Optimization by Record Dynamics.

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Abstract
Large dynamical changes in thermalizing glassy systems are triggered by trajectories crossing record sized barriers, a behavior revealing the presence of a hierarchical structure in configuration space. The observation is here turned into a novel local search optimization algorithm dubbed Record Dynamics Optimization, or RDO. RDO uses the Metropolis rule to accept or reject candidate solutions depending on the value of a parameter akin to the temperature, and minimizes the cost function of the problem at hand through cycles where its ‘temperature’ is raised and subsequently decreased in order to expediently generate record high (and low) values of the cost function. Below, RDO is introduced and then tested by searching the ground state of the Edwards-Anderson spin-glass model, in two and three spatial dimensions. A popular and highly efficient optimization algorithm, Parallel Tempering (PT) is applied to the same problem as a benchmark. RDO and PT turn out to produce solution of similar quality for similar numerical effort, but RDO is simpler to program and additionally yields geometrical information on the system’s configuration space which is of interest in many applications. In particular, the effectiveness of RDO strongly indicates the presence of the above mentioned hierarchically organized configuration space, with metastable regions indexed by the cost (or energy) the transition states connecting them.

1 Introduction
Built on analogies with physical or biological processes, heuristic optimization techniques are widely used in science [1, 2, 3, 4, 5, 6, 7]. Of present interest is Simulated Annealing (SA), a well known local search algorithm based on the Metropolis algorithm, which minimizes the cost
of candidate solutions in a way similar to a physical system minimizing its free energy under cooling \cite{8, 9}. In SA, a proposed solution is first generated by locally modifying the current solution. Changes lowering the cost are accepted and others are accepted with probability \( \exp(-\Delta E/T) \), where \( \Delta E > 0 \) is the additional cost incurred and where the parameter \( T \) is conventionally called temperature. Ideally, a cooling schedule gradually decreasing the temperature down to zero should reach the ground state, i.e. the desired solution of the optimization problem. However, in applications to hard combinatorial problems SA invariably gets stuck in one of the many suboptimal or metastable configurations which characterize these systems. Since available local configurational changes mainly get rejected, a larger partial randomization is required to obtain further improvements.

Large changes leading a thermalizing complex system from a metastable configuration to another are often triggered by thermal energy fluctuations of record magnitude \cite{10, 11, 12, 13}. It is then natural to hypothesize that visiting configurations of record-high cost, or energy, similarly helps a ‘thermal’ optimization algorithm of the SA type to escape suboptimal solutions.

The configuration space, or energy landscape, of the Edward-Anderson spin glass \cite{14} was previously investigated using Extremal Optimization \cite{7}, an optimization and exploration algorithm indifferent to energy barriers, and by the Waiting Time Method \cite{15}, a kinetic Monte Carlo algorithm with no rejections. The analysis led to the conclusion that, in order to achieve a lower BSF value, which is desirable in optimization, the barrier \( B(t) \) must previously reach a new high record. Importantly, this property is not associated to the algorithms used, but pertains to all energy landscapes which can be coarse-grained into inverted binary trees where nodes represent metastable configurations \cite{16} and height represents the energy. Motivated by the above considerations, the Record Dynamics Optimization (RDO) algorithm introduced below dynamically generates a non-monotonic SA schedule where heating and cooling phases alternate. Each heating phase terminates once a record high ‘barrier’ (defined below) is encountered and each cooling phase terminates once a state of record low cost is found. Möbius et al.\cite{17} earlier introduced a non monotonic annealing schedule where temperature oscillations are controlled by a tunable parameter instead of being determined by intrinsic geometrical properties of the landscape.

For demonstration purposes, RDO is used to search for the ground state of a three dimensional Edwards-Anderson (EA) spin glass \cite{14}, a standard NP hard optimization problem. For completeness, it is further applied to the two dimensional EA model. RDO performance is then compared to that of a carefully optimized version of Parallel Tempering (PT). The numerical effort needed to obtain results of compa-
rable quality is similar for the two methods. However, RDO has fewer tunable parameters and is more easily implemented. Secondly, RDO provides, at no extra cost, some information on configuration space structure which might be of interest in landscape explorations.

