Learning Iterative Reasoning through Energy Minimization

Yilun Du ¹  Shuang Li ¹  Joshua Tenenbaum ¹  Igor Mordatch ²

Abstract

Deep learning has excelled on complex pattern recognition tasks such as image classification and object recognition. However, it struggles with tasks requiring nontrivial reasoning, such as algorithmic computation. Humans are able to solve such tasks through iterative reasoning – spending more time thinking about harder tasks. Most existing neural networks, however, exhibit a fixed computational budget controlled by the neural network architecture, preventing additional computational processing on harder tasks. In this work, we present a new framework for iterative reasoning with neural networks. We train a neural network to parameterize an energy landscape over all outputs, and implement each step of the iterative reasoning as an energy minimization step to find a minimal energy solution. By formulating reasoning as an energy minimization problem, for harder problems that lead to more complex energy landscapes, we may then adjust our underlying computational budget by running a more complex optimization procedure. We empirically illustrate that our iterative reasoning approach can solve more accurate and generalizable algorithmic reasoning tasks in both graph and continuous domains. Finally, we illustrate that our approach can recursively solve algorithmic problems requiring nested reasoning. Code and additional information is available at https://energy-based-model.github.io/iterative-reasoning-as-energy-minimization/.

1. Introduction

Human thinking is often characterized in terms of mechanisms for two distinct modes of cognitive processing: SYSTEM 1 mechanisms for fast, habitual, and associative processing, and SYSTEM 2 mechanisms for slower, more deliberate and controlled, symbolic reasoning (Kahneman, 2011). Neural networks have excelled at habitual SYSTEM 1-style processing in familiar environments and task contexts, such as mapping images of familiar objects to their semantic classes, or familiar locations to the corresponding routes of movement. However, confronted with a novel environment or task demanding a more flexible response, humans can invoke controlled SYSTEM 2 processes, often in the form of iteratively reasoning about relationships between observed entities that builds on past experiences primarily through shared abstractions and algorithms rather than direct re-use of concrete, habitual responses. Such flexible processing of novel inputs is difficult for neural networks, with even large pretrained language models (Radford et al., 2019) that have proven effective in many zero-shot generalization contexts failing to extrapolate the simplest algorithmic operations, such as addition, to more complex inputs.

Iterative reasoning, the ability to repeatedly apply an underlying computation to the outputs of previous reasoning steps, is a crucial component of scalable SYSTEM 2-style processing. We may take abstractions learned from simpler variants of a problem, and iteratively apply them to solve harder variants of potentially unbounded complexity. As an example, having learned how to compute shortest paths on small graphs, we might generalize and apply our algorithmic intuition sequentially to substantially larger graph problems.
problems at test time through iterative application of learned computations. Such iterative processing enables humans to solve more challenging tasks, such as question answering, arithmetic calculation, or proof writing, even in novel or unfamiliar domains.

We present a new neural approach towards iterative reasoning, which we formulate as an energy minimization process on a learned energy landscape (Figure 1). By representing individual steps of reasoning as an optimization process, we may iteratively reason for longer on harder problems by running additional steps of optimization on the induced energy landscapes. Simultaneously, by monitoring the geometric energy landscape surrounding an optimized solution, we may automatically determine the completion of the algorithmic computation (by checking the presence of a local energy minimum). We refer to our underlying reasoning framework as Iterative Reasoning as Energy Minimization (IREM).

To evaluate and benchmark the effectiveness of iterative reasoning using IREM, we propose and construct a suite of different algorithmic reasoning tasks on both graph and continuous domains. Effective algorithmic reasoning requires repetitive application of underlying algorithmic computations, dependent on problem complexity, and thus serves as a natural benchmark for iterative reasoning. We compare IREM with past works on our algorithmic benchmark for iterative reasoning and find that IREM outperforms prior works in performance and generalization.

Our contributions in this paper are threefold. First, we present IREM, a new framework for iterative reasoning and analyze why it is beneficial to use such a framework for reasoning. Second, we present a benchmark for iterative algorithmic reasoning, both on graphs and continuous vector inputs, and show that IREM significantly outperforms prior approaches in both performance and generalization. Finally, we show how our approach can recursively solve algorithmic computation requiring nested reasoning. Our results point to IREM as a promising new approach towards iterative reasoning.

2. Related Work

Iterative Reasoning A variety of recent works have explored the integration of iterative reasoning into neural networks. One branch of works implements iterative reasoning by constructing neural programmatic operations (Graves et al., 2014; Reed & De Freitas, 2015; Banino et al., 2021) which are repeatedly executed till halting. Another branch of work implements iterative reasoning through recurrent computation (Graves, 2016; Bolukbasi et al., 2017; Chung et al., 2016; Schwarzschild et al., 2021). A key challenge with both types of approaches lies in the halting time of computation. Existing approaches learn halting policies through reinforcement learning (Chen et al., 2020; Chung et al., 2016), heuristic policies (Bolukbasi et al., 2017), or variational inference (Graves, 2016; Banino et al., 2021). Such approaches are unstable in nature (Banino et al., 2021), and many of them require manual hyper-parameter specification. We present an orthogonal approach towards implementing iterative reasoning as an energy minimization procedure on a learned energy landscape. By determining when a local energy minimum has been found, our approach provides a natural mechanism for terminating computation.

Algorithmic Reasoning with External Memory Several approaches towards iterative reasoning utilize an external memory scratchpad for algorithmic computation. Such a scratchpad enables models to store intermediate algorithmic computations, and thus boosts the underlying performance of the algorithm (Graves et al., 2014; Reed & De Freitas, 2015; Cai et al., 2017; Kaiser & Sutskever, 2015). In the Appendix B, we illustrate a manner through which we may utilize an external memory with IREM to improve underlying further improve reasoning performance.

