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From Near to Eternity: Spin-glass planting, tiling puzzles, and constraint satisfaction problems

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We present a methodology for generating Ising Hamiltonians of tunable complexity and with a priori known ground states based on a decomposition of the model graph into edge-disjoint subgraphs. The idea is illustrated with a spin-glass model defined on a cubic lattice, where subproblems, whose couplers are restricted to the two values \{-1, +1\}, are specified on unit cubes and are parametrized by their local degeneracy. The construction is shown to be equivalent to a type of three-dimensional constraint satisfaction problem known as the tiling puzzle. By varying the proportions of subproblem types, the Hamiltonian can span a dramatic range of typical computational complexity, from fairly easy to many orders of magnitude more difficult than prototypical bimodal and Gaussian spin glasses in three space dimensions. We corroborate this behavior via experiments with different algorithms and discuss generalizations and extensions to different types of graphs.

I. INTRODUCTION

Hard optimization problems are ubiquitous throughout the sciences and engineering, and have consequently been the subject of intense theoretical and practical study. While it is generally considered highly unlikely that problems classified as NP-hard can be solved efficiently for all members of their class, numerous algorithms have been devised that either strive for an approximate solution (e.g., simulated annealing or stochastic local searches) or solve the problems exactly in exponential time, but through mathematical insight (e.g., branch-and-bound or branch-and-cut), the algorithms increase the practically-feasible sizes over what can be achieved using brute force. Evaluation of such algorithms often requires access to benchmarking problems of various types; ideally their difficulty should also be a controllable (or tunable) property. Ideas of generating test instances with planted solutions, that is whose optimizing values are known to the problem constructor, have been explored in various fields for decades [1–3]. A method of planting solutions to hard random Boolean satisfiability (SAT) problems based on statistical mechanics was first proposed in Ref. [4]. In contrast, doing so for topologically-structured problems is considerably less charted territory, as the correspondence between random SAT and the diluted spin glass disappears; thus replica symmetry breaking analysis [5] no longer applies. As such, generating hard problems with a known solution for nearest-neighbor Ising models on a hypercube is a relatively new field of study.

While the task of creating such problems is of theoretical interest due to their potential assistance in answering open questions about the nature of spin glasses, the pace of research has been hastened in recent years by practical motivations, in particular, the availability of quantum annealing [6, 7] and related analog devices (e.g., optical [8]) physically implementing Ising Hamiltonian minimization heuristics. Such Hamiltonians are usually constrained by device manufacturing considerations to having short-ranged interaction terms [9]; while more general objectives can typically be encoded onto their native graphs, this may require many system variables for each objective function variable, limiting the problem sizes that can be studied. Hence, a way of encoding topology-native problems is desirable.

In this work, we present a methodology for generating short-range Ising problems with a known ground state. Our primary focus is on the three-dimensional lattice with periodic boundary conditions, primarily because the three-dimensional spin glass is a prototypical complex system and of tremendous interest to condensed matter physicists, but also because the regular structure over the lattice allows a description of our key contributions in a natural and transparent manner. We stress, however, that the method is adaptable to different graph topologies.

By carefully decomposing the underlying graphical structure and selecting Ising subproblems over the resultant components, whose spin-spin interactions are restricted to being bimodal, from classes designed to have specific features influencing hardness, we obtain a factor graph representation having the topology of a body-centered cubic lattice (see Fig. 1). Following a Voronoi tessellation of this lattice, we note that the resultant problem is equivalent to a specific type of three-dimensional tile matching puzzle, a generalization of the two-dimensional edge-matching puzzle [10]. In their
FIG. 1. Cutaway diagram showing a decomposition of the cubic lattice into edge-disjoint induced subgraphs, the colored unit cube cells. A cell in the right foreground is omitted for clarity. The collection of vertices specifying each colored cube is defined as \( C \). Under a periodic boundary, each spin appears in exactly two subgraphs. Ising subproblems with \( J_{ij} \in \{-1, 1\} \), i.e., requiring a single bit of precision, are specified for the cells such that one of each subproblem's ground states is consistent with its neighbors'. Over the full problem, the concatenation of these partial states forms the planted solution. As discussed in the text, the unit cell Hamiltonians' lack of shared edges is a crucial property enabling large, tunable changes in complexity while guaranteeing a known ground state.

two-dimensional form, these puzzles are constraint satisfaction problems requiring placement of tiles from a given set in permissible spatial locations so that patterns on the tile edges match those of their neighbors. A well-known example is Eternity II [11], a to date unsolved problem for whose solution a $2$ million prize was once offered; the prize deadline expired in 2010. Our construction yields polyhedral rather than square puzzle pieces; in particular the units are truncated octahedra whose facets are of various color patterns (see Fig. 6 below). By varying the pattern properties via manipulation of the underlying Ising subproblems, we quickly and efficiently induce a tremendous variation in problem difficulty observed over all examined heuristic algorithms; hundreds of problems at any hardness level within our attainable range can be obtained in seconds with modest computing resources. Hard problems within our class are observed to be many orders of magnitude more difficult than randomly generated spin-glass problems with bimodal disorder, i.e., where the spin-spin interactions are randomly drawn from \( \{\pm 1\} \). Despite our restriction to using problem parameters in \( \{\pm 1\} \), we can directly generate problems with a known solution that are vastly more difficult than the Gaussian spin glass, which is itself known to be considerably harder than the random bimodal class. We thus consider our construction technique as giving rise to a novel type of disordered system, and discuss our results within the context of hardness phase transitions for combinatorial problems.

The paper is structured as follows. In Sec. II we present an overview of recently-introduced planting techniques, followed by an outline of the planting approach presented in this work in Sec. III. Sections IV and V discuss the relationship of the presented problems to constraint satisfaction problems. Section VI presents numerical experiments using different algorithms highlighting the tunability of typical computational complexity for the problems with planted solutions, followed by discussions, generalizations to other graphs, as well as concluding remarks.

