Experimental quantum annealing: case study involving the graph isomorphism problem

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Quantum annealing is a proposed combinatorial optimization technique meant to exploit quantum mechanical effects such as tunneling and entanglement\textsuperscript{1}. Real-world quantum annealing-based solvers require a combination of annealing and classical pre- and post-processing; at this early stage, little is known about how to partition and optimize the processing. This article presents an experimental case study of quantum annealing and some of the factors involved in real-world solvers, using a 504-qubit D-Wave Two machine and the graph isomorphism problem. To illustrate the role of classical pre-processing, a compact Hamiltonian is presented that enables a reduced Ising model for each problem instance. On random $N$-vertex graphs, the median number of variables is reduced from $N^2$ to fewer than $N \log_2 N$ and solvable graph sizes increase from $N = 5$ to $N = 13$. Additionally, error correction via classical post-processing majority voting is evaluated. While the solution times are not competitive with classical approaches to graph isomorphism, the enhanced solver ultimately classified correctly every problem that was mapped to the processor and demonstrated clear advantages over the baseline approach. The results shed some light on the nature of real-world quantum annealing and the associated hybrid classical-quantum solvers.

Quantum annealing (QA) is a proposed combinatorial optimization technique meant to exploit quantum mechanical effects such as tunneling and entanglement\textsuperscript{1}. Machines purportedly implementing a type of quantum annealing have recently become available\textsuperscript{2}. While the extent of “quantumness” in these implementations is not fully understood, some evidence for quantum mechanical effects playing a useful role in the processing has been appearing\textsuperscript{3–5}. Aside from the debate over quantumness, there are interesting questions regarding how to effectively solve a real-world problem using a quantum annealer. Quantum annealing-based solvers require a combination of annealing and classical pre- and post-processing; at this early stage, little is known about how to partition and optimize the processing. For instance, current quantum annealers have severe practical limitations on the size of problems that can be handled. Can the pre-processing algorithms be modified in order to improve scalability? A second question involves post-processing. Quantum annealers provide solutions to an “embedded” version of a problem involving physical qubits. Post-processing is generally required for translating these to solutions to the original problem involving logical qubits (aka variables). Occasionally, a chain of physical qubits representing a single variable resolves to an inconsistent state, a scenario known as a broken chain. Studies are needed regarding broken chains and the possibility of classical error correction during post-processing.

This article presents an experimental case study of quantum annealing and some of the factors involved in real-world solvers, using a 504-qubit D-Wave Two machine. An example of parsimonious pre-processing is considered, along with post-processing majority voting. Through experiments on a 504-qubit D-Wave Two machine, we quantify the QA success probabilities and the impact of the methods under study. We use the graph isomorphism (GI) problem as the problem of focus. The GI problem is to determine whether two input graphs $G_{1,2}$ are in essence the same, such that the adjacency matrices...
can be made identical with a relabeling of vertices. This problem is an interesting candidate for several reasons. First, an accurate quantum annealing-based solver for GI has never been implemented. Second, quantum approaches can sometimes provide new insight into the structure of a problem, even if no speedup over classical approaches is achieved or even expected. Third, the GI problem is mathematically interesting; though many sub-classes of the problem can be solved in polynomial time by specialized classical solvers, the run time of the best general solution is exponential and has remained at \(O(N^{\log N})\) since 1983. The classical computational complexity of the problem is currently considered to be NP-intermediate, and the quantum computational complexity of the problem is unknown. Graph isomorphism is a non-abelian hidden subgroup problem and is not known to be easy in the quantum regime. Lastly, the GI problem is of practical interest. It appears in fields such as very large-scale integrated circuit design, where a circuit’s layout graph must be verified to be equivalent to its schematic graph, and in drug discovery and bio-informatics, where a graph representing a molecular compound must be compared to an entire database, often via a GI tool that performs canonical labeling.

