Interpolating greedy and reluctant algorithms

Pierluigi Contucci\textsuperscript{a}, Cristian Giardinà\textsuperscript{a}, Claudio Giberti\textsuperscript{b}, Francesco Unguendoli\textsuperscript{c} and Cecilia Vernia\textsuperscript{c}

\textsuperscript{a} Dipartimento di Matematica, Università di Bologna
Piazza di Porta S.Donato 5, 40127 Bologna, Italy
\{contucci, giardina\}@dm.unibo.it

\textsuperscript{b} Dipartimento di Informatica e Comunicazione, Università dell’Insubria,
via Mazzini 5, 21100 Varese, Italy
claudio.giberti@uninsubria.it

\textsuperscript{c} Dipartimento di Matematica Pura ed Applicata, Università di Modena
e Reggio Emilia, via Campi 213/B, 41100 Modena, Italy
\{unguendoli, vernia\}@unimore.it

Abstract

In a standard NP-complete optimization problem we introduce an interpolating algorithm between the quick decrease along the gradient (greedy dynamics) and a slow decrease close to the level curves (reluctant dynamics). We find that for a fixed elapsed computer time the best performance of the optimization is reached at a special value of the interpolation parameter, considerably improving the results of the pure cases greedy and reluctant.

1 Introduction and results

Combinatorial optimization stands as one of the most fruitful fields in the intersection of applied and pure mathematics: it connects the theory of the computational hardness to the techniques widely used in the search for global minima for complex functionals, i.e. functions with many local minima. Over the years there have been many strategies proposed to solve efficiently hard computational problems \[\square \square \square \]. Among them the
statistical mechanics approach [4, 5] has opened new interesting perspectives. In this paper we study the interpolation between two algorithms, the greedy and the reluctant. The first is the standard decrease along the deepest descent direction, while the second is the closest decrease to the level lines. In previous works [6, 7] we have studied and compared the two algorithms focusing on relaxation time and minimum reached level. We observed, moreover, how a simple convex interpolation between them could improve the performances of both of them in large size regime. In this work we push further such analysis introducing a smooth interpolation between greedy and reluctant depending on a parameter $\lambda$: small $\lambda$ plays the role of the greedy algorithm, while large $\lambda$ that of reluctant. This allows us a better tuning among the two and especially a parameter optimization. The newly introduced algorithm is tested against a model which has become the standard of NP complete problems: the Sherrington Kirkpatrick model of the mean field spin glass. Our results confirm and extend those in [6, 7]: we find that the relaxation time grows linearly when the algorithm is close to the greedy regime and quadratically when it is close to the reluctant one with a progressive condensation for large values of $\lambda$. The dynamics is then tested in the search for low energy configurations for fixed values of initial conditions, where the reluctant dynamics works substantially better than any other. The main result of this work is then the minimization at fixed elapsed computer time. In this case, in fact, we find that in the small size regime (compared to total search time) the greedy component performs better than any other due the short relaxation time and the fact that, basically, the dynamics is able to find the ground state or, at least, to get very close to it. Moreover and more interestingly, we find that increasing the system size does not lead to a uniform deteriorating of the greedy performance toward an improvement of the reluctant one. We find, in fact, an optimal value of $\lambda \sim 10$, for which the lowest energies are reached against a rather poor performance of the greedy ($\lambda \sim 1$) and reluctant algorithm ($\lambda \sim 100$). This optimal value appears to be independent of the size.

2 The model and the algorithm

Let us consider the Sherrington-Kirkpatrick model [8] defined by the Hamiltonian

$$H(J, \sigma) = -\frac{1}{2} \sum_{i,j=1}^{N} J_{ij} \sigma_i \sigma_j$$

(1)
where $\sigma_i = \pm 1$ for $i = 1, \ldots, N$ are spin variables which interact through an $N \times N$ symmetric matrix with $J_{ij}$ independent, identically distributed gaussian random couplings ($J_{ij} = J_{ji}, J_{ii} = 0$) with zero mean and variance $1/N$. We focus our attention on a stochastic energy-decreasing dynamics that, starting from any initial spin configuration at time $t = 0$ (which we choose at random with uniform distribution), ends up on a local energy minimum. The evolution rule is:

1. Let $\sigma(t) = (\sigma_1(t), \ldots, \sigma_N(t))$ be the spin configuration at time $t$.

2. Calculate the spectrum of energy change obtained by flipping the spin in position $i$, for $i = 1, \ldots, N$:

$$\Delta E_i = \sigma_i(t) \sum_{j \neq i} J_{ij} \sigma_j(t). \quad (2)$$

If $\Delta E_i > 0$, $\forall i = 1, \ldots, N$, then the algorithm stops ($\sigma(t)$ is a local minimum).