Optimization Based Computational Blocks Prior works have explored optimization as a computation block to solve different tasks. In (Brockett, 1991), a dynamical system is constructed that can solve various algorithmic tasks. Optimization has since been used as an intermediate neural network computation block for quadratic programs (Amos & Kolter, 2017; Donti et al., 2017) and submodular programs (Djolonga & Krause, 2017; Wilder et al., 2019) for flexible neural networks. Most similar to our work, Bai et al. (2019) utilizes equilibrium energy minimization as an intermediate computation block for memory-efficient neural networks. In contrast, we explore how direct optimization over a learned energy landscape can enable generalizable iterative reasoning. Concurrent to our work, (Rubanova et al., 2021) utilizes energy minimization to simulate physical dynamics.

Energy-Based Models Our work is related to works in Energy-Based Models (EBMs) (LeCun et al., 2006). Most recent works using EBMs have focused on learning probabilistic models over data (Du & Mordatch, 2019; Nijkamp et al., 2019; Grathwohl et al., 2020; Du et al., 2020; 2021b; Arbel et al., 2020; Li et al., 2020; Xiao et al., 2020). Instead of using EBMs as a probabilistic model, we use EBMs to define an energy landscape for solving iterative reasoning problems.

Learning Optimizers Our work utilizes backpropagation through intermediate optimization steps to train our energy function. Prior work has explored a similar idea of backpropagation through optimization to learn meta optimizers (Andrychowicz et al., 2016; Ravi & Larochelle, 2016; Bengio et al., 1995; Schmidhuber, 1992; Hochreiter
We next discuss how to learn IREM. Given an input problem \( x_i \), with a unique solution \( y_i \), the simplest method to learn \( E_\theta(x, y) \) is to directly supervise the minimal energy state \( \arg\min_y E_\theta(x_i, y) \) with \( y_i \) through regression

\[
\mathcal{L}_{\text{MSE}}(\theta) = \| \arg\min_y E_\theta(x_i, y) - y_i \|^2. \tag{4}
\]

However, in practice, during training time, it is computationally expensive to compute \( \arg\min_y E_\theta(x, y) \), as the underlying energy landscape may be complex.

As a fast approximation to \( \arg\min_y E_\theta(x, y) \), we may approximate the \( \arg\min \) operation with respect to \( y \), via \( N \) steps of gradient optimization. An approximate optimum \( y_i^N \) is obtained by:

\[
y_i^N = y_i^{N-1} - \lambda \nabla_y E_\theta(x_i, y_i^{N-1}). \tag{5}
\]

In Equation 5, we initialize \( y_i^0 \) using uniform noise, with \( \lambda \) denoting the step size for each gradient step, and sample \( y_i^N \) corresponding to the result after \( N \) steps of gradient descent. We then minimize the corresponding loss:

\[
\mathcal{L}_{\text{MSE}}(\theta) = \| y_i^N - y_i \|^2, \tag{6}
\]

where we may directly differentiate through the underlying optimization procedure (Finn et al., 2017). To alleviate the computational burden of computing second-order gradients across optimization steps, we empirically found that simply truncating back-propagation to the last optimization step maintained good performance at a significantly faster training speed as illustrated in Table 2.

An underlying difficulty of utilizing \( y_i^N \) as an approximation of \( \arg\min_y E_\theta(x_i, y) \) is that since only a small finite number of steps of gradient descent is applied, \( y_i^N \) may be far from reaching \( \arg\min_y E_\theta(x_i, y) \). As a result, when running a larger number of iterative reasoning steps at test time to more precisely compute \( \arg\min_y E_\theta(x, y) \), our underlying solution may degrade.

To remedy this issue, we maintain a replay buffer of previously optimized samples \( y_i^N \), and initialize \( y_i^0 \) either from previously optimized values \( y_i^N \) or uniform noise. By initializing gradient descent optimization with previously optimized samples, we ensure that these samples are closer to \( \arg\min_y E_\theta(x_i, y) \). A similar application of replay buffers has been utilized to train EBMs for consistent probability landscapes (Du & Mordatch, 2019).

We provide pseudocode for training IREM in Algorithm 1 and executing algorithmic reasoning with IREM in Algorithm 2, where an energy minimum is determined at test time once the energy value of a solution does not change for certain iterations. A fixed learning rate, \( \lambda = 100 \), is used for training IREM. At test time, \( \lambda \) is empirically tuned so that energy values decrease smoothly across iterations of optimization (harder problems require smaller \( \lambda \) to optimize). In Appendix B, we discuss how we may utilize optimization...
We construct a class of algorithmic reasoning tasks that are well-known in complexity theory that constructing Algorithm 1 IREM training algorithm

**Input:** Data Dist \( P_D(x, y) \), Replay Buffer \( B \), Step Size \( \lambda \), Number of Steps \( N \), EBM \( E_0(\cdot) \), Uniform Distribution \( U(-1, 1) \)

