtain an external input $I_e = 1$ during this transition (otherwise $I_e = 0$). In the navigation scenario, the reward signal $R$ was inversely proportional to the distance between neighboring locations $d_{k,l}$, i.e., $R = 1 - \beta d_{k,l}$ (here, we used $\beta = 1$). Thus, smaller distances between locations lead to larger reward values and larger weight updates, and vice versa. In the navigation learning scenario, initially, all weights were set to 0, and the learning rate was set to $\alpha = 0.02$. The learning and planning processes were repeated many times, where the learning process was performed for 100 iterations, and then, a path from the source (bottom-left corner) to the destination location (top-right corner) was planned using current state of the network.

We used two navigation learning cases. In the first case, a static environment with multiple obstacles was used where obstacles were present during the whole experiment [see Fig. 8(a)], whereas in the second case, a dynamic environment was simulated, where one big obstacle was used for some period of time and then was removed some time later [see Fig. 8(b)].

Results for navigation learning in a static environment are shown in Fig. 8(a), where we show the development of the network’s connectivity (black lines; line thickness is proportional to the weight strength) and the planned path (red trajectory) based on the network’s connectivity during the learning process. After 100 learning iterations, the environment was only partially explored. Therefore, the full path could not yet be planned. After 200 iterations, only a suboptimal path was found, since the environment was still not completely explored. As exploration and learning process proceeds, connections between different locations get stronger (note thicker lines in the second row), and eventually, the planned path converges to the SP (after 400 iterations).

In Fig. 8(b), we show results for navigation in a dynamic environment. Here, we first run the learning process until the path has converged (see the top row) and then remove the obstacle after 300 learning iterations. As expected, after obstacle removal, the middle part of the environment is explored, and new connections between neurons are built. Eventually, as the learning proceeds, the output of the planning network converges to the SP (after 1200 iterations).
Due to the learning rule used, in both cases, systems converge to the shortest Euclidean distance paths.

2) Sequence Learning: In the second learning scenario, we were dealing with sequence learning, where the task was to learn executing a predefined sequence of arbitrary events, which, in our study, were denoted by letters. For instance, this could correspond to an action sequence in order to execute some task by a robot and could be used for task planning in robotics to generate optimal plans. Note that sequences do not necessarily have to be predefined (as in our example), but optimal sequences can be learned based on obtained rewards during action execution.

In this case, we used a fully connected network (without self-connections) where neurons correspond to letters. In total, we used six different events (letters), i.e., A–F [see Fig. 9(a)]. Initially, all weights were set to random values from a uniform distribution $U(0, 0.5)$. Here, the planning and learning procedures are performed in epochs, where we execute the sequence based on the planner’s outcome always starting with letter A and ending with letter F and perform weight updates for each pair in the planned sequence. Suppose that, at the beginning, the planner generates a sequence $ABF$, then there will be two learning iterations in this epoch, i.e., weight update for the transition from A to B and from B to F. As in the navigation learning scenario, pre- and postsynaptic neurons will receive external input $I_e = 1$ whenever a certain transition from one event to the next event happens, e.g., from A (presynaptic neuron) to B (postsynaptic neuron).

Different from the navigation scenario, here, we used positive and negative rewards, i.e., $R = 1$ if a sequence pair is correct and $R = -1$ otherwise. In case of our previous $ABF$ example, this would lead to an increase of the synaptic weight between neurons A and B (correct) and to a decrease of the synaptic weight between neurons B and F (incorrect; after B should be C). The learning procedure is repeated for several learning epochs until convergence. Here, we used the relatively high learning rate $\alpha = 0.9$, which allows fast convergence to the correct sequence.

We performed two sequence learning cases, where in the first case, we were learning a sequence consisting of all possible six events $A B C D E F$, and in the second case, a shorter sequence $ACDF$ had to be learned.

Results for learning of the full sequence $A B C D E F$ and of the partial sequence $ACDF$ are presented in Fig. 9(a) and (b), respectively. As in the navigation learning example, we show the development of the network’s
connectivity during the learning process. In Fig. 9(a), we can see that before learning (epoch 0), due to random weight initialization, the generated sequence is \(ABF\), which is neither correct nor complete. However, after the first learning epoch, we can see that the connection weight between neurons \(A\) and \(B\) was increased (note thicker line between \(A\) and \(B\) as compared with epoch 0), whereas the connection weight between neurons \(B\) and \(F\) was decreased, which then led to a different sequence \(ABDF\). After epoch 3, the connection weight between neurons \(E\) and \(F\) was increased, since in previously generated sequence \(AEF\) (epoch 2), the transition from \(E\) to \(F\) was correct. Finally, after learning epoch 7, the network generates the correct sequence.

Similarly, in case of learning of the partial sequence \(ACDF\) [see Fig. 9(b)], learning converges to the correct sequence already after epoch 5, since the sequence is shorter.

