
Discrete Neural Algorithmic Reasoning

Gleb Rodionov¹ Liudmila Prokhorenkova²

Abstract

Neural algorithmic reasoning aims to capture computations with neural networks by training models to imitate the execution of classic algorithms. While common architectures are expressive enough to contain the correct model in the weights space, current neural reasoners struggle to generalize well on out-of-distribution data. On the other hand, classic computations are not affected by distributional shifts as they can be described as transitions between discrete computational states. In this work, we propose to force neural reasoners to maintain the execution trajectory as a combination of finite predefined states. To achieve this, we separate discrete and continuous data flows and describe the interaction between them. Trained with supervision on the algorithm’s state transitions, such models are able to perfectly align with the original algorithm. To show this, we evaluate our approach on multiple algorithmic problems and achieve perfect test scores both in single-task and multitask setups. Moreover, the proposed architectural choice allows us to prove the correctness of the learned algorithms for any test data.

1. Introduction

Learning to capture algorithmic dependencies in data and to perform algorithmic-like computations with neural networks is a core problem in machine learning, long studied using various approaches (Roni Khardon, 1994; Graves et al., 2014; Zaremba & Sutskever, 2014; Reed & De Freitas, 2015; Kaiser & Sutskever, 2015; Veličković et al., 2020b).

Neural algorithmic reasoning (Veličković & Blundell, 2021) is a research area focusing on building models capable of executing classic algorithms. Relying on strong theoretical guarantees of algorithms to work correctly on any input of

any size and distribution, this setting provides unlimited challenges for out-of-distribution generalization of neural networks. Prior work explored this setup using the CLRS-30 benchmark (Veličković et al., 2022), which covers classic algorithms from the Introduction to Algorithms textbook (Cormen et al., 2009) and uses graphs as a universal tool to encode data of various types. Importantly, CLRS-30 also provides the decomposition of classic algorithms into subroutines and simple transitions between consecutive execution steps, called hints, which can be used during training in various forms.

The core idea of the CLRS-30 benchmark is to understand how neural reasoners generalize well beyond the training distribution, namely on larger graphs. Classic algorithms possess strong generalization due to the guarantee that correct execution steps never encounter ‘out-of-distribution’ states, as all state transitions are predefined by the algorithm. In contrast, when encountering inputs from distributions that significantly differ from the training data, neural networks are usually not capable of robustly maintaining internal calculations in the desired domain. Consequently, due to the complexity and diversity of all possible data that neural reasoners can be tested on, the generalization performance of such models can vary depending on particular test distributions (Mahdavi et al., 2023).

Given that, it is becoming important to interpret internal computations of neural reasoners to find errors or to prove the correctness of the learned algorithms (Georgiev et al., 2021).

Interpretation methods have been actively developing recently due to various real-world applications of neural networks and the need to debug and maintain systems based on them. Especially, the Transformer architecture (Vaswani et al., 2017) demonstrates state-of-the-art performance in natural language processing and other modalities, representing a field for the development of interpretability methods (Elhage et al., 2021; Weiss et al., 2021; Zhou et al., 2024; Lindner et al., 2024). Based on active research on a computational model behind the transformer architecture, recent works propose a way to learn models that are fully interpretable by design (Friedman et al., 2023).

We found the ability to design models that are interpretable in a simple and formalized way to be crucial for neural

¹Yandex Research, Moscow, Russia ²Yandex Research, Amsterdam, The Netherlands. Correspondence to: Gleb Rodionov <rodionovgleb@yandex-team.ru>.

algorithmic reasoning as it is naturally related to the goal of learning to perform computations with neural networks.

In this paper, we propose to force neural reasoners to follow the execution trajectory as a combination of finite predefined states, which is important for both generalization ability and interpretability of neural reasoners. To achieve that, we start with an attention-based neural network and describe three building blocks to enhance its generalization abilities: feature discretization, hard attention and separating discrete and continuous data flows. In short, all mentioned blocks are connected:

- State discretization does not allow the model to use complex and redundant dependencies in data;
- Hard attention is needed to ensure that attention weights will not be annealed for larger graphs. Also, hard attention limits the set of possible messages that each node can receive;
- Separating discrete and continuous flows is needed to ensure that state discretization does not lose information about continuous data.

Then, we build fully discrete neural reasoners for different algorithmic tasks and demonstrate their ability to perfectly mimic ground-truth algorithm execution. As a result, we achieve perfect test scores on multiple algorithmic tasks with guarantees of correctness on any test data. Moreover, we demonstrate that a single network is capable of executing all covered algorithms in a multitask manner with perfect generalization too.

In summary, we consider the proposed blocks as a crucial component for robust and interpretable neural reasoners and demonstrate that discretized models, trained with hint supervision, perfectly capture the dynamics of the underlying algorithms and do not suffer from distributional shifts.

2. Background

2.1. Algorithmic Reasoning

Performing algorithmic-like computations usually requires the execution of sequential steps and the number of such steps depends on the input size. To imitate such computations, neural networks are expected to be based on some form of recurrent unit, which can be applied to a particular problem instance several times (Zaremba & Sutskever, 2014; Kaiser & Sutskever, 2015; Vinyals et al., 2015; Veličković et al., 2020b).

The CLRS Algorithmic Reasoning Benchmark (CLRS-30) (Veličković et al., 2022) defines a general paradigm of algorithmic modeling based on Graph Neural Networks (GNNs), as graphs can naturally represent different input types and

manipulations over such inputs. Also, GNNs are proven to be well-suited for neural execution (Xu et al., 2020; Dudzik & Veličković, 2022).

The CLRS-30 benchmark covers different algorithms over various domains (arrays, strings, graphs) and formulates them as algorithms over graphs. Also, CLRS-30 proposes to utilize the decomposition of the algorithmic trajectory execution into simple logical steps, called hints. Using this decomposition is expected to better align the model to desired computations and prevent it from utilizing hidden non-generalizable dependencies of a particular train set. Prior work demonstrates a wide variety of additional inductive biases for models towards generalizing computations, including different forms of hint usage (Veličković et al., 2022; Bevilacqua et al., 2023), biases from standard data structures (Jain et al., 2023; Jürß et al., 2024), knowledge transfer and multitasking (Xhonneux et al., 2021; Ibarz et al., 2022; Numeroso et al., 2023), etc. Also, recent studies demonstrate several benefits of learning neural reasoners end-to-end without any hints at all (Mahdavi et al., 2023; Rodionov & Prokhorenkova, 2023).

The recently proposed SALSA-CLRS benchmark (Minder et al., 2023) enables a more thorough OOD evaluation compared to CLRS-30 with increased test sizes (up to 100-fold train-to-test scaling, compared to 4-fold for CLRS-30) and diverse test distributions. Despite significant gains in the performance of neural reasoners in recent work, current models still struggle to generalize to out-of-distribution (OOD) test data (Mahdavi et al., 2023; Georgiev et al., 2023; Minder et al., 2023). While de Luca & Fountoulakis (2024) prove by construction the ability of the transformer-based neural reasoners to perfectly simulate graph algorithms (with minor limitations occurring from the finite precision), it is still unclear if generalizable and interpretable models can be obtained via learning. Importantly, the issues of OOD generalization are induced not only by the challenges of capturing the algorithmic dependencies in the data but also by the need to carefully operate with continuous inputs. For example, investigating the simplest scenario of learning to emulate the addition of real numbers, Klindt (2023) demonstrates the failure of some models to exactly imitate the desired computations due to the nature of gradient-based optimization. This limitation can significantly affect the performance of neural reasoners on adversarial examples and larger input instances when small errors can be accumulated.

2.2. Transformer Interpretability and Computation Model

Transformer (Vaswani et al., 2017) is a neural network architecture for processing sequential data. The input to the transformer is a sequence of tokens from a discrete vocabulary. The input layer maps each token to a high-dimensional

embedding and aggregates it with the positional encoding. The key components of each layer are attention blocks and MLP with residual connections. Providing a detailed description of mechanisms learned by transformer models (Elhage et al., 2021) is of great interest due to their widespread applications.