3. Generate a random number $D$ with probability density

$$f(x) = \begin{cases} \lambda e^{\lambda x} & \text{if } x \leq 0 \\ 0 & \text{if } x > 0 \end{cases}, \quad \lambda > 0. \quad (3)$$

4. Select the site $i^*$ associated with the closest energy change to the value $D$, i.e.:

$$i^* = \left\{ i \in \{1, \ldots, N\} : \Delta E_{i^*} = \min_{i \in \{1, \ldots, N\}} \{|\Delta E_i - D| : \Delta E_i < 0\} \right\}. \quad (4)$$

5. Flip the spin on site $i^*$:

$$\sigma_i(t + 1) = \begin{cases} -\sigma_i(t) & \text{if } i = i^* \\ \sigma_i(t) & \text{if } i \neq i^*. \end{cases} \quad (5)$$

This algorithm generates a dynamics that, following a 1-spin flip decreasing energy trajectory, arrives at a 1-spin flip stable configuration, that is a configuration whose energy cannot be decreased by a single spin-flip. The speed of convergence to local energy minima is tuned by $\lambda$, the control parameter in the probability distribution function for the move acceptance. Of course, the larger is $\lambda$, the bigger it is the probability of doing small energy-decreasing steps, so that the trajectory will follow an evolution path.
close to level curves (reluctant). On the other hand, small values of $\lambda$ enrich the probability of large negative energy steps, which will quickly drive the dynamic to the end-point (greedy). In the next Section, by varying the control parameter $\lambda$, we study the efficiency of the algorithm by measuring the average time to reach a metastable configuration and the lowest energy value found as the system size is increased.

3 Results

We performed a set of experiments for different values of $N$, starting from $N$ initial conditions (for a system of size $N$) and averaging the data on $nreal = 1000$ disorder realizations. We probed basically two quantities to measure the performance of the algorithm:

- the average time (i.e. the number of spin flips) to reach a minimum energy level

$$ \tau = \frac{1}{M} \sum_{i=1}^{M} t_i, $$

(6)

with $M = N \cdot nreal$ and $t_i, i = 1, \ldots, M$ the time for each initial condition

- the lowest energy found (averaged over disorder)

$$ H_N = \left\langle \frac{\min_{\sigma} H_N(J, \sigma)}{N} \right\rangle_{nreal}, $$

(7)

where $\min_{\sigma} H_N(J, \sigma)$ is the minimum value of the energy of the metastable states attained starting from $N$ initial conditions.

In Fig.1 we represent $\tau$ as a function of $N$ in the range $[25, 300]$ for six distinct values of $\lambda$ ($\lambda = 1, 10, 25, 45, 70, 100$), together with the best numerical fits. Because of high computational costs (which increase with $\lambda$), the case $N = 300$ is studied in details only for $\lambda = 1$ and $\lambda = 10$. On the other hand, the average time has good self-averaging properties so that, in order to have the trend for “large” $\lambda$ we focused on the case $N = 300, \lambda = 100$ with $nreal = 140$ disorder realizations instead of $nreal = 1000$. From Fig.1 we observe the progressive increase of the slope in log-log scale from an almost linear law in $N$ for $\lambda = 1$ (○) to an almost quadratic one for $\lambda = 100$ (∗). Thus, this algorithm behaves as a “smooth interpolation” between the two deterministic dynamics: “greedy”, that we obtain for $\lambda = 1$, and “reluctant”, here represented by $\lambda = 100$. More in detail, the numerical fits of
Figure 1: Average time $\tau$ to reach a metastable configuration as a function of $N$ for $\lambda = 1$ (⋄), $\lambda = 10$ (+), $\lambda = 25$ (□), $\lambda = 45$ ($\times$), $\lambda = 70$ (△), $\lambda = 100$ (∗).