This article relates to previous works as follows. A pre-print by King and McGeoch discusses tuning of quantum annealing algorithms, including the use of low-cost classical post-processing majority voting similar to what is evaluated in this article. Our study goes further regarding pre-processing (designing a Hamiltonian to generate compact Ising models) and covers graph isomorphism rather than problems such as random not-all-equal 3-SAT. A work by Rieffel et al. maps real-world problems such as graph coloring to a D-Wave quantum annealer. Regarding the graph isomorphism problem in particular, multiple attempts have been made using adiabatic quantum annealing. One of the first attempts assigned a Hamiltonian to each graph and conjectured that measurements taken during each adiabatic evolution could be used to distinguish non-isomorphic pairs. A subsequent experimental study using a D-Wave quantum annealer found that using quantum spectra in this manner was not sufficient to distinguish non-isomorphic pairs. A second approach converted a GI problem to a combinatorial optimization problem whose non-negative cost function has a minimum of zero only for an isomorphic pair. The approach required \(N/\log N\) problem variables and additional ancillary variables. It was numerically simulated up to \(N=7\) but not validated on a quantum annealing processor. An alternative GI Hamiltonian was proposed by Lucas.

Proposed Solver

Preliminaries. Before proceeding, we briefly define some key concepts. In quantum annealing, a system is first initialized to a superposition of all possible states as dictated by Hamiltonian \(H_{\text{init}}\). Then slowly evolved such that the initializing function becomes weaker and the energy function of interest, defined by the problem Hamiltonian \(H_p\), becomes dominant. The time-dependent combination of the two Hamiltonians is referred to as the system Hamiltonian \(H(t)\):

\[
H(t) = A(t)H_{\text{init}} + B(t)H_p, \quad t \in [0, T],
\]

where \(A(t)\) and \(B(t)\) are monotonic functions representing the annealing schedule up to time \(T\). In this work, we focus on problem Hamiltonians \(H_p\) of the problem Hamiltonian represents an Ising model, where each spin variable \(s_i \in \{-1, 1\}\) is subject to a local field \(h_i\), and where variables interact pairwise with coupling strength \(J_{ij}\):

\[
H_p = \sum_i h_i s_i + \sum_{ij} J_{ij} s_i s_j.
\]

Problems can alternatively be represented as quadratic unconstrained binary optimization (QUBO) problems, with binary variables \(x_i \in \{0, 1\}\) instead of spin variables, and with the following cost function:

\[
f_{\text{qubo}}(x) = \sum_{ij} Q_{ij} x_i x_j.
\]

In some cases, a QUBO representation is a more convenient starting point. A QUBO problem can be converted to an Ising problem using the relation \(x_i = \frac{(s_i + 1)}{2}\).

It is often the case that an Ising problem \([h, J]\) involves variable interactions not supported by a quantum annealing processor graph. An example is a variable with degree higher than the maximum supported by the D-Wave Chimera architecture (six). One common strategy is to find a minor embedding of the problem graph in the processor graph, in which a set of physical qubits is used to represent each variable. Each qubit is strongly coupled to at least one other qubit in the set, in an effort to keep the entire set in a consistent state. These sets are commonly referred to as chains. We refer to a minor-embedded version of a problem as an embedded Ising problem \([h', J']\).

Baseline Hamiltonian. We first describe a baseline penalty Hamiltonian for the GI problem, building upon the Hamiltonian described in Lucas. The problem input is a pair of simple, undirected \(N\)-vertex graphs \(G_{1,2}\). Penalties are applied such that the ground state energy is zero if the pair is isomorphic and
greater than zero otherwise. The intent is for an energy minimization process (such as quantum annealing) to provide a solution to this decision problem.

The baseline Hamiltonian includes binary variables \( x_{ui} \in \{0, 1\} \) for every possible mapping of a vertex \( u \) in \( G_i \) to a vertex \( i \) in \( G_2 \); in a solution, the variable is 1 if \( u \) is mapped to \( i \), and 0 otherwise. There are two types of penalties that can be applied to a vertex set mapping. One type penalizes mappings that are not bijective. For instance, a mapping in which vertices 1 and 2 in \( G_i \) are both mapped to vertex 1 in \( G_2 \) is invalid; the term \( x_{11}x_{21} \) assigns a positive penalty \( C_1 \) such that if \( x_{11} \) and \( x_{21} \) both resolve to 1, the solution energy will necessarily be greater than zero. The second penalty type involves edge inconsistencies. An example of an edge inconsistency is when vertices \( u,v \) exist in one input graph and not the other (e.g. \( uv \notin E_i \) while \( ij \notin E_i \)). Edge inconsistencies are assigned a positive penalty \( C_2 \). Additional details regarding this style of Hamiltonian can be found in Lucas17.

