# Combinatorial Optimization with Physics-Inspired Graph Neural Networks

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Combinatorial optimization problems are pervasive across science and industry. Modern deep learning tools are poised to solve these problems at unprecedented scales, but a unifying framework that incorporates insights from statistical physics is still outstanding. Here we demonstrate how graph neural networks can be used to solve combinatorial optimization problems. Our approach is broadly applicable to canonical NP-hard problems in the form of quadratic unconstrained binary optimization problems, such as maximum cut, minimum vertex cover, maximum independent set, as well as Ising spin glasses and higher-order generalizations thereof in the form of polynomial unconstrained binary optimization problems. We apply a relaxation strategy to the problem Hamiltonian to generate a differentiable loss function with which we train the graph neural network and apply a simple projection to integer variables once the unsupervised training process has completed. We showcase our approach with numerical results for the canonical maximum cut and maximum independent set problems. We find that the graph neural network optimizer performs on par or outperforms existing solvers, with the ability to scale beyond the state of the art to problems with millions of variables.

### I. INTRODUCTION

Optimization is ubiquitous across science and industry. Specifically, the field of combinatorial optimization—the search for the minimum of an objective function within a finite but often large set of candidate solutions—is one of the most important areas in the field of optimization, with practical (vet notoriously challenging) applications found in virtually every industry, including both the private and public sectors, as well as in areas such as transportation and logistics, telecommunications, and finance [1-5]. While efficient specialized algorithms exist for specific use cases, most optimization problems remain intractable, especially in real-world applications where problems are more structured and thus require additional steps to make them amenable to traditional optimization techniques. Despite remarkable advances in both algorithms and computing power, significant yet generic improvements have remained elusive, generating an increased interest in new optimization approaches that are broadly applicable and radically different from traditional operations research tools.

In the broader physics community, the advent of quantum annealing devices such as the D-Wave Systems Inc. quantum annealers [6–9] has spawned a renewed interest in the development of heuristic approaches to solve discrete optimization problems. On the one hand, recent advances in quantum science and technology have inspired the development of novel classical algorithms, sometimes dubbed nature-inspired or physics-inspired

algorithms (e.g., simulated quantum annealing [10, 11 running on conventional CMOS hardware) that have raised the bar for emerging quantum annealing hardware; see, for example, Refs. [12–15]. On the other hand, in parallel to these algorithmic developments, substantial progress has been made in recent years on the development of programmable special-purpose devices based on alternative technologies, such as the coherent Ising machine based on optical parametric oscillators [16, 17], digital MemComputing machines based on self-organizing logic gates [18, 19], and the ASIC-based Fujitsu Digital Annealer [20–22]. Some of these approaches face severe scalability limitations. For example, in the coherent Ising machine there is a trade off between precision and the number of variables and the Fujitsu Digital Annealer — baked into an ASIC — can currently handle at most 8192 variables. Thus, it is of much interest to find new alternate approaches to tackle large-scale combinatorial optimization problems, going far beyond what is currently accessible with quantum and nature-inspired approaches alike.

In the deep learning community, graph neural networks (GNNs) have seen a burst in popularity over the last few years [23–30]. In essence, GNNs are deep neural network architectures specifically designed for graph structure data, with the ability to learn effective feature representations of nodes, edges, or even entire graphs. Prime examples of GNN applications include classification of users in social networks [31, 32], the prediction of future interactions in recommender systems [33], and the prediction of certain properties of molecular graphs [34, 35]. As a convenient and general framework to model a variety of real-world complex structural data, GNNs have successfully been applied to a broad set of problems, including recommender systems in

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social media and e-commerce [36, 37], the detection of misinformation (fake news) in social media [38], and various domains of natural sciences including event classification in particle physics [39, 40], to name a few. While several specific implementations of GNNs exist [29, 41, 42], at their core typically GNNs iteratively update the features of the nodes of a graph by aggregating the information from their neighbors (often referred to as message passing [43]) thereby iteratively making local updates to the graph structure as the training of the network progresses. Because of their scalability and inherent graph-based design, GNNs present an alternate platform to build large-scale combinatorial heuristics.

In this work we present a highly-scalable GNNbased solver to (approximately) solve combinatorial optimization problems with up to millions of variables. The approach is schematically depicted in Fig. 1, and works as follows: First, we identify the Hamiltonian (cost function) H that encodes the optimization problem in terms of binary decision variables  $x_{\nu} \in \{0,1\}$  and we associate this variable with a vertex  $\nu \in \mathcal{V}$  for an undirected graph  $\mathcal{G} = (\mathcal{V}, \mathcal{E})$  with vertex set  $\mathcal{V} =$  $\{1, 2, \ldots, n\}$  and the edge set  $\mathcal{E} = \{(i, j) : i, j \in \mathcal{V}\}$ capturing interactions between the decision variables. We then apply a relaxation strategy to the problem Hamiltonian to generate a differentiable loss function with which we perform unsupervised training on the node representations of the GNN. The GNN follows a standard recursive neighborhood aggregation scheme [43, 44], where each node  $\nu = 1, 2, \dots, n$  collects information (encoded as feature vectors) of its neighbors to compute its new feature vector  $\mathbf{h}_{\nu}^{k}$  at layer k=1 $0, 1, \ldots, K$ . After k iterations of aggregation, a node is represented by its transformed feature vector  $\mathbf{h}_{\nu}^{k}$ , which captures the structural information within the node's khop neighborhood [28]. For binary classification tasks we typically use convolutional aggregation steps, followed by the application of a nonlinear softmax activation

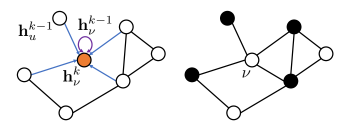


FIG. 1. Schematic illustration of the graph neural network approach for combinatorial optimization presented in this work. Following a recursive neighborhood aggregation scheme, the graph neural network is iteratively trained against a custom loss function that encodes the specific optimization problem (e.g., maximum cut). At training completion, we project the final values for the soft node assignments at the final graph neural network layer back to binary variables  $x_i = 0, 1$ , providing the solution bit string  $\mathbf{x} = (x_1, x_2, \ldots)$ . Further details are given in the text.

function to shrink down the final embeddings  $\mathbf{h}_{\nu}^{K}$  to one-dimensional soft (probabilistic) node assignments  $p_{\nu} = \mathbf{h}_{\nu}^{K} \in [0, 1].$  Finally, once the unsupervised training process has completed, we apply a projection heuristic to map these soft assignments  $p_{\nu}$  back to integer variables  $x_{\nu} \in \{0,1\}$  using, for example,  $x_{\nu} =$  $int(p_{\nu})$ . We numerically showcase our approach with results for canonical NP-hard optimization problems such as maximum cut (MaxCut) and maximum independent set (MIS), showing that our GNN-based approach can perform on par or even better than existing wellestablished solvers, while being broadly applicable to a large class of optimization problems. Further, the scalability of our approach opens up the possibility of studying unprecedented problem sizes with hundreds of millions of nodes when leveraging distributed training in a mini-batch fashion on a cluster of machines as demonstrated recently in Ref. [45].