\( B \leftarrow \emptyset \)

while not converged do
  \triangleright Sample data and candidate solutions from \( p_D \) and replay buffer \( B \)
  \[ x_i, y_i \sim P_D, \ y_i^0 \sim U(-1, 1) \]
  \[ x_i, y_i^0, y_i^1 \sim B \]
  \[ x_i, y_i^0, y_i^1 \leftarrow x_i \cup x_i^0 \cup y_i \cup y_i^0 \cup y_i^1 \]
  \triangleright Generate low energy solutions through optimization:
  for sample step \( n = 1 \) to \( N \) do
    \[ \tilde{y}_i^n \leftarrow \tilde{y}_i^{n-1} - \lambda \nabla_{\tilde{y}_i} E_0(x_i, \tilde{y}_i^{n-1}) \]
  end for
  \triangleright Optimize objective \( L_{\text{MSE}} \) wrt \( \theta \):
    \[ \Delta \theta \leftarrow \nabla_{\theta} \sum_{i=0}^{n} || \tilde{y}_i^n - y_i ||^2 \]
  Update \( \theta \) based on \( \Delta \theta \) using Adam optimizer

\triangleright Update replay buffer \( B \)
  \[ B \leftarrow B \cup (x_i, y_i, \tilde{y}_i^N) \]
end while

Algorithm 2 IREM prediction algorithm

**Input:** Data Dist \( P_D(x) \), Step Size \( \lambda \), Number of Steps \( K \), EBM \( E_0(\cdot) \), Uniform Distribution \( U(-1, 1) \)

\triangleright Sample input from \( P_D \) and initialize candidate solution
  \[ x_i \sim P_D \]
  \[ y_i \sim U(-1, 1) \]

while Not at Energy Minima do
  \triangleright Optimize candidate solution \( y_i \) with gradient descent:
    \[ y_i \leftarrow y_i - \lambda \nabla_{y_i} E_0(x_i, y_i) \]
end while

\triangleright Final predicted solution:
  \[ y = \tilde{y} \]



to further incorporate an external scratchpad into iterative reasoning using IREM.

As IREM directly trains an energy landscape by optimizing samples to regress solutions, there is no guarantee that a smooth underlying energy landscape \( E(x, y) \) is learned. Empirically, as seen in Figure 1 and Figure 5, we find that our objective does lead to consistent energy landscapes similar to past work (Du et al., 2021a).

### 3.3. Analysis

In this section, we provide complexity-theoretic motivation for representing iterative reasoning as an energy minimization is advantageous. We consider two approaches to represent reasoning:

**Feedforward Computation.** We consider reasoning as a feedforward function \( f \) of the form \( f(x) \) where \( f(x) : \mathbb{R}^O \rightarrow \mathbb{R}^M \) maps a input \( x \) to a predicted solution \( y \).

**Energy Minimization.** We next consider reasoning as a energy minimization problem, \( \arg \min_y E(x, y) \), where \( E(x, y) \) is a function from \( \mathbb{R}^O \times \mathbb{R}^M \rightarrow \mathbb{R} \), mapping a input \( x \) and solution \( y \) into an energy. This corresponds to the computation represented by IREM.

We construct a class of algorithmic reasoning tasks that are easier to learn using an energy function \( E(x, y) \) as opposed to the feedforward function \( f(x) \). Our result is based on the intuition that learning the energy function \( E(x, y) \) corresponds to learning a solution verifier, which assigns minimal energy to the correct solution, and high energy to all the other solutions. In contrast, learning a function \( f(x) \) corresponds to explicitly generating a solution. We rely on the well-known theorem in complexity theory that constructing a solution verifier is easier than a solution generator to show that learning energy minimization is easier than feedforward computation.

To analyze \( E(x, y) \), we consider solving the 3-SAT (Impagliazzo & Paturi, 1999) problem. We construct a energy function \( E(x, y) \) to verify the 3-SAT formula. Given a 3-SAT formula \( \phi \) with \( D \) variables and \( K \) clauses, we construct an energy function to represent \( \phi \) as \( E(x, y) := \sum_{1 \leq k \leq K} \epsilon_k(x, y) \), where \( x \) encodes the clauses in a 3-SAT problem, \( y \) corresponds to boolean assignments to each variable, and \( \epsilon_k \) verifies whether the boolean assignments in \( y \) satisfies the \( k \)th clause. To encode a set of \( K \) clauses using \( x \), we utilize an ordinal representation (e.g. \( x_1 = [1, 2, 3] \) to represent a particular clause \((y_1 \land y_2 \land y_3))\). The energy function \( \epsilon_k \) is then constructed by taking in clause \( x_i \) and outputting 0 if the corresponding entries of \( y \) satisfy the encoded clause and 1 otherwise (polynomial time to compute). We assume the Exponential Time Hypothesis (ETH) (Impagliazzo & Paturi, 1999), which states that checking the satisfiability of a 3-SAT formula takes time exponential in the sum of the number of variables and the number of clauses.

**Remark 1.** There exists a 3-SAT problem which may be encoded in an energy function \( E(x, y) \) which can be evaluated at any input in time polynomial in the number of dimensions of \( x \) but for which the computational complexity of encoding a feedforward solution \( f(x) \) which may be evaluated at any input is (worse-case) exponential in the number of dimensions of \( x \).

**Proof.** We encode the 3-SAT energy function \( E(x, y) \) as defined above, which is evaluated in time polynomial in the number of dimensions of \( x \). In contrast, ETH directly implies that constructing \( f(x) \), which corresponds to solving the 3-SAT problem, is exponential in dimension of \( x \).