In Lucas17, couplings can incur zero, one, or two penalties. Here, we require that couplings be penalized no more than once, in order to achieve a simple set of coupler values (e.g. \([0, 1]\) instead of \([0, 1, 2]\)) amenable to quantum annealing. Specifically, an edge-related penalty is applied to a coupling only if there is not a vertex mapping penalty. As an example, if vertices \( u,v \) are mapped to \( i,j \), then the bijection has been violated and coupling \( x_{ui}x_{vj} \) will incur a vertex mapping penalty \( (C_1) \); therefore, an additional edge inconsistency penalty \( (C_2) \) need not be applied. Edge inconsistency penalties are only applied if \( i \neq j \) and \( u \neq v \). The set of required coupler values can be further simplified by setting \( C_1 = C_2 \geq 0 \). The complete Hamiltonian is

\[
H_1 = C_1 \sum_{u} \left( 1 - \sum_{i} x_{ui} \right)^2 + C_1 \sum_{i} \left( 1 - \sum_{u} x_{ui} \right)^2 + C_2 \sum_{ij \in E_i} \sum_{u,v \in E_2} x_{ui}x_{vj} + C_2 \sum_{ij \in E_i} \sum_{u,v \in E_2} x_{ui}x_{vj}
\]

where the \( i \neq j \) and \( u \neq v \) conditions represent the main modifications to Lucas17.

**Compact Hamiltonian.** The approach embodied in the baseline Hamiltonian \( H_1 \) suffers from a severe lack of scalability. For \( N \)-vertex input graphs, it requires \( N^2 \) logical variables. Moreover, due to the limited direct connections between qubits in the D-Wave Chimera architecture, problems are often given a minor embedding into the processor working graph. This typically involves replicating variables across multiple qubits. Thus, the qubit requirements can reach \( O(N^4) \). Problems mapped in this way to a ~500-qubit processor tend to be limited to \( N = 5 \) or 6. We now investigate whether a more effective Hamiltonian can be designed. The idea is that many variables and interactions are unnecessary, and information indicating so can be leveraged up front during the requisite pre-processing. Note that an isomorphic mapping requires the vertices in each matched pair to have the same degree. Thus, degree information can be leveraged up front during the requisite pre-processing. Note that an isomorphic mapping of such vertices

\[
H_2 = \sum_{u} \left( 1 - \sum_{i} x_{ui} \right)^2 + \sum_{i} \left( 1 - \sum_{u} x_{ui} \right)^2 + \sum_{ij \in E_i} \sum_{u,v \in E_2} x_{ui}x_{vj} + \sum_{ij \in E_i} \sum_{u,v \in E_2} x_{ui}x_{vj}
\]

Figure 1 shows an example of problem instances generated using the baseline and the compact Hamiltonians on the same input; it illustrates how the number of variables and the number of non-zero interactions—as seen in the off-diagonal entries in the associated QUBO matrix \( Q \)—can be much smaller when using the compact Hamiltonian. The variable scaling of each Hamiltonian is quantified and compared in Results.

\( H_2 \) was validated using a software solver (D-Wave System's ising-heuristic version 1.5.2) that provides exact results for problems with low tree width. In exhaustive testing of all \( 2^{12} N = 4 \) pairs and \( 2^{20} N = 5 \) pairs, the ground state energy of \( H_2 \) was confirmed to be zero for isomorphic cases and greater than zero for non-isomorphic.

