Analysis of Random Noise and Random Walk Algorithms for Satisfiability Testing

Bhaskar Krishnamachari\textsuperscript{1}, Xi Xie\textsuperscript{1}, Bart Selman\textsuperscript{2}, and Stephen Wicker\textsuperscript{1}

\textsuperscript{1} School of Electrical Engineering, Cornell University, Ithaca, NY 14853
\{bhaskar, xie, wicker\}@ee.cornell.edu
\textsuperscript{2} Department of Computer Science, Cornell University, Ithaca, NY 14853
selman@cs.cornell.edu

Abstract. Random Noise and Random Walk algorithms are local search strategies that have been used for the problem of satisfiability testing (SAT). We present a Markov-chain based analysis of the performance of these algorithms. The performance measures we consider are the probability of finding a satisfying assignment and the distribution of the best solution observed on a given SAT instance. Although the analysis is restricted to SAT problems with a limited number of variables as it requires complete knowledge of the search space, the insights gained are pertinent to the performance of these algorithms on real world problems. We examine the effect of \( p \), the probability of making non-greedy moves, on these algorithms and provide a justification for the practice of choosing this value empirically.

1 Introduction

Local search algorithms such as GSAT, Random Walk and Random Noise search have been shown to be good at solving CNF satisfiability (SAT) problems [4], [13]. Such methods perform better than systematic search algorithms on large satisfiability problems involving thousands of variables. They may be used for the problem of maximum satisfiability (finding a truth assignment that satisfies as many clauses as possible) as well as complete satisfiability (satisfying all clauses).

However, due to the complex interactions between the problem instance and algorithm implementation details, it is hard to predict the performance of these algorithms. Researchers have, therefore, mainly relied upon empirical studies for this purpose [5], [14]. Although this approach provides very useful results, it is still desirable to have some theoretical understanding of algorithm performance.

A large portion of the literature on theoretical analysis of local search algorithms for other problems has been devoted to determining the convergence of search algorithms to the global optimum using Markov models [2], [3], [7], [8], [9], [11]. The rates of convergence to the optimum have also been discussed assuming various properties of cost functions and search spaces [15], [16]. Some work in the area of complexity theory has been focused on studying PLS (polynomial
local search) problems regarding the time required to locate local optima [6], [10].

In this paper, we show how the Random Walk and Random Noise algorithms can be modeled using discrete Markov chains. We present a procedure to determine the probability of finding the global optimum as well as complete statistics of the best solution observed in a given number of iterations. The former measure of algorithm performance is most relevant to maximum satisfiability problems, while the latter is the statistic of interest when considering complete satisfiability. These measures are relevant because they tell us quantitatively how the algorithm will perform on a given problem in limited computational time.

The procedure presented requires complete knowledge of the search space. Hence, it can only be carried out for small-scale satisfiability problems. Still, the insights we get from this analysis are pertinent to the performance of algorithms on real world problems. Both Random Noise and Random Walk algorithms are characterized by the probability \( p \) of making non-greedy moves – a measure of the randomness of the search. The value of \( p \) that provides optimum algorithm performance is of great interest. We find theoretical support for the practice of empirically choosing an optimal value for this parameter.

The rest of the paper is organized as follows: section 2 reviews the definitions of the Random Noise and Random Walk algorithms. Section 3 shows how these algorithms can be modeled as discrete Markov chains and presents a procedure for determining the performance statistics for these algorithms. The method of determining the one-step state transition matrix for these algorithms is described in section 4. Section 5 presents and discusses results obtained using this procedure. Concluding comments are presented in section 6.