RASP (Weiss et al., 2021) is a programming language proposed as a high-level formalization of the computational model behind transformers. The main primitives of RASP are elementwise sequence functions, *select* and *aggregate* operations, which conceptually relate to computations performed by different blocks of the model. Later, Lindner et al. (2024) presented Tracr, a compiler for converting RASP programs to the weights of the transformer model, which can be useful for evaluating interpretability methods.

While RASP might have limited expressibility, it supports arbitrarily complex continuous functions that in theory can be represented by the transformer architecture, but are difficult to learn. Also, RASP is designed to formalize computations over sequences of fixed length. Motivated by that, Zhou et al. (2024) proposed RASP-L, a restricted version of RASP, which aims to formalize the computations that are easy to learn with transformers in a size-generalized way. The authors also conjecture that the length-generalization of transformers on algorithmic problems is related to the ‘simplicity’ of solving these problems in the RASP-L language.

Another recent work (Friedman et al., 2023) describes Transformer Programs: constrained transformers that can be trained using gradient-based optimization and then automatically converted into a discrete, human-readable program. Built on RASP, transformer programs are not designed to be size-invariant.

3. Discrete Neural Algorithmic Reasoning

3.1. Encode-Process-Decode Paradigm

Our work follows the encode-process-decode paradigm (Hamrick et al., 2018), which is usually employed for step-by-step neural execution.

All input data is represented as a graph G with an adjacency matrix A and node and edge features that are first mapped with a simple linear encoder to high-dimensional vectors of size h . Let us denote node features at a time step t ($1 \leq t \leq T$) as $X^t = (x_1^t, \dots, x_n^t)$ and edge features as $E_t = (e_1^t, \dots, e_m^t)$. Then, the processor, usually a single-layer GNN, recurrently updates these features, producing node and edge features for the next step:

$$X^{t+1}, E^{t+1} = \text{Processor}(X^t, E^t, A).$$

The processor network can operate on the original graph defined by the task (for graph problems) or on the fully

connected graph. For the latter option, the information about the original graph can be encoded into the edge features.

The number of processor steps T can be defined automatically by the processor or externally (e.g., as the number of steps of the original algorithm). After the last step, the node and edge features are mapped with another linear layer, called the decoder, to the output predictions of the model.

If the model is trained with hint supervision, the changes of node and edge features at each step are expected to be related to the original algorithm execution. In this sense, the processor network is aimed to mimic the algorithm’s execution in the latent space.

3.2. Discrete Neural Algorithmic Reasoners

In this section, we describe the constraints for the processor that allow us to achieve a fully interpretable neural reasoner. We start with Transformer Convolution (Shi et al., 2020) with a single attention head.

As mentioned above, at each computation step t ($1 \leq t \leq T$), the processor takes the high-dimensional embedding vectors for node and edge features as inputs and then outputs the representations for the next execution step.

Each node feature vector x_i is projected into *query* (Q_i), *key* (K_i), and *value* (V_i) vectors via learnable parameter matrices W_Q , W_K , and W_V , respectively. Edge features e_{ij} are projected into *key* (K_{ij}) vector with a matrix W_K^E . Then, for each directed edge from a node j to a node i in the graph G , we compute the attention coefficient

$$\alpha_{ij} = \frac{\langle Q_j, K_i + K_{ij} \rangle}{\sqrt{h}},$$

where $\langle a, b \rangle$ denotes the dot product. Then, each node i normalizes all attention coefficients across its neighbors using the softmax function with temperature τ and receives the aggregated message:

$$\hat{\alpha}_{ij} = \frac{\exp(\alpha_{ij}/\tau)}{\sum_{k \in \mathcal{N}(i)} \exp(\alpha_{ik}/\tau)}, \quad M_i = \sum_{k \in \mathcal{N}(i)} \hat{\alpha}_{ik} V_k, \quad (1)$$

where $\mathcal{N}(i)$ denotes the set of all incoming neighbors of the node i and M_i is the message sent to the i -th node.

For undirected graphs, we consider two separate edges in each direction. Also, for each node, we consider a self-loop connecting the node to itself. For multi-head attention, each head l separately computes the messages M_i^l which are then concatenated.

Similar to Transformer Programs, we enforce attention to be hard attention. We found this property important not only for interpretability but also for size generalization, as hard attention allows us to overcome the annealing of the

attention weights for arbitrarily large graphs and strictly limits the set of messages that each vertex can receive.

After the message computation, node and edge features are updated depending on the current values and received messages using feed-forward MLP blocks:

$$\begin{aligned}\hat{x}_i^{t+1} &= \text{FFN}_{nodes}([x_i^t, M_i^t]), \\ \hat{e}_{ij}^{t+1} &= \text{FFN}_{edges}([e_{ij}^t, \hat{\alpha}_{ji} V_i, \hat{\alpha}_{ij} V_j]).\end{aligned}$$

We also enforce all node and edge features to be from a fixed finite set, which we call *states*. We ensure this property by adding discrete bottlenecks at the end of the processor block:

$$\begin{aligned}x_i^{t+1} &= \text{Discretize}_{nodes}(\hat{x}_i^{t+1}), \\ e_{ij}^{t+1} &= \text{Discretize}_{edges}(\hat{e}_{ij}^{t+1}).\end{aligned}$$

We implement discretization by projecting the features to the vectors of size k (states logits) which we then use to produce a one-hot vector corresponding to a discrete state. During training with hints, we use teacher forcing and directly use ground truth states as inputs for the next processor step. For no-hint learning, we apply Gumbel-Softmax (Jang et al., 2017) with annealing temperature to states logits. During inference, we simply use the argmax function over the states logits dimension.

3.3. Continuous Inputs

Clearly, most algorithmic problems operate with continuous or unbounded inputs (e.g., weights on edges). Usually, all input data is encoded into node and edge features and the processor operates over the resulting vectors. The proposed discretization of such vectors would lead to the loss of information necessary for performing correct execution steps. One possible option to operate with such inputs (we will call them *scalars*, meaning both continuous and size-dependent integer inputs, such as node indexes) is Neural Execution Engines (Yan et al., 2020), which enables operating with bit-wise representations of integer and (in theory) real numbers. Such representations are bounded by design, but fully discrete and interpretable.

We propose another option: to maintain scalar inputs (denote them by S) separately from the node and edge features and use them only as edge priorities s_{ij} in the attention block. If scalars are related to the nodes, we assign them to edges depending on the scalar of the sender or receiver node. Now, we can consider the hard attention block as a selector which for each node selects the best edge based on an ordered pair of ‘states priority’ (attention weights described above, which depend only on states of the corresponding nodes and edges) and s_{ij} . We note that this selector is related to the theoretical primitive *select.best* from RASP. We implement this simply by augmenting the key vectors K_{ij} of each edge

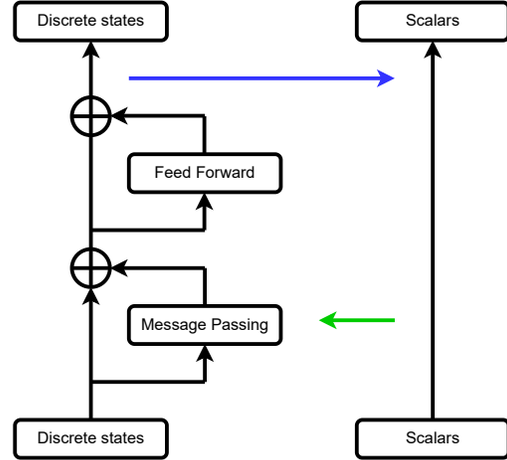


Figure 1. An illustration of the proposed separation between discrete and continuous data flows. Scalars can only affect the attention weights (Green) and can be modified with actions via ScalarUpdate (Blue).

with the indicator whether the given edge has the ‘best’ (min or max) scalar among the other edges to node j . Thus, scalars affect only the attention weights, not the messages or the node states.

