The Randomness Recycler: A New Technique for Perfect Sampling

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Abstract

For many probability distributions of interest, it is quite difficult to obtain samples efficiently. Often, Markov chains are employed to obtain approximately random samples from these distributions. The primary drawback to traditional Markov chain methods is that the mixing time of the chain is usually unknown, which makes it impossible to determine how close the output samples are to having the target distribution. Here we present a new protocol, the randomness recycler (RR), that overcomes this difficulty. Unlike classical Markov chain approaches, an RR-based algorithm creates samples drawn exactly from the desired distribution. Other perfect sampling methods such as coupling from the past use existing Markov chains, but RR does not use the traditional Markov chain at all. While by no means universally useful, RR does apply to a wide variety of problems. In restricted instances of certain problems, it gives the first expected linear time algorithms for generating samples. Here we apply RR to self-organizing lists, the Ising model, random independent sets, random colorings, and the random cluster model.

1 Introduction

The Markov chain Monte Carlo (MCMC) approach to generating samples has enjoyed enormous success since its introduction, but in certain cases it is possible to do better. The “randomness recycler” technique we introduce here (and whose name is explained in Section 2) works for a variety of problems without employing the traditional Markov chain. Our approach is faster in many cases, generating in particular the first algorithms that have expected running time linear in the size of the problem, under certain restrictions.

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In classical MCMC approaches, small random changes are made in the observation until the observation has nearly the stationary distribution of the chain. The Metropolis [13] and heat bath algorithms utilize the idea of reversibility to design chains with a stationary distribution matching the desired distribution. Unfortunately, this standard Markov chain approach does have problems.

The samples generated by MCMC will not be drawn exactly from the stationary distribution, but only approximately. Moreover, they will not be close to the stationary distribution until a number of steps larger than the mixing time of the chain have been taken. Often the mixing time is unknown, and so the quality of the sample is suspect.

Recently, Propp and Wilson have shown how to avoid these problems using techniques such as coupling from the past (CFTP) [15]. For some chains, CFTP provides a procedure that allows perfect samples to be drawn from the stationary distribution of the chain, without knowledge of the mixing time. However, CFTP and related approaches have drawbacks of their own. These algorithms are non-interruptible, which means that the user must commit to running such an algorithm for its entire (random) running time even though that time is not known in advance. Failure to do so can introduce bias into the sample. Other algorithms, such as FMMR [3], are interruptible (when time is measured in Markov chain steps), but require storage and subsequent rereading of random bits used by the algorithm. The method we will present is both interruptible and "read-once," with no storage of random bits needed.

In addition, algorithms like CFTP and FMMR require an underlying Markov chain, and can never be faster than the mixing time of this underlying chain. Often these chains make changes to parts of the state where the state has already been suitably randomized. This leads to wasted effort when running the algorithm that often adds a log factor to the running time of the algorithm.

The randomness recycler (RR) is not like any of these perfect sampling algorithms. In fact, the RR approach abandons the traditional Markov chain entirely. This is what allows the algorithm in several cases to reach an expected
running time that is linear, the first for several problems of interest. The RR technique gives interruptible, read-once perfect samples.

In the next section we illustrate the randomness recycler for the problem of finding random independent sets of a graph. After this example we present in Section 3 the general randomness recycler procedure and present a (partial) proof of correctness. In Section 4 we present other applications and in Section 5 we review the results of applying our new approach to several different problems.

2 Weighted Independent Sets

We begin by showing how the randomness recycler technique applies to the problem of generating a random independent set of a graph. This will illustrate the key features of RR and lay the groundwork for the more general procedure described in the next section.

Recall that an independent set of a graph is a subset of vertices no two of which share an edge. We will represent an independent set as a coloring of the vertices from \( \{0, 1\} \), denoted generically by \( x \). Set \( x(v) = 1 \) if \( v \) is in the independent set, and \( x(v) = 0 \) if \( v \) is not in the independent set. This implies that \( \sum_v x(v) \) is the size of the independent set.

We wish to sample from the distribution

\[
\pi(x) = \frac{\lambda \sum_v x(v)}{Z_\lambda},
\]

where \( \lambda \) (called the fugacity) is a parameter of the problem, and \( Z_\lambda \) is the normalizing constant needed to make \( \pi \) a probability distribution.

This distribution is known as the hard core gas model in statistical physics, and also has applications in stochastic loss networks [11]. When \( \lambda \) is large the sample tends to be a large independent set, and if \( \lambda > 28/\Delta \) where \( \Delta \) is the maximum degree of the graph, it is known that generating samples from this distribution cannot be done in polynomial time unless \( NP = RP \) [1].