The paper is structured as follows. In Sec. II we provide some context for our work, discussing recent developments at the cross-section between machine learning and combinatorial optimization. Section III summarizes the basic concepts underlying our approach, as well as information on the class of problems that this approach can solve. Section IV outlines the implementation of the proposed GNN-based optimizer, followed by numerical experiments in Sec. V. In Sec. VI we discuss potential real-world applications in industry. In Sec. VII we draw conclusions and give an outlook on future directions of research.

### II. RELATED WORK

In this Section we briefly review relevant existing literature, with the goal to provide a detailed context for our work. Broadly speaking, our work makes a physics-inspired contribution to the emerging crossfertilization between combinatorial optimization and machine learning, where the development of novel deep learning architectures has sparked a renewed interest in heuristics for solving NP-hard combinatorial optimization problems using neural networks, as extensively reviewed in e.g., Refs. [46, 47]. Leaving alternative, non-graph-based approaches as presented for example in Ref. [48] aside, in the following short survey we focus on graph-based optimization problems—where modern deep learning architectures such as sequence models, attention mechanisms, and GNNs provide a natural tool set [46]—and we primarily distinguish between approaches based on supervised learning, reinforcement learning, or unsupervised learning. This categorization can be refined further with respect to the typical size of a problem solved by a specific approach and the scope of the solver (special-purpose vs generalpurpose).

**Supervised Learning.** The majority of neural-network-based approaches to combinatorial optimization

are based on supervised learning, with the goal to approximate some (typically complex, non-linear) mapping from an input representation of the problem to the target solution, based on the minimization of some empirical, handcrafted loss function. Early work was based on pointer networks which leverage sequence-to-sequence models to produce permutations over inputs of variable size, as, for example, relevant for the canonical traveling salesman problem (TSP) [49]. Since then, numerous studies have fused GNNs with various heuristics and search procedures to solve specific combinatorial optimization problems, such as quadratic assignment [50], graph matching [51], graph coloring [52], and the TSP [53, 54]. As pointed out in Ref. [55], however, the viability and performance of supervised approaches critically depends on the existence of large, labelled training data sets with previously optimized hard problem instances, resulting in a problematic chicken-and-egg scenario, that is further amplified by the fact that it is hard to efficiently sample unbiased and representative labeled instances of NP-hard problems [56].

Reinforcement Learning. The critical need for training labels can be circumvented with Reinforcement Learning (RL) techniques that aim to learn a policy with the goal of maximizing some expected reward function. Specifically, optimization problems can typically be described with a native objective function that can then serve as a reward function in an RL approach [46]. Motivated by the challenges associated with the need for optimal target solutions, Bello et al. extended the pointer network architecture [49] to an actor-critic RL framework to train an approximate TSP solver, using a recurrent neural network encoder scheme and the expected tour length as a reward signal [57]. Using a general RL framework based on a graph attention network architecture [42], significant improvements in accuracy on two-dimensional euclidean TSP have subsequently been presented in Ref. [58], getting close to optimal results for problems up to 100 nodes. Moreover, TSP variants with hard constraints have been analyzed in Ref. [59], with the help of a multi-level RL framework in which each layer of a hierarchy learns a different policy, and from which actions can then be sampled. Finally, while the majority of the RL-based approaches have focused on the TSP or variants thereof, Dai et al. proposed a combination of RL and graph embedding to learn efficient greedy meta-heuristics to incrementally construct a solution, and showcased their approach with numerical results for Minimum Vertex Cover, MaxCut, and TSP as test problems, for graphs with up to  $\sim 1000 - 1200$  nodes

Unsupervised Learning. Conceptually, our work is most similar to those that aim to train neural networks in an unsupervised, end-to-end fashion, without the need for labelled training sets [55]. Specifically, Toenshoff *et al.* have recently used a recurrent GNN architecture—dubbed RUN-CSP—to solve optimization problems that

can be framed as maximum constraint satisfaction problems [61]. For other types of problems, such as the maximum independent set problem, the model relies on empirically-selected hand-crafted loss functions. Using the language of constraint satisfaction problems, where the system size is expressed in terms of both the number of variables and the number of constraints. the authors solve problem instances of Maximum 2satisfiability, 3-colorability, MaxCut and Maximum Independent Set with up to 5000 nodes, showing that RUN-CSP can compete with traditional approaches like greedy heuristics or semi-definite programming. Finally, by either optimizing a smooth relaxation of the cut objective or applying a policy gradient, Yao et al. trained a GNN to specifically solve the MaxCut problem, albeit at relatively small system sizes with up to 500 nodes [62], and without any details on runtime.

Here, we present a highly-scalable, physics-inspired framework that uses deep-learning tools in the form of GNNs to approximate solutions to hard combinatorial optimization problems with up to millions of variables. Our GNN optimizer is based on a direct mathematical relation between prototypical Ising spin Hamiltonians [63], the Quadratic Binary Unconstrained Optimization (QUBO) and Polynomial Binary Unconstrained Optimization (PUBO) formalism and the differentiable loss function with which we train the GNN, thereby providing one unifying framework for a broad class of combinatorial optimization problems, and opening up the powerful toolbox of statistical physics to modern deep-learning approaches. Fusing concepts from statistical physics with modern machine learning tooling, we propose a simple, generic, and robust solver that does not rely on hand-crafted loss functions. Specifically, we show that the same GNN optimizer can solve different QUBO problems, without any need to change the architecture or loss function, while scaling to problem instances orders of magnitude larger than what many traditional QUBO solvers can handle [6, 12, 64, 65].

## III. PRELIMINARIES

To set up our notation and terminology we start out with a brief review of both combinatorial optimization, and graph neural networks.

Combinatorial Optimization. The field of combinatorial optimization is concerned with settings where a large number of yes/no decisions must be made and each set of decisions yields a corresponding objective function value, like a cost or profit value, that is to be optimized [1]. Canonical combinatorial optimization problems include, among others, the maximum cut problem (MaxCut), the maximum independent set problem (MIS), the minimum vertex cover problem, the maximum clique problem and the set cover problem. In all cases exact solutions are not feasible for sufficiently-large systems due to the exponential growth

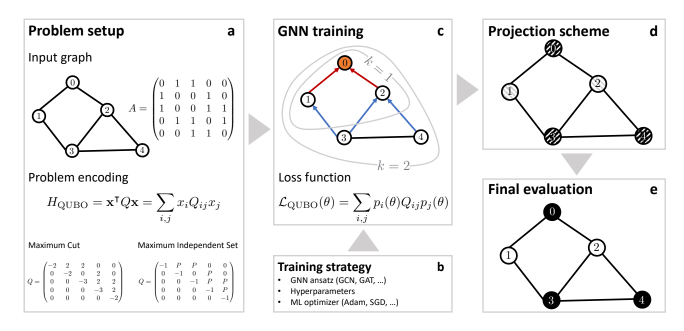


FIG. 2. Flow chart illustrating the end-to-end workflow for the proposed physics-inspired GNN optimizer. (a), The problem is specified by a graph  $\mathcal{G}$  with associated adjacency matrix A, and a cost function as described (for example) by the QUBO Hamiltonian  $H_{\text{QUBO}}$ . Within the QUBO framework the cost function is fully captured by the QUBO matrix Q, as illustrated for both MaxCut and MIS for a sample (undirected) graph with five vertices and six edges. (b), The problem setup is complemented by a training strategy that specifies the GNN Ansatz, a choice of hyperparameters and a specific ML optimizer. (c), The GNN is iteratively trained against a custom loss function  $\mathcal{L}_{\text{QUBO}}(\theta)$  that encodes a relaxed version of the underlying optimization problem as specified by the cost function  $H_{\text{QUBO}}$ . Typically, a GNN layer operates by aggregating information within the local one-hop neighbourhood (as illustrated by the k=1 circle for the top node with label 0). By stacking layers one can extend the receptive field of each node, thereby allowing distant propagation of information (as illustrated by the k=2 circle for the top node with label 0). (d)-(e), The GNN generates soft node assignments which can be viewed as class probabilities. Using some projection scheme, we then project the soft node assignments back to (hard) binary variables  $x_i = 0, 1$  (as indicated by the binary black/white node coloring), providing the final solution bit string  $\mathbf{x}$ .