For multiple different scalar inputs (e.g., weighted edges and node indexes), we use multi-head attention, where each head operates with separate scalars.

Thus, the interface of the proposed processor can be described as

$$X^{t+1}, E^{t+1} = \text{Processor}(X^t, E^t, A, S),$$

where X^t, X^{t+1} and E^t, E^{t+1} are from fixed sets. State sets are independent of the execution step t and the input graph (including scalar inputs S).

3.4. Manipulations over Continuous Inputs

The proposed selector offers a read-only interface to scalar inputs, which is not expressive enough for most algorithms. However, we note that the algorithms can be described as *discrete* manipulations over input data. For example, the Dijkstra algorithm (Dijkstra, 1959) takes edge weights as inputs and uses them to find the shortest path distances. Computed distances can affect the subsequent execution steps. We note that such distances can be described as the sum of the weights of the edges that form the shortest path to the given node. In other words, the produced scalars depend only on input scalars and discrete execution states.

To avoid the challenges of learning continuous updates with high precision (Klindt, 2023), we propose to learn discrete

manipulations with scalars. The updated scalars can then be used with the described selector in the next steps.

In our experiments, we use a scalar updater capable of incrementing, moving, and adding scalars depending on discrete node/edge states:¹

$$s_i^{t+1} = \text{inc}(x_i^t) + \text{keep}(x_i^t) \cdot s_i^t + \sum_{j \in \mathcal{N}(i)} \text{push}(e_{ji}^t) \cdot s_{ji}^t,$$

$$s_{ij}^{t+1} = \text{inc}(e_{ij}^t) + \text{keep}(e_{ij}^t) \cdot s_{ij}^t + \text{push}(x_i^t) \cdot s_i^t,$$

where s_i are node-related scalars, s_{ij} are edge-related scalars, and inc, keep, push are 0-1 functions representing whether a scalar in each node/edge should be incremented, kept, or pushed to any of its neighbors. We implement these functions as simple linear projections of node/edge features with subsequent discretization.

Finally, our proposed method (see Figure 1) can be described as:

$$X^{t+1}, E^{t+1} = \text{Processor}(X^t, E^t, A, S^t),$$

$$S^{t+1} = \text{ScalarUpdate}(X^{t+1}, E^{t+1}, A, S^t).$$

The proposed neural reasoners are fully discrete and can be interpreted by design. Moreover, the proposed selector block guarantees the predictable behavior of the message passing for any graph size, as it compares discrete state importance and uses continuous scalars only to break ties between equally important states.

4. Experiments

In this section, we perform experiments to evaluate how the proposed discretization affects the performance of neural reasoners on diverse algorithmic reasoning tasks. Our main questions are:

1. Can the proposed discrete neural reasoners capture the desired algorithmic dynamics with hint supervision?
2. How does discretization affect OOD and size-generalization performance of neural reasoners?
3. Is the proposed model capable of multi-task learning?

Additionally, we are interested whether discrete neural reasoners can be learned without hints and how they tend to utilize the given number of node and edge states to solve the problem. We discuss no-hint experiments separately in Section 6.

Finally, we investigate whether the training size can be reduced without compromising generalization performance; see Appendix B for details.

¹We additionally investigate if the operation set can be extended, see Appendix D.

4.1. Datasets

We perform our experiments on the problems from the recently proposed SALSA-CLRS benchmark (Minder et al., 2023), namely BFS, DFS, Prim, Dijkstra, Maximum Independent Set (MIS), and Eccentricity. We believe that the proposed method is not limited by the covered problems, but we leave the implementation of the required data flows (e.g., edge-based reasoning (Ibarz et al., 2022), graph-level hints, interactions between different scalars) for future work.

The train dataset of SALSA-CLRS consists of random graphs with at most 16 nodes sampled from the Erdős-Rényi (ER) distribution with parameter p chosen to be as low as possible such that graphs remain connected with high probability. The test set consists of sparse graphs of sizes from 16 to 1600 nodes.

We slightly modify the hints from the benchmark without conceptual changes (e.g., we have modified the hints for the DFS problem to remove graph-level hints and node discovery/finish times). Discrete states are fully described by the non-scalar hints and scalars are exactly the hints of the *scalar* type (we refer to Veličković et al. (2022) for the details on hint design).

4.2. Baselines and Evaluation

We compare the performance of our proposed discrete model with two baseline sparse models, GIN (Xu et al., 2019) and Pointer Graph Network (Veličković et al., 2020a). We report both node-level and graph-level metrics for the baselines and our model. Also, we compare our model with Triplet-GMPNN (Ibarz et al., 2022) and two recent approaches, namely Hint-ReLIC (Bevilacqua et al., 2023) and G-ForgetNet (Bohde et al., 2024), which demonstrate state-of-the-art performance in hint-based neural algorithmic reasoning. However, as these methods are evaluated on the CLRS-30 benchmark and their code is not yet publicly available, we can only compare them on the corresponding tasks (BFS, DFS, Dijkstra, Prim) and CLRS-30 test data, namely ER graphs with $p = 0.5$ of size 64. Note that this test data is denser than that of SALSA-CLRS, meaning shorter roll-outs for the given tasks. Also, only node-level metrics have been reported for these methods.

4.3. Model Details

For our experiments, we use the model described in Section 3. We use one attention head for each scalar value. The number of processor steps is defined externally as the length of the ground truth algorithm trajectory, which is consistent with prior work. We use one architecture (except the task-dependent encoders/decoders), including the ScalarUpdate module for all the problems.

We recall that neural reasoners can operate either on the base graph (which is defined by the problem) or on more dense graphs with the original graph encoded into edge features. SALSA-CLRS proposes to enhance the size-generalization abilities of neural reasoners with increased test sizes (up to 100-fold train-to-test scaling), so we use the base graph for message-passing, similar to the SALSA-CLRS baselines. We also add a virtual node that communicates with all nodes of the graph.

4.4. Training Details and Hyperparameters

We train each model using the Adam optimizer, learning rate $\eta = 0.001$, using teacher forcing, batch size of 32 graphs with 1000 optimization steps, and evaluate the resulting model. We anneal softmax temperatures in the discrete bottlenecks (attention weights, ScalarUpdate operations) geometrically from 3.0 to 0.01, decreasing the temperature at each training step. We report all hyperparameters in the source code.²

During training, we minimize the standard hints and output losses: scalar hints are optimized with MSE loss, and other types of hints are optimized with cross-entropy and categorized cross-entropy losses (Veličković et al., 2022; Ibarz et al., 2022). Note that we do not supervise any additional details in model behavior, e.g., selecting the most important neighbor in the attention block, the exact operations with scalars, etc.

For multitask experiments, we follow the setup proposed by Ibarz et al. (2022) and train a single processor with task-dependent encoders/decoders to imitate all covered algorithms simultaneously. We make 10000 optimization steps on the accumulated gradients across each task and keep all hyperparameters the same as in the single task.

Our models are trained on a single A100 GPU, requiring less than 1 hour for single-task and 5-6 hours for multitask training.

4.5. Results

We found learning with teacher forcing suitable for discrete neural reasoners, as discretization blocks allow us to perform the exact transitions between the states. Trained with step-wise hint supervision, discrete neural reasoners are able to perfectly align with the original algorithm and generalize on larger test data without any performance loss. We report the evaluation results in Tables 1 and 2. Also, our multitask experiments show that the proposed discrete models are capable of multitask learning and demonstrate the perfect generalization scores in a multitask manner too.

Recall that we have three key components of our contribu-

Table 1. Node \ graph level test scores for the proposed discrete reasoner and the baselines on SALSA-CLRS test data. Scores are averaged across 5 different seeds, standard deviation is omitted. For the eccentricity problem, only the graph-level metric is applicable.