As a part of a quantum annealing-based solver, an algorithm can be employed that accepts a graph pair as input and uses the Hamiltonian to generate an associated QUBO problem (later converted to...
Figure 1. Illustration of problem instances generated using baseline Hamiltonian $H_1$ and compact Hamiltonian $H_2$ on the same input. For many inputs, $H_2$ requires fewer variables and non-zero interactions.

Figure 2. Graph isomorphism solver flow.
an Ising problem). The proposed algorithm using the compact Hamiltonian $H_2$ is presented (in pseudo code) in Algorithm 1.

**Algorithm 1 | High-level algorithm for generating a QUBO problem using the proposed compact Hamiltonian $H_2$.**

```
Input: adjacency matrices $G_{1,2}$
Output: QUBO matrix

// Define QUBO variables
for each node $v$ in $G_2$
do
  for each node $i$ in $G_1$
do
    if $\deg(v) = \deg(i)$ and $\deg(i) > 0$ then
      create QUBO variable $x_{v,i}$
    end
  end
end

// Populate QUBO matrix with penalty terms
for each pair of different QUBO variables $x_{u,i}$ and $x_{v,j}$ do
  // Assign penalty for node mapping conflict
  if $i = j$ then
    penalize mapping $u \rightarrow i$, $v \rightarrow j$ // Two nodes in $G_2$ map to the same node in $G_1$
  end
  else if $u = v$ then
    penalize mapping $u \rightarrow i$, $v \rightarrow j$ // A node in $G_2$ maps to two nodes in $G_1$
  end
  // Assign penalty for edge discrepancy
  else if $G_1(i, j) \neq G_2(u, v)$ then
    penalize mapping $u \rightarrow i$, $v \rightarrow j$ // Edge in one graph, non-edge in other
  end
end
for each QUBO variable do
  assign value along diagonal of $Q$
end
```

**Complete Solver Flow.** The high-level flow of the proposed GI solver is shown in Fig. 2. The problem input is a graph pair $G_{1,2}$. In this article, the graph types considered are random, simple, undirected graphs. Graphs are generated using the Erdős-Rényi model $G(N, p)$ where we set the probability of an edge being present $p = 0.5$. An advantage of $G(N, 0.5)$ graphs for an initial study is that all graphs are equally likely. This type has been used in classical graph isomorphism work as well. In step 1, the input graphs and the Hamiltonian formulation of interest (e.g. $H_1$ or $H_2$) are used to generate a QUBO problem which is then converted to an Ising problem $[h, J]$. An example of an algorithm for generating the QUBO problem is shown in Algorithm 1. The Ising problem is then compiled to a specific quantum annealing processor in step 2. A main task is to find sets of physical qubits to represent the problem variables (aka logical qubits); this is achieved by providing the $J$ matrix and the processor working graph to the D-Wave findEmbedding() heuristic. Subsequently, the parameters of the embedded Ising problem $[h', J']$ are set following certain strategies such as the use of a random spin gauge (see Methods). The embedded Ising problem, sometimes referred to as a machine instruction, is submitted to the quantum annealing machine along with several job parameters. The quantum annealing job is executed in step 3 and solutions are returned in the form of strings of two-valued variables. These solutions and energies are associated with the embedded problem, not the original Ising problem. Therefore a post-processing step is necessary (step 4), in which the state of each qubit chain is plugged into the cost function of the Ising problem. A difficulty arises when the states of the qubits in a chain are inconsistent, a case referred to as a broken chain. In the proposed solver, broken chains can be handled by either discarding the associated solution, or by performing majority voting over each chain. The two strategies are compared empirically in Results. Given a solution to the original Ising problem, the solution energy can be calculated. If the
lowest energy is zero, then the input pair can be declared isomorphic and no further jobs are necessary. Otherwise, a decision must be made whether to repeat the process from step 2 or to stop and declare that isomorphism could not be established.