Our remark shows that it is computationally advantageous to represent an energy function \( E(x, y) \) as opposed to a feedforward decoder \( f(x) \). In particular, learning neural networks \( E_0(x, y) \) and \( f_0(x) \) to approximate either \( E(x, y) \) or \( f(x) \), our remark implies that a larger network is necessary to represent the exponential computations of \( f(x) \).
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| Task          | Method          | Same Size | Larger Size | Task          | Method          | Same Diff. | Harder Diff. |
|---------------|-----------------|-----------|-------------|---------------|-----------------|------------|--------------|
| Edge Copy     | Feedforward     | 0.3016    | 0.3124      | Add           | Feedforward     | 0.0448    | 0.7029       |
|               | Recurrent       | 0.3015    | 0.3113      |               | Recurrent       | 0.3610    | 2.6133       |
|               | Programmatic    | 0.3053    | 0.4409      |               | Programmatic    | 0.0111    | 0.3446       |
|               | Iterative Feedforward | 0.6163 | 0.6498      |               | Iterative Feedforward | 0.0144 | 0.1577       |
|               | IREM (Ours)     | 0.0019    | 0.0019      |               | IREM (Ours)     | 0.0003    | 0.0021       |
| Connected Components | Feedforward     | 0.1796    | 0.3460      | Matrix Completion | Feedforward     | 0.0203    | 0.2720       |
|               | Recurrent       | 0.1794    | 0.2766      |               | Recurrent       | 0.0266    | 0.3285       |
|               | Programmatic    | 0.2338    | 3.1381      |               | Programmatic    | 0.0203    | 0.2637       |
|               | Iterative Feedforward | 0.4908 | 1.2064      |               | Iterative Feedforward | 0.0253 | 0.2102       |
|               | IREM (Ours)     | 0.1424    | 0.2171      |               | IREM (Ours)     | 0.0183    | 0.2074       |
| Shortest Path | Feedforward     | 0.1233    | 1.4089      | Matrix Inverse | Feedforward     | 0.0112    | 0.2150       |
|               | Recurrent       | 0.1259    | 0.1083      |               | Recurrent       | 0.0109    | 0.2123       |
|               | Programmatic    | 0.1375    | 0.1290      |               | Programmatic    | 0.0124    | 0.2209       |
|               | Iterative Feedforward | 0.4588 | 0.7688      |               | Iterative Feedforward | 0.0270 | 0.5250       |
|               | IREM (Ours)     | 0.0274    | 0.0464      |               | IREM (Ours)     | 0.0108    | 0.2083       |

Table 1. Algorithmic Reasoning with IREM– IREM is a general framework for iterative reasoning which can learn algorithmic computation on both graph (left) and continuous (right) inputs. IREM generalizes at test time to both larger (left) and harder (right) algorithmic problems through iterative computation. Error reported on each task using elementwise mean square error. Approaches on the left are trained on graphs with ten nodes and evaluated on larger graphs with fifteen nodes. IREM significantly outperforms comparisons.

Such a network $f_\theta(s)$ would require either exponentially deeper or wider layers, if there is no iterative computation. With iterative computation, we may parameterize $f_\theta(x)$ and $E_\theta(x, y)$ with a similar number of parameters, but then it would be necessary for $f_\theta(x)$ to be iteratively applied an exponential number of times. Direct recurrent backpropagation through such a number of iterative computations has been proven to be unstable to train. While in principle computing $\arg\min_y E(x, y)$, would require a similar number of computations, we may train $E(x, y)$ with a simple inexact energy minimization procedure and run a more extensive minimization procedure at test time. We next analyze the computational complexity of representing $E(x, y)$ and $f(x)$ as a function of problem size.

**Remark 2.** As the underlying number of variables in 3-SAT problem increases, we may construct $E(x, y)$ which can be evaluated at any input in time polynomial in the number of variables, but for which the computational complexity of encoding a feedforward solution $f(x)$ which may be evaluated at any input is (worse-case) exponential in the number of underlying variables.

**Proof.** Our constructed energy function $E(x, y)$ above may be evaluated in time polynomial in the number of input variables. In contrast, ETH implies that constructing $f(x)$, which corresponds to solving the 3-SAT problem, is exponential in the number of underlying variables. □

Similar to our previous remark, this result implies that representing $f(x)$ requires exponentially more computation as the underlying problem size of a 3-SAT problem increases. In particular, our remark has implications for generalization. If we learn $f_\theta(x)$ and $E_\theta(x, y)$ on smaller problem instances of $x$, our neural networks to generalize correctly to larger problems instances, the underlying computation executed by the neural network must increase polynomially for $E_\theta(x, y)$ and exponentially for $f_\theta(x)$. Existing architectures, such as transformers and graph networks, adaptively increase computation polynomially for larger inputs. In contrast, few architectures can adaptively increase computation exponentially for larger problem instances. To realize $f(x)$, we thus require iterative approaches which can execute exponentially longer on larger inputs. As seen in Figure 2, this is difficult for existing approaches.

4. Experiments

4.1. Experimental Setup

We compare IREM with feedforward and iterative baselines for reasoning. We discuss each approach in detail below.

**Feedforward Computation.** First, we compare with (one-step) feedforward computation, where we train a neural network that directly outputs the values of solutions.

**Recurrent Network Computation.** Next, we compare our approach with methods utilizing a recurrent neural network to execute iterative computation. Recurrent architectures have been shown to successfully execute reasoning recently (Schwarzschild et al., 2021). We use a LSTM network to represent iterative computation.

**Learned Programmatic Computation.** We compare our method with past works which construct iterative computation through building programmatic structures with neural networks. We compare our method with the recent architecture and training objective of PonderNet (Banino et al., 2021), which variationally learns both a halting probability and individual computation step networks.

**Iterative Feedforward Computation.** We further compare our approach with direct iterative application of a feed-
We report the underlying elementwise mean squared error between predictions from models and their associated ground truth outputs. We evaluate performance on the three algorithmic reasoning tasks. All iterative methods are trained with 5 steps of iterative computation on size 10 graphs, while a recurrent network fails to do so.

forward computation. We train the iterative feedforward computation using an iterative denoising objective (Sohl-Dickstein et al., 2015).