TASK	SIZE	GIN	PGN	DNAR (OURS)
BFS	16	98.8 \ 92.5	100. \ 100.	100. \ 100.
	80	95.3 \ 59.4	99.8 \ 88.1	100. \ 100.
	160	95.1 \ 37.8	99.6 \ 66.3	100. \ 100.
	800	86.9 \ 0.9	98.7 \ 0.2	100. \ 100.
	1600	86.5 \ 0.0	98.5 \ 0.0	100. \ 100.
DFS	16	41.5 \ 0.0	82.0 \ 19.9	100. \ 100.
	80	30.4 \ 0.0	38.4 \ 0.0	100. \ 100.
	160	20.0 \ 0.0	26.9 \ 0.0	100. \ 100.
	800	19.5 \ 0.0	24.9 \ 0.0	100. \ 100.
	1600	17.8 \ 0.0	23.1 \ 0.0	100. \ 100.
SP	16	95.2 \ 49.8	99.3 \ 89.5	100. \ 100.
	80	62.4 \ 0.0	94.2 \ 3.3	100. \ 100.
	160	53.3 \ 0.0	92.0 \ 0.0	100. \ 100.
	800	40.4 \ 0.0	87.1 \ 0.0	100. \ 100.
	1600	36.9 \ 0.0	84.5 \ 0.0	100. \ 100.
PRIM	16	89.6 \ 29.7	96.4 \ 69.9	100. \ 100.
	80	51.6 \ 0.0	79.7 \ 0.0	100. \ 100.
	160	49.5 \ 0.0	75.6 \ 0.0	100. \ 100.
	800	45.0 \ 0.0	69.5 \ 0.0	100. \ 100.
	1600	43.2 \ 0.0	66.8 \ 0.0	100. \ 100.
MIS	16	79.9 \ 3.3	99.8 \ 98.6	100. \ 100.
	80	79.9 \ 20.0	99.4 \ 88.9	100. \ 100.
	160	78.2 \ 0.0	99.4 \ 76.2	100. \ 100.
	800	83.4 \ 0.0	98.8 \ 18.0	100. \ 100.
	1600	79.2 \ 0.0	98.9 \ 5.2	100. \ 100.
ECC.	16	25.3	100.	100.
	80	23.8	100.	100.
	160	26.1	100.	100.
	800	17.1	100.	100.
	1600	16.0	83.0	100.

tion: feature discretization, hard attention, and separating discrete and continuous data flows. To evaluate the importance of each component for generalization capabilities of the proposed models, we conduct an ablation study, the details can be found in Appendix A. In short, we demonstrate that removing each of these components yields the model without provable guarantees of perfect generalization. However, these components differ in terms of the impact on the performance. In particular, using regular attention instead of hard attention yields perfect test scores for given datasets, but it is possible to construct adversarial examples with large neighborhood sizes where performance drops. On the other hand, removing discretization from the scalar updater significantly affects the performance even on the small test graphs.

²<https://github.com/yandex-research/dnar>

Table 2. Node-level test scores for the proposed discrete reasoner and the baselines on CLRS-30 test data. Test graphs are of size 64. Scores are averaged across 5 different seeds.

TASK	HINT-RELIC	G-FORGETNET	DNAR (OURS)
BFS	99.00 \pm 0.2	99.96 \pm 0.0	100. \pm 0.0
DFS	100. \pm 0.0	74.31 \pm 5.0	100. \pm 0.0
SP	97.74 \pm 0.5	99.14 \pm 0.1	100. \pm 0.0
PRIM	87.97 \pm 2.9	95.19 \pm 0.3	100. \pm 0.0

5. Interpretability and Testing

In addition to the empirical evaluation of the trained models on diverse test data, the discrete and size-independent design of the proposed models allows us to interpret and test them manually. The main idea is to show that the model will perform the exact discrete state transitions (including discrete operations with scalars) as the original algorithm.

First, we note that due to the hard attention, each node receives exactly one message. Also, the message depends only on the discrete states of the corresponding nodes and edges. Thereby, as each node and edge states after a single processor step depend only on the current states and received message, all the possible options can be directly enumerated and tested if all states change to the correct ones.

The only remaining part to fully interpret the whole model is the attention block. We note that our implementation of the *select.best* selector (Section 3.3) does not necessarily produce the top-1 choice over the ordered pairs of ‘states priority’ and scalars s_{ij} as it simply augments key vectors with indicators if the given edge has the best scalar among others. For example, it may happen that for some state, the maximum attention weight is achieved for an edge without the indicator. However, given the finite number of discrete states, we can manually check if the mentioned “best” indicator increases the attention weight between every pair of states. Combined with hard attention, this would imply that the attention block attends depending on the predefined states and uses scalar priorities only to break ties. The particular example with more detailed explanation for the BFS problem can be found in Appendix C.

Given that, we can unit-test all possible state transitions and attention blocks. With full coverage of such tests, we can guarantee the correct execution of the desired algorithm for any test data. We tested our trained models from Section 4 manually verifying state transitions. As a result, we confirm that the attention block indeed operates as *select.best* selector, as the model actually uses these indicators to increase the attention weights. Thus, we can guarantee that for any graph size, the model will mirror the desired algorithm, which is correct for any test size.

6. Towards No-Hint Discrete Reasoners

In this section, we discuss the challenges of training discrete reasoners without hints, which can be useful when tackling new algorithmic problems.

Training deep discretized models is known to be challenging: without hyperparameter search, discrete models are only slightly improved over the untrained models. Therefore, we focus only on the BFS algorithm, as it is well-aligned with the message-passing framework, has short roll-outs, and can be solved with small states count (note that for no-hint models node/edge states count is a hyperparameter due to the absence of the ground truth states trajectory).

We recall that the output of the BFS problem is the exploration tree pointing from each node to its parent. Each node chooses as a parent the neighbor from the previous distance layer with the smallest index.

We perform hyperparameter search over the training sizes (using ER graphs with $p = 0.5$ and $n \in [4, 16]$), discrete node states count (from 2 to 6 states), softmax temperature annealing schedules ($[3, 0.01]$, $[3, 0.1]$, $[3, 1]$). For each hyperparameter choice, we train 5 models with different seeds. We validate the resulting models on the graphs of size 16. The best resulting model is obtained with the training size 5 and 4 node states. The trained models never achieved the perfect validation scores, see Table 3 for the results.

Table 3. BFS node/graph level scores of the *best_no_hint_model* for different graph sizes.

	5	16	64
<i>best_no_hint_model</i>	99 / 86	94 / 34	79 / 0

Then, we select the best-performing models and try to analyze the mistakes of the resulting models and reverse-engineer how they utilize the given states. First, we look at the node states after the last step of the processor and note that the states correspond to the distances from the starting node. More formally, we note that the model with four states uses the first state for the starting node, the second state for its neighbors, the third state for nodes at distance two from the starting node, and the last state for all other nodes and such states-based classification of distance has accuracy $> 98\%$ when tested on 1000 random graphs with 16 nodes. Then, we note that for the nodes that are from the first four distance layers from the starting node, the pointers are predicted with 100% accuracy and these pointers are computed layer-by-layer as in the ground truth algorithm (we refer to Appendix E for illustrations). The mistakes of the model are on the distance ≥ 4 from the starting node (we did not reverse-engineer the specific logic of computations on larger distances).

We found this behavior well-aligned to the BFS algorithm and indicating the possibility of achieving the perfect validation score with enough states count. However, this algorithm does not generalize since it fails at distances larger than those encountered during training.

On the other hand, one can demonstrate that for a small enough state count (for BFS, it is two node states and two edge states) and diverse enough validation data, the perfect validation performance implies that the learned solution will generalize to any graph size.

Therefore, we highlight the need to achieve perfect validation performance with models that use as few states as possible, which corresponds to the minimum description length (MDL) theory (Myung, 2000; Rissanen, 2006) and is related to the notion of Kolmogorov complexity (Kolmogorov, 1963).