Results

Ising Model Scaling. To compare the resource requirements of the two proposed Hamiltonians, 100 pairs of graphs are used as inputs to Step 1 of the solver flow (Fig. 2), where 50 pairs are isomorphic and 50 are non-isomorphic for each size up to $N = 100$. Since $H_1$ models a variable for each possible vertex pair, $N^2$ variables are required by definition. Ising problems generated using $H_2$ are found to use fewer variables than $H_1$; scaling of the median problems fits to $0.748 N^{1.45}$. Incidentally, this indicates that most problems have fewer variables than with the Gaitan et al. approach, which entails $N \log_2 N$ plus ancillary variables\textsuperscript{16}. The variable scaling is illustrated in Fig. 3. In addition to the number of variables, a second resource metric is the number of non-zero interactions between variables; dense interactions make the minor embedding problem more difficult. We find that the scaling of variable interactions has been improved from $O(N^4)$ for $H_1$ to $O(N^{2.9})$ for $H_2$ (where $R^2 = 0.9991$).

Embeddability. Next, we compare the embeddability of the two approaches, in other words the extent to which Ising problems can be minor-embedded in a given processor graph. The processor of choice is the D-Wave Two Vesuvius-6 processor housed at USC ISI. At the time of this writing, the working graph contains 504 qubits and 1427 couplers. Embedding is attempted using the D-Wave findEmbedding() heuristic\textsuperscript{21} with default parameter values such as 10 “tries” per function call. As shown in Fig. 4a, embeddings are found for the majority of problems only for sizes $N \leq 6$ when using $H_1$, but sizes $N \leq 14$ with $H_2$ (Fig. 4a). The median number of qubits across all problems scales as $O(N^{4.22})$ for $H_1$ and has been reduced to $O(N^{3.29})$ for $H_2$ (Fig. 4b).

Experimental Quantum Annealing for Graph Isomorphism. The accuracy of the solver described in the previous section was measured via trials conducted on a D-Wave Two Vesuvius quantum annealing processor. Several alternative strategies were compared—the use of Hamiltonians $H_1$ vs. $H_2$, running a single job per problem vs. multiple jobs, and the use of chain majority voting during post-processing. Note that by construction of the Ising models using a penalty Hamiltonian, problems with non-isomorphic input graphs cannot achieve a zero energy state, regardless of annealing results. The main challenge for the solver is to find the zero energy state for isomorphic pairs. Thus, we first focus on the isomorphic case. One hundred isomorphic pairs were input into the solver for each size $N$ from 3 to 20.

For one strategy in particular the zero energy state was always eventually achieved—the use of Hamiltonian $H_2$ combined with multiple jobs and chain majority voting. Thus, with this strategy there were no false negatives and classification accuracy reached 100% of the embeddable problems, as shown in Table 1. For the most difficult problem, the zero energy state was achieved on the 9th job. All other
Variable requirements decreased from \( \log_2 N \) to fewer than \( N \) discarded, false negatives occurred. For instance, at \( N \) without this form of error correction (in other words, when all solutions containing a broken chain were for which the zero energy ground state solution was only achieved when using this post-processing; \( H_2 \) and in [Lucas] 17) blindly creates QUBO problem, the baseline approach (embodied in Hamiltonian \( H_1 \)) and in \( H_2 \) fits to 0.0723 \( N^{0.29} \) (\( R^2 = 0.9955 \)). A missing data point indicates no embedding was found for the median instance. The 65th and 35th percentiles are marked with bars (not always visible).

strategies incurred false negatives. For the successful strategy, the expected total annealing time was calculated (as described in Methods). Results are shown in Fig. 5.

For completeness, non-isomorphic pairs were run as well, using Hamiltonian \( H_2 \) and chain majority voting. Since in the worst case nine jobs were required to correctly classify the isomorphic pairs above, nine jobs were submitted for each non-isomorphic problem. One hundred non-isomorphic \( G(N, 0.5) \) problems were tested at each size between \( N = 3 \) to 14; of the 1200 problems, embeddings were found for 1186. In addition, pairs of isospectral non-isomorphic graphs (PINGs) were tested. All \( N = 5 \) PINGs were tested (150 permutations), as well as 100 random \( N = 6 \) PINGs. As expected, none of the non-isomorphic problems achieved a zero energy state and thus none were classified as isomorphic. In other words, there were no false positives.