of the solution space as the number of variables n increases. Bespoke (approximate) algorithms to solve these problems can typically be identified, at the cost of limited scope and generalizability. Conversely, in recent years the QUBO framework has resulted in a powerful approach that unifies a rich variety of these NP-hard combinatorial optimization problems [1–3, 66]. The cost function for a QUBO problem can be expressed in compact form with the following Hamiltonian

$$H_{\text{QUBO}} = \mathbf{x}^{\mathsf{T}} Q \mathbf{x} = \sum_{i,j} x_i Q_{ij} x_j, \tag{1}$$

where  $\mathbf{x} = (x_1, x_2, \dots)$  is a vector of binary decision variables and the QUBO matrix Q is a square matrix of constant numbers that encodes the actual problem to solve. Without loss of generality, the Q-matrix can be assumed to be symmetric or in upper triangular form [1]. We have omitted any irrelevant constant terms, as well as any linear terms as these can always be absorbed into the Q-matrix because  $x_i^2 = x_i$  for binary variables  $x_i \in \{0,1\}$ . Problem constraints, as relevant for many real-world optimization problems, can be accounted for with the help of penalty terms entering the objective function (rather than being explicitly

imposed), as detailed in Ref. [1]. The significance of QUBO problems is further illustrated by the close relation to the famous Ising model, which is known to provide mathematical formulations for many NP-complete and NP-hard problems, including all of Karp's 21 NP-complete problems [66]. As opposed to QUBO problems, Ising problems are described in terms of binary spin variables  $z_i \in \{-1,1\}$ , that can be mapped straightforwardly to their equivalent QUBO form, and vice versa, using  $z_i = 2x_i - 1$ . By definition, both the QUBO and the Ising models are quadratic, but can be naturally generalized to higher order PUBO problems, as described by the N-local Hamiltonian

$$H_{\text{PUBO}} = \sum_{k=0}^{N} \sum_{\langle i_1, i_2, \dots, i_k \rangle} Q_{i_1 i_2 \cdots i_k} x_{i_1} x_{i_2} \cdots x_{i_k},$$
 (2)

with real-numbered coefficients  $Q_{i_1i_2\cdots i_k}$ , for some  $N\geq 3$ , and  $\langle i_1,i_2,\ldots,i_k\rangle$  indicating a group of k binary variables (or spins in the Ising formulation). Terms containing a product of k variables, of the form  $Q_{i_1i_2\cdots i_k}x_{i_1}x_{i_2}\cdots x_{i_k}$ , are commonly referred to as k-local interactions with  $Q_{i_1i_2\cdots i_k}$  being the coupling constant. As we exemplify below for some canonical problems, graph (hypergraph)

problems can be naturally framed as QUBO (PUBO) problems. To this end, given an undirected graph  $\mathcal{G} = (\mathcal{V}, \mathcal{E})$ , we simply associate a binary variable  $x_i$  with every vertex  $i \in V$ , and then express the (node classification) objective as a QUBO problem, where the specific assignment  $\mathbf{x}$  can be visualized as a specific twotone (e.g., light and dark) coloring of the graph [67]; see Fig. 1.

Graph Neural Networks. On a high level, GNNs are a family of neural networks capable of learning how to aggregate information in graphs for the purpose of representation learning. Typically, a GNN layer is comprised of three functions [35]: (i) a message passing function that permits information exchange between nodes over edges, (ii) an aggregation function that combines the collection of received messages into a single, fixed-length representation, and (iii) a (typically nonlinear) update activation function that produces node-level representations given the previous layer representation and the aggregated information. While a single-layer GNN encapsulates a node's features based on its immediate or one-hop neighborhood, by stacking multiple layers, the model can propagate each node's features through intermediate nodes, analogous to the broadening the receptive field in downstream layers of convolutional neural networks. Formally, at layer k=0, each node  $\nu \in \mathcal{V}$  is represented by some initial representation  $\mathbf{h}_{\nu}^{0} \in \mathbb{R}^{d_{0}}$ , usually derived from the node's label or given input features of dimensionality  $d_0$  [68]. Following a recursive neighborhood aggregation scheme, the GNN then iteratively updates each node's representation, in general described by some parametric function  $f_{\theta}^{k}$ , resulting in

$$\mathbf{h}_{\nu}^{k} = f_{\theta}^{k} \left( \mathbf{h}_{\nu}^{k-1}, \{ \mathbf{h}_{u}^{k-1} | u \in \mathcal{N}_{\nu} \} \right), \tag{3}$$

for the layers  $k=1,\ldots,K$ , with  $\mathcal{N}_{\nu}=\{u\in\mathcal{V}|(u,\nu)\in\mathcal{E}\}$  referring to the local neighborhood of node  $\nu$ , i.e., the set of nodes that share edges with node  $\nu$ . The total number of layers K is usually determined empirically as a hyperparameter, as are the intermediate representation dimensionality  $d_k$ . Both can be optimized in an outer loop. While a growing number of possible implementations for GNN architectures [30] exists, here we use a graph convolutional network (GCN) [29] for which Eq. (3) reads explicitly as

$$\mathbf{h}_{\nu}^{k} = \sigma \left( \mathbf{W}_{k} \sum_{u \in \mathcal{N}(\nu)} \frac{\mathbf{h}_{u}^{k-1}}{|\mathcal{N}(\nu)|} + \mathbf{B}_{k} \mathbf{h}_{\nu}^{k-1} \right), \tag{4}$$

with  $\mathbf{W}_k$  and  $\mathbf{B}_k$  being (shared) trainable weight matrices, the denominator  $|\mathcal{N}(\nu)|$  serving as normalization factor (with other choices available as well) and  $\sigma(\cdot)$  being some (component-wise) nonlinear activation function such as sigmoid or ReLU. While GNNs can be used for various prediction tasks (including node classification, link prediction, community detection, network similarity, or graph classification), here we focus

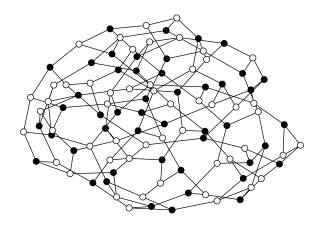


FIG. 3. Example solution to MaxCut for a random 3-regular graph with n=100 nodes. After training completion, the GNN provides a binary bit string  ${\bf x}$  that assigns one of two possible colors (e.g., black or white) to each vertex. An edge is said to be cut when it connects two vertices of different colors. For a given graph, the optimization problem is to assign the colors in a way that as many edges as possible can be cut at the same time (corresponding to the antiferromagnetic ground-state of the system).

on node classification, where usually the last (K-th) layer's output is used to predict a label  $y_{\nu}$  for every node  $\nu \in \mathcal{V}$ . To this end, we feed the (parametrized) final node embeddings  $\mathbf{z}_{\nu} = \mathbf{h}_{\nu}^{K}(\theta)$  into a problem-specific loss function and run stochastic gradient descent to train the weight parameters.

