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Improved extremal optimization for the Ising spin glass

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A version of the extremal optimization (EO) algorithm introduced by Boettcher and Percus is tested on 2D and 3D spin glasses with Gaussian disorder. EO preferentially flips spins that are locally “unfit”; the variant introduced here reduces the probability to flip previously selected spins. Relative to EO, this adaptive algorithm finds exact ground states with a speed-up of order $10^9 (10^{10})$ for $16^2$ (8$^3$) spin samples. This speed-up increases rapidly with system size, making this heuristic a useful tool in the study of materials with quenched disorder.

Exploring the low temperature behavior of disordered materials, such as spin glasses and other random magnets [1], is quite challenging due to the very phenomena, glassy dynamics and multiple metastable states, that are important in such materials. Scaling arguments [2, 3] indicate that many properties of the glassy state, including the scaling of the energy of excitations and correlation functions, can be found by studying the ground state and its response to perturbations. Significant effort has been invested in identifying models whose ground states can be computed in time polynomial in the system size [4]. Where no polynomial-time algorithm is known, exact and heuristic methods which take exponential in system size are used. This enterprise is intimately connected with concepts developed in computer science, especially the distinction between P and NP-hard optimization problems [5].

The Ising spin glass (ISG) is a prototypical example of a disordered magnet. NP-hard problems such as the 3D ISG are, of course, particularly challenging. Exact methods for the 3D ISG with Gaussian bond weights can solve $12^3$-spin samples with open boundary conditions [7]. Such sizes have not proven to be sufficiently large to decide between alternate pictures for the low-temperature behavior. Heuristic genetic methods mix configurations and can therefore generate large scale “moves”; such methods are used for samples with $14^3$ spins for $\pm J$ couplings [8]. Heuristics with local moves generally have difficulty finding the exact ground state, due to the large barriers separating metastable states. Techniques such as flat histogram methods [9] can partially lower free energy barriers between metastable states.

In this Communication, I study a modified version of extremal optimization (EO) [10]. EO is a local search algorithm that preferentially flips spins with low “fitness”. The version presented here, “jaded” extremal optimization (JEO) increases the fitness of a spin by an amount proportional to the number of times it has been flipped. The goal of this adjustment is to reduce the repetition in exploring paths in configuration space, so that more possibilities can be quickly explored. Empirically, this simple change dramatically increases the effectiveness of the EO algorithm for finding ground states of two- and three-dimensional spin glass samples. As exact ground states are needed for studies of excitations and scaling, the algorithm is, for the most part, stringently tested by demanding that it find the ground states computed by exact methods. Both EO and JEO take time exponential in the system size to find the exact ground state, but the rate of growth is slower for JEO. Though JEO introduces an extra parameter, large improvements are achieved with only modest tuning.

I. EXTREMAL OPTIMIZATION AND EXTENDED ALGORITHM

A principle motivation for applying EO is to explore the energy landscape near the trial configuration by unconditionally modifying “unfit” variables. Preferentially (but not exclusively) changing variables with low fitness tends to raise the expected fitness while maintaining large fluctuations. The algorithm differs some from traditional Monte Carlo algorithms that conditionally select variables according to the expected improvement. In EO, the potential moves are selected according to their rank by fitness, rather than a Boltzmann distribution by weight.

A correspondence can be defined between fitness and the Hamiltonian for the Ising spin glass [10]. The Hamiltonian for spins $s_i$ indexed by position $i$, in a $d$-dimensional ISG of linear size $L$ is

$$H = -\sum_{ij} J_{ij} s_i s_j,$$  

where $J_{ij}$ are random bond strengths each chosen with probability $P(J_{ij}) = e^{-J_{ij}/2}/\sqrt{2\pi}$ for nearest neighbor spins with $1 \leq i, j \leq N = L^d$. When $d = 2$, algorithms with running times polynomial in $N$ are available [11] to find the ground state. When $d \geq 3$, finding the ground state energy is NP-hard, so that finding ground states for the worst-case choice of $J_{ij}$ is expected to take time exponential in $N$. In the context of EO, one choice for the fitness variable $\lambda_i$ for a spin variable $s_i$ is

$$\lambda_i = \lambda_i^0 \equiv s_i \left( \sum_{j \in U_i} J_{ij} s_j \right),$$  

where $U_i$ are the set of unsatisfied bonds ($s_i J_{ij} s_j < 0$) containing $s_i$. (Allowing for site-dependent constant shifts $\lambda_i \rightarrow \lambda_i^0 + \kappa_i$ as in Ref. [12] did not affect the comparisons here.) The configuration energy is related
to the fitness by $H = -\frac{1}{2} \sum_{i} \lambda_{i}^{0} + \sum_{ij} |J_{ij}|$. Any increase in the fitness decreases the total energy.