V. CONCLUSION AND OUTLOOK

Finding the optimal path in a connected graph is a classical problem, and, as discussed in Section II-A, many different algorithms exist to address this. Here, we proposed a neural implementation of the BF algorithm. The main reason, as we would think, for translating BF into a network algorithm is the now-arising possibility to directly use network learning algorithm on a path-finding problem. To do this with BF or other cost-based algorithm, you would have to switch back and forth between a cost- and a weight-based representation of the problem. This is not needed for NN-BF. Given that BF and NN-BF have the same algorithmic complexity, computational speed of the path-finding step is similar, too, where learning epochs can be built into NN-BF as needed. The examples shown above rely on a local learning rule that alters the weights only between directly connected nodes. Globally acting rules could, however, also be used to address different problems, too, and—as far as we see—there are no restrictions with respect to possible network learning mechanisms, because NN-BF is a straightforward architecture for activity propagation without any special limitations.

Hence, network algorithm with NN-BF allows for new and different applications where learned path augmentation is directly coupled with path finding in a natural way.

So far, we only investigated the performance of our method on specific environments and tasks. Some of these task addressed the problem of unseen environments using our learning methods. In our future work, we will investigate the capability of our approach to generalize across different environments and/or different tasks, and we can also extend the experiments into a deeper investigation of complex unknown environments to even more strongly emphasize the power of being able to combine path finding directly with learning.