## IV. COMBINATORIAL OPTIMIZATION WITH GRAPH NEURAL NETWORKS

We now detail how to use GNNs to solve combinatorial optimization problems, as schematically outlined in Fig. 2. To this end, we frame combinatorial optimization problems as unsupervised node classification tasks, without the need for any labelled data. Because the nodes do not carry any inherent features, in our setup the node embeddings  $\mathbf{h}_{\nu}^{0}$  are initialized randomly. Warmstarting the training process with pre-training (transfer learning) will be left for future research. The class of Hamiltonians described above are not differentiable and cannot be used straightforwardly within the GNN training process. Therefore, for a given problem Hamiltonian H and graph  $\mathcal{G}$ , we generate a differentiable loss function  $\mathcal{L}(\theta)$ , as required for standard backpropagation, by promoting the binary decision variables  $x_i \in \{0,1\}$  to continuous (parametrized) probability parameters  $p_i(\theta)$  with the following (heuristic) relaxation approach

$$x_i \longrightarrow p_i(\theta) \in [0, 1].$$
 (5)

The soft assignments  $p_i$  can be viewed as class probabilities. They are generated by our GNN Ansatz as final node embeddings  $p_i = \mathbf{h}_i^K \in [0, 1]$  at layer K, after the application of a non-linear softmax activation function. Then, they are used as input for the loss function  $\mathcal{L}(\theta)$ . In particular, for QUBO-type problems:

$$H_{\text{QUBO}} \longrightarrow \mathcal{L}_{\text{QUBO}}(\theta) = \sum_{i,j} p_i(\theta) Q_{ij} p_j(\theta),$$
 (6)

which is differentiable with respect to the parameters of the GNN model  $\theta$ , and similarly for PUBO problems on hyper-graphs with higher-order terms of the form  $p_i p_j p_k$  etc., thereby establishing a straightforward, general connection between combinatorial optimization problems, Ising Hamiltonians and GNNs. For training with gradient descent, standard ML optimizers such as ADAM can be used. Once the (unsupervised) training process has completed, we apply projection heuristics to map these soft assignments  $p_i$  back to integer variables  $x_i = 0, 1$ , using for example simply  $x_i = \text{int}(p_i)$ . The application of other, more sophisticated projection schemes will be left for future research. Note that any projection heuristics can be applied throughout training after every epoch, thereby increasing the pool of solution candidates, at no additional computational cost. With the GNN guiding the search through the solution space, one can then book keep all solution candidates identified throughout training and simply pick the best solution found.

Our general GNN approach features several hyperparameters, including the number of layers K, the dimensionality of the embedding vectors  $\mathbf{h}_i^k$ , and the learning rate  $\beta$ , with details depending on the specific architecture and optimizer used. These can be fine-tuned and optimized in an outer loop, using, e.g., standard techniques such as grid search or more advanced Bayesian optimization methods.

Our GNN-based approach can be readily implemented with open-source libraries such as PyTorch Geometric [69] or the Deep Graph Library [70]. The core of the corresponding code is displayed in the supplemental material for a GCN with two layers and a loss function for any QUBO problem. For illustration, an example solution to the archetypal MaxCut problem (as implemented with this Ansatz) for a 3-regular graph with n=100 vertices is shown in Fig. 3. Here, the cut size achieved with our GNN method amounts to 132. Further details are provided below.

### V. NUMERICAL EXPERIMENTS

We perform numerical experiments using MaxCut and MIS benchmark problems. Before providing details on these numerical experiments, we first describe our GNN model architecture as it is consistent across the *d*-regular MaxCut and MIS problem instances described below. It is certainly possible that better solutions can be found by

fine-tuning the hyper-parameters for every given problem instance. However, one of our goals is to design a robust and scalable solver that is able to solve a large sample of instances efficiently without the need of hand-tuning the parameters on an instance-by-instance base.

**GNN Architecture.** We use a simple two-layer GCN architecture based on PyTorch GraphConv units. The first convolutional layer is fed the node embeddings of dimension  $d_0$  and outputs a representation of size  $d_1$ . Next, we apply a component-wise, non-linear ReLU transformation. The second convolutional layer is then fed this intermediate representation and outputs the output layer of size  $d_2$ , which is then fed through the component-wise sigmoid transformation to provide a soft probability  $p_i \in [0,1]$  for every node  $i \in \mathcal{V}$ . We find that the following simple heuristic for determining the hyper-parameters  $d_0$  and  $d_1$  works well: if the number of nodes is large  $(n \ge 10^5)$ , then we set  $d_0 = \operatorname{int}(\sqrt{n})$ , else we set  $d_0 = \operatorname{int}(\sqrt[3]{n})$ , and we take  $d_1 = \operatorname{int}(d_0/2)$ . Because we solve for binary classification tasks, we set the final output dimension as  $d_2 = 1$ . However, for multi-color problems this could be extended to C > 2classes by passing the output layer through a softmax transformation (instead of a sigmoid) and taking the argmax. Note that as the graph size scales beyond  $\sim 10^5$  nodes, memory becomes a concern, and so we further reduce the representations to allow the GNN to be trained on a single GPU. Distributed training leveraging a whole cluster of machines will be discussed below in Sec. VII. With the GNN's output depending on the random initialization of the hidden feature vectors there is a risk of becoming stuck in a local optimum where the GNN stops learning. To counter this issue, one can take multiple shots (i.e., run the GNN training multiple times for different random seeds and choose the best solution), thereby boosting the performance at the cost of extended runtime. In our numerical experiments we limited the number of shots per instance to five, only re-running the training when an obviously sub-optimal solution was detected. Finally, we set the learning rate to  $\beta = 10^{-4}$ and allow the model to train for up to  $\sim 10^5$  epochs, with a simple early stopping rule set to an absolute tolerance of  $10^{-4}$  and a patience of  $10^3$ .