Given the fitness variables $\lambda_{i}^{0}$, there are a variety of strategies one could employ to attempt to improve the total fitness. The simplest version of EO takes “greedy” steps: the algorithm repeatedly flips the least fit variable until a static state is achieved. The greedy method converges quite rapidly, but in a spin glass the convergence is to a local minimum that is generally quite far from the optimal solution, both in configuration of the $\{s_{i}\}$ and often in energy per degree of freedom $H/N$. Similar greedy approaches for decision problems such as SAT, which seeks truth assignments for Boolean formula so that all clauses contain a true value, can be quite successful for given ensembles of problems [13].

An improved method, $\tau$-EO [10], sorts the spins by $\lambda_{i}$ and chooses the $m$th spin in the list with probability proportional to $m^{-\tau}$. This favors the choice of spins with low fitness, but allows for the occasional choice of sites with very high fitness. Fluctuations arising from the stochastic choice among spins with low fitness and the ranking of spins by the total weight of broken bonds, rather than energy improvement, allow the search to escape metastable states. It is argued [10] that for large systems, the optimal choice of $\tau$ approaches $\tau = 1$.

The extension considered in this paper (JE-EO) adjusts the fitness by an amount proportional to the number of times $k_{i}$ that a site $i$ has been previously chosen, that is,

$$\lambda_{i} = \lambda_{i}^{0} + \Gamma k_{i},$$

where $\Gamma$ is a site-independent “aging” parameter. The variables are sorted by $\lambda_{i}^{0}$ and then selected by rank as in $\tau$-EO. The $\tau$-EO algorithm corresponds to the choice $\Gamma = 0$. Setting $\Gamma \neq 0$ reduces the probability of selecting moves that have been flipped many times before. For configurations near (or in) the ground state, it is favorable for some spins to have low fitness, in order that a number of other spins can maximize their fitness. When $\Gamma = 0$, these spins, which are actually in their ground state orientation relative to the other spins, will be flipped in utility. Shifting the $\lambda_{i}$ during the algorithm also breaks the finite set of offsets between fitnesses of distinct spins that exist at $\Gamma = 0$ (due to the finite number of bond configurations at each site). This adaptive scheme has similarities to a variety of methods for solving problems such as SAT (satisfiability of sets of logical constraints) that disfavor repeated selection of the same move, such as Novelty [14] and variants of WALKSAT and GSAT [13, 16]. In contrast with these other schemes, the selection process in JEO is combined with the power law distribution for selecting ranked moves. Spin glasses with continuous disorder differ from SAT problems as they have less local degeneracy but also possess a global up-down symmetry, so that distinct methods may be appropriate.

In order to select spins quickly, I used the approximate selection method described in Ref. [12]. The spins are stored in a heap structure [14] according to their current fitness. This structure is a tree that is relatively cheap to maintain ($O(\log N)$ total cost to select a spin and update the tree). Each spin has a parent (except for the root) and at most two children. Each child is more fit than its parent and the root of the tree contains the least fit spin. This structure does not guarantee any other interlevel sorting, so that a spin $i$ that is deeper in the tree than, but not a direct descendant of, a given spin $i'$, may have a lower fitness. The heap structure does maintain a useful approximate sorting, though. To select a spin to flip, a level $\ell$ is selected with probability proportional to $2^{-\ell(\tau-1)/\tau}$ and then a random spin within level $\ell$ is chosen. The spin at this site is then inverted. The fitness of the neighboring spins is adjusted and the heap is updated using standard methods [17].

EO does not take advantage of the special structure of the 2D problem: it is not necessary or even expected that it will find the solution in time polynomial in the system size. Polynomial-time solvable problems have been used to study algorithms, for example, for hard mean-field problems [18]. For some classes of problems, heuristics can find solutions in polynomial time [13, 19]. In the 2DISG, large low-energy excitations may make local algorithms especially inefficient.

II. PERFORMANCE OF THE ALGORITHM

In this section, I compare the performance of the extended EO algorithm, JEO, against $\tau$-EO as applied Ising spin glasses with Gaussian disorder. When feasible, comparisons with ground states found using exact methods provide a precise and direct test for convergence.

Two-dimensional spin glass. The 2DISG models are on a square lattice with $L^{2}$ spins and open boundary conditions. To determine the 2D ground state, each sample is mapped to a general weighted matching problem. The matching problem for a graph is to find a set of edges with minimal total weight such that each vertex belongs to exactly one edge. The weighted graph for a 2DISG sample has edges dual to the lattice bonds, with weight $|J_{ij}|$ for an edge that crosses a bond with weight $J_{ij}$, and extra edges of weight zero that ensure that the frustration of each plaquette is maintained: unfrustrated (frustrated) plaquettes give an even (odd) number of the bonds dual to the edges of the plaquette in the matching. To find the minimum weight matching and hence the ground state energy for a 2DISG sample, I used the Blossom IV algorithm developed by Cook and Rohe [20].