2 The RDO algorithm

First some notation: A sweep in a MC run comprises a number of elementary moves or queries, i.e. the generation and acceptance or rejection of a candidate move, equal to the number of independent variables of the problem. The number of sweeps carried out up to a certain point is dubbed time and denoted by the symbol $t$. Each query generates a putative solution or state, and the ordered sequence of states sampled in $[0, t]$ is called a trajectory. The cost associated to a state is called its energy $E$. The Best So Far energy, $\text{BSF}(t)$, is the lowest energy sampled in a single trajectory in $[0, t]$. The barrier $B(t)$ associated to a state sampled at time $t$ is $B(t) \overset{\text{def}}{=} E(t) - \text{BSF}(t)$. Lower case symbols are used for quantities scaled by system size, i.e. in the example considered $b(t)$ is the barrier energy per spin. We stress that the BSF and barrier functions are stochastic processes and that inherent geometrical properties of the landscape can only be estimated by averaging over a suitably large ensemble of independent trajectories.

The RDO algorithm comprises an initial phase followed by a succession of cooling and a heating phases controlled by record events. Each of these phases involves decreasing or increasing the temperature within a set of 22 predefined and equidistant values in the temperature range $[T_{\text{MIN}}, T_{\text{MAX}}]$. Several preliminary simulations showed that $T_{\text{MAX}} = 1.2$, slightly above the critical temperature of the 3d model is a good choice. Furthermore, $T_{\text{MIN}} = 0.3$ was chosen as BSF values are rarely, if at all, found below $T = 0.3$. We let the system cool and heat $ad libitum$ since each cooling or heating phase produces gradually lower extremal values. Once the minimal temperature is reached, and no further BSF is found, the algorithm stops.

1. Initialization of BSF and barrier values: Any short naive optimization at a constant temperature typically slightly below the critical temperature $T_g$ will produce the first BSF value, $\text{BSF}_0$. The first high barrier value $b_0 > \text{BSF}_0$ is found by running the algorithm at a slightly higher constant temperature. For $i = 1, 2, \ldots$, the ‘barrier’ $B(t) = E(t) - \text{BSF}_i$ is used to control the algorithm. The highest barrier overcome in heating phase $i$ is called $B_i$.

2. Cooling: Let $S_{B,i}$ be the configuration corresponding to $B_i$. Starting from $S_{B,i}$ run SA with decreasing temperature until a lower
BSF value is found. If no lower BSF is found, cooling stops after $N_{\text{step}} = 50000$ sweeps.

3. Running at constant $T$: the Metropolis algorithm at constant $T$ is used until either $m$ new BSF values have been found or the preset max time is exceeded. In practice $m$ is a small integer, i.e. $m = 3$ in the present simulations. This step ensures that once the correct region of configuration space is identified, some time is spent exploring it. BSF$_{i+1}$ is the lowest BSF value identified during this phase.

4. Heating: starting from $S_{i+1}$, the configuration corresponding to BSF$_{i+1}$, heat the system until $B(t) = E(t) - \text{BSF}_{i+1} > B_i$. The achieved record value of $B(t)$ defines $B_{i+1}$.

5. Set $i+1 \rightarrow i$, go to step 2 and repeat ad libitum.

3 Parallel tempering

Parallel Tempering (PT) avoids trapping by independently searching a number $N_T$ of identical replicas of the problem at hand. The $m$’th replica is explored by a conventional Metropolis algorithm run at a temperature $T_m$. Additional configurational swaps between replicas, also controlled by the Metropolis criterion in order to ensure detailed balance, provide the sought escape route from suboptimality. A successful PT implementation requires consideration of the temperatures at which the replicas are run and a compromise between the number of attempted swaps and the number of standard queries within the replicas. The reader is referred to [18, 19, 20] for a in-depth discussion of PT. The brief summary provided below describes the implementation presently used to benchmark RDO.