2 Random Noise and Random Walk Algorithms for Satisfiability

The Random Noise and Random Walk algorithms are both based on GSAT, a greedy local search procedure for satisfiability which works as follows [12]:

Procedure GSAT

for \( i := 1 \) to MAX-TRIES
  T := a randomly generated truth assignment
  for \( j := 1 \) to MAX-FLIPS
    if T satisfies expression then return T
    Flip any variable in T that results in greatest decrease
    (could even be 0) in the number of unsatisfied clauses
  end for
end for
return "No satisfying assignment found"

The success of GSAT depends on its ability to make either strictly improving or "sideways" moves (moves to assignments with an equal number of unsatisfied
clauses. When the algorithm finds itself at a non-optimal point in the search space where no further improvements are possible, it is essentially trapped in a region which is a local minimum and needs to be restarted with a random new assignment. Another mechanism for escaping such local minima that is widely used is to permit the search to make uphill moves occasionally. Random Noise and Random Walk algorithm are both closely related in the way they allow for the possibility of uphill moves [13]:

**Random Noise**

With probability $p$, pick any variable at random and flip its truth assignment.
With probability $1-p$, follow the standard GSAT scheme, i.e., make the best possible local move

**Random Walk**

With probability $p$, pick a variable occurring in some unsatisfied clause and flip its truth assignment.
With probability $1-p$, follow the standard GSAT scheme, i.e., make the best possible local move

Experimental results comparing the basic GSAT algorithm, Simulated Annealing, Random Walk and Random Noise strategies on a test suite including randomly-generated CNF problems and Boolean encodings of circuit synthesis, planning and circuit diagnosis problems can be found in [13]. The authors of this paper found that the Random Walk strategy significantly out-performed the other algorithms on these problems.

### 3 Modeling and Analysis

Most local search algorithms are memoryless in the sense that the process of selecting the next point in the search space depends iteratively on the current point. This allows us to model these search algorithms as Markov chains. Such models for two widely used local search algorithms – Simulated Annealing and Genetic Algorithms, can be found in [1] and [9] respectively. In the context of satisfiability problems, each point in the search space corresponds to a different truth assignment.

**Theorem 1:** The sequence of points visited by the Random Noise (also Random Walk) algorithm forms a Homogeneous Markov Chain.

**Proof:** To prove that this sequence forms a Markov chain, it suffices to show that the point visited at the $(k + 1)^{th}$ iteration depends only upon which point was visited at the $k^{th}$ iteration. This can be seen as follows: by the definition
of both these algorithms, the truth assignment at the \((k + 1)^{th}\) iteration differs from the truth assignment at the \(k^{th}\) iteration at exactly one variable. The set of variables that may be flipped at the \(k^{th}\) iteration depends only upon whether we are considering the Random Noise or Random Walk algorithm and not the points visited at any previous iteration. Finally, the probability of flipping each variable in this set also depends only on the value of \(p\) and not the points visited in the first \((k - 1)\) iterations. The Markov chain will be homogeneous because the state transition probabilities will only be a function of \(p\) which is assumed to be constant for the duration of the search. Q.E.D.

If \(N\) is the number of variables, the search space consists of a total of \(|X| = 2^N\) possible truth assignments. Let \(x_{jd}\) be a point in the search space. The cost \(f(x_{jd})\) is the number of unsatisfied clauses in the corresponding truth assignment. For simplicity of analysis and description, we assume that the points in the search space are sorted in non-decreasing order of costs, i.e. \(j < k \Rightarrow f(x_{jd}) \leq f(x_{jd})\). The search space may contain two points with the exact same cost function value. We represent the sorted list of costs using a row vector \(\bar{f}\) of size \(|X|\) such that \(\bar{f}_{jd} = f(x_{jd})\).

Let \(x_i\) be the random variable describing which point the search is at during iteration \(i\). The pmf of \(x_i\) is represented by a row vector \(\pi_i\) of size \(|X|\) such that \(\pi_{i,m} = P(x_i = x_i^m)\). A homogeneous Markov chain based local search algorithm can then be described by a one-step state transition matrix \(P\) such that:

\[
\pi_i = \pi_{i-1}P
\]

The performance statistics of interest (probability of finding the global optimum within a given number of iterations, best solution observed to date) require us to incorporate the search history as well. For this purpose, it is necessary to fully describe the state that the search algorithm is in at a given iteration. This description should include a) the probability mass function (pmf) describing the probability that the search algorithm is at any given point in the search space, and b) the conditional pmf's of the best (lowest) cost seen up to the current iteration given that the search is currently at a certain point in the search space. Both these pmf's can be iteratively calculated at each step as they depend only on the pmf's of the previous iteration and on the search algorithm being analyzed.