**Discussion**

Several observations can be made from this case study. First, the formulation of the cost function (Hamiltonian) can have a noticeable impact on quantum annealing results. For the graph isomorphism problem, the baseline approach (embodied in Hamiltonian \( H_1 \) and in [Lucas]17) blindly creates QUBO variables for every possible vertex pair, whereas the proposed Hamiltonian \( H_2 \) is more parsimonious. Variable requirements decreased from \( N^2 \) to fewer than \( N \log N \) (Fig. 3) on the graph type under study, allowing larger problems to be solved (Fig. 4 and Table 1). Along with Rieffel13, this is one of the first quantum annealing studies to experimentally quantify the effect of alternative Hamiltonian formulations. One of the impacts of this observation is increased appreciation for the fact that all quantum annealing-based solvers are actually classical-quantum hybrids and that focus must be placed on effectively partitioning the processing and optimizing the classical portion. A caveat is in order—if the classical side is made to do too much work then the quantum annealing aspect becomes trivial and of little value. Further work is needed in identifying the specific strengths of annealing processors, and in leveraging the two sides appropriately.

A second observation is that using chain majority voting during post-processing can in some cases provide a benefit. Previously, such majority voting was evaluated for a different set of problems (scheduling) and was not found to provide a significant benefit13. In our context, there were many problems for which the zero energy ground state solution was only achieved when using this post-processing; without this form of error correction (in other words, when all solutions containing a broken chain were discarded), false negatives occurred. For instance, at \( N = 12, 53 \) of 83 embedded problems were solved on the first job without using chain majority voting, and an additional 12 problems were solved by applying chain majority voting (Table 1). Classical error correction strategies other than majority voting should be explored and assessed in future studies, and their costs quantified.

To our knowledge, the evaluated solver is the first validated, experimental implementation of a QA-based graph isomorphism solver. While it ultimately classified every embeddable problem correctly and demonstrated clear advantages over the baseline approach, it has serious limitations as a graph

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**Figure 4. Embeddability when targeting the USC-LM Vesuvius processor’s 504-qubit, 1427-coupler working graph.** The inputs are 50 isomorphic and 50 non-isomorphic \( G(N, 0.5) \) pairs at each size. (a) Fraction of problems for which an embedding was found within 10 tries, using the D-Wave findEmbedding() heuristic. (b) Median number of utilized qubits among all 100 problems. \( H_1 \) data points fit to \( y = 0.235 N^{0.22} \) (dotted line; \( R^2 = 0.9996 \)); \( H_2 \) fits to 0.0723 \( N^{0.29} \) (\( R^2 = 0.9955 \)). A missing data point indicates no embedding was found for the median instance. The 65th and 35th percentiles are marked with bars (not always visible).
The problem sizes are not competitive with those handled by classical solvers, which can handle $G(N, 0.5)$ graphs with thousands of vertices and even for the hardest graph types can handle hundreds of vertices before running into difficulty. Similarly, the scaling of the total annealing times (Fig. 5) is not competitive with classical scaling. Ultimately, new approaches are likely needed if quantum annealing is to contribute to graph isomorphism theory or practice. Fortunately, the case study provides some new insight into experimental quantum annealing, and contributes methods that have relevance beyond the GI problem. It is hoped that the experimental evaluation of alternative Hamiltonian formulations adds to the understanding of the factors affecting quantum annealing performance, and that the demonstration of majority voting raises new questions about the role of post-processing for a variety of problems.
Methods

Quantum annealing experiments were performed on the D-Wave Two machine housed at USC ISI and operated by the USC-Lockheed Martin Quantum Computing Center. Experiments were conducted in October and November, 2014. The working graph of the machine’s Vesuvius-6 quantum annealing processor consisted of 504 qubits and 1427 couplers during this period. The pattern of working qubits is shown in Figure 6. The qubit temperature was estimated by the manufacturer to be $16 \pm 1$ mK. Additional processor specifications include a maximum anti-ferromagnetic mutual inductance of 1.33, and $f_1/\text{amplitude}$ of $7.5 \pm 1 \mu \phi_0/\sqrt{\text{Hz}}$.