1. $N_T$ different copies of the system are updated in parallel at temperatures $T_m > T_{m+1}$, $m = 1...N_T$ through one or more Monte Carlo sweeps.

2. A proposed swap between configuration $C_m$ and $C_{m+1}$ is accepted or rejected according to the Metropolis criterion. Defining $\beta_m = 1/T_m$, and

$$
\Delta S = \left[ \beta_{m+1} E(C_m) + \beta_m E(C_{m+1}) \right] - \left[ \beta_m E(C_m) + \beta_{m+1} E(C_{m+1}) \right],
$$

the exchange is accepted with probability $\min(1, e^{-\Delta S})$.

3. Further exchanges between the configurations associated with $\beta_{m+1}$ and $\beta_{m+2}$ are accepted or rejected in the same way, eventually exploring the whole set of temperatures.
4. Go to step 1 and repeat *ad libitum*.

After a number of exploratory simulations, the highest temperature was chosen as \( T_{\text{max}} = 1.6 \), a value higher than the critical temperature of the Edwards-Anderson spin glass i.e. \( T_c \approx 0.95 \) [21]. The lowest temperature is dynamically determined as discussed below. A suitable number of temperatures for PT is generally estimated to be \( N_T \approx \sqrt{N_{\text{spin}}} \) [19]. In the following, \( N_T = 30, 50 \) and 90 are used for \( L = 30, 50 \) and 100 in the 2d simulations, while \( N_T = 30, 40 \) and 80 are used in the 3d case for \( L = 8, 14 \) and 20.

To accept an exchange between copies with probability \( \approx 0.5 \), a value considered to be optimal [20], the \( T_m \) values are treated as dynamical variables using the recursive method described in Ref. [19]. Initially, the inverse temperatures \( \beta_m \) are set to

\[
\beta_m = \beta_1 + (\beta_M - \beta_1) \frac{m-1}{M-1} \tag{2}
\]

with \( M = N_T \). The updated set \( \{ \beta'_m \} \) is obtained using the sampled exchange rates \( p_m \) between configurations at inverse temperatures \( \beta_m \) and \( \beta_{m-1} \):

\[
\beta'_1 = \beta_1 \\
\beta'_m = \beta'_{m-1} + (\beta_m - \beta_{m-1}) \frac{p_m}{c} \text{ with } m = 2, ..., M
\]

\[
c = \frac{1}{M-1} \sum_{m=2}^{M} p_m \tag{3}
\]

While in Ref. [19] temperatures are only updated initially to reach the constant values used in the simulation, we found it more convenient to update them during the simulation itself, at logarithmically equidistant times \( 2^n \times 100 \) MC sweeps, with \( n = 1, 2, ..., N \).

Two different benchmarks for RDO are provided. The first, our ‘fast’ PT, has \( N = 10 \) and \( N_{\text{step}} = 102400 \) sweeps per replica. Adding the computational effort for all replicas, PT is eight time faster than RDO but produces results of somewhat lesser quality. The second version, ‘slow’ PT, has \( N = 13 \) and \( N_{\text{step}} = 819200 \), with the total number of sweeps approximately corresponding to that used in our RDO implementation. Both versions of the PT algorithm include a final quench to \( T = 0 \), a step omitted in RDO. Importantly, the PT versions implemented are carefully optimized and based on the recent literature on the subject.
4 Results