Maximum Cut. MaxCut is an NP-hard combinatorial optimization problem with practical applications in machine scheduling [71], recognition [72], and electronic circuit layout design [73]. In the current era of noisy intermediate-scale quantum devices, with the advent of novel hybrid quantum-classical algorithms such as the Quantum Approximate Optimization Algorithm (QAOA) [74], the MaxCut problem has recently attracted considerable attention as a potential use case of pre-error-corrected quantum devices, see Refs. [75–80]. MaxCut is a graph partitioning problem defined as follows: given a graph with vertex set  $\mathcal{V}$  and edge set  $\mathcal{E}$ , we seek a partition of V into two subsets with maximum cut, where a cut refers to edges connecting two nodes from different vertex sets. Intuitively, that means we score a point whenever an edge connects two nodes of different colors. To formulate MaxCut mathematically, we introduce binary variables satisfying  $x_i = 1$  if vertex i is in one set and  $x_i = 0$  if it is in the other set. It is then easy to verify that the quantity  $x_i + x_j - 2x_ix_j = 1$  if the edge (i,j) has been cut, and 0 otherwise. With the help of the adjacency matrix  $A_{ij}$  with  $A_{ij} = 0$  if edge (i,j) does not exist and  $A_{ij} > 0$  if a (possibly weighted) edge connects node i with j, the MaxCut problem is described by the following quadratic Hamiltonian

$$H_{\text{MaxCut}} = \sum_{i < j} A_{ij} (2x_i x_j - x_i - x_j) \tag{7}$$

that falls into the broader class of QUBO problems described by Eq. (1); we provide the explicit Q-matrix for a sample MaxCut problem in Fig. 2. Up to an irrelevant constant, the MaxCut problem can equivalently by described by the compact Ising Hamiltonian  $H_{\text{MaxCut}} = \sum_{i < j} J_{ij} z_i z_j$  with  $J_{ij} = A_{ij}/2$ , favoring antiferromagnetic ordering of the spins for  $J_{ij} > 0$ , as expected intuitively based on the problem definition. As our figure of merit, we denote the largest cut found as cut<sup>\*</sup> =  $-H_{\text{MaxCut}}(\mathbf{x}^*)$ , with  $\mathbf{x}^*$  referring to the corresponding bit string.

The complexity of MaxCut depends on the regularity and connectivity of the underlying graph. Following an existing trend in the community [76], we first consider the MaxCut problem on random (unweighted) d-regular graphs, where every vertex is connected to exactly dother vertices. We perform the benchmarks as follows. For graphs with up to a few hundred nodes, we compare our GNN-based solver to the (approximate) polynomialtime Goemans-Williamson (GW) algorithm [81], which provides the current record for an approximate answer within some fixed multiplicative factor of the optimum (referred to as approximation ratio  $\alpha$ ), using semidefinite programming and randomized rounding. Specifically. the GW algorithm achieves a guaranteed approximation ratio of  $\alpha \sim 0.878$  for generic graphs. This lower bound can be raised for specific graphs such as unweighted 3-regular graphs where  $\alpha \sim 0.9326$  [82]. implementation of the GW algorithm is based on the open-source CVXOPT solver, with CVXPY as modeling interface. For very large graphs with up to a million nodes, numerical benchmarks are not available, but we can compare our best solution cut\* to an analytical result derived in Ref. [83], where it was shown that with high probability (in the limit  $n \to \infty$ ) the size of the maximum cut for random d-regular graphs with n nodes is given by cut\* =  $(d/4 + P_*\sqrt{d/4} + \mathcal{O}(\sqrt{d}))n + \mathcal{O}(n)$ . Here,  $P_* \approx 0.7632$  refers to an universal constant related to the ground-state energy of the Sherrington-Kirkpatrick model [84, 85] that can be expressed analytically via Parisi's formula [83]. We thus take  $cut_{ub} = (d/4 +$  $P_*\sqrt{d/4}$ )n as an upper-bound estimate for the maximum cut size in the large-n limit. We complement this upper bound with a lower bound as achieved by a

simple, randomized 0.5-approximation algorithm that (on average) cuts half of the edges, yielding a cut size of  $\operatorname{cut}_{\operatorname{rnd}} \approx (d/4)n$  for a d-regular graph with  $|\mathcal{E}| = (d/2)n$ . Our results for the achieved cut size as a function of the number of vertices n are shown in Fig. 4. All results are bootstrapped estimates of the mean, with error bars denoting twice the bootstrapped standard deviations, sampled across 20 random d-regular graphs for every data point. For graphs with up to a few hundred nodes. we find that a simple two-layer GCN architecture can perform on par with the GW algorithm, while showing a runtime advantage compared to GW starting at around  $n \approx 100$  nodes. For large graphs with  $n \approx 10^4$  to  $10^6$ nodes, we find that our approach consistently achieves high-quality solutions with  $cut^* \gtrsim 0.9 \cdot cut_{ub}$  for both d=3 and d=5, respectively (i.e., much better than any naive randomized algorithm). As expected for dregular graphs, we find cut\* to scale linearly with the number of nodes n, i.e., cut\*  $\approx \gamma_d n$ , with  $\gamma_3 \approx 1.28$  and  $\gamma_5 \approx 1.93$  for d=3 and d=5, respectively. Moreover, utilizing modern GPU hardware, we observe a favorable runtime scaling at intermediate and large system sizes that allows us to solve instances with  $n = 10^6$  nodes in approximately 10 minutes (which includes both GNN model training and post-processing steps). Specifically, as shown in Fig. 4, we observe an approximately linear scaling of total runtime with  $\sim n$ , for large d-regular graphs with  $10^5 \le n \le 10^6$ ; contrasted with the observed GW algorithm scaling as  $\sim n^{3.5}$  for problem sizes in the range  $n \lesssim 250$ , thereby showing the (expected) time complexity  $\tilde{O}(n^{3.5})$  of the interior-point method (as commonly used for solving the semidefinite program underlying the GW algorithm) that dominates the GW algorithm runtime [86, 87].

To complement our work on random d-regular graphs, we have performed additional experiments on standard Max-Cut benchmark instances, with published results, based on the publicly available Gset data set [91] commonly used for testing Max-Cut algorithms. provide benchmark results for seven different graphs, with thousands of nodes, including (i) two Erdös-Renyi graphs with uniform edge probability, (ii) two graphs where the connectivity gradually decays from node 1 to n, (iii) two 4-regular toroidal graphs, and (iv) one of the largest Gset instances with  $n = 10^4$ . results are displayed in Tab. I. Here, we report cut sizes achieved with our physics-inspired GNN solver (PI-GNN), together with results sourced from Refs. [61, 88– 90]; the latter include an SDP solver using dual scaling (DSDP) [90], a combination of local search and adaptive perturbation referred to as Breakout Local Search (BLS) [89] (providing the best known solutions for the Gset data set), a Tabu Search metaheuristic (KHLWG) [88], and a recurrent GNN architecture for maximum constraint satisfaction problems (RUN-CSP) [61]. We assess the solution quality achieved with PI-GNN with the relative error  $\epsilon = (\operatorname{cut}_{\operatorname{best}} - \operatorname{cut}^*)/|\mathcal{E}|$  quantifying the gap to the best known solution, normalized by the number of

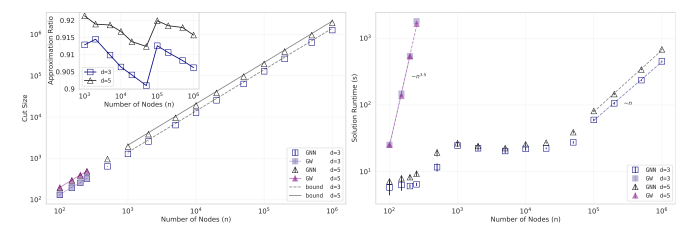


FIG. 4. Numerical results for MaxCut. **Left panel:** Average cut size for d-regular graphs with d=3 and d=5 as a function of the number of vertices n, bootstrap-averaged over 20 random graph instances, for both the GNN-based method and the Goemans-Williamson (GW) algorithm. On each graph instance, the GNN solver is allowed up to five shots, and the GW algorithm takes 100 shots. Solid lines for  $n \ge 10^3$  represent theoretical upper bounds, as described in the main text. Inset: The estimated relative approximation ratio defined as cut\*/cut<sub>ub</sub> shows that our approach consistently achieves high-quality solutions. **Right panel:** Algorithm runtime in seconds for both the GNN solver and the GW algorithm. Error bars refer to twice the bootstrapped standard deviations, sampled across 20 random graph instances for every data point.