The exact ground state energy of each 2DISG sample was input to the $\tau$-EO and JEO codes. When the heuristic codes found this energy, the codes terminated. The primary results from these computations were the distributions of the running times, measured in number of spin flips, to find the true ground state. The time to solution is a function of both the seed used to generate the sample and an independent “algorithm seed” used to generate the random initial configuration and to select spin flips. In a given sample, the distribution of times to find a ground
state was roughly Poissonian. This suggests that restarting the algorithm with different initial configurations or seeds for selecting flips does not significantly decrease the mean running time. This conclusion was consistent with empirical trials of restarting the algorithm: the algorithm does not get stuck in history dependent traps. Given a sample $k$, the median $t_{m}^k$ of the running time was estimated from the solution time for 100 algorithm seeds. The results reported here are for $t_{m}$, the sample mean of $t_{m}^k$. The $\Gamma = 0$ data is in agreement with previously results for $\tau$-EO, with $t_{m}$ minimal at $\tau \approx 1.5$.

The results for the mean solution time $t_{m}$ for optimal $\tau$ and $\Gamma$ are summarized in Fig. 1. As suggested by the data plotted in Fig. 1, $t_{m}$ is not very sensitive to the exact choice of parameters, as long as $\tau$ is in the range $1.5 < \tau < 2.5$ and the optimal $\Gamma$ (on the order of $10^{-3}$ to $10^{-1}$) is found to within a factor of about 2, for the sizes studied here. The best running times for $\tau$-EO grow much more rapidly than those for JEO. For $L = 16$, JEO is of the order $10^4$ times faster than $\tau$-EO. Extrapolation suggests that the advantage of JEO increases significantly with $L$. For comparison, an exponential dependence $t_{m} = 15 \cdot 2^L$ is shown in Fig. 1. This function does a good job of describing the JEO data for $L = 4$ through $L = 32$.

In separate runs, for comparison, the heuristic algorithm was terminated when the energy was within 1% of the exact ground state energy. These approximate solutions were found much more rapidly than exact solutions ($\approx 10^5$ times faster for $L = 32$).

Three-dimensional spin glass. A similar comparison was carried out for 3DISG samples with Gaussian disorder. The $L^3$ spins in the 3DISG samples lie on a cubic lattice with periodic boundary conditions. For 3DISG samples of size up to $6^3$, the spin glass server at the University of Köln [21] (which applies branch-and-cut [3]) was used to generate exact solutions. The termination condition of the algorithm was modified, as exact ground states for the larger samples were not readily available. All samples were simulated in parallel with $n = 10$ algorithm seeds. When the minimal record energy for eight (8) of the samples were identical, the algorithm was terminated. This criterion produced configurations equal to the exact solutions for all $L = 4, 6$ samples (45 at each size). This suggests that true ground states were found with a high probability for $L = 8$ and possibly also $L = 10$. The summary results are plotted in Fig. 2. Given the termination criterion, JEO was of the order of $10^2$ times faster than $\tau$-EO in converging to a potential solution for $L = 8$ samples. Very roughly, $L = 6$ samples were solved in $\approx 10$s on average both on the Köln spin glass server (a 400 MHz Sun Ultra) and using JEO (on a 1 GHz Intel P5). Further studies would be needed to provide better estimates of the confidence in the ground states and how to improve such confidence.

III. DISCUSSION

JEO extends the extremal optimization algorithm of Boettcher and Percus by adaptively reducing the frequency of flipping previously selected spins. As a local move can lead to avalanche-like behavior, due to induced changes in the fitness of neighbors, this modification also reduces the frequency of flipping larger domains. This extension of EO does add a parameter, the aging parameter $\Gamma$. However, a near-optimal value for $\Gamma$ for each
problem type at a given size can be found quickly and less tuning of the parameter $\tau$ is required than for $\tau$-EO.

One possible avenue of exploration is to check whether avalanche regions correspond to important domains or excitations in the sample. Possible modifications of JEO include using a selection distribution with sharp cutoffs \cite{23}, rather than power-law distributions. Other schemes for reducing the fitness of frequently repeated moves could be considered, such as modifying the fitness using non-linear functions of the number of flips at a spin.

Regardless of the exact details of the role of domains and possible improvements, empirical testing shows that the aging of the spins during state-space exploration greatly reduces the time for EO to find the ground state of the ISG in two and three dimensions. Though the 2D model was used to make a precise comparison with exact results, the exponential equilibration times for the 2DISG using extremal optimization are consistent with those that would be seen for an NP-hard optimization problem with a similar local solution strategy. It may be useful to use an algorithm like JEO to locally improve the configurations formed by whole sample crossover in genetic algorithms \cite{23}. As exact solutions for small samples can be found with confidence in a relatively small number of steps, in machine time very similar to that for branch-and-cut, this simple algorithm also provides a very convenient way to study small 3D samples.

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