The model

The EA model[14] with Gaussian interactions deals with a set of spins $\sigma_i = \pm 1$ which are placed on a d-dimensional discrete grid with linear size $L$ and periodic boundary conditions. The spins interact via a coupling matrix $J$ which is symmetric, has diagonal elements all equal zero and off-diagonal elements $J_{ij}$ likewise equal zero, unless spins $i$ and $j$ reside on neighboring grid points. In this case, and for $i < j$, their values are independently drawn from a Gaussian distribution with zero average and unit variance. The energy of a configuration $\alpha$ is then given by

$$E^\alpha = \frac{1}{2} \sum_{i,j} J_{ij} \sigma_i^\alpha \sigma_j^\alpha. \quad (4)$$

Determining the energy of the ground state of the 3d Edwards-Anderson spin glass problem in its different guises is an NP hard combinatorial problem which has been attacked using a variety of techniques. For future reference we note that Pal [22] combined a genetic algorithm with local search and found that the ground state energy per spin is $e_{gs} = -1.699926 + 2.1373L^{-3}$, where the first term is the thermodynamic limit and the second term describes finite size corrections. More recently, Romá et al. [23] used Parallel Tempering and considered three different finite size corrections to the thermodynamic limit, one of which is $e_{gs} = -1.7000 + 2.01L^{-2.94}$. The same authors find $e_{gs} = -1.3149 + 1.3L^{-2.28}$ for the 2d system.

In this work, the EA model is used to test how RDO works and to benchmark it against PT. As there is no ambition to improve on existing estimates of the model’s ground state energy, we limit ourselves to three different system sizes in both 2d and in 3d, and offer no analysis of finite size corrections. Most results are averages over $N_{\text{sample}}$ of independent realizations of $J$ and, unless otherwise specified, the combination $N_{\text{sample}} = 200, 50$ and 10 is either used for 2d systems of linear size $L = 30, 50$ and 100 or for 3d systems of linear size $L = 8, 14$ and 20. Since the RDO algorithm produces trajectories with varying number of steps and with BSF($t$) values obtained at different times, coarse-graining each trajectory is required to average different trajectories. Accordingly, within each trajectory, the measured values were averaged every 750000 steps, as long as possible. This choice fits the worst case encountered for every size studied, i.e. provides a partition of the trajectory with the slowest decay of the BSF energy. Faster trajectories were padded with the last BSF energy value achieved. In this way, the computational time used by the RDO algorithm to achieve a certain averaged BSF energy value is slightly overestimated.
Temperature, barriers and energies in RDO

Since alternating heating and cooling phases are characteristic features of the algorithm, the time dependence of the temperature and the related time dependence of the record barriers which control the switch from heating to cooling are discussed first. Second, we discuss the time dependence of the BSF energy per spin, which, at each stage of the calculation, provides the RDO estimate of the ground state energy. Finally, the randomizing effect of a temperature cycle is discussed in terms of Hamming distances.

The left panel of Fig. 1 depicts the average barrier value per spin, \( b(t) \), with the initial value subtracted. The trend is in all cases logarithmic, but the slope is lesser in 3d than in 2d. The data collapse obtained for sufficiently large sizes shows that barriers are extensive, in contrast to barriers reached in an isothermal relaxation process, which are sub-extensive\(^2\). Hence, heating the system up is a far more efficient way to partial configurational randomization than isothermal relaxation.

In the right panel of Fig. 1, the algorithm’s temperature \( T \) is plotted vs. time for a single 2d system with \( L = 100 \). Similar behavior is observed for all other systems considered. In the curve, each local minimum corresponds to the temperature \( T_{\text{min}}(t) \) for which a new BSF energy value is found at stage 3 of the RDO algorithm, while each local maximum corresponds to a temperature \( T_{\text{max}}(t) \) reached at stage 4, for which a new record sized barrier is found. The upper and lower envelopes of the curves are least square fits of the form \( T_{\text{min}}(t) = T(t_0) - a_{\text{min}} \sqrt{t} \) and \( T_{\text{max}}(t) = T(t_0) - a_{\text{max}} \log(t) \), where \( a_{\text{min}} \) and \( a_{\text{max}} \) are numerical constants. We first note that even though the barrier values increase in time, the (un-shifted) energy of the corresponding barrier states decreases, i.e. the RDO algorithm explores regions of configuration space of gradually lower energy. The decreasing trend of \( T_{\text{max}}(t) \) matches the logarithmic decrease of the energy of the different barrier states, see Fig. 2. The square root term in \( T_{\text{min}}(t) \) indicates that the BSF energy minima belonging to different regions of the landscape decrease in a roughly linear fashion from one region to the next and are reached via the diffusion-like process associated to the constant temperature search in the third phase of the RDO algorithm.