Finally, we note that for sequential problems, such as DFS, obtaining a good no-hint model can be even more challenging and can require additional effort. One possible way to overcome this limitation is to implement a curriculum learning setup, which we leave for future work.

7. Limitations and Future Work

Limitations In this work, we propose a method to learn robust neural reasoners that demonstrate perfect generalization performance and are interpretable by design. In this section, we describe some limitations of our work and important directions for future research.

First, several proposed design choices strictly reduce the expressive power of the model. For example, due to the hard attention, the proposed model is unable to compute the mean value from all the neighbors in a single message-passing step, which is trivial for attention-based models (note that this can be computed in several message-passing steps). Thus, the model in its current form is unable to express transitions between hints for some algorithms from the CLRS-30 in a single processor step.

However, we believe that the expressivity of the proposed model can be enhanced with additional architectural modifications (e.g., edge-based reasoning (Ibarz et al., 2022), global states, interactions between different scalars) that can be combined with the proposed discretization ideas. Additionally, there are several potential ways to balance between expressive power of the model and strong generalization guarantees, e.g. removing hard attention, removing feature discretization (while keeping the discretization inside the ScalarUpdate module).

Second, while we report the perfect scores for the covered tasks, we cannot guarantee that the training will converge to the correct model for any initialization/training data dis-

tributions. However, we empirically found the proposed method to be quite robust to various architecture/training hyperparameter choices.

Future work Our method is based on the particular architectural choice and actively utilizes the attention mechanism. However, the graph deep learning field is rich in various architectures exploiting different inductive biases and computation flows. The proposed separation between discrete states and continuous inputs may apply to other models, however, any particular construction can require additional efforts.

Also, we provide only one example of the ScalarUpdate block. We believe that utilizing a general architecture (e.g., some form of discrete Neural Turing Machine (Graves et al., 2014; Gulcehre et al., 2016)) capable of executing a wider range of manipulation is of interest for future work.

With the development of neural reasoners and their ability to execute classic algorithms on abstract data, it is becoming more important to investigate how such models can be applicable in real-world scenarios according to the Neural Algorithmic Reasoning blueprint (Veličković & Blundell, 2021) and transfer their knowledge to high-dimensional noisy data with intrinsic algorithmic dependencies. While there are several examples of NAR-based models tackling real-world problems (Beurer-Kellner et al., 2022; Numeroso et al., 2023), there are no established benchmarks for extensive evaluation and comparison of different approaches.

Lastly, we leave for future work a deeper investigation of learning interpretable neural reasoners without hints, which we consider essential from both theoretical perspective and practical applications, e.g., combinatorial optimization.

8. Conclusion

In this paper, we force neural reasoners to maintain the execution trajectory as a combination of finite predefined states. To achieve that, we separate discrete and continuous data flows and describe the interaction between them. The obtained discrete reasoners are interpretable by design. Moreover, trained with hint supervision, such models perfectly capture the dynamic of the underlying algorithms and do not suffer from distributional shifts. We consider discretization of hidden representations as a crucial component for robust neural reasoners.

Impact Statement

This paper presents work whose goal is to advance the field of Machine Learning. There are many potential societal consequences of our work, none which we feel must be specifically highlighted here.

References

- Beurer-Kellner, L., Vechev, M., Vanbever, L., and Veličković, P. Learning to configure computer networks with neural algorithmic reasoning. *Advances in Neural Information Processing Systems*, 35:730–742, 2022.
- Bevilacqua, B., Nikiforou, K., Ibarz, B., Bica, I., Paganini, M., Blundell, C., Mitrovic, J., and Veličković, P. Neural algorithmic reasoning with causal regularisation. In *International Conference on Machine Learning*, 2023.
- Bohde, M., Liu, M., Saxton, A., and Ji, S. On the markov property of neural algorithmic reasoning: Analyses and methods. In *The Twelfth International Conference on Learning Representations*, 2024.
- Cormen, T. H., Leiserson, C. E., Rivest, R. L., and Stein, C. *Introduction to Algorithms*. MIT press, 2009.
- de Luca, A. B. and Fountoulakis, K. Simulation of graph algorithms with looped transformers. In *Forty-first International Conference on Machine Learning*, 2024.
- Dijkstra, E. W. A note on two problems in connexion with graphs. *Numerische Mathematik*, 1:269–271, 1959.
- Dudzik, A. J. and Veličković, P. Graph neural networks are dynamic programmers. *Advances in Neural Information Processing Systems*, 35:20635–20647, 2022.
- Elhage, N., Nanda, N., Olsson, C., Henighan, T., Joseph, N., Mann, B., Askell, A., Bai, Y., Chen, A., Conerly, T., DasSarma, N., Drain, D., Ganguli, D., Hatfield-Dodds, Z., Hernandez, D., Jones, A., Kernion, J., Lovitt, L., Ndousse, K., Amodei, D., Brown, T., Clark, J., Kaplan, J., McCandlish, S., and Olah, C. A mathematical framework for transformer circuits. *Transformer Circuits Thread*, 2021.
- Friedman, D., Wettig, A., and Chen, D. Learning transformer programs. *Advances in Neural Information Processing Systems*, 2023.
- Georgiev, D., Barbiero, P., Kazhdan, D., Veličković, P., and Lio, P. Algorithmic concept-based explainable reasoning. In *AAAI Conference on Artificial Intelligence*, 2021.
- Georgiev, D., Lio, P., Bachurski, J., Chen, J., and Shi, T. Beyond erdos-renyi: Generalization in algorithmic reasoning on graphs. In *NeurIPS 2023 Workshop: New Frontiers in Graph Learning*, 2023.
- Graves, A., Wayne, G., and Danihelka, I. Neural turing machines. *arXiv preprint arXiv:1410.5401*, 2014.
- Gulcehre, C., Chandar, S., Cho, K., and Bengio, Y. Dynamic neural turing machine with soft and hard addressing schemes. *arXiv preprint arXiv:1607.00036*, 2016.
- Hamrick, J. B., Allen, K. R., Bapst, V., Zhu, T., McKee, K. R., Tenenbaum, J. B., and Battaglia, P. W. Relational inductive bias for physical construction in humans and machines. *arXiv preprint arXiv:1806.01203*, 2018.
- Ibarz, B., Kurin, V., Papamakarios, G., Nikiforou, K., Benani, M., Csordás, R., Dudzik, A. J., Bošnjak, M., Vitvitskyi, A., Rubanova, Y., et al. A generalist neural algorithmic learner. In *Learning on Graphs Conference*. PMLR, 2022.
- Jain, R., Veličković, P., and Liò, P. Neural priority queues for graph neural networks. *arXiv preprint arXiv:2307.09660*, 2023.
- Jang, E., Gu, S., and Poole, B. Categorical reparameterization with gumbel-softmax. *International Conference on Learning Representations*, 2017.
- Jürß, J., Jayalath, D. H., and Veličković, P. Recursive algorithmic reasoning. In *Learning on Graphs Conference*. PMLR, 2024.
- Kaiser, Ł. and Sutskever, I. Neural GPUs learn algorithms. *arXiv preprint arXiv:1511.08228*, 2015.
- Klindt, D. A. Controlling neural network smoothness for neural algorithmic reasoning. *Transactions on Machine Learning Research*, 2023. ISSN 2835-8856.
- Kolmogorov, A. N. On tables of random numbers. *Sankhya: The Indian Journal of Statistics*, 25, 1963.
- Lindner, D., Kramár, J., Farquhar, S., Rahtz, M., McGrath, T., and Mikulik, V. Tracr: Compiled transformers as a laboratory for interpretability. *Advances in Neural Information Processing Systems*, 36, 2024.
- Mahdavi, S., Swersky, K., Kipf, T., Hashemi, M., Thrampoulidis, C., and Liao, R. Towards better out-of-distribution generalization of neural algorithmic reasoning tasks. *Transactions on Machine Learning Research*, 2023. ISSN 2835-8856.
- Minder, J., Grötschla, F., Mathys, J., and Wattenhofer, R. SALSA-CLRS: A sparse and scalable benchmark for algorithmic reasoning. *arXiv preprint arXiv:2309.12253*, 2023.
- Myung, I. J. The importance of complexity in model selection. *Journal of mathematical psychology*, 44(1):190–204, 2000.
- Numeroso, D., Bacciu, D., and Veličković, P. Dual algorithmic reasoning. In *International Conference on Learning Representations*, 2023.
- Reed, S. and De Freitas, N. Neural programmer-interpreters. *arXiv preprint arXiv:1511.06279*, 2015.