REFERENCES

[1] R. Bellman, “On a routing problem,” *Quart. Appl. Math.*, vol. 16, no. 1, pp. 87–90, 1958.
[2] L. R. Ford, “Network flow theory,” Rand Corp., Santa Monica, CA, USA, Tech. Rep. P-923, 1956.
[3] J. S. Baras and G. Theodorakopoulos, “Path problems in networks,” *Synth. Lectures Commun. Netw.*, vol. 3, no. 1, pp. 1–77, Jan. 2010.
[4] D. O. Hebb, *The Organization of Behavior*. New York, NY, USA: Wiley, 2005.
[5] L. Kusmierz, T. Isomura, and T. Toyoizumi, “Learning with three factors: Modulating Hebbian plasticity with errors,” *Current Opinion Neurol.*, vol. 46, pp. 170–177, Oct. 2017.
[6] P. Porr and F. Würgötter, “Learning with ‘relevance’: Using a third factor to stabilize Hebbian learning,” *Neural Comput.*, vol. 19, no. 10, pp. 2694–2719, 2007.
[7] E. F. Moore, “The shortest path through a maze,” in *Proc. Int. Symp. Theory Switching*, 1959, pp. 285–292.
[8] D. Merrill, M. Garland, and A. Grimshaw, “Scalable GPU graph traversal,” *ACM SIGPLAN Notices*, vol. 47, no. 8, pp. 117–128, Sep. 2012.
[9] E. W. Dijkstra, “A note on two problems in connexion with graphs,” *Nieuw Arch. Wisk.*, vol. 1, no. 1, pp. 269–271, Dec. 1959.
[10] P. Hart, N. Nilsson, and B. Raphael, “A formal basis for the heuristic determination of minimum cost paths,” *IEEE Trans. Syst. Sci. Cybern.*, vol. SSC-4, no. 2, pp. 100–107, Jul. 1968.
[11] R. E. Korf, “Depth-first iterative-deepening: An optimal admissible tree search,” *Artif. Intell.*, vol. 27, no. 1, pp. 97–109, 1985.
[12] S. Koenig, M. Likhachev, and D. Furcy, “Lifelong planning A∗,” *Artif. Intell.*, vol. 155, nos. 1–2, pp. 93–146, 2004.
[13] X. Sun, S. Koenig, and W. Yeoh, “Generalized adaptive A∗,” in *Proc. 7th Int. Conf. Auto. Agents Multiagent Syst.*, 2008, pp. 469–476.
[14] D. Harabor and A. Grastien, “Online graph pruning for path finding on grid maps,” in *Proc. AAAI*, 2011, pp. 1114–1119.
[15] R. W. Floyd, “Algorithm 97: Shortest path,” *Commun. ACM*, vol. 5, no. 6, p. 345, Jun. 1962.
[16] S. Warshall, “A theorem on Boolean matrices,” *J. ACM*, vol. 9, no. 1, pp. 11–12, Jan. 1962.
[17] E. F. Johnson, “Efficient algorithms for shortest paths in sparse networks,” *J. ACM*, vol. 24, no. 1, pp. 1–13, Jan. 1977.
[18] M. K. M. Ali and F. Kamoun, “Neural networks for shortest path computation and routing in computer networks,” *IEEE Trans. Neural Netw.*, vol. 4, no. 6, pp. 941–954, Nov. 1993.
[19] D.-C. Park and S.-E. Choi, “A neural network based multi-destination routing algorithm for communication network,” in *Proc. IEEE Int. Joint Conf. Neural Netw.*, vol. 2, May 1998, pp. 1673–1678.
[20] F. Araujo, B. Ribeiro, and L. Rodrigues, “A neural network for shortest path computation,” *IEEE Trans. Neural Netw.*, vol. 12, no. 5, pp. 1067–1073, Sep. 2001.
[21] J. Häkkinen, M. Lagerholm, C. Peterson, and B. Söderberg, “A Potts neuron approach to communication routing,” *Neural Comput.*, vol. 10, no. 6, pp. 1587–1599, Aug. 1998.
[22] R. Glasius, A. Komoda, and S. C. A. M. Gielen, “Neural network dynamics for path planning and obstacle avoidance,” *Neural Netw.*, vol. 8, no. 1, pp. 125–133, Jan. 1995.
[23] R. Glasius, A. Komoda, and S. C. A. M. Gielen, “A biologically inspired neural net for trajectory formation and obstacle avoidance,” *Biol. Cybern.*, vol. 74, no. 6, pp. 511–520, Jun. 1996.
[24] N. Bin, C. Xiong, Z. Liming, and X. Wendong, “Recurrent neural network for robot path planning,” in *Proc. Int. Conf. Parallel Distrib. Comput., Appl. Technol.*, 2004, pp. 188–191.
[25] S. X. Yang and M. Meng, “Neural network approaches to dynamic collision-free trajectory generation,” *IEEE Trans. Syst., Man Cybern., B*, vol. 31, no. 5, pp. 302–318, Jun. 2001.
[26] H. Qu, S. X. Yang, A. R. Willoms, and Z. Yi, “Real-time robot path planning based on a modified pulse-coupled neural network model,” *IEEE Trans. Neural Netw.*, vol. 20, no. 11, pp. 1724–1739, Nov. 2009.
[27] J. Ni, L. Wu, P. Shi, and S. X. Yang, “A dynamic bioinspired neural network-based real-time path planning method for autonomous underwater vehicles,” *Comput. Intell. Neurosci.*, vol. 2017, pp. 1–16, Feb. 2017.
[28] T. Kulvicius, S. Herzog, M. Tamosiunaite, and F. Wörgötter, “Finding optimal paths using networks without learning—Unifying classical approaches,” *IEEE Trans. Neural Netw. Learn. Syst.*, vol. 33, no. 12, pp. 8778–8787, Dec. 2022.
[29] A. H. Qureshi, M. J. Bency, and M. C. Yip, “Motion planning networks,” *Artif. Intell.*, vol. 24, no. 1, pp. 1–13, Jan. 1977.
[30] N. Bin, C. Xiong, Z. Liming, and X. Wendong, “Recurrent neural network for robot path planning,” in *Proc. Int. Conf. Parallel Distrib. Comput., Appl. Technol.*, 2004, pp. 188–191.
[31] S. X. Yang and M. Meng, “Neural network approaches to dynamic collision-free trajectory generation,” *IEEE Trans. Syst., Man Cybern., B*, vol. 31, no. 5, pp. 302–318, Jun. 2001.
[32] H. Qu, S. X. Yang, A. R. Willoms, and Z. Yi, “Real-time robot path planning based on a modified pulse-coupled neural network model,” *IEEE Trans. Neural Netw.*, vol. 20, no. 11, pp. 1724–1739, Nov. 2009.
[33] J. Ni, L. Wu, P. Shi, and S. X. Yang, “A dynamic bioinspired neural network-based real-time path planning method for autonomous underwater vehicles,” *Comput. Intell. Neurosci.*, vol. 2017, pp. 1–16, Feb. 2017.
[34] T. Kulvicius, S. Herzog, M. Tamosiunaite, and F. Wörgötter, “Finding optimal paths using networks without learning—Unifying classical approaches,” *IEEE Trans. Neural Netw. Learn. Syst.*, vol. 33, no. 12, pp. 8778–8787, Dec. 2022.
[35] A. H. Qureshi, M. J. Bency, and M. C. Yip, “Motion planning networks,” *Artif. Intell.*, vol. 24, no. 1, pp. 1–13, Jan. 1977.
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[32] T. Kulvicius, S. Herzog, T. Luderbeck, M. Tamosiunaite, and F. Wörgötter, “One-shot multi-path planning using fully convolutional networks in a comparison to other algorithms,” Frontiers Neuroinformatics, vol. 14, p. 115, Jan. 2021.

[33] M. J. Bency, A. H. Qureshi, and M. C. Yip, “Neural path planning: Fixed time, near-optimal path generation via Oracle imitation,” 2019, arXiv:1904.11102.

[34] R. S. Sutton and A. G. Barto, Reinforcement Learning: An Introduction.