Let \(x_i^*\) denote the point with the lowest cost function seen up to iteration \(i\). We can use a matrix \(D_i^*\) to represent the conditional probability of the lowest cost seen to date being \(m\) given that the current point is \(n\), i.e. \(D_{i,mn}^* = P(x_i^* = m|x_i = n)\). Note that \(D_0^* = I\). For entries representing equal value points, it does not matter how the weight is distributed among them as long as the total probability remains the same.
The distribution $\mathbf{\bar{\pi}}_i$ can be calculated from $\mathbf{\bar{\pi}}_{i-1}$ using equation 1. The following formulae\(^1\) can be used in sequence to calculate $\mathbf{D}_i^*$ from $\mathbf{\bar{\pi}}_i$ and $\mathbf{D}_{i-1}$:

$$\mathbf{B}_i^* = \mathbf{D}_{i-1}^* \text{diag}(\mathbf{\bar{\pi}}_{i-1}) \mathbf{P}$$

$$
C_{i,j}^* = \begin{cases} 
\frac{1}{k} & j < k \\
\frac{1}{k} \sum_{b=j}^{k} B_{i,j}^* & j = k \\
0 & j > k 
\end{cases}
$$

$$\mathbf{D}_i^* = \mathbf{C}_i^* (\text{diag}(\mathbf{\bar{\pi}}_i))^{-1}$$

Thus, given the initial state distribution $\mathbf{\bar{\pi}}_0$ and the state transition matrix $\mathbf{P}$, we can derive $\mathbf{D}_n^*$ – the conditional pmf's of the lowest cost function value seen to date, and $\mathbf{\bar{\pi}}_n$ – the distribution of costs at the $n^{th}$ iteration\(^2\). It is typically assumed that each point in the search space is equally likely to be picked as the starting point (uniform initial distribution). Once $\mathbf{D}_n^*$ and $\mathbf{\bar{\pi}}_n$ are known, the expectation and variance of the best-to-date cost can then be readily calculated:

$$E[f(x_n^*)] = \bar{\mathbf{\bar{\pi}}} D_n^* \mathbf{\bar{\pi}}_n$$

$$VAR[f(x_n^*)] = \bar{\mathbf{\bar{\pi}}} \text{diag}(\mathbf{\bar{\pi}}) D_n^* \mathbf{\bar{\pi}}_n - (\bar{\mathbf{\bar{\pi}}} D_n^* \mathbf{\bar{\pi}}_n)^2$$

The expectation and variance of best-to-date cost are useful measures if we are interested in the problem of maximum satisfiability. For complete satisfiability, we would like to know the probability $P[f(x_n^*) = f^*]$ of achieving the global optimum $f^*$ within $n$ iterations. This can be calculated as follows:

$$P[f(x_n^*) = f^*] = \mathbf{\bar{\pi}} D_n^* \mathbf{\bar{\pi}}_n$$

where $\mathbf{\bar{\pi}} = [1 \ 0 \ 0 \ 0 \ldots \ 0]$, consisting of a 1 followed by $(|X| - 1)$ zeros.

4 Determining the State Transition Matrix

We have shown how the performance statistics of interest may be obtained once the state transition matrix $\mathbf{P}$ is known. This matrix depends upon both the

\(^1\) In these formulae, $\text{diag}(\mathbf{\bar{\pi}})$ represents the diagonal matrix derived from a vector $\mathbf{\bar{\pi}}$; $\mathbf{B}_i^*$ and $\mathbf{C}_i^*$ are temporary matrices used during this updating process.