We scale network sizes to ensure that each individual baseline has roughly the same number of parameters. The architectural details of each model are provided in Appendix D of the paper. We utilize MLP neural networks for continuous algorithmic reasoning tasks and graph neural networks for graph algorithmic reasoning tasks. All iterative methods are trained with 5 steps of reasoning. We provide comparisons with each baseline on graphical algorithmic reasoning in Section 4.2 and on continuous algorithmic reasoning in Section 4.3. We provide additional benchmark comparisons of IREM on an existing iterative image denoising task in Appendix A.

4.2. Graphical Algorithmic Reasoning

Setup. We first evaluate our approach on graphical algorithmic reasoning. We train models on fully connected graphs with between 2 to 10 nodes, and evaluate performance on larger fully connected graphs with size 15 nodes. We report the underlying elementwise mean squared error between predictions from models and their associated ground truth outputs. We evaluate performance on the three different graphical algorithmic reasoning tasks, aiming to capture different aspects of reasoning, which we detail below, with additional details about each dataset in Appendix C.

1. **Edge Copy**: We first test the ability of models to copy and output the values of all edges in a dense graph that is given as input. This task serves as a simple test for iterative reasoning, and requires a method to sequentially copy over input edge values to the output.

2. **Connected Components**: Next, we evaluate the ability of models to infer the underlying connected components of a graph. We construct a sparse graph, where 5% of all fully connected edges exist, and ask models to predict a binary indicator on whether a node of a graph is connected to another for all pairs of nodes in a graph. This task tests structural discovery, an aspect of cognitive reasoning (Kemp & Tenenbaum, 2008).

3. **Shortest Path**: Finally, we evaluate the ability of models to compute the shortest path distances between all pairs of nodes in a graph. This task tests for planning, and the underlying calculation necessary to compute the shortest paths between nodes is analogous to that of planning.

Quantitative Results. We present quantitative results on graphical algorithmic reasoning in the left column of Table 1. During test time, we evaluate on problems with either similar or larger sizes than those seen during training. Across all three tasks, IREM outperforms all the compared baselines. This difference in performance is magnified when evaluating on larger graphs during testing. Approaches other than IREM fail even on the relatively simple edge copy task. This is due to the fact that the small size of graph networks (hidden size 128), requires methods to iteratively copy different subsets of input edges to output predictions. While IREM successfully learns this iterative computation, the compared baselines were unable to do so.

Adaptive Computation. Next, we analyze the ability of IREM and baselines to adapt its underlying computational budget to different larger problem instances. In Figure 2, we illustrate shortest path computation error on different input graphs sizes at test time. While all methods are only trained with 5 steps of iterative computation on size 10 graphs, we find that IREM can generalize iterative computation to size 30 graphs. In contrast, the learned iterative baselines, such as the recurrent network in Figure 2, fail to do so. We find similar behavior across other graphical algorithmic reasoning tasks.

Ablations. We run an ablation analysis on the impact of utilizing a replay buffer to train IREM, as well as the effect of truncating backpropagation to the last step of optimization. In Table 2, we find that utilizing a replay buffer significantly improves the performance of IREM. We further find that truncating backpropagation to the last step of optimization has a limited impact on IREM and greatly reduces the overall memory cost of training, as well as slightly

**Table 2. Ablations** – Ablations of proposed components of IREM on test performance on the shortest path algorithmic reasoning task. The use a replay buffer boosts the underlying performance of IREM and truncating gradient backpropagation to the last step of optimization reduces both training time and memory cost.

| Ablations | Buffer | Truncate | Test Performance | Training Speed | Memory Usage |
|-----------|--------|----------|------------------|----------------|--------------|
| No        | No     | 0.0491   | 87%              | 295%           |
| Yes       | No     | 0.0287   | 83%              | 295%           |
| Yes       | Yes    | 0.0274   | 100%             | 100%           |
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Figure 3. Computation Steps vs Problem Difficulty – Illustration of MSE error of prediction as a factor of the test time difficulty (harder difficulty right) of the task and computation steps applied. Each model is trained with 5 steps of iterative computation. Models missing in plots have errors greater than the range displayed in the plot. The error of IREM improves with the number of underlying algorithmic computation steps, with larger number of computation steps benefiting performance on harder algorithmic tasks.

improving the training speed of IREM.

4.3. Continuous Algorithmic Reasoning

Data Setup. We next evaluate IREM and baselines on continuous algorithmic reasoning tasks. We apply algorithmic operations on input vectors of size 400 (resized to 20×20 matrices for matrix operations). We report the underlying MSE error between the predictions and the associated ground truth outputs on test problem instances. We evaluate different methods on the following three tasks, aiming to capture different aspects of reasoning, with additional details in the Appendix C.

1. Addition: We first evaluate the algorithmic computation of addition. We train networks to add entries in two separate input vectors (element-wise). We construct harder variants of the addition problems at test time by feeding input vectors with larger magnitudes. This task aims to test simple arithmetic reasoning.

2. Matrix Completion: Next, we evaluate the algorithmic computation of matrix completion. We mask out 50% of the entries of a low-rank input matrix constructed from two separate low-rank matrices $U$ and $V$, and train networks to reconstruct the original input matrix. We construct harder variants of the matrix completion problem at test time by increasing the complexity of $U$ and $V$. This task aims to test both structural and analogical reasoning, with both shown to be equivalent to matrix completion (Lampinen et al., 2017).