- Rissanen, J. Information and complexity in statistical modeling. In *Information Theory Workshop*, 2006.
- Rodionov, G. and Prokhorenkova, L. Neural algorithmic reasoning without intermediate supervision. *Advances in Neural Information Processing Systems*, 2023.
- Roni Khardon, D. R. Learning to reason. In *Proceedings of the 12th National Conference on Artificial Intelligence, Seattle, WA, USA, July 31 - August 4, 1994, Volume 1*, pp. 682–687. AAAI Press / The MIT Press, 1994.
- Shi, Y., Huang, Z., Feng, S., Zhong, H., Wang, W., and Sun, Y. Masked label prediction: Unified message passing model for semi-supervised classification. *arXiv preprint arXiv:2009.03509*, 2020.
- Vaswani, A., Shazeer, N., Parmar, N., Uszkoreit, J., Jones, L., Gomez, A. N., Kaiser, Ł., and Polosukhin, I. Attention is all you need. *Advances in neural information processing systems*, 30, 2017.
- Veličković, P. and Blundell, C. Neural algorithmic reasoning. *Patterns*, 2(7):100273, 2021.
- Veličković, P., Buesing, L., Overlan, M., Pascanu, R., Vinyals, O., and Blundell, C. Pointer graph networks. *Advances in Neural Information Processing Systems*, 33: 2232–2244, 2020a.
- Veličković, P., Ying, R., Padovano, M., Hadsell, R., and Blundell, C. Neural execution of graph algorithms. In *International Conference on Learning Representations*, 2020b.
- Veličković, P., Badia, A. P., Budden, D., Pascanu, R., Bannino, A., Dashevskiy, M., Hadsell, R., and Blundell, C. The CLRS algorithmic reasoning benchmark. In *International Conference on Machine Learning*, pp. 22084–22102. PMLR, 2022.
- Vinyals, O., Fortunato, M., and Jaitly, N. Pointer networks. *Advances in neural information processing systems*, 28, 2015.
- Weiss, G., Goldberg, Y., and Yahav, E. Thinking like transformers. In *International Conference on Machine Learning*, pp. 11080–11090. PMLR, 2021.
- Xhonneux, L.-P., Deac, A.-I., Veličković, P., and Tang, J. How to transfer algorithmic reasoning knowledge to learn new algorithms? *Advances in Neural Information Processing Systems*, 34:19500–19512, 2021.
- Xu, K., Hu, W., Leskovec, J., and Jegelka, S. How powerful are graph neural networks? In *The Seventh International Conference on Learning Representations*, 2019.
- Xu, K., Li, J., Zhang, M., Du, S. S., Kawarabayashi, K.-i., and Jegelka, S. What can neural networks reason about? In *International Conference on Learning Representations*, 2020.
- Yan, Y., Swersky, K., Koutra, D., Ranganathan, P., and Hashemi, M. Neural execution engines: Learning to execute subroutines. *Advances in Neural Information Processing Systems*, 33:17298–17308, 2020.
- Zaremba, W. and Sutskever, I. Learning to execute. *arXiv preprint arXiv:1410.4615*, 2014.
- Zhou, H., Bradley, A., Littwin, E., Razin, N., Saremi, O., Susskind, J. M., Bengio, S., and Nakkiran, P. What algorithms can transformers learn? A study in length generalization. In *The Twelfth International Conference on Learning Representations*, 2024.

A. Ablation Study

Recall that we have three key components of our contribution: feature discretization, hard attention, and separating discrete and continuous data flows. In this section, we study the importance of these components for generalization capabilities of the proposed models.

Discrete bottlenecks First, we evaluate the model without all discrete bottlenecks: the result is a simple Transformer Convolution processor, which performs comparable to other baseline models, see Table 4.

Table 4. Node \ graph level test scores for our base model without all discrete bottlenecks. Scores are averaged across 5 different seeds, standard deviation is omitted.

SIZE	16	80	160	800	1600
BFS	99.9 \ 99.3	99.7 \ 88.2	99.5 \ 57.9	98.4 \ 0.0	97.2 \ 0.0
DFS	79.2 \ 6.8	41.1 \ 0.0	28.1 \ 0.0	24.7 \ 0.0	21.9 \ 0.0
SP	99.3 \ 88.7	94.1 \ 12.4	90.3 \ 0.0	86.9 \ 0.0	82.4 \ 0.0
PRIM	95.1 \ 72.7	82.6 \ 0.0	79.7 \ 0.0	68.1 \ 0.0	66.0 \ 0.0
MIS	99.8 \ 98.6	99.6 \ 86.1	99.2 \ 69.0	97.1 \ 11.9	96.3 \ 0.0
ECC	79.2	41.1	28.1	24.7	21.9

Hard attention To highlight the importance of the hard attention for strong size generalization, we train the proposed model but with the regular attention mechanism on the BFS task. The resulting model also demonstrates the perfect scores for the given test data. However, the standard test data (the Erdős-Renyi graphs with low edge probability) does not contain nodes with a large number of neighbors, while such nodes can be problematic due to the annealing of the attention weights. Thus, we additionally test the resulting model on the complete bipartite graphs $K_{2,n-2}$ for different n . For each n , we assign the starting node from the smaller component and test if the second node in this component correctly selects its parent in the BFS tree, see Figure 2.

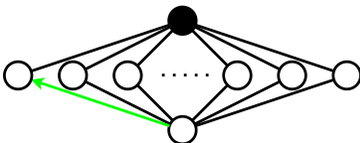


Figure 2. Complete bipartite graphs $K_{2,n-2}$ used to evaluate the effect of the attention weights annealing. The black node is the starting node. The highlighted edge (Green) is the ground truth pointer from the bottom node to its parent from the BFS tree.

Our experiments demonstrate that the model without hard attention fails to predict the correct pointer for larger graphs due to the attention weight annealing, see Table 5. We note that the models from Section 4 are provably correct on any test data.

Table 5. Attention weights for the ground truth pointer (green pointer from Figure 2) for different graph sizes; (+/-) denotes if the correct pointer was predicted.

SIZE	16	80	160	800	1600
ATTENTION WEIGHT	0.97 (+)	0.86 (+)	0.76 (+)	0.38 (-)	0.24 (-)

Scalar updater As demonstrated in Klindt (2023), simple neural networks trained to sum two real numbers fail to learn the structure of the task and struggle to extrapolate beyond the training data distributions. In this section, we study how these limitations affect the overall performance of neural reasoners to highlight the importance of the proposed discrete manipulations with scalars.

First, we recall that usually all input data is encoded into node and edge features and the processor operates over the resulting vectors. Then, hints of type scalar are directly predicted from the node/edge features. To evaluate the effect of non-discrete ScalarUpdate modules, we simply replace the proposed discrete ScalarUpdate module with a single-layer

transformer convolution network, which inputs scalars and node/edge states and outputs the scalars for the next step, keeping the remaining architecture the same as in the main experiments. We train the resulting model on Dijkstra and MST problems.