Simple undirected $N$-vertex graphs were constructed according to the Erdős-Rényi $G(n, p)$ model with $n = N$ and with the probability $p$ of including each edge equal to 0.5. Non-isomorphic pairs were generated by creating two graphs as above and checking for non-isomorphism using the MATLAB graphisomorphism() function. Isomorphic pairs were generated by generating a single graph then applying a random permutation to arrive at the second graph. For each pair of input graphs, an Ising model was created using equation (4) or (5). Programming was performed using MATLAB R2014a win64 and the D-Wave MATLAB pack 1.5.2-beta2. The current version of the D-Wave sapiFindEmbedding() function cannot directly embed Ising models with more than one connected component (i.e. a set of variables that interact only with each other and not any of the remaining variables); therefore, models with this characteristic were not included in the input data. When attempting to generate 100 input pairs for each size, such disconnected models occurred no more than 4 times for each size $N \geq 14$. Similarly, the heuristic cannot accept models with fewer than two variables, so in the rare case of a trivial Ising problem with fewer than two variables (e.g. a non-isomorphic pair with no matching degrees), dummy variables were added to the problem.

The $h_i$ values of the Ising problem were split evenly across each qubit in the associated chain in the embedded Ising problem. The $J_{ij}$ values of the Ising problem were assigned to a single coupler connecting two variable chains in the embedded problem. The magnitudes of the embedded $h'$ and $J'$ were scaled together such that the maximum magnitude reached 20% of the full range supported by the processor; the range of the embedded $h'$ values was $[-0.4, 0.4]$ and the range of the embedded $J'$ values coupling different variables was $[-0.2, 0.2]$. This 20% value was determined empirically to provide good performance on the median difficulty problem at the largest sizes. Subsequently, the $J'$ values connecting physical qubits within a chain were set to the maximum ferromagnetic value ($−1$). A single random spin gauge transformation$^2$ was then applied to each embedded problem, with a gauge factor $a_i \in \{-1, 1\}$

Figure 6. Physical layout of the working qubits in the USC-LM D-Wave Two Vesuvius-6 processor as of October 10, 2014. 504 working qubits (gray), 8 non-working (black). All 1427 couplers connecting working qubits are part of the working graph.
associated with each qubit and transformation \( h'_i \rightarrow a, h'_i; J'_i \rightarrow a, J'_i \). One job was submitted to the quantum annealer per embedded problem; some Ising problems were associated with multiple embedded problems and jobs. After each programming cycle, the processor was allowed to thermalize for 10 ms (the maximum supported by the machine). The annealing time was set to the minimum value of 20\( \mu \)s. The number of annealing and readout cycles per programming cycle was 40000, which allowed the total job time to be within the limits of the machine (1 s). The readout thermalisation time was set to the default value of 0. Regarding error correction through majority voting of chains of physical qubits, ties were broken by choosing the spin up state. The probability of achieving the zero energy state on job \( k \) is denoted

\[
P_{0,k} = \frac{\text{number of annealing cycles achieving zero energy}}{\text{number of annealing cycles}}
\]

When multiple jobs are required, we calculate the geometric mean in the style of Boixo et al.:

\[
P_0 = 1 - \prod_{k=1}^{K} (1 - P_{0,k})^{1/K}.
\]

The total annealing time required to reach 0.99 probability of success was calculated by multiplying the annealing time by the expected number of annealing cycles (repetitions \( R \)) using the formula:

\[
R = \left\lfloor \frac{\ln(1 - 0.99)}{\ln(1 - P_0)} \right\rfloor.
\]

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Author Contributions
K.Z. and O.S. conceived the research; O.S. performed mathematical modeling and developed an algorithm for generating an Ising model; K.Z. designed and implemented the solvers, conducted experiments, and...
wrote the manuscript text; M.F. and K.Z. supervised the research; M.F. reviewed the manuscript and contributed revisions.

**Additional Information**

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