graph	nodes	edges	BLS	DSDP	KHLWG	RUN-CSP	PI-GNN	relative error $\epsilon$
G14	800	4694	3064	2922	3061	2943	3026	0.81%
G15	800	4661	3050	2938	3050	2928	2990	1.29%
G22	2000	19990	13359	12960	13359	13028	13181	0.89%
G49	3000	6000	6000	6000	6000	6000	5918	1.37%
G50	3000	6000	5880	$\boldsymbol{5880}$	5880	5880	5820	1.00%
G55	5000	12468	10294	9960	10236	10116	10138	1.25%
G70	10000	9999	9541	9456	9458	_	9421	1.20%

TABLE I. Numerical results for MaxCut on Gset instances. We report cut sizes achieved with our physics-inspired GNN solver (PI-GNN), together with results sourced from Refs. [61, 88–90]. Best known results are marked in bold. The last column specifies the relative error  $\epsilon$  comparing PI-GNN to the best known cut size. Further details are provided in the main text. GNN model configurations are detailed in the supplemental material.

edges  $|\mathcal{E}|$ , thereby giving the fraction of uncut edges as compared to the best known solution. We find that our general-purpose approach is competitive with other solvers and typically within  $\sim 1\%$  of the best published results.

Maximum Independent Set. The MIS problem is a prominent combinatorial optimization problem with practical applications in network design [93] and finance [94], and is closely related to the maximum clique, minimum vertex cover, and set packing problems. In the quantum community, the MIS problem has recently attracted significant interest [95] as a potential target use case for novel experimental platforms based on neutral atom arrays [96]. The MIS problem reads Given an undirected graph  $\mathcal{G} = (\mathcal{V}, \mathcal{E})$ , as follows. an independent set is a subset of vertices that are not connected with each other. The MIS problem is then the task to find the largest independent set, with its (maximum) cardinality typically denoted as the independence number  $\alpha$ . To formulate the MIS problem

mathematically, for a given graph  $\mathcal{G} = (\mathcal{V}, \mathcal{E})$ , one first associates a binary variable  $x_i \in \{0,1\}$  with every vertex  $i \in V$ , with  $x_i = 1$  if vertex i belongs to the independent set, and  $x_i = 0$  otherwise. The MIS can then be formulated in terms of a Hamiltonian that counts the number of marked (colored) vertices and adds a penalty to nonindependent configurations (when two vertices in this set are connected by an edge). It is given by

$$H_{\text{MIS}} = -\sum_{i \in \mathcal{V}} x_i + P \sum_{(i,j) \in \mathcal{E}} x_i x_j, \tag{8}$$

with a negative pre-factor to the first term (because we solve for the largest independent set within a minimization problem), and the penalty parameter P > 0 enforcing the constraints. Note that the numerical value for P is typically set as P = 2 [97], but can be further optimized in an outer loop. Energetically, the Hamiltonian  $H_{\rm MIS}$  favors each variable to be in the state  $x_i = 1$  unless a pair of these are connected by an

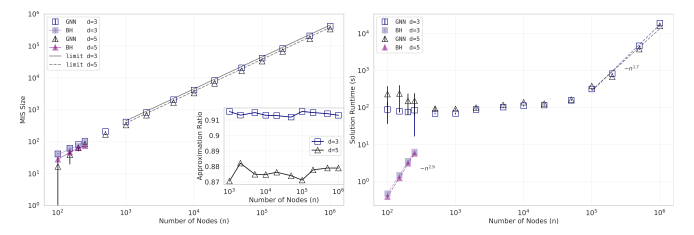


FIG. 5. Numerical results for the MIS problem. Left panel: Average independence number  $\alpha$  for d-regular graphs with d=3 and d=5 as a function of the number of vertices n, (bootstrap-)averaged over 20 random graph instances, for both the GNN-based method and a traditional MIS algorithm [92]. Solid lines for  $n \geq 10^3$  refer to theoretical upper bounds as described in the main text. Inset: The estimated relative approximation ratio comparing the achieved independence number  $\alpha$  against known theoretical upper bounds shows that our approach consistently achieves high-quality solutions. Right panel: Algorithm runtime in seconds for both the GNN solver and the Boppana-Halldorsson algorithm. Error bars refer to twice the bootstrapped standard deviations, sampled across 20 random graph instances for every data point.

edge. Again, the Hamiltonian  $H_{\text{MIS}}$  is quadratic and falls into the broader class of QUBO problems described by Eq. (1); again we provide the explicit Q-matrix for a sample MIS problem in Fig. 2.

The MIS problem is known to be strongly NPhard, making the existence of an efficient algorithm for finding the maximum independent set on generic graphs unlikely. In addition, the MIS problem is even hard to approximate. In general, the MIS problem cannot be approximated to a constant factor in polynomial time (unless P = NP). Again we study the MIS problem on random unweighted d-regular graphs. Because in our approach the independence constraint is enforced with soft penalty terms  $\sim P$ —just like in any QUBO-based model—the predicted set may violate the independence condition (i.e., the set may contain nodes connected by an edge). Setting P=2, we have observed these violations only in very few cases. If present, as part of our postprocessing, we have enforced the independence constraint by greedily removing one of the nodes of each induced edge from the set, and only reporting results after this correction. For small graphs with up to a few hundred nodes, we compare the GNN-based results to results obtained with the Boppana-Halldorsson algorithm built into the Python NetworkX library [92]. For very large graphs with up to a million nodes (where benchmarks are not available) we resort to analytical upper bounds for random d-regular graphs as presented in Ref. [98]. Here, the best known bounds on the ratio  $\alpha_d/n$  are reported as  $\alpha_3/n = 0.45537$  and  $\alpha_5/n = 0.38443$  for d=3 and d=5, respectively, as derived using refined versions of Markov's inequality [99]. Our results for the achieved independence number as a function of the number of vertices n are shown in Fig. 5. All results are bootstrapped estimates of the mean, with error bars

denoting twice the bootstrapped standard deviations, sampled across 20 random d-regular graphs for every data point. Our numerical results for MIS are similar to the observations we have made for MaxCut: for graphs with up to a few hundred nodes, we find that a simple two-layer GCN architecture can perform on par with (or better than) the traditional solver, with the GNN solver showing a favorable runtime scaling. For large graphs with  $n \approx 10^4$  to  $10^6$  nodes we find that our approach consistently achieves high-quality solutions with  $\alpha_3/n \approx$ 0.416 and  $\alpha_5/n \approx 0.338$  for d=3 and d=5, respectively, resulting in estimated numerical approximation ratios of  $0.416/0.45537 \sim 0.92$  and  $0.338/0.38443 \sim 0.88$ , respectively. Finally, as shown in Fig. 5, we observe a moderate, super-linear scaling of the total runtime as  $\sim$  $n^{1.7}$  for large d-regular graphs with  $n \gtrsim 10^5$ , as opposed to the Boppana-Halldorsson solver with a runtime scaling of  $\sim n^{2.9}$  in the range  $n \lesssim 500$ . Note that the GNN model training alone displays sub-linear runtime scaling as  $\sim n^{0.8}$ , in line with our MaxCut results, while the aggregate runtime (including post-processing to enforce the independence condition) scales as  $\sim n^{1.7}$  in the regime  $n \sim 10^5 - 10^6$ .