Fig.1 are: $\tau_{\lambda=1}(N) \sim N^{1.027}$ (⋄), $\tau_{\lambda=10}(N) \sim N^{1.263}$ (+), $\tau_{\lambda=25}(N) \sim N^{1.600}$ (□), $\tau_{\lambda=45}(N) \sim N^{1.796}$ ($\times$), $\tau_{\lambda=70}(N) \sim N^{1.911}$ (△), $\tau_{\lambda=100}(N) \sim N^{1.932}$ (∗). The fits are quite good for all the cases but $\lambda = 10$ and $\lambda = 25$. In these cases the quality of the fit is enhanced excluding the data corresponding to $N = 25$ and $N = 40$. So we obtain $\tau_{\lambda=10}(N) \sim N^{1.184}$ and $\tau_{\lambda=25}(N) \sim N^{1.488}$. 
Next, we measured the lowest energy found by the algorithm. In Fig. 2 we represent $H_N$ as a function of $N$ for different values of $\lambda$ and for a fixed number of $N$ initial conditions. While, for small $N$, the ground state is believed to be closely approximated for all values of $\lambda$ (in fact, varying $\lambda$, the lowest energy $H_N$ undergoes a relative change of $2.5 \cdot 10^{-4}$ for $N = 25$ and of $8.8 \cdot 10^{-4}$ for $N = 40$ and 50), the best results for large $N$ ($> 50$) are obtained for $\lambda = 100$ which corresponds to deterministic reluctant dynamics. Therefore, this confirms that, for a fixed number of initial spin configurations, the algorithm that makes moves corresponding to the “smallest” possible energy decrease is the most efficient in reaching low-energy states. In other words, the slower ... the better! However, this is reflected in an increasing cost for the computational time.

Indeed, when the analysis is focused on the performances for a fixed elapsed time, the situation changes drastically. In Fig. 3 we compare the minimum energy values $H_N$, obtained considering different system sizes

Figure 2: Lowest energy value $H_N$ as a function of $N$ obtained using a protocol with $N$ initial conditions for 1000 disorder realizations for $\lambda = 1$ (○), $\lambda = 10$ (+), $\lambda = 25$ (□), $\lambda = 45$ (×), $\lambda = 70$ (△), $\lambda = 100$ (∗).
and, for each of them, five different parameter values ($\lambda = 1, \lambda = 10, \lambda = 25, \lambda = 45, \lambda = 100$) for an elapsed time of 50 h of CPU on a IBM SP4. The system size $N = 350$ is studied only for $\lambda = 10$, $\lambda = 25$ and $\lambda = 45$. Each run (i.e. for fixed $N$ and $\lambda$) consists of 1000 disorder realizations, with the same CPU time length (3 min.) assigned to each sample. For $N \leq 100$ we believe to find the ground state of the system, since varying $\lambda$ the values of $H_N$ coincide, within our numerical accuracy ($10^{-10}$). The best result is obtained for the case $\lambda = 10$, which seems to be the best compromise to obtain a dynamical trajectory that is able to arrive deep enough with respect to energy levels but without wasting all the time in the search for the slower of possible path. We note that this finding is in good agreement with the result of previous analysis [6], where a convex linear combination of reluctant (with probability $P$) and greedy (with probability $1 - P$) dynamics was considered. The optimal value $\lambda \sim 10$ is the one for which the relaxation time $\tau \sim N^\alpha$ grows with a scaling exponent $\alpha$ which is the closest, among the others, to the value $\alpha = 1.26$ of the optimal convex combination with $P = 0.1$.

Improvements of the greedy and reluctant algorithms is presently under study [9], by permitting also increase in energy with exponential decrease in time, in the very same spirit of the well-known Simulated Annealing strategies.

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References


Figure 3: Lowest energy value $H_N$ as a function of $N$ for a fixed CPU time of 50 h on a IBM SP4 for $\lambda = 1$ (○), $\lambda = 10$ (+), $\lambda = 25$ (□), $\lambda = 45$ (×), $\lambda = 100$ (∗).