3. Matrix Inverse: Finally, we evaluate the algorithmic computation of matrix inverse. We train networks to compute the matrix inverse of an input matrix. We construct harder matrix inverse problems by considering less well-conditioned input matrices. This task aims to test the numerical reasoning, with matrix inversion a crucial operation across various numerical algorithms.

Quantitative Results. We present quantitative results on each of our three continuous algorithmic reasoning problems on the right side of Table 1 on test problems with either the same or harder difficulty than training problems. Across all three tasks, we find that IREM outperforms baselines, with the underlying difference magnified on harder, out-of-distribution test problems. In particular, on the task of addition, we find that IREM is able to nearly perfectly solve the underlying task even on harder, out-of-distribution test problems. In contrast, all other evaluated iterative and
We further qualitatively visualize the underlying iterative computation learned by IREM and baselines. In Figure 3, we illustrate the prediction error of different methods as a function of the number of computation steps applied. As we increase the number of iterative computation steps, the underlying performance of IREM continues to improve. In particular, iterative computation helps more substantially on harder variants of the algorithmic problem (right column), such as addition. In contrast, several iterative methods show significant degradation of performance with increased iterative computation.

We further qualitatively visualize the underlying iterative computation learned by IREM. In Figure 4, we visualize the element-wise mean square error of the predicted solution as a function of the number of iterative reasoning steps applied on the addition task. We find that energy minimization gradually refines a predicted solution to the ground truth additive answer, with different elements of the solution exhibiting different convergence rates.

**Energy Landscape.** IREM parameterizes an energy landscape across all possible solutions for a given problem. Such an energy landscape enables us to assess the relative quality of solutions dependent on their associated energies, and further gives a natural objective to terminate iterative computation when an underlying local energy minimum is reached. In Figure 5, we visualize the predicted energy of different candidate solutions and their corresponding MSE distances from the ground truth answer. We find that across different continuous algorithmic tasks, the underlying energy value assigned to a candidate solution is well correlated with its distance from the ground truth answer, with low energy solutions close to the ground truth.

**Sensitivity to Step Size.** We assess the performance of IREM under different values of step size \( \lambda \) using during energy optimization at training. We consider the continuous addition task in Table 3. We find that the underlying performance is not sensitive to hyperparameter choice for step size, and utilize a fixed step size of 100 across our experiments.

### 4.4 Recursive Algorithmic Computation

**Setup.** We further evaluate the ability of algorithms represented by IREM to be recursively applied on inputs. Recursively nesting algorithms enable complex computations, but require learned networks to be robust to out-of-distribution outputs from prior algorithmic execution.

We consider recursive applications of a learned algorithmic operator \( \text{Alg}_{\theta} (\cdot) \) representing addition as introduced in Section 4.3. We evaluate the element-wise mean square error of the predicted output \( \hat{y} \) of recursively applying the learned algorithmic operator.

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Table 3. Ablation Analysis of Step Size in IREM – Analysis of training step size of IREM performance on the continuous addition task.

| Step Size | Same Difficulty | Harder Difficulty |
|-----------|-----------------|-------------------|
| 10        | 0.0003          | 0.0021            |
| 30        | 0.0003          | 0.0020            |
| 100       | 0.0003          | 0.0021            |
| 300       | 0.0004          | 0.0023            |
| 1000      | 0.0004          | 0.0025            |

Table 4. Algorithmic Composition – Test performance when composing multiple instances of the addition operation. Error is reported using element-wise mean square error. IREM is able to generalize well when composing algorithmic computations.

```
Method       2  5  10
Feedforward  0.0445 0.2717 0.8898
Recurrent    2.1377 3.1861 4.8706
Programmatic 0.0203 0.1068 0.4587
Iterative Feedforward 0.0826 0.5930 3.6004
IREM (Ours)  0.0014 0.0078 0.0422
```
algorithmic operator $k$ times

$$\hat{y}^k = \text{Alg}_\theta(\hat{y}^{k-1}, y_k),$$  \quad (7)$$

with the corresponding ground truth solution being $\sum_k y_k$, for different values $k$ of recursive application.

**Quantitative Results.** We report the results of recursively applying each learned algorithmic operator between two to ten times in Table 4. We find that IREM supports the most stable recursive application of algorithmic operators. IREM exhibits significantly lower error than all compared baseline, due to its ability to utilize iterative computation to deal with out-of-distribution inputs and to accurately compute intermediate algorithmic outputs.

**Qualitative Results.** We illustrate error as a factor of the number of applied algorithmic operations in Figure 7, and find that the error of predictions from IREM rises slowly in comparison to other baselines. We further visualize the nested algorithmic predictions from IREM. We illustrate the inferred array sum predicted by IREM when four separate inputs are summed in Figure 6. As seen above, our approach enables us to closely approximate the addition of four input matrices (with the first nine entries shown).

5. Conclusion and Limitations

In this paper, we present IREM, a new approach towards iterative computation, by formulating it as an energy minimization process. We illustrate, on both continuous and graphical domains, how iterative computation utilizing IREM enables better algorithmic performance, as well as generalization to more complex instances of problems. We further illustrate how the underlying algorithmic computation learned by IREM may be nested to implement more complex algorithmic computations.

Iterative reasoning with IREM has several limitations. First, while IREM substantially outperforms existing iterative methods on tasks where output solutions have high dimensionality, limited gains are obtained when IREM is executed on problems with lower dimensionality solutions (such as parity prediction). Second, as training and inferring solutions with IREM relies on continuous gradient optimization, IREM struggles when output solutions have discrete values. An interesting line of future work would be to explore how discrete optimization could be integrated with training IREM to solve such discrete valued problems. Finally, since the training procedure of IREM requires backpropagation across gradient optimization steps, it is computationally expensive. An interesting line of future work could be exploring alternative ways to train an energy function for reasoning, such as utilizing gradient-free optimization.