Additionally, we evaluate the non-discrete ScalarUpdate module in a more straightforward setup. Similarly to Klindt (2023), we train a 2-layer MLP to add two real numbers and use the resulting model as a ScalarUpdate module for the Dijkstra algorithm. We manually use the learned addition module when the node distances are updated (e.g., the distance of the node u is updated with the sum of the distance of v and the edge (v, u) cost), and use the ground truth scalars for other scalar updates. Our experiments demonstrate that the resulting model outperforms the baselines on the test size of 16 nodes, but does not generalize well on larger graphs, see the evaluation results in Table 6.

Table 6. Node \ graph level test scores for the proposed model with ScalarUpdate replaced by a regular attention-based network trained to predict hints of type scalar. ‘Addition only’ means that ScalarUpdate is replaced by a 2-layer MLP trained to predict the sum of two numbers (other values are taken from the ground truth).

SIZE	16	80	160	800	1600
DIJKSTRA	99.3 \ 94.6	60.7 \ 0.0	42.8 \ 0.0	19.0 \ 0.0	11.8 \ 0.0
MST	99.8 \ 98.1	98.2 \ 54.1	97.2 \ 28.1	95.5 \ 0.0	91.73 \ 0.0
DIJKSTRA (ADDITION ONLY)	99.8 \ 96.6	95.3 \ 71.0	86.5 \ 46.3	41.6 \ 3.1	22.2 \ 0.0

We note that all the resulting models demonstrate perfect scores when evaluated with teacher-forced ground truth scalars. Thus, all state transitions are learned correctly and imperfect test scores are fully described by the errors in manipulations with continuous values.

To summarize, small errors in manipulations with scalars (even restricted on the simplest addition sub-task) strictly affect the overall performance of the model, highlighting the importance of the proposed discrete manipulations with scalars.

B. Minimal Training Sizes

In this section, we highlight an important property of the proposed models: the correct state transitions can be learned even from a trivially small size.

For example, for the BFS problem, it is enough to use graphs with only 3 nodes to observe that a not visited node becomes visited or not depending on the received message. However, the subtask of selecting the parent from the multiple visited neighbors requires at least 4 nodes (where the minimum sufficient example is the complete bipartite graph $K(2, 2)$).

To demonstrate that, we conducted additional experiments to empirically find the smallest training size for perfect fitting of each covered algorithm. For this experiment we used training with hints: we train our models for each problem on $ER(n, 0.5)$ graphs for different n and test the resulting models on the graphs with 160 nodes. Results are presented in Table 7. Note that the empirical bound is around 4-5 nodes.

Table 7. Node-level test scores for the proposed discrete reasoner on test graphs with 160 nodes, across different training sizes.)

Algorithm	3 Nodes	4 Nodes	5 Nodes
BFS	41	100	100
DFS	38	100	100
Dijkstra	13	26	100
MST	11	14	100
MIS	79	100	100
Ecc.	45	100	100

C. Interpretability and Testing Details

In this section, we provide additional details on interpretability and testing of the proposed discrete reasoners.

As an example, consider the BFS algorithm. First, recall the pseudocode of the algorithm:

```

Starting_node ← visited
All_other_nodes ← not_visited
for step in range(T) do
  for node  $U$  in a graph do
    if  $U$  is visited on previous steps then
      continue
    end if
    if  $U$  has a neighbor  $P$  that visited on previous steps then
       $U \leftarrow$  visited on this step
       $U$  select the smallest-indexed such neighbor  $P$  as parent:
      Edge  $(U, P) \leftarrow$  pointer
      Self-loop  $(U, U) \leftarrow$  not_pointer
    end if
  end for
end for
return a BFS tree described by pointers

```

Now let us describe how we can verify that the trained DNAR model will perfectly imitate this algorithm for any test data.

First, we note that for each node U , the node state on the step $t + 1$ (denoted by U_{t+1}) is the function of U_t and V_t , where V is the node that sends the message to U on step t :

$$U_{t+1} = \text{StateUpdate}(U_t, \text{message_from_}V_t)$$

How does the node U select a node that will send a message to it? For any node V connected to U , the node U computes attention scores depending on discrete states of V of each node and a discrete indicator if each node has the smallest (or largest) scalar among all neighbors of U with the same discrete state as V . Then, the node U selects the node V with the largest attention score.

In our (slightly simplified) case, the attention scores only depend on the tuples $(U_{state}, V_{state}, \text{indicator_if_}u\text{ has the smallest index})$ and there are only 8 such tuples. We can directly compute these attention scores and verify the required invariants, e.g.,

$$\text{Attention}(\text{not_visited}, \text{visited}, \text{smallest}) > \text{Attention}(\text{not_visited}, *any_other*),$$

which would imply that the not_visited node will receive the message from the smallest-indexed visited neighbor if such exists independently of the graph size and distribution. If there is no such neighbor, the node U will receive the message from another not_visited node (or from itself).

After verifying the correctness of the message flows, we need to ensure that the state updates are computed correctly, e.g.,

$$\begin{aligned}
 \text{visited} &= \text{StateUpdate}(\text{not_visited}, \text{message_from_visited}), \\
 \text{visited} &= \text{StateUpdate}(\text{visited}, *any*), \\
 \text{not_visited} &= \text{StateUpdate}(\text{not_visited}, \text{message_from_not_visited}).
 \end{aligned}$$

The main idea is that due to the finite states count and discrete manipulations with scalars, there are only finite amounts of such checks that can cover all possible state transitions and all of them should be evaluated only once.

D. Extended ScalarUpdate Module

In this section, we investigate if the proposed ScalarUpdate module can be successfully extended to support more complex manipulations with scalars.

First, we note that simple manipulations with scalars cover a significant part of the classic algorithms. In this work, we use the minimum set of the required functions, but it can be directly extended by other functions. Importantly, as ScalarUpdate can be viewed as a separate module, we can separately check if it is possible to train it with any given set of predefined manipulations for any problem only with supervision on the results.

Let us formalize the problem: each input of the `ScalarUpdate` module can be described as an object with a discrete state s_i (from a fixed predefined set) and several scalar values (we consider two scalars x_i and y_i). Note that we omit the separation between nodes and edges and consider objects with several scalar values. For each discrete state s_i , there exists a ground truth update of scalars, e.g., $f(s, x, y) = x + \cos(y)$. The output of the scalar updater can be viewed as a sum:

$$\text{ScalarUpdate}(s, x, y) = \sum_{g \in \mathcal{OPS}} g(x, y) \cdot \text{activate}_g(s),$$

where \mathcal{OPS} is a predefined set of operations and activate_g is a 0-1 function representing if a specific operation should be applied. Note that activate_g depends only on a discrete state of the input.

For our additional experiment, we train several different `ScalarUpdate` modules with the extended set of operations:

- $g_0(x, y) = 1$;
- $g_1(x, y) = x$;
- $g_2(x, y) = \cos(x)$;
- $g_3(x, y) = x \cdot y$;
- $g_4(x, y) = \text{atan2}(x, y)$

to learn the following set of the ground truth updates simultaneously:

- $f_0(x, y) = x$;
- $f_1(x, y) = \cos(x)$;
- $f_2(x, y) = \cos(x) + x \cdot y$;
- $f_3(x, y) = \text{atan2}(x, y)$;
- $f_4(x, y) = 1 + x + \text{atan2}(x, y)$.

In particular, we consider a set of 16 discrete states (numbered from 0 to 15 and sampled uniformly) and the ground truth scalar update is derived from these states by taking the remainder of the division by 5 (updates count).

The learnable parameters of the `ScalarUpdate` are states’ embeddings and linear projections for each indicator. We train `ScalarUpdate` to minimize the MSE loss between the ground truth and predicted outputs with 5000 optimization steps. Additionally, we train a non-discrete scalar updater (2-layer MLP), similar to our ablation experiments. We refer to the source code for the experiment details.

Inspired by [Klindt \(2023\)](#), we generate training scalars X and Y from $\text{Uniform}[0.5, 1.0]$ and generate test set by sampling scalars from the $\text{Uniform}[0., 0.5]$ distribution.