\(^2\) Rigorously, the inverse of $\text{diag}(\mathbf{\bar{\pi}}_i)$ does not exist if any of the elements of $\mathbf{\bar{\pi}}_i$ are 0 – although this only happens when using purely greedy search. However the notation used in equation 4 is convenient and the difficulty can be overcome by treating these 0 elements as arbitrarily small values $\epsilon$. 

specific problem instance, as well as the algorithm used. We discuss here via an example how the matrix can be obtained for Random Noise algorithms if the satisfiability instance is known.

Table 1 shows a randomly generated 3-SAT instance with 3 variables and 15 clauses\(^3\). The cost for each truth assignment, i.e. the number of unsatisfied clauses as well as the neighboring truth assignments are shown in figure 1. From the figure, it is easy to see that the assignment 001 is a local minimum and that the global minima 111 and 100 are satisfying assignments.

Given a problem instance, the one-step transition matrix \( P \) can be determined for the Random Noise algorithm as follows:

- Determine the transition matrix \( P_{\text{greedy}} \) for the GSAT algorithm (\( p = 0 \)).
- Determine the transition matrix \( P_{\text{random}} \) for the random noise algorithm with \( p = 1 \).
- \( P = (1 - p) P_{\text{greedy}} + p P_{\text{random}} \)

For the SAT instance presented in figure 1, the corresponding transition matrices for the Random Noise algorithm: \( P_{\text{greedy}} \) and \( P_{\text{random}} \) are shown in figure 2. The \( P_{\text{random}} \) matrix is constructed by assigning equal transition probabilities to each neighbor of a given truth assignment (elements corresponding to non-neighbor points are 0). The \( P_{\text{greedy}} \) is constructed by assigning equal transition probabilities from any given truth assignment to the neighbor(s) which have the greatest decrease in cost (0 or more). The procedure is nearly identical for obtaining the \( P \) for the Random Walk algorithm, with the only

\(^3\) It may be seen that a number of these clauses are identical in this example, but this is due to the small number of variables used for illustration.
difference being in the construction of $\mathbf{P}_{\text{random}}$. To construct $\mathbf{P}_{\text{random}}$ for the Random Walk algorithm, assign equal state transition probabilities to each neighbor of a given truth assignment that can be obtained by flipping a variable involved in unsatisfied clauses.

$$
\mathbf{P}_{\text{random}} = \frac{1}{3}
\begin{bmatrix}
0 & 0 & 0 & 1 & 0 & 0 & 1 & 1 \\
0 & 0 & 0 & 1 & 1 & 0 & 1 & 0 \\
0 & 0 & 0 & 1 & 0 & 1 & 1 & 0 \\
1 & 1 & 0 & 0 & 0 & 1 & 0 & 0 \\
0 & 1 & 1 & 0 & 0 & 1 & 0 & 0 \\
0 & 0 & 0 & 1 & 1 & 0 & 0 & 1 \\
1 & 1 & 1 & 0 & 0 & 0 & 0 & 0 \\
1 & 0 & 1 & 0 & 0 & 1 & 0 & 0 \\
\end{bmatrix}
$$

$$
\mathbf{P}_{\text{greedy}} = \frac{1}{2}
\begin{bmatrix}
2 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 2 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 2 & 0 & 0 & 0 & 0 & 0 \\
1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 2 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 2 & 0 & 0 & 0 & 0 & 0 \\
1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
2 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\end{bmatrix}
$$

**Fig. 2.** Constructing State Transition Probability Matrices

5 Results

Using the method of analysis presented in this paper, it is possible to investigate the effect of $p$, the non-greedy move probability, on the performance of random noise algorithms. One significant result is the following:

**Theorem 2:** $E[f(x^*_n)]$, the expected best cost seen by a random noise or random walk algorithm after $n$ iterations, is a polynomial in $p$ of order at most $n$.

**Proof:** See appendix A.

**Corollary 1:** The variance of the best cost $VAR[f(x^*_n)]$ is a polynomial in $p$ of order at most $2n$.

**Corollary 2:** $P[f(x^*_n) = f_\star]$, the probability of having found the global best assignment after $n$ iterations, is a polynomial in $p$ of order at most $n$.