II. RELATED WORK

Our methodology addresses several shortcomings of recently-proposed techniques for solution planting in short-range Ising models. In Ref. [12], a construction method was presented in which frustrated loops with known solution were employed as subproblem units. While the idea is appealing, the resultant problems typically have coupler strengths spanning a large range, posing serious issues for physical devices with precision limitations. An ad hoc limited-precision variant was introduced in Ref. [13]. Both approaches, however, bear the more serious and systematic disadvantage of tending to generate sets of problems whose members are not of sufficient hardness; we contend that this is a consequence of adding subproblem couplers. It is not hard to see that when partial Hamiltonians having consistent ground states are added, couplers in common which in the individual minima are either both satisfied or both violated will increase in magnitude. In a ground state, however, satisfied couplers are more common than violated ones, hence for any given shared coupler, it is relatively likely that the resultant (added) coupler tends to more strongly prefer alignment with the shared ground state. When this occurs over many edges, the problem's degrees of freedom are "softly" constrained. The effect is the introduction of "hints" facilitating an optimization algorithm's task.

In Ref. [14], a method was presented for generating difficult instances with bimodal couplers by iteratively chang-
ing the couplers’ signs to maximize the time required by a given solver to reach the (hypothesized) ground state. This is a promising approach as it both avoids the coupler addition issue and makes modest precision demands, however it generally does not allow the ground state to be known with certainty and hence, is difficult to scale to larger systems. Furthermore, the approach requires a Monte-Carlo-like bond moving algorithm that will likely not easily scale up to much larger problem sizes. The authors do discuss a variant enabling solution planting, but as it again relies on adding couplers, the precision issue returns and, they report, the gains in hardness are modest compared to the mode not controlling the ground state. While the authors provide a useful analysis of post hoc empirical correlates of hardness, for example the parallel tempering empirical mixing time, first proposed in Ref. [15], the generation procedure is ultimately a local search technique with respect to solver computational time, and hence does not yield or use physical or algorithmic insight at the problem level about what tunes difficulty. In other words, one could not, merely by reference to the resultant Hamiltonian, predict whether the problem is easy or hard. Finally, to design problems based on the time-to-solution of a given solver, assuming this will carry over to other optimization techniques, requires theoretical backing that is missing to date.

A recent paper [16] introduced the method of patch planting, in which subgraphs with known solution are coupled to each other satisfying all interactions and thus planting an overall ground state. An advantage is that low-precision and tunable problems can be readily created for a given application domain. The authors report, however, that the resultant problems are observed to be less difficult than random problems, simply because frustrated loops typically do not extend beyond the subgraph building blocks, making the ground state less frustrated than in the fully random case. This problem can be slightly alleviated by post-processing mining of the data [16].

III. PROBLEM CONSTRUCTION

Consider a cubic lattice of linear size \( L \) with periodic boundary conditions. Let the underlying graph be \( G = (V, E) \) with \( V \) and \( E \) its vertex and edge sets, respectively. Define \( U \) to be the vertices of \( V \) whose coordinates \((i_x, i_y, i_z)\) consist of integers of the same parity, for example, \((1,3,1)\) or \((0,2,4)\). For each \( u \in U \), define \( C \) to be the vertices of the cubic unit cell implied when \( u \) is opposite to the vertex with coordinates \((i_x + 1, i_y + 1, i_z + 1)\), where the addition is modulo \( L \) to account for the periodic boundary. Let \( C \) be the collection of all such cell vertex sets. This construction is partially illustrated in Figure 1, where each colored unit cell represents a subgraph \( G[C] \) induced by a vertex set \( C \in C \) as defined. It should be clear that first, each vertex in \( V \) appears in two and only two unit cells, and second, that the unit cell subgraphs do not share any edges. In graph terminology, the family \( C \) partitions \( G \) into edge-disjoint induced subgraphs.

We are concerned here with constructing Ising problems on the lattice with zero field, i.e., whose Hamiltonians are of the form

\[
H(\mathbf{s}) = \sum_{ij \in E} J_{ij} s_i s_j, \tag{1}
\]

Due to the disjointness of the cell edge sets, we can regroup the Hamiltonian into terms each dependent only on the couplings within a unit cell in \( C \):

\[
H(\mathbf{s}) = \sum_{C \in \mathcal{C}} H_C(\mathbf{s}), \tag{2}
\]

where

\[
H_C(\mathbf{s}) \triangleq \sum_{ij \in E[C]} J_{ij} s_i s_j. \tag{3}
\]

The terms \( \{H_C(\mathbf{s})\} \) are called the unit cell subproblem Hamiltonians. Specifying each subproblem’s Hamiltonian is sufficient to imply a full Hamiltonian over the lattice. A straightforward consequence of the relation between summation and minimization, exploited in Ref. [12], is that if the subproblems share the same minimizing configuration \( \mathbf{s}^* \), then \( H \) is also minimized at \( \mathbf{s}^* \). This, in turn, suggests a natural construction procedure for a problem \( H \) with a known ground state. We note a seemingly small but in fact deep and far-reaching difference between prior methodologies and what we propose is that subproblem couplers are never added, avoiding the issue discussed in Section II.

It may appear surprising at first that any interesting behavior can result from linking such apparently simple unit cell subproblems, but this impression turns out to be quite false. Indeed, by restricting our attention to simple classes of subproblems whose couplers belong to the set \( \{\pm 1\} \), in other words representable with a single bit of precision, we can effect dramatic changes in problem complexity, from trivial to many orders of magnitude harder than, e.g., Gaussian spin glasses.

Without loss of generality, we focus on planting the ferromagnetic ground state, i.e., \( \mathbf{s}^* = (+1, +1, \ldots, +1) \) and its \( \mathbb{Z}_2 \) image, because once such a problem has been generated, the structure of the construction procedure can be concealed by gauge randomization from a would-be adversary seeking
the solution. Specifically, to translate to an arbitrarily chosen ground state $t^*$, one would transform the initially-determined couplings $\{J_{ij}\}$ via

$$J'_{ij} \leftarrow J_{ij}t_i^*t_j^*.$$  \hspace{1cm} (4)

We thus turn our attention to the set of unit cell Hamiltonians having couplers of magnitude 1 and ground state $s^*$. Here we disregard the trivial subproblem comprised uniquely of ferromagnetic bonds; those remaining can be naturally partitioned into three types according to their number of frustrated facets. We call these types $F_2$, $F_4$, and $F_6$, as they contain unit cells with 2, 4, and 6 frustrated facets, respectively. These types can be further partitioned into members equivalent under action of the cube graph automorphism group. More precisely, problems within a given equivalence class can be arrived at from one another via transformation from $O_h$, the 48 octahedral symmetries comprised of rotation and reflection leaving a (generic) cube invariant. It turns out that $F_2$ and $F_4$ contain two such classes (the orbits of $O_h$ acting on $F_2$ and $F_4$, called $F_{21}$, $F_{22}$ and $F_{41}$, $F_{42}$) while all members of $F_6$ are equivalent under $O_h$. Figure 2 illustrates an arbitrarily-chosen set of equivalence class representatives for the three problems types, while in Figure 4 a few members of type $F_6$ are shown.