We report the evaluation results in Table 8. The proposed discrete `ScalarUpdate` module successfully learned the correct operations for updates f_1, \dots, f_4 for all seeds and 3 times out of 5 for f_0 (note that the model was trained to predict different manipulations for different states simultaneously). For unsuccessful runs, when f_0 was not learned correctly, the learned operation for f_0 was g_3 for some states (i.e., $x \cdot y$ instead of x), which can be explained by optimization challenges as the distribution of y is close to 1.

Our experiment demonstrates that the proposed `ScalarUpdate` module can be extended to support a wider range of manipulations with scalars. We note that this complicates the optimization problem of selecting the correct operations/operands from the operations results (e.g., such decomposition might not be unique).

Table 8. MSE for train/test distributions for the discrete and non-discrete ScalarUpdate modules and different operations.

	f_0	f_1	f_2	f_3	f_4
discrete	0.01 / 0.1	0. / 0.	0 / 0	0 / 0	0 / 0
non-discrete	$5.7e-6$ / 0.03	$5.1e-6$ / 0.001	$1.1e-5$ / 0.007	$1.0e-5$ / 0.08	$2.0e-5$ / 0.03

E. State Usage for No-Hint Models

In this section, we provide several illustrations of the node states and the dynamics of the pointer prediction updates for the BFS algorithm (Figures 3-5). Our analysis suggests that no-hint models with K states tend to use states as distances from the starting node, with the distances $\geq K$ merged into the same state. Also, pointer predictions for the first K BFS layers are correct and computed layer-by-layer as in the ground truth algorithm. The mistakes of the model are at the later layers and some pointers at the later layers are computed before that in the ground truth algorithm (Figure 5c). For simplicity of illustrations, we use the model with 3 discrete states.

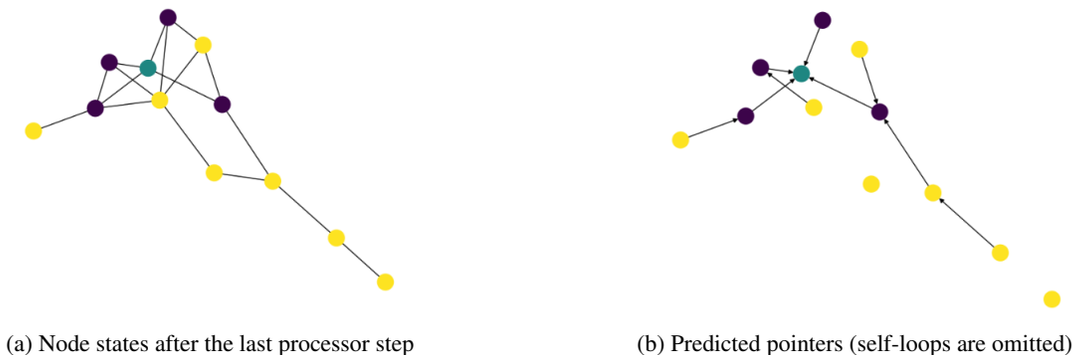


Figure 3. Node states and the predicted pointers after the last processor step of the DNAR model (with 3 states), trained without hints. Different colors represent different states. The green node is the starting node.

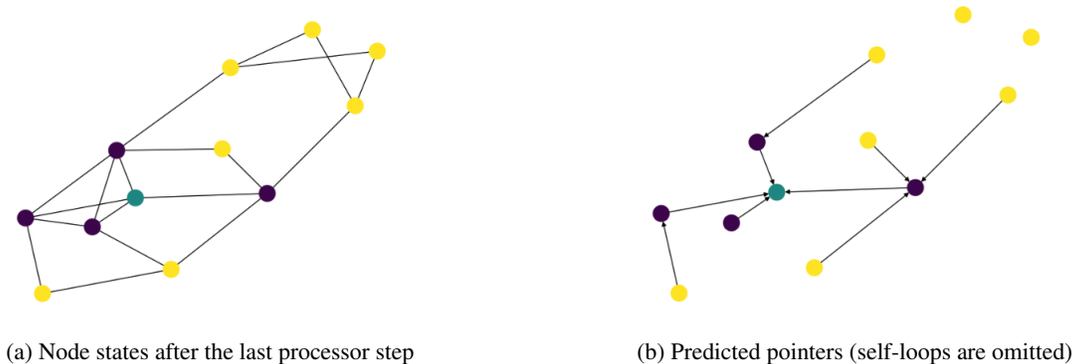


Figure 4. Node states and the predicted pointers after the last processor step of the DNAR model (with 3 states), trained without hints. Different colors represent different states. The green node is the starting node.

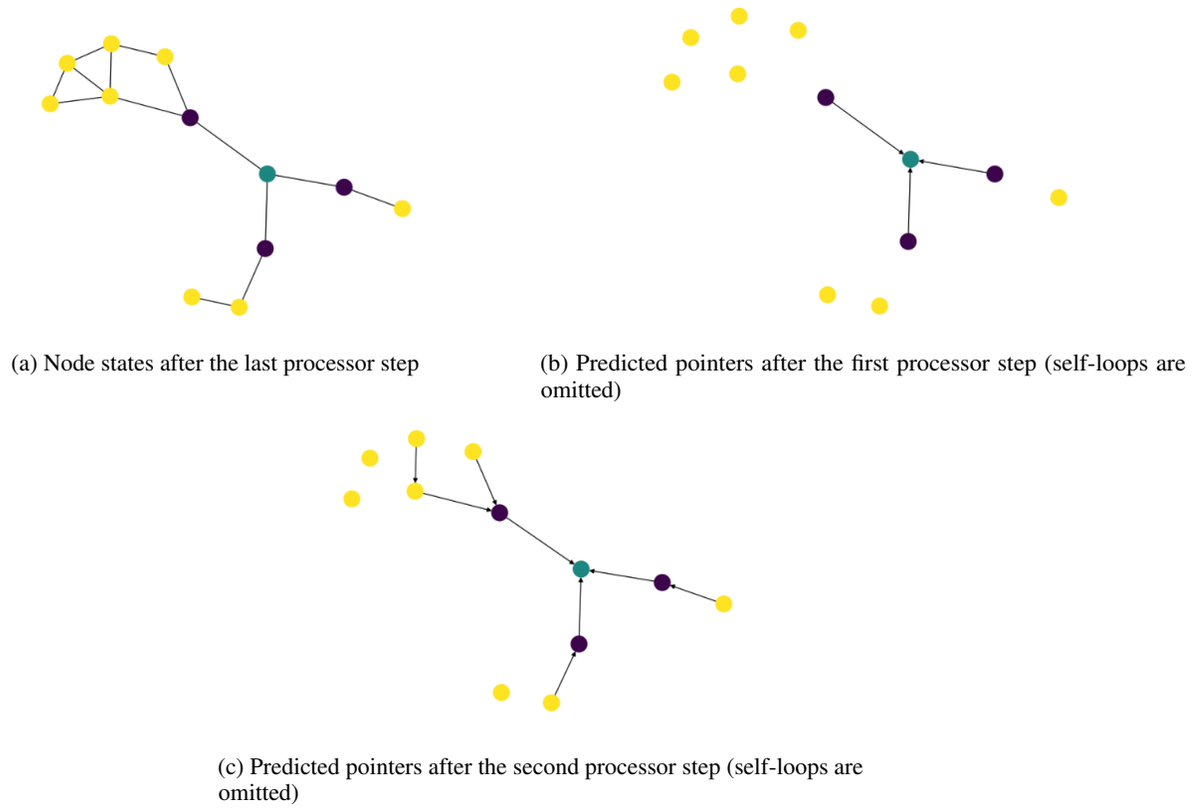


Figure 5. Node states and the dynamics of the pointer prediction updates of the DNAR model (with 3 states), trained without hints. Different colors represent different states. The green node is the starting node.