We applied the procedure described in equations (1) through (7) on randomly generated 3-SAT instances to determine the performance of Random Walk and Random Noise algorithms. Table 2 shows a typical instance with 7 variables and 30 clauses. Figure 3 shows the effect of $p$ on Random Noise (3a,3b) and Random Walk (3c,3d) algorithms for this instance.

The data in Figure 3 is for 20 values of $p$ ranging from 0 to 1, for the first 20 iterations of the algorithm. Figures 3a and 3c show the expected best cost. The first line on top corresponds to iteration 0, the starting point of the search. As the iteration number increases, the expected cost goes down at each step and is
indicated by the successively lower curves. Figure 3b and 3d show the probability of having found the global optimum (in this case, a satisfying assignment with 0 cost). This probability increases with iteration and is hence represented by successively higher curves. In all these graphs, for any given iteration, there is some \( p = p_{\text{best}} \) for which the algorithm achieves the best value. In figures 3b and 3d, the \( p_{\text{best}} \) points for each iteration (subject to the resolution of the \( p \) values tested) are connected, forming a locus.

The performance statistics for the two algorithms are different, and this can be seen more clearly in figure 4, where the probability of having found the global minimum for both algorithms is compared for \( p = 0, 0.5, \text{and} 1 \). When \( p = 0 \), as noted earlier, both algorithms are identical to the GSAT algorithm and hence their performance is the same. For the other two values of \( p \), it is seen that Random Walk algorithm out-performs the Random Noise algorithm. We have noticed this on other instances as well. This has also been observed empirically on large-scale problems [14],[13].

6 Conclusion

Local search algorithms can be modeled as Markov chains. We have shown how the Random Noise and Random Walk algorithms for satisfiability testing can also be described using such a model. Based on this modeling, we are able to derive the probability of finding a satisfying assignment, and the statistics of the best solution observed within a given number of iterations. The former measure of algorithm performance is useful when considering problems of complete satisfiability, while the latter is more relevant to the problem of maximum satisfiability and related optimization problems.

For real world problems, it is almost always the case that a value of \( p = p_{\text{best}} \in (0,1) \) offers the best performance on a given problem. The results obtained for randomly generated 3-SAT instances using our analysis also show this behavior. The value of \( p_{\text{best}} \) depends upon the specific problem instance, as well as the iteration number. We observed that the performance measures vary slowly with respect to \( p \). Further, we have proved that these performance measures are polynomial (hence continuous) functions of \( p \), the probability of making non-greedy moves. Therefore, for real world problems, if the value of \( p \) chosen via experiments is close to \( p_{\text{best}} \), it will result in near-optimal performance. In nearly

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Table 2. sample 3-SAT instance with 7 variables, 30 clauses
Fig. 3. Performance for Random Noise and Random Walk algorithms with respect to $p$ for a randomly generated 3-SAT instance with 7 variables.
all the 3-SAT instances we tested, the Random Walk algorithm out-performed the Random Noise algorithm. This merits further study.

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Appendix A

Remark – Properties of the Transition Matrix: Since the sum of any row of
the transition matrix P is 1, P \( \mathbf{T}^r \) = \( \mathbf{T}^r \), where \( \mathbf{T} \) is a 1 \times 2^N vector of 1’s.
The elements of P are polynomials in p of degree either 0 or 1.

Definition: If a matrix A has elements that are polynomials in p, \( \mathcal{O}(A) \) is
defined as the highest possible degree of these polynomials.

Theorem 2: For a given neighborhood, \( E[f(x_n^*)] \), the expected best cost seen
by a random noise algorithm after n iterations, is a polynomial in p of order at
most n.