While the subproblems can all readily be verified to have $s^*$ as a GS, they clearly differ in the number of other minimizers they possess. Up to $Z_2$ symmetry, members of $F_{21}$ have a unique ground state while those of $F_{22}$ have 4. All problems in $F_4$ and $F_6$ have 2 and 8 global minima, respectively. Figure 2 shows the 4 ground states of a specific member of $F_{22}$. As we shall see, this variation in subproblem ground-state degeneracy turns out to exert a crucial effect on the computational difficulty of the final planted problem.

We are now ready to straightforwardly specify our problem construction procedure. First, a probability distribution is chosen over the problem classes $\{F_{ij}\}$. A naïve example, which in fact turns out to be a poor choice for generating hard problems, is to sample them uniformly. Next, a distribution over members of the classes is specified. In our examples this was always uniform. The randomness in generating the problems is thus decomposed into what we call subproblem disorder (over the classes) and $O_h$-disorder (over the members); the latter so called because it arises equivalently by sampling symmetries of the octahedral group. Subproblems over the unit cell set $C$ are then sampled from the defined distributions.

Note that $O_h$-disorder is sufficient to induce considerable richness in the types of final problems, even when the cells in $C$ are all assigned from the same subproblem class. For example, it may appear at first that restricting all subproblems to the class $F_6$ (a regime which turns out to be highly interesting) will result in a so-called fully-frustrated model [17], i.e., where all chordless cycles (or plaquettes) are frustrated in the final Hamiltonian, because problems in $F_6$ are themselves fully-frustrated. That, however, would be incorrect; it is easy to check that cycles forming the facets of unit cubes not in $C$ may or may not be frustrated depending on the random orientation of the bounding $C$-cell problems.

It should be pointed out that while the method does ensure
IV. THE PUZZLE OF SPIN GLASSES

The proposed problem generation formalism enables translation to a specific type of geometrically-appealing constraint satisfaction problem (CSP) called a tiling puzzle. While these problems are usually defined over two-dimensional domains, we shall see that our three-dimensional technique yields a polyhedral generalization of an edge-matching puzzle, an example of which is Eternity II [11]. The idea is to abstract the Ising problem into a generic CSP representation known as a factor graph [19]. While the Ising Hamiltonian itself can always be interpreted as a factor graph, under our planting technique additional structure is present, namely the restriction to certain subsets of states. Following construction of the CSP exploiting this structure, the natural connection to a tiling puzzle is observed.

Figure 5 illustrates the necessary steps for the mapping. For clarity, a direct two-dimensional analog of the three-dimensional case is shown, in which the unit cells induced by the decomposition into \( \mathcal{C} \) are the cycles forming a checkerboard pattern rather than unit cubes as displayed in Figure 1.

We first consider the construction of a graph with respect to \( \mathcal{C} \) which we call the subproblem lattice, denoted by \( \tilde{G} \). Suppose a vertex is placed at the centroid of each unit cell. The subproblem lattice is defined by considering two such vertices to be adjacent if and only if their corresponding unit cells share a corner vertex. Clearly, this lattice may be 2-colored, that is partitioned into two subsets none of whose members are neighbors. This in turn implies a partitioning of the cells in \( \mathcal{C} \) into two disjoint sets \( \mathcal{W} \) and \( \mathcal{B} \), whose elements are labeled with colors \( W \) and \( B \), respectively. Figure 5(a) demonstrates the two-dimensional construction of \( \tilde{G} \) and its canonical coloring. We observe that the planted state is in fact a ground state, as the couplers are selected from the \( \{ \pm 1 \} \) class, it is in principle possible for other ground states to exist [18]. Our experimental validation in Sec. VII is thus concerned with the difficulty of finding any ground state. Methods of removing the degeneracy are certainly conceivable and are discussed briefly in Sec. VII yet will be the subject of future study.

We proceed next to a further exploration of the problem structure, discussing its connection to tiling puzzles.
vertex coloring, where yellow (light shading) and blue (dark shading) may represent labels \( W \) and \( B \) or vice versa. In three space dimensions, each yellow (light shading) cell is surrounded by eight blue (dark shading) cells instead of four.

We next construct the factor graph CSP representation equivalent to the problem of minimizing \( H \). The idea is to derive an objective consisting of independent variables subject to equality constraints along the edges of \( G \). First, each vertex in \( G \), mapping to cell \( C \) in the original lattice, is identified with a variable \( s^{W}_{i} \) or \( s^{B}_{i} \), over 8 Ising variables (4 in two space dimensions) depending on its label in the 2-coloring. Because each vertex of the initial lattice is a shared corner of two unit cells in \( C \) of opposite color, in the concatenation of the states \( \prod_{C \in W} s^{W}_{C} \prod_{C' \in B} s^{B}_{C'} \) variables \( s^{W}_{i} \) and \( s^{B}_{i} \) will occur exactly once for each vertex \( i \in V \). The domain of each subproblem lattice variable is simply the ensemble of its subproblem ground states, the \( \text{ground-state set} \ \Gamma_{C} \), so that the full configuration space of the new problem is the Cartesian product

\[
\Gamma^{W,B} \triangleq \prod_{C \in W} \Gamma_{C} \prod_{C' \in B} \Gamma_{C'}.
\]

To be a feasible solution to the original problem however, equality of neighboring variables must be enforced, that is \( s^{W}_{i} = s^{B}_{i} \forall i \). If we define the problem

\[
\max_{s^{W,B}} \prod_{i \in V} \delta(s^{W}_{i}, s^{B}_{i}),
\]

where \( \delta(x,y) = 1 \) if \( x = y \) and zero otherwise, we readily see that the solution is obtained for the overall planted ground state \( s^{B} = s^{W} = s^{*} \). To make the graphical connection, we note that each delta function in Eq. (6) can be interpreted as a factor over neighboring subproblem lattice variables (identified with \( C, C' \), i.e., \( \psi_{C,C'}(s^{W}_{C}, s^{B}_{C'}) \triangleq \delta(s^{W}_{i}, s^{B}_{i}) \) with \( i = C \cap C' \)). The final maximization objective is then the product of all factors over subproblem lattice variables

\[
f(s^{W,B}) = \prod_{C,C' \in E} \psi_{C,C'}(s^{W}_{C}, s^{B}_{C'}),
\]

where the product index refers to edges of \( \tilde{G} \) by the cells in \( C \) mapping to their endpoint vertices. The factor graph corresponding to the two-dimensional lattice in Fig. 5 (a) appears in Fig. 5 (b), with variables represented by circular vertices (whose colors identify their labels as \( W \) or \( B \)) and factors by the squares along the edges. In three dimensions, each variable is involved in eight factors rather than four.