We will show that for \( \lambda < 4/(3\Delta - 4) \) the randomness recycler approach gives an algorithm with expected running time linear in the size of the graph, the first such result for this problem.

The RR approach is to start not with the entire graph, but rather with a small graph where we can easily find an independent set from this distribution. For example, if a graph has only a single vertex, finding an independent set is easy. Starting from a single vertex, we attempt to add vertices to the graph, building up until we are back at our original problem. Sometimes we fail in our attempt to build up the graph, and indeed we need to remove vertices that we had previously added. The set \( V_t \) will comprise those vertices we have built up by the end of time step \( t \). After step \( t \), the vector \( x_t \) will hold an independent set which has the correct distribution over (the subgraph induced by) the vertices in \( V_t \).

**Randomness Recycler for Independent Sets**

Set \( V_0 \leftarrow \emptyset, x_0 \leftarrow 0, t \leftarrow 0 \)

Repeat

Set \( x_{t+1} \leftarrow x_t \)

Choose any \( v \in V \setminus V_t \)

Set \( V_{t+1} \leftarrow V_t \cup \{v\} \)

Draw \( U \) uniformly at random from \([0, 1]\)

If \( U \leq 1/(1 + \lambda) \)

Let \( x_{t+1}(v) \leftarrow 1 \)

Else

Let \( x_{t+1}(v) \leftarrow 0 \)

If a neighbor \( w \) of \( v \) has \( x_{t+1}(w) = 1 \)

Let \( w \) be the lowest-numbered such neighbor

Set \( x_{t+1}(w) \leftarrow 0, x_{t+1}(v) \leftarrow 0 \)

Remove from \( V_{t+1} \) the vertices \( v \) and \( w \),

all neighbors of \( w \), and all neighbors of \( v \)

with numbers less than that of \( w \)

Set \( t \leftarrow t + 1 \)

Until \( V_t = V \)

(In advance of running the algorithm, choose and fix a numbering of the vertices.) The algorithm proceeds inductively as follows. At the outset of step \( t + 1 \), we begin with an independent set \( x_t \) of \( V_t \) chosen with the correct probability. Then we choose a vertex \( v \) not in \( V_t \) to attempt to add. This vertex may be chosen in any fashion desired (randomly, or according to some fixed order, but not depending on the independent set \( x_t \)). Because the desired probability of choosing an independent set \( x \) is proportional to \( \lambda \sum_v \pi(v) \), putting \( x_{t+1}(v) = 1 \) has \( \lambda \) times the weight of putting \( x_{t+1}(v) = 0 \). Therefore we select \( x_{t+1}(v) = 1 \) with probability \( \lambda/(1 + \lambda) \) and \( x_{t+1}(v) = 0 \) with probability \( 1/(1 + \lambda) \) (these are the heat bath probabilities).

Unfortunately, the vector \( x_{t+1} \) resulting from this selection may fail to correspond to an independent set. At line 11 of the pseudocode, we check whether some neighbor of \( v \) was already colored 1 (in the independent set). Note that we cannot simply remove \( v \). Prior to the step, we knew that \( x_t \) was an independent set of \( V_t \). If we observe that \( x_t(w) = 1 \) for some lowest-numbered neighbor \( w \) of \( v \), then \( x_t \) is an independent set on \( V_t \) conditioned on this knowledge.

Our solution is this: In line 14 we "undo" the knowledge gained by removing from \( V_{t+1} \) the vertices \( v \) and \( w \), all the neighbors of \( w \), and all the neighbors of \( v \) with number less than that of \( w \). On the remaining vertices of \( V_{t+1} \), \( x_{t+1} \) will continue to be an independent set from the correct distribution. We will say that an RR step of this type preserves the correct distribution.

Note that although \( V_{t+1} \) is made smaller than \( V_t \) in the case of a conflict, we are able to salvage most of the vertices.
in \(V_i\). In other words, we “recycle” the randomness built up in all of the vertices except \(v\) and \(w\) and some neighbors. This is where our approach gets its name, and “recycling” is the key new feature that enables us to construct similar practicable algorithms for a wide variety of problems.

We repeat until \(V_i = V\). Because each step preserves the correct distribution, we know that \(z_t\) will have the correct distribution \(\pi\) at the end. This is proved formally in the next section; here we concentrate on bounding the running time of our procedure.

**Theorem 1** If \(\lambda < 1/(2\Delta - 1)\), then the expected running time of the above randomness recycling procedure for random independent sets is \(O(n)\).

A more careful statement of Theorem 1 is given following the proof.