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Appendix

In this appendix we provide additional evaluation and details on IREM. First, we illustrate how IREM may be applied to an existing image iterative reasoning task in Appendix A. Next we discuss how we may utilize optimization to execute IREM with an external scratchpad in Appendix B. We further discuss additional experimental details on our evaluated algorithmic tasks in Appendix C. Finally, we discuss individual model architectures used in Appendix D.

A. Image Denoising

Setup We compare IREM with existing approaches on an existing image based iterative reasoning benchmark from (Chen et al., 2020). The benchmark task is to denoise images with various levels of Gaussian noise corruption added. Harder denoising tasks are constructed at test time by adding larger amounts of noise to input images. We directly compare with baselines and numbers for UNLNet, DnCNN and DnCNN-stop from (Chen et al., 2020) and utilize the authors provided training code to train IREM with five steps of iterative reasoning.

Quantitative Results We quantitatively evaluate the performance of IREM and baselines in terms of PSNR (numbers for baselines directly from (Chen et al., 2020)) in Table 5. While we found that IREM performed similarly to existing approaches when the test noise corruption is similar to that of training, IREM significantly outperformed the compared baselines when evaluated on harder images.

While we trained IREM with 5 steps of reasoning at noise level \( \sigma = 65 \) improved performance, while running up to 100 steps of reasoning at noise level \( \sigma = 75 \) further improved performance.

Qualitative Results We qualitatively illustrate IREM being applied to an image corrupted with an unseen noise level \( \sigma = 75 \) (significantly larger than what is seen during training) in Figure 8. As seen in Figure 8, images continue to become clearer, even after 50 steps of iterative computation using IREM (with dress in the top row and floor in the bottom row becoming clearer).

B. Iterative Reasoning with a Scratchpad

We next discuss how we may incorporate an external computational scratchpad when executing an iterative computation with IREM. To enable the processing of an external scratchpad, \( z \in \mathbb{R}^D \), we construct a EBM, \( E_\theta(x, y, z) : \mathbb{R}^N \times \mathbb{R}^M \times \mathbb{R}^D \to \mathbb{R} \), which takes as input an input problem \( x \), a candidate solution \( y \), and a scratchpad state \( z \).

We then define our predicted solution

\[
\hat{y} = \arg \min_y \min_z E_\theta(x, y, z),
\]

where optimize over both output solutions \( y \) and an external scratchpad \( z \) at prediction time. Analogous to the training procedure in the main paper, we may obtain a fast approximation of \( \hat{y} \) as \( y^N \) training time by jointly optimizing \( y \) and \( z \)

\[
\begin{align*}
    y_i^n &= y_i^{n-1} - \lambda \nabla_y E_\theta(x, y_i^{n-1}, z_i^{n-1}) \\
    z_i^n &= z_i^{n-1} - \lambda \nabla_z E_\theta(x, y_i^{n-1}, z_i^{n-1}).
\end{align*}
\]

We then analogously minimize the corresponding loss:

\[
\mathcal{L}_{\text{MSE}}(\theta) = \|y_i^N - y_i\|^2,
\]

We provide the overall pseudocode for training IREM with external memory in Algorithm 3.

C. Experimental Details

Graphical Algorithmic Computation Models were trained in approximately 2 hours on a single Nvidia Titan X GPU using a training batch size of 64 and the Adam optimizer with learning rate 1e-4. Each model was trained for 10,000 iterations and evaluated on 1000 test problems.
Algorithm 3 IREM Training with External Memory

**Input:** Data Dist $p_d(x, y)$, Replay Buffer $B$, Step Size $\lambda$, Number of Steps $N$, EBM $E_0()$, Uniform Distribution $U(-1, 1)$

1. **while not converged do**
   2. **Sample data and candidate solutions from $p_d$ and replay buffer $B$**
   3. $x_i, y_i \sim p_d, \tilde{y}_i^0, \tilde{z}_i^0 \sim U(-1, 1)$
   4. $x_i, y_i, \tilde{y}_i^0, \tilde{z}_i^0 \sim B$
   5. $x_i, y_i, \tilde{y}_i^0, \tilde{z}_i^0 \leftarrow x_i \cup x_i^1 \cup y_i \cup \tilde{y}_i^0 \cup \tilde{y}_i^0 \cup \tilde{z}_i^0 \cup \tilde{z}_i^0$
   6. **Generate low energy solutions through optimization:**
   7. **for** sample step $n = 1$ to $N$ **do**
   8. $\tilde{y}_i^n \leftarrow \tilde{y}_i^{n-1} - \lambda \nabla_y E_0(x_i, \tilde{y}_i^{n-1}, \tilde{z}_i^{n-1})$
   9. $\tilde{z}_i^n \leftarrow \tilde{z}_i^{n-1} - \lambda \nabla_z E_0(x_i, \tilde{y}_i^{n-1}, \tilde{z}_i^{n-1})$
   10. **end for**
   11. **Optimize objective $L_{MSE}$ with $\theta$:**
   12. $\Delta \theta \leftarrow \nabla_\theta \sum_{i=1}^N \| \tilde{y}_i^N - y_i \|^2$
   13. **Update $\theta$ based on $\Delta \theta$ using Adam optimizer**
   14. **Update replay buffer $B$**
   15. $B \leftarrow B \cup (x_i, y_i, \tilde{y}_i^0, \tilde{z}_i^N)$
   16. **end while**

Each model was trained with five steps of iterative computation, with PonderNet trained with a halting geometric distribution of 0.8. Below, we provide additional numerical details about each of the evaluated algorithmic tasks.