Consider now the subproblem lattice with its vertices lying at their “natural” points in Euclidean space, i.e., at the centroids of the unit cells, rather than as a generic graph. A Voronoi tessellation \([20] \) is a partitioning of the space surrounding the vertices into convex polytopes, usually called tiles, but for reasons that will soon be clear, we refer to as tiling locations, such that all points within a polytope are closest (typically in the \( L_2 \) norm sense) to the enclosed vertex. It is easy to see in Fig. 5 (a) that in two dimensions, the subproblem lattice forms a periodic pattern, sometimes called the quincunx, of squares of length 2 each with a central vertex. The appropriate tessellation of \( \tilde{G} \) is a partitioning into tilted square regions centered at each vertex. Neglecting for an instant the significance of the colors, Fig. 5 (c) shows 5 full and 8 partial tiling squares in the section of lattice drawn. Returning to the CSP in which the task was to assign the variables from their respective domains (ground-state sets) so that factor constraints are satisfied, we now observe the equivalence to a tile-matching puzzle. This is comprised of a board (the space embedding \( \tilde{G} \)), segmented into octagons (the cells in the Voronoi tessellation), each of which is endowed with a repertoire of tiles (the ground-state set associated with the location). The tile faces could be in one of two colors (the spin values, not to be confused with the 2-coloring introduced to construct the CSP). The puzzle task is to select the tiles so that no two adjacent tile faces have a different color. If orange (light) and purple (dark) represent spin values, say 1 and \(-1\), respectively, then Fig. 5 (c) shows a hypothetical tiling assignment. Here, a violation occurs where the tile colors on both sides of a factor differ.

In three dimensions, the puzzle is analogous, but with a few twists. In generalization of the two-dimensional case, the subproblem lattice \( \tilde{G} \), will, when respecting Euclidean coordinates, form a pattern of cubes of length 2, each with a vertex at its centroid. Such a formation is known as the body-centered cubic (BCC) lattice. The corresponding Voronoi tessellation is the space-filling \( \text{bitruncated cubic honeycomb} \) whose base unit is the \( \text{truncated octahedron} \) \([20, 21] \). These polyhedral cells, which now define the three-dimensional tiling locations, each consist of eight \( \text{hexagonal} \) facets, through which the edges of our CSP factor graph pass, and six square facets, defining the boundaries between \( \tilde{G} \) vertices at distance 2, but through which no CSP factors pass. The ground-state sets associated with each vertex of \( \tilde{G} \) thus map to coloring configurations on the hexagonal facets of the enclosing Voronoi cell, defining a set of tiles, and again the task is to place the tiles while meeting all coloring constraints. The construction is illustrated in Fig. 6.
FIG. 5. Steps in the derivation of a tile-matching puzzle from the unit cell planting methodology, shown in two dimensions with a free boundary for clarity. Note that the two-dimensional cells induced by the decomposition are the unit cycle (plaquette) subgraphs arranged in a checkerboard pattern rather than the unit cubes shown in Fig. 1 in three dimensions. In panel (a), the subproblem lattice $\tilde{G}$ is constructed by placing a vertex at each subproblem cell’s centroid. As discussed in the text, the vertices are two-colored to yield a convenient representation of the puzzle. In three space dimensions, each yellow (light shading) unit cell contacts 8 rather than 4 blue (dark shading) neighbors; the corresponding $\tilde{G}$ is the body-centered cubic lattice. The factor graph associated with $\tilde{G}$ appears in panel (b), with factors shown as squares. Each vertex is involved in 8 factors in three space dimensions. A Voronoi tessellation of $\tilde{G}$ yields the set of tiling locations. In two space dimensions, these are the areas within the thick lines in panel (c). The factor graph domains, determined by the Ising subproblems, specify a set of allowable tile patterns at each location. The figure shows a hypothetical tile placed at each site. The factor constraints require adjacent colors to agree; hence, the central tile shown has 1 violation among its 4 neighbors. The puzzle for the three-dimensional tessellation is shown in Fig. 6.

latter, each tile from the given set may be placed at more than one board location (in an allowable orientation) while presently, the ground-state set associated with each site constrains the choices. This fact, along with the polynomial-time solvability [22] of the original ground-state problem in two dimensions, could lead one to suspect that the puzzles we have contrived may not be overly interesting. In three space dimensions, however, solution by graph matching is no longer applicable, and as we shall see, the problems exhibit a rich range of behavior intimately tied to the known properties of generic constraint satisfaction problems.

V. THE PHASES OF SATISFACTION

Hardness phase transitions in combinatorial problems have been an active area of study for several decades now [23]. Of particular interest is the emergence of difficulty divergencies in random satisfiability (SAT) problems as parameters guiding instance generation are varied [24]. While the tiling CSPs we have proposed here can certainly be expressed as SAT problems [25], the resultant formulae do not, however, follow the typical assumptions of random SAT. Most notably, the variables are not free to appear in any clause with some probability, but follow the highly-structured topological constraints of the puzzle. Of course, the subproblem and $O_h$-disorder impose stochasticity over the factor graph variable domains $\{\Gamma_C\}$, but this is a localized randomness, violating key assumptions in the analysis of random SAT. Nonetheless, we expect that commonly-known facts and intuitions regarding the relation between how constrained a problem is and its empirical hardness will apply to our situation.