All curves show a logarithmic dependence on time. The common asymptotic limit of the curves, which corresponds to the predicted ground state energy value is, as expected, nearly independent of system size and in agreement with current numerical estimates \(^2\). In contrast, the plot of the local energy maxima along trajectories which is depicted Fig. 3 shows that \( e_{\text{max}}(t) = e(t) + b(t) \) retains a system size dependence throughout the simulation. A glance to Fig. 1 shows that the latter mainly stems from the system size dependence of the initial
Figure 1: (Color online) Left: the average barrier per spin $b(t)$ with the initial value subtracted and for all the systems considered, is plotted versus time on a logarithmic horizontal scale. Right: the temperature $T(t)$ vs. time for a single trajectory of a 2d system of linear size $L = 100$. The lower and upper lines show fits to the local minima and maxima $T_{\text{min}}(t)$ and $T_{\text{max}}(t)$, respectively.

Figure 2: (Color online) The disorder averaged BSF energy per spin $e$ is plotted vs. time for 2d (left panel) and 3d systems using a logarithmic horizontal scale.
barrier value. We stress that both $e_{\text{max}}$ and $e(t)$ decrease logarithmically in time while the barrier $b(t)$ correspondingly increases.

The Hamming distance between configurations $\alpha$ and $\beta$ is $H(\alpha, \beta) = 1 - \frac{1}{N_{\text{spin}}} \sum_i \sigma_i^\alpha \sigma_i^\beta$. In the left panel of Fig. 4 $\alpha$ denotes the configuration corresponding to the ‘current’ BSF energy value, and $\beta$ is its immediate predecessor along a trajectory. In the right panel of the same figure, $\alpha$ is the initial configuration and $\beta$ the current one. In other words, the left panel describes the configurational change occurring in a single thermal cycle, while the right panel describes the change accumulated from the beginning of the evolution of the system. All data are disorder averaged as earlier explained. In the 3d systems, $H_c$, the Hamming distance between consecutive minima, has a clear decreasing trend, interspersed by some oscillations. Hence the randomizing effect of crossing a record-sized barrier gradually tapers off, which is a desirable property in an optimization setting. The 2d case has much more pronounced oscillations, some of which correspond to system size configurational changes.

The right panel shows that, as expected, the Hamming distance of the current minimum to the initial configuration nearly remains constant in time. This constant is near one in the 2d case, implying that low energy configurations successively identified are all nearly orthogonal to the initial configuration. In the 3d case the constant is much smaller, indicating a higher degree of persistent correlation and a lingering memory of the past. In Fig. 4 the disorder averaged BSF energy per spin, $e$, is plotted vs. time on a logarithmic horizontal scale for 2d (left panel) and 3d systems.
4.1 Comparing RDO and PT

The RDO and PT algorithms are compared in terms of the disorder averaged BSF energy per spin, respectively average energy per spin as a function of temperature obtained using the two methods. To simplify the notation, the same symbol $e$ is used for both quantities, both converging for large times to the desired ground state energy, the target of the search.

We considered two types of comparison: in the first, we use a ‘fast’ PT, where minimizing the execution time is a priority, but where the results are in some case of lesser quality than those obtained by RDO. In the second, we use a ‘slow’ version of PT algorithm, with parameters tuned to obtain better, i.e. lower, energy values. As mentioned, slow PT requires the same computational effort as RDO, while fast PT is 8 time faster than RDO. All our results lie, as expected, slightly above the thermodynamic limit of the model’s ground state energy, see the previous model discussion and the original references [22, 23].