## VI. APPLICATIONS IN INDUSTRY

While our previous analysis has focused on canonical graph optimization problems such as maximum cut and maximum independent set, in this section we discuss real-world applications in industry for which our solver could provide solutions, in particular at potentially unprecedented problem scales. We focus on applications of the QUBO formalism, even though our methodology is not limited to this modeling framework. We first review

the existing literature, providing relevant references across a wide stack of problem domains. Thereafter, we explicitly show how to distill combinatorial QUBO problems for a few select real-world use cases, from risk diversification in finance to sensor placement problems in water distribution networks. Once in QUBO format, the problem can be plugged into our general-purpose physics-inspired GNN solver, as outlined above.

As extensively reviewed in Refs. [1], [2], and [66], the QUBO (or, equivalently, Ising) formalism provides a comprehensive modeling framework encompassing a vast array of optimization problems, including knapsack problems, task (resource) allocation problems, and capital budgeting problems, among others. Specifically, the applicability of the QUBO representation has been reported for problem settings involving circuit board layouts [100], capital budgeting in financial analysis [101], computer aided design (CAD) [102], electronic traffic management [103, 104], cellular radio channel allocation [105], molecular conformation [106], and the prediction of epileptic seizures [107], among others. As mentioned earlier, practical applications of the MaxCut problem can be found in machine scheduling [71], image recognition [72], and electronic circuit layout design [73]. Similarly, in what follows we discuss in detail three select use cases and how they can be cast in QUBO form and thus made amenable to our solver.

Risk diversification. Graphs offer a convenient framework to model portfolio management problems in finance. Specifically, here we outline a risk diversification strategy, but similar considerations apply for the implementation of hedging strategies [108]. We consider a (potentially very large) universe of n assets, for which we are given a vector  $\boldsymbol{\mu} \in \mathbb{R}^n$  describing expected future returns, and the covariance matrix  $\Sigma \in \mathbb{R}^{n \times n}$  capturing volatility through the correlations among assets. minimize the volatility of returns of our portfolio, our goal is to select a subset of uncorrelated assets with the largest possible diversified portfolio. To this end we consider a graph  $\mathcal{G}$  with n nodes, with every node representing one asset. Correlations can be described in graph form, either by directly taking the cross-correlation matrix as a weighted adjacency matrix, or by creating a binary adjacency matrix A through thresholding. We set  $A_{i,j} = 1$  if and only if the absolute value of the correlation between assets i and j is greater than some user-specific threshold parameter  $\lambda$ , and  $A_{i,j} = 0$  otherwise [108]. Accordingly, in our model pairs of assets are classified as correlated or uncorrelated, based on whether or not the corresponding correlation coefficient exceeds a minimum level. Overall, the risk diversification strategy outlined above can be cast as an optimization problem in QUBO form, with the Hamiltonian

$$H_{\text{wMIS}} = -\sum_{i \in \mathcal{V}} \mu_i x_i + P \sum_{(i,j) \in \mathcal{E}} x_i x_j, \tag{9}$$

in a straightforward extension of Eq. (8) from the standard MIS problem to the weighted MIS problem.

Accordingly, by minimizing the first term of  $H_{\rm wMIS}$  we pick large-return assets subject to the independent set constraint (as captured by the second term). This diversification model is reminiscent of the mean-variance Markowitz model [109], albeit in discretized form. Given our results for the standard MIS problem with small tweaks to the loss function, such a problem could be readily plugged into our solver, for example as part of a larger, two-stage portfolio management pipeline where first a subset of assets is selected from a larger universe of assets using our solver, and then capital is allocated within a smaller, sparsified basket of assets using off-the-shelf solvers.

Interval scheduling. Here we consider a scenario involving the scheduling of tasks with given start and end times, as relevant for example in the context of algorithm design in computer science [110]. Specifically, we face n resource requests, each represented by an interval specifying the time in which it needs to be processed by some machine. Typically some requests will overlap in time leading to request clashes that cannot be satisfied by the same machine (resource). Conversely, a subset of intervals is deemed compatible if no two intervals overlap on the machine. As commonly done in resource allocation problems and scheduling theory, this situation can conveniently be described with the help of an undirected interval graph  $\mathcal{G}$  in which we introduce a vertex for each request and edges between vertices whose requests overlap. With the goal to maximize the throughput (i.e., to execute as many tasks as possible on a single machine), the interval scheduling maximization problem is then to find the largest compatible set, that is, a set of non-overlapping intervals of maximum size. This use case is equivalent to finding the maximum independent set in the corresponding interval graph  $\mathcal{G}$  with n nodes. While inexpensive (special-purpose) algorithms exist for interval graphs [111], we can then solve the underlying MIS problem, as described by Eq. (8), on this interval graph, in the same way as any other QUBO problem, using our general-purpose GNNbased approach, following the methodology outlined above.

Sensor placement  $_{
m in}$ water distribution Optimal sensor placement is key to networks. the detection and isolation of fault events—such as water leaks—in water distribution networks (WDN) [112]. As detailed in Ref. [112], the problem of optimally placing pressure sensors on a WDN can be efficiently cast as QUBO problem. Specifically, a WDN can be readily mapped to a graph  $\mathcal{G} = (\mathcal{V}, \mathcal{E})$ , with the nodes  $i \in \mathcal{V}$  referring to tanks or junctions and edges  $(i, j) \in \mathcal{E}$ representing pipes, respectively. We then associate a binary variable  $x_i = 0,1$  with every node, and set  $x_i = 1$  if node i hosts a sensor, and  $x_i = 0$  otherwise. The problem of covering the WDN with the smallest possible number of pressure sensors then maps onto the minimum vertex cover problem [112], as described by

the Hamiltonian

$$H_{\text{MVC}} = \sum_{i \in \mathcal{V}} c_i x_i + P \sum_{(i,j) \in \mathcal{E}} (1 - x_i - x_j + x_i x_j).$$
 (10)

Here,  $c_i \geq 0$  denotes the cost of node i hosting a sensor, the first term describes the overall cost of any potential sensor placement strategy, while the second (penalty) term with P > 0 ensures the constraint  $x_i + x_j \geq 1$  for all edges  $(i,j) \in \mathcal{E}$  (i.e., at least one of the endpoints of each edge will be in the cover [1]). Potential tweaks to this model are detailed in Ref. [112], however, variations can all be represented as a QUBO problem. Along the lines of our previous analysis for the MaxCut or MIS problem, the Hamiltonian  $H_{\text{MVC}}$  (or some variation thereof) can be straightforwardly mapped to a relaxed loss function with which we can train our solver and then solve the corresponding sensor placement use case.