Because the tiling puzzle color factors are invariant to the distribution over cell problems, constrainedness depends exclusively on the features of the ground-state sets. As we alluded to earlier, a dominant correlate of problem hardness is the average size of the ground-state sets, $\mathbb{E} |\Gamma|$. A highly-constrained regime is one in which the subproblems tend to have relatively small degeneracy, and vice versa for a loosely constrained one. A subproblem demonstrating a maximal constraint level is any member of class $F_{21}$ (with a unique ground state up to $Z_2$), while one constrained minimally, which was not considered in the experiments, is the zero $J$ subproblem, in which all 256 states are ground states. Within the class
These extremes strongly suggest that for some intermediate state using simple strategies to escape from local minima. The state space, while for local search, the energy landscape encounter relatively small branching factors when traversing choices from the ground-state sets, a tree search method will problems are also typically easy: Because there are few local ground-state sets. At the other extreme, highly-constrained interactions (or without trivially removing all $J_{ij}$), an interesting question, probed experimentally in the sequel is: Given the proposed classes of unit-cell subproblems ($F_2$, $F_4$, and $F_6$) and all possible mixture distributions over the classes, did the hypothesized peak occur for some interior point of the distribution set? An affirmative answer would suggest that the corresponding CSP class is among the hardest of all tile-matching puzzles of the type we have introduced, i.e., for which the locations’ tile sets may be chosen arbitrarily, not by constraining them to map to subproblem ground states. Conversely, if the hardness maximum within the set of mixtures occurs at the set boundary, it would point to the existence of more difficult tiling puzzles not representable within our framework.

VI. EXPERIMENTS

We now numerically study the typical complexity of certain subproblem classes to illustrate their tunability. This section focuses on three-dimensional cubic lattices, followed by a demonstration that the approach works well also in two space dimensions (Sec. VI.B). We conclude with a discussion of generalizations to other graph types.

While we would ideally have performed a rigorous numerical study for a discrete “grid” representing all subproblem types we have proposed, this is computationally prohibitive. Consequently, we focus on two illustrative regimes demonstrating interesting variations in problem difficulty over three algorithms, and the role played by subproblem degeneracy. The simulation results show that highly complex problems with known ground state, and more difficult than conventional spin glasses with both Gaussian and random bimodal couplings, are accessible via our methodology, but they also strongly suggest that the hardness maximum over the distribution set occurs at a boundary, namely when the distribution is a point mass concentrated on class $F_6$. Perhaps unsurprisingly then, we conclude that tiling puzzles equivalent to Ising problems constructed using the classes $\{F_2, F_4, F_6\}$ are likely not the hardest among those where the tile sets are arbitrarily specifiable.

All problems are defined on three-dimensional lattices of size $8 \times 8 \times 8$ with periodic boundary conditions. We consider two classes of problems corresponding to one-dimensional slices of the problem mixture parameter space. The first class, gallus_26, allows subproblems to belong solely to classes $F_6$ and $F_{22}$, while in the second, gallus_46, they
are constrained to $F_6$ and $F_{42}$. Both instance classes are parametrized by $p_6$, the probability of selecting class $F_6$ instead of the alternative [26]. The classes each contained 200 instances generated at 30 uniformly-spaced values of $p_6 \in [0.8, 1]$, for a total of 6000 instances per class. All sub-problems are subject to uniform $O_4$-disorder. The selected range of $p_6$ values is interesting as within it, problems are more difficult than conventional spin glasses with Gaussian or random bimodal couplings. We note that the problems continue to become easier as $p_6$ decreases. Using the values of $|\Gamma|$ shown in Fig. 2, the expected ground-state set sizes in terms of $p_6$ are $\mathbb{E} |\Gamma| = 8p_6 + 4(1 - p_6)$ for gallus_{26} and $\mathbb{E} |\Gamma| = 8p_6 + 2(1 - p_6)$ for gallus_{46}.

Problem difficulty is assessed through performance of three different algorithms designed for problems with rough energy landscapes. The measures of difficulty are in strong agreement across the methods, providing corroboration that the observed difficulty trends should persist robustly across a fairly large class of heuristic algorithms, including backtrack-based search [23].

Simulated annealing (SA) [27] is the most basic algorithm considered. We use the optimized implementation developed by Isakov et al. [28] with $\beta_{\text{min}} = 0.01$ and $\beta_{\text{max}} = 1$. While some form of optimized time to solution (TTS) measure, which considers the best tradeoff between simulation length and number of simulations, would rather have been used, SA runs were too time consuming on the hard problems to generate the requisite runtime histograms reliably. Consequently, we selected a fixed run length of $N_S = 8192$ sweeps for all problems with a single sweep per temperature. Each problem is simulated for three-dimensional problem classes gallus_{26} and gallus_{46}. SA and ICM TTS are measured in Monte Carlo sweeps and seconds, respectively. Within each problem class, every point in a given color (symbol) jointly shows the two corresponding measures for one of the 30 subclasses described in the text (averages are computed over 200 instances from each subclass). The data show that PAMC $\langle \log \rho_S \rangle$ exhibits a strong linear correlation with the other two algorithms’ $\log$ TTS metrics, confirming its power as a measure of hardness and landscape roughness. Furthermore, as relative hardness measures for these problems, the three quantities studied here can be used more or less interchangeably. Error bars omitted for clarity.

FIG. 7. Simulated annealing (SA) $\langle \log_{10} \text{TTS} \rangle$ (right y-axes, red triangles) and parallel tempering with isoenergetic cluster moves (ICM) $\langle \log_{10} \text{TTS}_{\text{opt}} \rangle$ (left y-axes, blue circles) as a function of population annealing Monte Carlo (PAMC) $\langle \log_{10} \rho_S \rangle$ for three-dimensional problem classes gallus_{26} (a) and gallus_{46} (b). SA and ICM TTS are measured in Monte Carlo sweeps and seconds, respectively. Within each problem class, every point in a given color (symbol) jointly shows the two corresponding measures for one of the 30 subclasses described in the text (averages are computed over 200 instances from each subclass). The data show that PAMC $\langle \log \rho_S \rangle$ exhibits a strong linear correlation with the other two algorithms’ $\log$ TTS metrics, confirming its power as a measure of hardness and landscape roughness. Furthermore, as relative hardness measures for these problems, the three quantities studied here can be used more or less interchangeably. Error bars omitted for clarity.