Proof: By substituting equation (4) into (5), we have:

\[
E[f(x^*_n)] = \mathbf{J} C^*_i (\text{diag}(\mathbf{\pi^*_i}))^{-1} \mathbf{\pi^*_i} = \mathbf{J} C^*_i \mathbf{T}^r
\]

If the elements of \( C^*_i \) be polynomials in p, then \( E[f(x^*_n)] \) is also a polynomial
in p, and

\[
\mathcal{O}(E[f(x^*_n)]) = \mathcal{O}(C^*_i \mathbf{T}^r)
\]

Hence it suffices to show that the elements of \( C^*_i \) are polynomials in p, and that
\( \mathcal{O}(C^*_i) = i \). This can be done inductively:

Base Case (n = 1)

From the fact that \( D^*_0 = I \) and equation (2), we get \( B^*_1 = \text{diag}(\mathbf{\pi^*_0})P \). Hence
the elements of \( B^*_1 \) are polynomials in p and \( \mathcal{O}(B^*_1) = 1 \). All the rows of \( B^*_1 \) add
up to one:

\[
\mathcal{O}(B^*_1 \mathbf{T}^r) = \mathcal{O}(\text{diag}(\mathbf{\pi^*_0})P \mathbf{T}^r) = \mathcal{O}(\text{diag}(\mathbf{\pi^*_0}) \mathbf{T}^r) = 0
\]
The operations in equation (3) consist of adding all elements of $B_i^*$ that are below the diagonal to the diagonal element in each column and then setting these below-diagonal elements to 0. Thus $C_i^*$ is an upper-triangular matrix such that all its elements are also polynomials in $p$ and $\overline{O}(C_i^*) = \overline{O}(B_i^*) = 1$. When each row of $C_i^*$ is summed, the order 1 terms will not necessarily cancel out, and hence $\overline{O}(C_i^* 1^T) = 1$.

**Inductive Hypothesis**

For any $k > 1$, the elements of $B_{k-1}^*$ and $C_{k-1}^*$ are polynomials in $p$. Further, $\overline{O}(C_{k-1}^*) = \overline{O}(B_{k-1}^*) = k - 1$, $\overline{O}(B_{k-1}^* 1^T) = k - 2$, and $\overline{O}(C_{k-1}^* 1^T) = k - 1$.

**Inductive Step**

This is similar to the verification of the base case. From equations (2) and (4), we get:

$$B_i^* = C_{i-1}^*(\text{diag}(\overline{x}_{i-1})^{-1}\text{diag}(\overline{x}_{i-1})P = C_{i-1}^*P$$

(11)

By this equation, the elements of $B_k^*$ are polynomial in $p$, and $\overline{O}(B_k^*) = \overline{O}(C_{k-1}^*) + \overline{O}(P) = k$. The operations in equation (3) ensure that $C_k^*$ is an upper-triangular matrix and that all its elements are polynomials in $p$ with $\overline{O}(C_k^*) = \overline{O}(B_k^*) = k$. Also from equation (11),

$$\overline{O}(B_k^* 1^T) = \overline{O}(C_{k-1}^*P 1^T) = \overline{O}(C_{k-1}^* 1^T) = k - 1$$

(12)

This means that when each row of $B_k^*$ is summed, any terms of degree $k$ all cancel out. After the below-diagonal elements of $B_k^*$ are moved to the diagonal terms in equation 3, when the rows of $C_k^*$ are summed, the terms of degree $k$ will not necessarily cancel. Hence $\overline{O}(C_k^* 1^T) = k$.

Therefore by induction, we have that the elements of $C_n^*$ are polynomials in $p$ and that $\overline{O}(C_n^* 1^T) = n$, $\forall n \geq 1$. Q.E.D.

**Corollary 1:** The variance of the best cost $VAR[f(x_n^*)]$ is a polynomial in $p$ of order at most $2n$.

This follows immediately from the result of Theorem 1 and equation 6.

**Corollary 2:** $P[f(x_n^*) = f^*]$, the probability of having found the global best assignment after $n$ iterations, is a polynomial in $p$ of order at most $n$.

To see this, compare equations (5) and (7). The properties of $E[f(x_n^*)]$ with respect to $p$ hold for $P[f(x_n^*) = f^*]$ as well.