**Proof** We will show that for \(\lambda\) this small, on average \(|V_i|\) increases at each step. If \(U \leq 1/(1 + \lambda)\), then the size of \(|V_i|\) goes up by 1, but if \(U > 1/(1 + \lambda)\), then the size of \(|V_i|\) may decrease by at most \(2\Delta - 1\) [removing \(v\) (not previously included), \(w\), and some neighbors]. Hence

\[
E[|V_{i+1}| - |V_i| \mid V_i, z_t] \geq \frac{1}{1 + \lambda}(1 - \frac{\lambda}{1 + \lambda}(2\Delta - 1)) \approx \frac{1}{1 + \lambda}[1 - (2\Delta - 1)\lambda],
\]

which is positive precisely when \(\lambda < 1/(2\Delta - 1)\). Given an increase of \(|V_i|\) on average at each step, standard martingale stopping theorems (see, e.g., [14]) show that after \(O(n)\) expected time the value of \(|V_i|\) will be \(n\), at which point \(V_i = V\) and the algorithm terminates. □

More carefully, if

\[
\frac{1}{1 + \lambda}[1 - (2\Delta - 1)\lambda] \geq \gamma \in (0, 1),
\]

i.e., if

\[
\lambda \leq \frac{2\Delta}{1 - \gamma - 1} - 1,
\]

then the expected value of the running time \(T\) (measured by number of iterations of the Repeat loop) satisfies

\[
ET \leq n/\gamma.
\]

Furthermore, a simple argument shows that the distribution of \(T\) has at worst geometrically thick tails:

\[
P(T \geq 2^m n) \leq 2^{-m}, \quad m = 1, 2, \ldots,
\]

Several tricks may be used to either improve our method or to improve our bounds on its performance. The first two we present by altering the algorithm, and the third gives a better analysis. First we concentrate on making sure that as few vertices as possible are removed in the rejection step. Note that we may assume that the graph is connected, since otherwise we simply work on each connected component separately. Therefore \(V \setminus V_i\) is connected, and by being slightly careful in how we choose \(v \in V \setminus V_i\), we can ensure that \(V \setminus V_i\) remains connected at each step. In every step (except when \(|V \setminus V_i| = 1\)), the vertex \(v\) is not adjacent to \(\Delta\) vertices in \(V_i\), but only to at most \(\Delta - 1\), so fewer vertices are removed during rejection. Since the vertices removed from \(V_i\) in case of rejection are connected to \(V \setminus V_i\), \(V \setminus V_{i+1}\) will also be connected whether we accept or reject, and no more than an extra constant amount of work is required at each step.

The second alteration concerns how we look for the neighbor of \(v\) that is colored 1 in case of rejection. Instead of starting at the lowest numbered neighbor and working our way up, we start at a random neighbor and continue looking in cyclic order until we find our \(w\) colored 1 and then the vertices that we remove are \(w\) and its neighbors and \(v\) and its neighbors encountered in the search prior to finding \(w\). On average (and together with the first trick), we need only look at \(\leq \Delta/2\) vertices in order to find \(w\).

Finally, in our analysis we keep track of a potential \(\phi(V_i, z_t) = |V_i|\) and showed that \(\phi\) increases on average. When we accept we sometimes add a vertex colored 1 to our set \(V_i\); but when we reject, precisely one vertex colored 1 (namely, \(w\)) is removed. This suggests that we modify \(\phi\) so that the acceptance and rejection phases both lead (in the worst case) to the same expected change in \(\phi\). We will consider

\[
\phi(V_i, z_t) = |V_i| - \alpha \sum_v z_t(v)
\]

and seek a suitable value of \(\alpha\). The expected change in \(\phi\) if no neighbor of \(v\) is colored 1 is \(1 - \alpha [\lambda/(1 + \lambda)]\). If some neighbor is colored 1, then the expected change is at least \(1 [1/(1 + \lambda)] + \left(\lambda/1 + \lambda\right) - (3\Delta/2 + \alpha)\). (The term \((3\Delta - 2)/2\) is an upper bound on the expected decrease in \(|V_i|\), since (see above) on average we lose at most \(\Delta/2\) neighbors of \(v\) and \(\Delta - 1\) neighbors of \(w\).)