In the left panel of Fig. 4 the average BSF energy per spin $e$ of the 2D system is plotted as a function of the temperature $T$. As explained in the figure text, the RDO and fast PT results are plotted for different system sizes. Interestingly, the average energy calculated by the RDO is consistently lower than its PT counterpart.

Also note that BSF energy obtained in the RDO at $T = 0.3$ is comparable or, in the case of the larger systems, even lower than the average energy obtained in PT at a lower temperature. On the right panel we compare RDO with our slow PT. Here PT finds lower energies...
Figure 5: (Color online) Left: the BSF energy, respectively average energy per spin $e$ vs. temperature $T$ for all 2d systems considered. Data obtained using RDO and fast PT, respectively. Right: the same RDO results compared with those of slow PT.

than RDO, but the difference is hardly significant.

Figure 6 is the 3-d analog Fig. 5 and similar observations apply to the trends and minima. Furthermore, the BSF energy given by RDO are almost coincident with the ground states at $T = 0$ given by slow PT. Table 1 summarizes the estimated ground state energy

|       | RDO          | Fast PT      | Slow PT      |
|-------|--------------|--------------|--------------|
| $L=30$ | -1.30824±0.00167 | -1.31306±0.00166 | -1.31501±0.00166 |
| $L=50$ | -1.30380±0.00202 | -1.30632±0.00200 | -1.30981±0.00202 |
| $L=100$ | -1.30403±0.00214 | -1.30441±0.00212 | -1.30815±0.00217 |
| $L=8$  | -1.69472±0.00236   | -1.69728±0.00236   | -1.69733±0.00276   |
| $L=14$ | -1.69114±0.00194   | -1.69416±0.00191   | -1.69597±0.00188   |
| $L=20$ | -1.69549±0.00194   | -1.69426±0.00214   | -1.69793±0.00215   |

Table 1: Ground-state energy estimates (energy per spin) obtained using RDO and the fast and slow PT algorithm. The first three rows pertain to 2d systems, and the last three to 3d systems. The errors are given as ±σ, where σ is the estimated standard error on the computed averages.

values obtained for all the systems considered using RDO and our fast and slow PT algorithm. The values quoted are ensemble averages and their standard (1σ) errors. The two methods very nearly produce results with overlapping error bars. The fast PT algorithm results for the largest systems are marginally inferior to the corresponding RDO results, while the slow PT results are marginally superior.
5 Discussion

Similarly to Simulated Annealing (SA), Record Dynamics Optimization is a ‘thermal’ optimization heuristics based on local search and on the Metropolis acceptance rule. Unlike SA, it features an alternation of heating and cooling phases, each delimited by the achievement of a record high ‘barrier energy’, and a lower Best So Far (BSF) energy, respectively. The current BSF energy provides an estimate of the solution of the optimization problem at hand. The physical idea behind RDO is that the configuration space of hard optimization problems explored by local searches can be coarse-grained into a hierarchy of nested sets. Starting from a poor solution, configurations of decreasing cost can only be accessed by scaling increasingly large barriers. Hence, quickly generating record high barriers provides an effective way to achieve better solutions. For a given application, the validity of a hierarchical description is buttressed whenever RDO works efficiently. In this way, which makes RDO into a landscape optimization tool.

RDO has a modicum of adjustable parameters. Most important are the cooling/heating rate, and the number of BSF energy values found when (briefly) searching at constant temperature following a cooling phase. The programming effort in RDO is similar to standard SA and considerably smaller than in PT. On our test problem, RDO seems to deliver marginally higher energy values than the slow PT algorithm on the largest systems considered. However, PT is a highly optimized algorithm with a long history of successes, while RDO is a new algorithm, which, we surmise, still has considerable potential for further improvements.
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