Furthermore, we have used a highly optimized implementation of parallel tempering Monte Carlo with isoenergetic cluster moves (ICM) [29], an adaptive hybrid parallel tempering (PT) [30][32] cluster algorithm [33]. Because ICM is considerably more efficient than SA, it allowed us to gather runtime statistics for each instance, allowing optimized TTSs to be computed. In contrast to SA, total ICM computational time is not merely a function of overall Monte Carlo sweeps, but includes the additional cost of constructing random-sized clusters. Consequently, to track aggregated ICM computational effort we record run times in seconds on hardware dedicated entirely to the simulations. If $P(\tau)$ is the empirically observed probability of finding the ground state in
time \( \tau \) or less, the optimized time to solution is defined as 
\[ \text{TTS}_{\text{opt}} = \min_{\tau} \frac{\tau \log(0.01)}{\log[1 - P(\tau)]}. \]
We note, however, that despite the efficiency of ICM, it commonly failed to find the solution for the harder problems within the maximum allowed 2^{24} total Monte Carlo sweeps, requiring 60 to 75 minutes of real time, when computing the runtime histograms. For difficult classes, these “timeouts” occurred at least half of the time within the 100 ICM repetitions used to tally the histograms. Fortunately, we were nonetheless able to infer an optimized TTS from the conditional histogram in which solutions were found. We used \( N_T = 30 \) temperatures spaced within \( T_{\text{max}} = 3 \) and \( T_{\text{min}} = 0.01 \); see Ref. \([29]\) for further details of ICM.

The final set of tests have used the sequential Monte Carlo \([34]\) method known as population annealing Monte Carlo (PAMC) \([35, 36]\). This algorithm is related to SA but differs crucially in its usage of weight-based resampling, which multiplies or eliminates members of a population according to the ratios of their Boltzmann factors at adjacent temperatures, maintaining thermal equilibrium at each step. Our simulations used \( N_T = 201 \) temperatures with \( 1/T = \beta \in [0.0, 5] \) and \( N_s = 10 \) sweeps per temperature, and a population size of \( R = 5 \times 10^5 \) replicas. In Ref. \([37]\), a PAMC-derived index of landscape roughness called the *entropic family size* \( \rho_S \) was proposed. If \( q_i \) is the fraction of replicas in the final population descended from initial replica \( i \), then \( \rho_S = \lim_{R \to \infty} R/e^{h[\eta]}, \) where \( h[\eta] = -\sum_i q_i \log q_i \). An energy landscape is thus deemed rough if \( h[\eta] \) is small, that is, if relatively few initial replicas survive to the final distribution, yielding a large value of \( \rho_S \). Note that \( \rho_S \) converges quickly in \( R \) and is easily estimated with finite populations. Thermal equilibration is ensured by requiring \( h[\eta] > \log 100 \); when this is not satisfied for an instance, it is rerun with a larger population size. \( \rho_S \) is known to covary strongly with the PT autocorrelation time \([37]\), and as can be seen from Fig. [7] does so for the TTS-based hardness metrics considered here as well. Figure [7] shows, respectively, the dependence of both SA and ICM \( \langle \log_{10} \text{TTS} \rangle \) measures on PAMC \( \langle \log_{10} \rho_S \rangle \) for the 30 subclasses of *gallus* \( 26 \) (a) and *gallus* \( 46 \) (b) studied, where the averages \( \langle \cdots \rangle \) at each point representing a subclass are computed over its 200 generated instances. The plots clearly show a distinct near-linear dependence of both
time-based measures on the PAMC-defined value. Larger ρs on average implies longer time to solution with respect to both SA and ICM algorithms, corroborating the former’s power as an objective measure of landscape roughness.

Results of the PAMC simulations are shown in Fig. 8. We observe a trend of increasing ρs (and hence complexity) for both problem classes as the fraction of subproblems from \( F_6 \) increases towards unity. For comparison, we display the mean log ρs value of two prototypical problems with rough landscapes on the same lattice, the random \( J_{ij} \in \{ \pm 1 \} \) and Gaussian \( J_{ij} \sim N(0, 1) \) \( [N(0, 1) \) a Normal distribution with zero mean and variance one] spin glass, computed using 1000 and 5099 instances of each type, respectively. It is clear that problems in most of the examined subclasses of gallus_26 and gallus_46 are more difficult than both of these widely-studied problem classes. Indeed, for subclasses of gallus_26 corresponding to \( p_6 \in [0.91, 1] \), problems are \( \sim 2 - 3 \) orders of magnitude harder than bimodal spin glasses. Perhaps more surprisingly, they are also \( \sim 2 - 3 \) orders more difficult than Gaussian spin glasses, despite the latter possessing continuous-valued couplings while our instances restrict couplers to \( \{ \pm 1 \} \) which are believed to generally be easier to minimize.

Setting \( p_6 = 1 \) yields the most complex problems of those considered. In fact, we conjecture, based on less comprehensive simulations, that these instances are the hardest among all those constructed with subproblem classes in \{ \( F_2 \), \( F_4 \), \( F_6 \) \}. We plan to perform a comprehensive analysis in the future. For this hard class, the unit cell subproblems, deriving exclusively from \( F_6 \), have 8 ground states each. Because this hardness peak occurs at the boundary of the problem parameter space, it seems plausible that one could instantiate yet more complex three-dimensional tiling puzzles, where the locations were allowed more than 8 (times 2) tile possibilities but not so many that the problem becomes underconstrained and easy to solve.

At first, the greater pointwise difficulty shown in Fig. 8 of an \{ \( F_6 \), \( F_{22} \) \} mixture over one made of \{ \( F_6 \), \( F_{32} \) \} appears to contradict intuition that greater frustration implies higher difficulty. After all, with four bounding frustrated facets, a member of \( F_{22} \) can be interpreted as more frustrated than one of \( F_{32} \), which has two. The story is somewhat more subtle though: While frustration certainly plays a role in tuning hardness in our problems, it appears to do so through its effect on constraint level, namely on the sizes of \( \Gamma_C \), with \( F_{22} \) inducing higher complexity than \( F_{32} \) because its ground-state set size is larger. This fact is displayed in Fig. 8(b), where log ρs for the two classes is plotted against \( \mathbb{E} |\Gamma| \) instead of \( p_6 \). The graph shows that for a given value of \( \mathbb{E} |\Gamma| \), gallus_26 and gallus_46 have a very similar roughness index, implying that mean subproblem degeneracy accurately predicts difficulty regardless of the underlying subproblem mixture.

For completeness, analogous plots displaying similar difficulty trends for SA’s log TTS and ICM’s log TTSopt are shown in Fig. 9 where they are again plotted against \( p_6 \).