## VII. CONCLUSION AND OUTLOOK

In summary, we have proposed and analyzed a versatile and scalable general-purpose solver that is powered by graph neural networks and draws from concepts Our approach is applicable in statistical physics. to any k-local Ising model, including canonical NPhard combinatorial optimization problems such as the maximum cut, maximum clique, minimum vertex cover or maximum independent set problems, among others Starting from a problem formulation in Ising form, we apply a relaxation strategy to the problem Hamiltonian by dropping integrality constraints on the decision variables in order to generate a differentiable loss function with which we perform unsupervised training on the node representations of the GNN. The GNN is then trained to generate soft assignments to predict the likelihood of belonging in one of two classes, for each vertex in the graph. To find a binary (twocolor) labelling consistent with the original problem formulation, simple projection heuristics are applied. Overall, we find that this approach can compete with existing special-purpose solvers, such as the Goemans-Williamson algorithm designed to solve the maximum cut problem, with the potential to tap into the rich toolbox of statistical physics, including, for example, the study of phase transitions. In the current noisy intermediate scale quantum era, our approach could be used as a broadly applicable, scalable benchmark for emerging quantum technologies, including special-purpose quantum [6] and quantum-inspired annealers [20], while not being resource constrained nor being limited to problem instances in QUBO form, as is also the case for coherent Ising machines [113].

Finally, we highlight possible extensions of research going beyond our present work. First, to better

understand the limitations of GNNs in the context of combinatorial optimization, further studies are in order, systematically benchmarking GNNs against state-of-theart solvers for a large class of optimization problems while leveraging the entire zoo of GNN implementations including, for example, GraphSAGE [27] or Graph Attention Networks (GATs) [42] to potentially boost the GNN Ansatz with an attention mechanism enabling vertices to weigh neighbor representations during the aggregation steps. Second, the presented GNN approach should be able to accommodate problems sizes with hundreds of millions of nodes when leveraging distributed training in a mini-batch fashion on a cluster of machines [45], thereby challenging the capabilities of several existing solvers. While we have solved individual problem instances from scratch, using a random initialization process for the initial node embeddings, in the future warm-starting the training process with pre-trained weights (transfer learning) could boost the time to solution. Moreover, one could potentially boost the performance of our optimizer by implementing randomized projection schemes (as opposed to the simple deterministic approach used here), or augment these strategies with simple greedy post-processing routines that check for local optimality with a sequence of local bit flips. Finally, as discussed in the main text, our approach can be generalized to PUBO problems on hyper-graphs where so-called hyper-edges may contain more than just two nodes, with no need for (typically) resource-intensive degree reduction schemes, as opposed to resourceconstrained QUBO solvers. Potential applications cover many real-world optimization problems involving multibody interactions, as found in scheduling problems [114] or chemistry [115, 116]. In conclusion, the proposed cross-fertilization between machine learning, operations research and physics opens up a number of interesting research directions, with the ultimate goal to further advance our ability to solve hard combinatorial optimization problems.

## Data availability

The data necessary to reproduce our numerical benchmark results are publicly available at https://web.stanford.edu/~yyye/yyye/Gset/. Random d-regular graphs have been generated using the open-source networkx library (https://networkx.org).

## Code availability

Code availability statement: An end-to-end open source demo version of the code implementing our approach has been made publicly available at https://github.com/amazon-research/co-with-gnns-example.

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Acknowledgments We thank Fernando Brandao, George Karypis, Michael Kastoryano, Eric Kessler, Tyler Mullenbach, Nicola Pancotti, Mauricio Resende, Shantu Roy, Grant Salton, Simone Severini, Ayinger Urweisse, and Jason Zhu for fruitful discussions.

Author contributions All authors contributed to the ideation and design of the research. M.J.A.S. and J.K.B. developed and ran the computational experiments, as well as wrote the initial draft of the the manuscript. H.G.K. supervised this work and revised the manuscript.

Competing interests M.J.A.S., J.K.B. and H.G.K. are listed as inventors on a US provisional patent application (number 7924-38500) on combinatorial optimization ith graph neural networks.

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# Supplemental Material for: Combinatorial Optimization with Physics-Inspired Graph Neural Networks

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### I. CORE GCN CODE BLOCK

Listing 1. Core code block of example script based on the DGL library. The first block defines a two-layer GCN architecture Ansatz; the second code block defines the loss function as described by Eq. (6). Further details can be found in the main text, as well as in Ref. [S117].

```
# Import required packages
import dgl
import torch
import torch.nn as nn
from dgl.nn.pytorch import GraphConv
# Define two-layer GCN
class GCN(nn.Module):
  def __init__(self ,in_feats ,hidden ,classes):
      super(GCN, self). __init__()
self.conv1 = GraphConv(in_feats, hidden)
      self.conv2 = GraphConv(hidden, classes)
  def forward (self,g,inputs):
      h = self.conv1(g,inputs)
      h = torch.relu(h)
      h = self.conv2(g,h)
      # binary classification
      h = torch.sigmoid(h)
      return h
# Define custom loss function for QUBOs
def loss_func(probs_,Q_mat):
  function to compute cost value for given
  soft assignments and predefined QUBO matrix
  \# minimize cost = x.T * Q * x
  cost = (probs .T @ Q mat @ probs ).squeeze()
  return cost
```

#### II. HYPERPARAMETERS FOR G-SET EXPERIMENTS

In this section, we provide details for the specific model configurations (hyperparameters) as used to solve the Gset instances with our physics-inspired GNN solver (PI-GNN). The results achieved with these model configurations are displayed in Tab. I; the corresponding hyperparameters are given in Tab. II. Our base GCN architecture with tunable number of layers K is specified in Listing 2.

graph	PI-GNN	embedding $d_0$	layers $K$	hidden dim $d_1$	hidden dim $d_2$	hidden dim $d_3$	learning rate $\beta$	dropout
G14	3026	369	1	5	_		0.00467	0.0
G15	2990	394	1	5	_	_	0.00587	0.0
G22	13181	419	2	1909	3401	_	0.00103	0.4498
G49	5918	2167	3	2338	1955	8	0.00058	0.3554
G50	5820	208	3	218	3582	566	0.00488	0.2365
G55	10138	278	3	8412	8352	5499	0.00161	0.1062
G70	9421	109	3	1233	7048	11869	0.00139	0.3912

TABLE II. Numerical results for MaxCut on Gset instances, with hyperparameters specified for the PI-GNN solver.

```
Listing 2. Base GCN architecture used for solving MaxCut on Gset problem instances.
# Define GNN object
class GCN_dev(nn.Module):
               _(self, in_feats, hidden_sizes, dropout, num_classes):
        super(GCN_dev, self).__init__()
# Combine all layer sizes into a single list
        all layers = [in feats] + hidden sizes + [num classes]
        # slice list into sub-lists of length 2
         self.layer_sizes = list(window(all_layers))
        # reference to ID final layer
         self.out layer id = len(self.layer_sizes) - 1
         self.dropout frac = dropout
         self.layers = OrderedDict()
         for idx, (layer_in, layer_out) in enumerate(self.layer sizes):
             self.layers[idx] = GraphConv(layer in, layer out).to(DEVICE)
    def forward(self, g, inputs):
         for k, layer in self.layers.items():
             if k == 0: # reference to ID final layer
                 h = layer(g, inputs)
                 h = torch.relu(h)
                 h = F.dropout(h, p=self.dropout_frac)
             elif 0 < k < self.out layer id: # intermediate layers
                 h = layer(g, h)
                 h = torch.relu(h)
                 h = F. dropout (h, p=self.dropout frac)
             else: # output layer
                 h = layer(g, h)
                 h = torch.sigmoid(h) # binary classification
        return h
```