1. **Edge Copy:** We randomly sample a value for each edge in a fully connected graph with a uniform value between -1 and 1. Models are then tasked with replicating the value of each individual edge in the graph in the final output prediction.

2. **Connected Components:** We randomly zero-out 95% of the edges in a fully connected graph. Models are then tasked with predicting the pairwise connectivity of all possible pairs of nodes in the graph.

3. **Shortest Path:** We randomly sample an edge distance between 0 and 1 for each edge in a fully connected graph. Models are then tasked with predicting the pairwise shortest distance between all possible pairs of nodes in the graph.

**Continuous Algorithmic Computation** Models were trained in approximately 2 hours on a single Nvidia Titan X GPU using a training batch size of 128 and the Adam optimizer with learning rate 1e-4. Each model was trained for 10,000 iterations and evaluated on 1000 test problems. Each model was trained with five steps of iterative computation, with PonderNet trained with a halting geometric distribution of 0.8. We further provide individual dataset details below.

1. **Addition:** We randomly construct two separate vectors, each with 400 elements, with each element in the vector randomly sampled between -1 and 1. Models are then tasked with summing up the elements in each vector element-wise. When constructing more difficult addition problems at test time, each element in the vector is randomly sampled between -2.5 and 2.5.

2. **Matrix Completion:** We randomly construct a low-rank matrix $M$ represented as the $M = U^T V + 0.1 N(0, 1)$, where $U$ and $V$ are $10 \times 20$ matrices, with each individual elements in $U$ and $V$ sampled from $N(0, 0.22)$ . Models are given 50% of the entries of $M$ are tasked with recovering all entries of $M$. When constructing more difficult matrix completion problems at test time, each element in $U$ and $V$ are sampled from $N(0, 0.47)$.

3. **Matrix Inverse:** We randomly construct a well conditioned invertible matrix $M = R + R^T + 0.5 * I$, where $R$ is a random matrix, with individual elements sampled between -1 and 1. Models are tasked with computing the matrix inverse of $M$. When constructing more difficult matrix inversion problems at test time, we make $M$ less well-conditioned by setting $M = R + R^T + 0.1 * I$.

**D. Model Architectures**

**Graphical Algorithmic Computation** For each iterative and feedforward method, we utilize the GINEConv layer from (Hu et al., 2019), where GINEConv$(128, 128)$ refers to a graph convolution operator with node features 128 and edge feature 128. An input problem instance $x$ consists of a set of nodes features $v$ and edge features $e$. To parameterize the EBM in IREM $E_0(x, y)$ we concatenate the optimized prediction $y$, with $e$, which then utilized in the GINEConv layer. To obtain edge predictions for baselines, we pairwise concatenate node features for the given edges, and apply an FC layer to obtain the corresponding prediction following (Zhang & Chen, 2018). We specify the architecture for IREM in Table 6, the architecture for feedforward and iterative feedforward baselines in Table 7, the architecture for recurrent baselines in Table 8, and the architecture for programmatic execution baselines in Table 9. All models have roughly the same number of underlying parameters.

| GINEConv$(128, 128)$ | GINEConv$(128, 128)$ |
|----------------------|----------------------|
| GINEConv$(128, 128)$ | GINEConv$(128, 128)$ |
| GINEConv$(128, 128)$ |
| Linear $128 \rightarrow 1$ |

Table 6. The model architecture for IREM on graphical tasks.

| GINEConv$(128, 128)$ | GINEConv$(128, 128)$ |
|----------------------|----------------------|
| GINEConv$(128, 128)$ |
| Linear $256 \rightarrow$ Output Dim |

Table 7. The model architecture for feedforward and iterative feedforward baselines on graphical tasks.
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| GINEConv(128, 128) | GINEConv(128, 128) |
|---------------------|---------------------|
| LSTM(128)           | GINEConv(128)       |
| GINEConv(128, 128)  | Linear 256 → Output Dim |

Table 8. The model architecture for recurrent baseline on graphical tasks.

| GINEConv(128, 128) | GINEConv(128, 128) |
|---------------------|---------------------|
| Linear 256 → Output Dim | Linear → 1 |

Table 9. The model architecture for PonderNet baseline on graphical tasks.

**Continuous Algorithmic Computation** For each iterative and feedforward method, we utilize a MLP to implement continuous algorithmic computation. To parameterize the EBM in IREM $E_\theta(x, y)$, we concatenate $x$ and $y$ together as input into the network. We utilize the ReLU activation in all networks except IREM, where we utilize the Swish activation. We specify the architecture for IREM in Table 10, the architecture for feedforward and iterative feedforward baselines in Table 11, the architecture for recurrent baselines in Table 12, and the architecture for programmatic execution baselines in Table 13. All models have roughly the same number of underlying parameters.

| Linear 512 |
|------------|
| Linear 512 |
| Linear 512 |
| Linear → 1 |

Table 10. The model architecture for IREM on continuous tasks.

| Linear 512 |
|------------|
| Linear 512 |
| Linear 512 |
| Linear 512 |
| Linear → Output Dim |

Table 11. The model architecture for feedforward and iterative feedforward baselines on continuous tasks.

| Linear 196 |
|------------|
| LSTM 196 |
| Linear → Output Dim |

Table 12. The model architecture for recurrent baseline on continuous tasks.

| Linear 512 |
|------------|
| Linear 512 |
| Linear 512 |
| Linear 512 |
| Linear → Output Dim |

Table 13. The model architecture for PonderNet baseline on continuous tasks.