In contrast to our results so far, which have considered instance-averaged difficulty measures, Fig. 10 shows histograms of optimized ICM \( \log_{10} \) TTS values for three subclasses of gallus_26 [Fig. 10(a)] and of gallus_46 [Fig. 10(b)] indexed by \( \mathbb{E} |\Gamma| \). The leftward shift in histogram support shows clearly that problems tend to become easier with decreasing \( \mathbb{E} |\Gamma| \). Given the histogram shapes, one may naturally suspect that TTSopt is log-normally distributed. As log TTS is unlikely to be precisely normal across the entire data range, we visualize correspondence with a Gaussian via normal probability plots [38], which relate the sample order statistics, obtained by sorting the data, with the theoretical means of the corresponding normal order statistics. Deviations from a linear relation signal lack of Gaussianity. Figure 11 shows the probability plots for the three subclasses used to generate the preceding histograms for the gallus_26 (a) and gallus_46 (b) classes, respectively. The relation is close to linear over the majority of the histogram support, implying in turn that the TTSopt is approximately log-normally distributed. This is consistent with quantities such as the parallel tempering Monte Carlo autocorrelation time and other roughness measures [15, 59] having the same property.

While we have argued that \( \mathbb{E} |\Gamma| \) is a good predictor of mean difficulty, the histogram results show that this value by no means provides a complete picture. Indeed, when \( \mathbb{E} |\Gamma| = 8 \) (i.e., \( p_6 = 1 \)) there is no variance in \( |\Gamma_C| \) as all subproblems have eight-fold degeneracy, yet the log TTS distributions in Fig. 10 nonetheless show considerable intraclass spread in difficulty. Therefore, there are certainly other factors at play in predicking the hardness, which will be analyzed in upcoming studies.

VII. DISCUSSION

In this Section we discuss generalizations of the planting approach using lattice animals. Furthermore, we present a case study on how the approach generalizes to other non-hypercubic lattices. Finally, we discuss the use of the planted problems for fundamental studies of spin glasses and related statistical systems.
FIG. 9. SA $\langle \log_{10} TTS \rangle$ (a) and ICM $\langle \log_{10} TTS_{opt} \rangle$ (b) plotted against parameter $p_6$ for 30 subclasses of gallus_26 and gallus_46 defined in the text. The problems show the same relative difficulty trends with respect to both algorithms and in accordance with the PAMC results shown in Fig. 8 (a).

FIG. 10. Histograms of (optimized) $\log_{10} TTS_{opt}$ relative to isoenergetic cluster moves for classes gallus_26 (a) and gallus_46 (b) for three subclasses characterized here by mean subproblem degeneracy $E|\Gamma|$: 200 problems within each subclass were used to obtain the histograms. The leftward trend in both sets of histograms shows clearly that the problems become easier with decreasing $E|\Gamma|$, and their shapes suggest that $TTS_{opt}$ is log-normally distributed.

A. Generalization via lattice animals

The proposed unit cell planting technique shows encouraging results and properties, and one may inquire how it may be generalized. In this Section, we present a natural extension of the idea, still assuming lattice-structured problems, where instead of defining subproblems on the unit cells of $C$, they are specified on subgraphs consisting of their unions. One can
verify that such subgraphs, called lattice animals or polyominoes, also partition the lattice into edge-disjoint subgrahps, meaning that subproblem couplers are still not added.

A two-dimensional example of decomposition into lattice animals is shown in Fig. 12. Generalization to three-dimensional polyominoes obtained by grouping cells from the decomposition shown in Fig. 1 is straightforward. Shown in non-gray colors are 6 lattice animals. Only the pink (top right) one is made of a single cell. The key difference from the basic method is that unit cells of a given color are no longer constrained to have their individual ground states agree; only the complete animal ground state is relevant. This extension thus considerably extends the types of subproblems that can be employed, and also introduces an additional mechanism of solution hiding, namely via randomization of the employed lattice animals, which would of course be unknown to the would-be adversary. While the tiling puzzle and CSP interpretation of the problem, suitably modified, continues to hold in this generalization, under lattice animal randomization, the adversary would in essence no longer be certain what puzzle they are even solving.

In this work, we have considered carefully chosen families of subproblems on three-dimensional unit cells. An exploration of extensions to more general lattice animals is outside the scope of the present work. We note, however, that if subproblem couplers are sampled from a given distribution, then provided that the subgraph tree widths are small, their ground states can be computed exactly [40] and gauged to the desired overall ground state.

Finally, we note that this lattice animal generalization is also key when attempting to reduce degeneracy in the planted problems. For example, by selecting the coupler values from a Sidon set [41–43] of the form \( \{ \pm(n - 2)/n, \pm(n - 1)/n, \pm1 \} \) with, e.g., \( n = 50 \) the degeneracy is drastically reduced. Similarly, one could select the couplers from a distribution of the form

\[
|J_{ij}| \in a + (1 - a)u,
\]

where \( u \in [0, 1) \) is a uniform random number and \( a \) close to 1 [44]. However, instead of using basic tiles, more complex lattice animals must be used to construct the problems, something we intend to tackle in the near future.

B. Generalization to arbitrary graphs

The need to benchmark both classical and quantum optimization heuristics has hastened the development of advanced planting techniques for solutions of spin-glass-like optimization problems. We have so far focused our attention on hypercubic lattice problems. However, quantum annealing machines typically have hardwired hardware graphs that are
FIG. 12. Generalization of unit cell planting using lattice animals, also known as polyominoes. Illustrations are in two dimensions for clarity. The analogous procedure in three space dimensions is similar. With lattice animal planting, the edge-disjoint subgraphs underlying the subproblems are no longer restricted to the unit cells in C (the checkerboard in two space dimensions and the decomposition in Fig. 1 for three space dimensions) but are now permitted to be connected subgraphs comprised of unions of cells from C. As before, subproblem couplers are not added. Above, six such subgraphs in non-gray colors are shown, of which only the pink one (top right) is comprised of a single cell. If the tree widths of the lattice animals are small, their ground states can be computed exactly and gauged to the overall target ground state. This extension considerably expands types of subproblems that can be employed and also introduces an additional mechanism of solution hiding via randomization of the employed lattice animals.