[35] K. Teru, E. Denis, and W. Hamilton, “Inductive relation prediction,” in Proc. Neural Inf. Process. Syst. (NIPS), Sep. 2017, pp. 31–36.

[36] A. I. Panov, K. S. Yakovlev, and R. Suvorov, “Grid path planning with deep reinforcement learning: Preliminary results,” Proc. Comput. Sci., vol. 45, no. 1, pp. 2129–2134, 2016.

[37] R. Zellers, M. Yatskar, S. Thomson, and Y. Choi, “Neural motifs: Scene graphs in artificial agents,” Nature, vol. 557, no. 7705, pp. 429–433, May 2018.

[38] T. Kulvicius, S. Herzog, T. Luddecke, M. Tamosiunaite, and KULVICIUS et al.: COMBINING OPTIMAL PATH SEARCH WITH TASK-DEPENDENT LEARNING 509

[39] L. Zhang et al., “TN-ZSTAD: Transferable network for zero-shot temporal activity detection,” IEEE Trans. Pattern Anal. Mach. Intell., vol. 45, no. 3, pp. 3918–3932, Mar. 2023.

[40] D. Xu, Y. Zhu, C. B. Choy, and L. Fei-Fei, “Scene graph generation by iterative message passing,” in Proc. IEEE Conf. Comput. Vis. Pattern Recognit. (CVPR), Jul. 2017, pp. 3097–3106.

[41] R. Zellers, M. Yatskar, S. Thomson, and Y. Choi, “Neural motifs: Scene graph parsing with global context,” in Proc. IEEE/CVF Conf. Comput. Vis. Pattern Recognit., Jun. 2018, pp. 5831–5840.

[42] W. Gao, Y. Zhu, W. Zhang, K. Zhang, and H. Gao, “A hierarchical recurrent approach to predict scene graphs from a visual-attention-oriented perspective,” Comput. Intell., vol. 35, no. 3, pp. 496–516, Aug. 2019.

[43] Y. Li, W. Ouyang, B. Zhou, J. Shi, C. Zhang, and X. Wang, “FactorizableNet: An efficient subgraph-based framework for scene graph generation,” IEEE Trans. Neural Netw. Learn. Syst., vol. 33, no. 5, pp. 1234–1253, Jun. 2018.

[44] J. Yang, J. Lu, S. Lee, D. Batra, and D. Parikh, “Graph R-CNN for scene graph generation,” in Proc. Eur. Conf. Comput. Vis. (ECCV), 2018, pp. 670–685.

[45] X. Chang, P. Ren, P. Xu, Z. Li, X. Chen, and A. Hauptmann, “A comprehensive survey of scene graphs: Generation and application,” IEEE Trans. Pattern Anal. Mach. Intell., vol. 45, no. 1, pp. 1–26, Jan. 2023.

[46] M. Zhang et al., “One-shot neural architecture search: Maximising diversity to overcome catastrophic forgetting,” IEEE Trans. Pattern Anal. Mach. Intell., vol. 43, no. 9, pp. 2921–2935, Sep. 2021.

[47] C. Li, G. Wang, B. Wang, X. Liang, Z. Li, and X. Chang, “DS-Net++: Dynamic weight slicing for efficient inference in CNNs and vision transformers,” IEEE Trans. Pattern Anal. Mach. Intell., vol. 45, no. 4, pp. 4430–4446, Apr. 2023.

[48] X. Gong, S. Chang, Y. Jiang, and Z. Wang, “AutoGAN: Neural architecture search for generative adversarial networks,” in Proc. IEEE/CVF Int. Conf. Comput. Vis. (ICCV), Oct. 2019, pp. 3223–3233.

[49] H. Wang and J. Huan, “GAN: Towards automated design of generative adversarial networks,” 2019, arXiv:1906.11080.

[50] C. Yan et al., “ZeroNAS: Differentiable generative adversarial networks search for zero-shot learning,” IEEE Trans. Pattern Anal. Mach. Intell., vol. 44, no. 12, pp. 9733–9740, Dec. 2022.

[51] Y. Xia and J. Wang, “A discrete-time recurrent neural network for shortest-path routing,” IEEE Trans. Autom. Control, vol. 45, no. 11, pp. 2129–2134, 2000.

[52] Y. Zhang and S. Li, “Distributed biased min-consensus with applications to shortest path planning,” IEEE Trans. Autom. Control, vol. 62, no. 10, pp. 5429–5436, Oct. 2017.

[53] S. Koenig and R. G. Simmons, Complexity Analysis of Real-Time Reinforcement Learning Applied to Finding Shortest Paths in Deterministic Domains. Pittsburgh, PA, USA: Carnegie-Mellon Univ., 1993.

[54] P. Pot and F. Wörgötter, “Isotropic sequence order learning,” Neural Comput., vol. 15, no. 4, pp. 831–864, Apr. 2003.

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