So far, we have focused primarily on planting subproblems on three-dimensional unit cells, in part because planar problems without a field are solvable in polynomial time \[22\], but as illustrated in Figs. 5 and 12, the two-dimensional analog...
is readily obtained. The analytical tractability of the planar lattice enables a deep exploration of physical and computational properties, a work which will be reported elsewhere \[47\]. Here we outline the idea and demonstrate that it does indeed perform well on planar lattices and nonplanar quasi-two-dimensional Chimera graphs using population annealing simulations. The two-dimensional subproblems, i.e., the analogs of the cells shown in Fig. 2, are partitioned into classes \( C_i \) for \( i \in \{1, \ldots, 4\} \), within which cells have \( i \) minimizing ground-state configurations each. To achieve the construction, first define two magnitudes \( J_x, J_l > 0 \) with \( J_l > J_x \); presently we take \( J_l = 2 \) and \( J_x = 1 \). A cell in class \( C_i \) is constructed by setting a random edge to be antiferromagnetic with magnitude \( J_x \), \( i - 1 \) of the remaining edges to be ferromagnetic with magnitude \( J_x \), and the leftover edges to be ferromagnetic with magnitude \( J_l \) \[48\]. It is readily verified that the subproblems do indeed have the specified number of local ground states which always include the ferromagnetic state.

As was done in three space dimensions, we consider two classes of problems each comprised of mixtures of two sub-problem classes. Problem class \( C_{2,4} \), consists of mixtures of \( C_2 \) and \( C_4 \) cells, while in \( C_{2,3} \) the cells may belong to \( C_2 \) or \( C_3 \). Both problem classes are parametrized by \( p_2 \), the probability of choosing a cell from \( C_2 \). The results on \( 24 \times 24 \) periodic lattices are shown in Fig. 14. Again, a wide range of landscape roughness is seen to be attainable, including a regime in which the problems are more difficult than random bimodal (±1 couplers) spin glasses. An interesting distinction from three-dimensional results is that the most difficult problems are not those in which cells exclusive belong to the maximally-degenerate class \( C_4 \), but rather the moderately constrained \( C_2 \) class. In fact, class \( C_4 \) is a highly underconstrained regime for this topology and gives rise to very easy problems. The fascinating connections between dimension, complexity, and phase behavior are the subject of ongoing study that goes beyond the scope of this paper.

If one wished to move beyond a regular lattice structure, the natural objective is a decomposition of the problem graph \( G \) into edge-disjoint subgraphs satisfying some constraint allowing tractable minimization. An example of a constrained decomposition is into subgraphs with given minimum vertex degree \[49\]. More directly applicable to the planting context would be a constraint on maximum subgraph tree width thereby allowing determination of the planted subproblem ground states. The edge-disjointness property is the key common aspect with the ideas presented in this paper, as it continues to circumvent the need for adding subproblem couplers. This idea is clearly a generalization of the lattice animal methodology discussed previously. While we have not presently considered planting using such generic subgraphs, one can readily envision a heuristic decomposition algorithm that greedily grows partitioning subgraphs until their tree widths exceed some criterion.

### C. Spin-glass physics

We hope researchers in the field embrace these planted problems to study physical properties of glassy systems beyond the benchmarking of optimization heuristics.

Having arbitrarily-large planted solutions for hypercubic lattices allows one to address different problems in the physics of spin glasses. For example, the computation of defect energies—intimately related to fundamental properties of these paradigmatic disordered systems—strongly depend on the knowledge of ground states (see, for example, Refs. \[50\]-\[55\]). Being able to plant problems would drastically reduce the computational effort in answering these fundamental questions.

Furthermore, by carefully tuning the different instance
classes, problems with different disorder and frustration can be generated. A systematic study of the interplay between disorder and frustration is therefore possible for nontrivial lattices beyond hierarchical ones. Similarly, being able to tune the fraction of frustrated plaquettes allows one to carefully study the emergence of chaotic effects in spin glasses (see, for example, Refs. [56–58] and references therein).

D. Application-based benchmark problems

It is well established that random benchmark problems \cite{59, 60} for classical and quantum solvers using spin-glass-like problems are computationally typically easy. Furthermore, the control over the hardness of the benchmark problems has been rather limited either because (post-) processing is expensive \cite{14, 42}, or because the benchmark generation approach does not give the user enough control over the problems to match, e.g., hardware restrictions \cite{12}.

Because application-based problems from industrial settings are highly structured, they pose additional challenges for the vast pool of optimization techniques designed, in general, for random unstructured problems. This has sparked the use of problems from industry to generate hard (and sometimes tunable) benchmark problems. Most notably, the use of circuit fault diagnosis \cite{61, 62} has produced superbly-hard benchmarks with small number of variables. However, the use of application-based problems for benchmarking is in its infancy and while circuit fault diagnosis shows most promise \cite{62}, many applications have produced benchmark problems that lack the richness needed to perform systematic studies; see, for example, Refs. \cite{63–66}.

The problems presented in this work are highly tunable and computationally easy to generate. Furthermore, they can be embedded in more complex graphs, as is, for example, commonly done on the D-Wave hardware for application benchmarks. Having this tunability thus should allow for not only the generation of problems that might elucidate quantum speedup in analog annealers, but it might also help gain a deeper understanding into quantum annealing for spin glasses in general. In parallel, having these tunable problems might elucidate the application scope of specific optimization techniques, both classical and quantum.

VIII. CONCLUSIONS

We have presented an approach for generating Ising Hamiltonians with planted ground-state solutions and a tunable complexity based on a decomposition of the model graph into edge-disjoint subgraphs. Although we have performed the construction for three-dimensional cubic lattices and illustrated the approach with the two-dimensional pendant, the approach can be generalized to other lattice structures, as shown in Sec. \ref{sec:VII-B} for the Chimera lattice. The construction allows for a wide range in computational complexity depending on the mix of the elementary building blocks used. We corroborate these results with experiments using different optimization heuristics. Subsequent studies will discuss constructions with controllable ground-state degeneracy, as well as the mapping of the complete complexity phase space.

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Referring to the subproblem classes though, a few moments’ reflection reveals that interestingly, free spin degeneracy is impossible in the planted state.
Triviality of the ground state

M. Palassini and A. P. Young, 


R. Harris et al., in preparation (2017).


Hence two bits are precision are required for these classes.


M. Palassini and A. P. Young, Triviality of the ground state structure in Ising spin glasses, Phys. Rev. Lett. 83, 5126 (1999).