These two expressions may be made equal by setting \(\alpha = 3\Delta/4\), and then the expected change in \(\phi\) will be positive when

\[
\lambda < \frac{4}{3\Delta - 4}.
\]

Note that \(\phi\) at time 0 equals 0, and can never be more than \(n\), and we have shown that \(\phi\) is expected to increase by a fixed positive amount at each step (when we formulate carefully as in the paragraph following the proof of Theorem 1). This fact together with standard martingale stopping theorems can then be used to show that the expected time needed for \(|V_i|\) to equal \(n\) is at most linear in \(n\).
2.1 Markov chain approaches

Several Markov chains for this problem exist [12] [2], together with techniques for using CFTP to obtain perfect samples [6] [7]. These Markov chains are known to mix in time \( O(n \ln n) \), and the corresponding perfect sampling algorithms are known to run in time \( O(n \ln n) \), when \( \lambda < 2/(\Delta-2) \), which is a larger range of \( \lambda \) than our method gives. However, when \( \lambda \) is small enough, the \( O(n) \) bound for our RR algorithm is smaller. It is hoped that with further refinement of the rejection step, the range of \( \lambda \) may be increased to where it matches the Markov chain analysis.

3 The Randomness Recycler

We now present a more general outline of the randomness recycler technique. Many state spaces \( \Omega \) of interest are of the form \( \Omega \subseteq C^V \), where \( C^V \) is the set of (proper or improper) colorings of a graph. Our goal is to sample from \( \Omega \) in expected time linear in \( |V| \). We have already seen how the independent sets of a graph may be encoded by coloring a vertex 1 if it is in the independent set and 0 otherwise. For another example, the set of permutations of \( n \) elements is a subset of \( \{1, \ldots, n\}^n \). Of course, the size of \( \Omega \) may be as large as \( |C|^{|V|} \), and this is in part what makes generating samples from these distributions difficult.

![Image](image)

The Randomness Recycler (Outline)

Set \( V_0 \leftarrow \emptyset \), \( X_0 \leftarrow \text{suitable} \ x_0 \), \( t \leftarrow 0 \)

Repeat

Set \( X_{t+1} \leftarrow X_t \)

Choose \( v \in V \setminus V_t \)

Randomly choose color \( c \) for \( v \)

Compute probability of accepting color \( c \) for \( v \)

If we accept

Set \( X_{t+1}(v) \leftarrow c \)

Set \( V_{t+1} \leftarrow V_t \cup \{v\} \)

Else

Set \( V_{t+1} \) and \( X_{t+1}\mid V\setminus V_{t+1} \) in a way that ‘undoes’ the effect of rejection

Set \( t \leftarrow t+1 \)

Until \( V_t = V \)

In an RR algorithm, a sample (i.e., one draw from \( \pi \)) is built up one vertex of \( V \) at a time until we include all of the vertices. Let \( V_t \) be the subset of vertices on which we have already built up a sample at time \( t \). On the vertices in \( V \setminus V_t \), the sample is fixed at some value, whereas on \( V_t \), the sample is random, and drawn exactly from the desired distribution. \( V_t \) starts out empty, and at each step of the algorithm we attempt to add a vertex to \( V_t \). Sometimes this is possible, and sometimes it is not. We continue in this fashion until \( V_t = V \), at which point we have a sample drawn exactly from the desired distribution. Let \( X_t \) denote the coloring of the graph \( V \) at time \( t \).

The way in which we randomly choose \( c \), compute the acceptance probability, and set \( V_{t+1} \) and \( X_{t+1}\mid V\setminus V_{t+1} \) in case of rejection will all depend on the target distribution \( \pi \). What differentiates this algorithm from an elementary stepwise rejection approach is our rejection step. Rather than starting over when rejection is faced, we keep as much of \( V_t \) as possible, "recycling" the coloring on \( V_{t+1} \).

At each time step \( t \) we keep track of the vertex set \( V_t \) together with the colors that are fixed on \( V \setminus V_t \). The state \( X_t \) is random over \( V_t \) while on \( V \setminus V_t \) it is deterministic. Let \( X_t^* = (V_t, X_t\mid V\setminus V_t) \), and for any possible value \( x^* = (S, x\mid V\setminus S) \) of \( X_t^* \), let \( \pi_\pi \) be \( \pi \) conditioned given that the colors of \( V \setminus S \) are as specified by \( x\mid V\setminus S \).

To achieve both the desired distribution and interruptibility, we want \( X_t \) to be random over \( V_t \) independent of the history \( X_t^* \) for \( t' < t \). In other words we want the identity

\[
P(X_t = x_t \mid X_t^* = x_t^*) = \pi_{x_t^*}(x_t),
\]

(1)
to hold. Indeed, if it does, then letting \( T \) denote the first time that \( V_T = V \), it follows easily that

\[
P(X_T = x \mid T = t) = \pi(x).
\]

Thus if (1) is satisfied for all \( t \), then at termination time \( T \) the RR algorithm returns a sample \( X_T \) that is distributed according to the desired distribution, and we have the interruptibility property that \( T \) and \( X_T \) are independent random variables.

Since \( V_0 \) is empty, it is easy to begin with \( X_0 \) from \( \pi x \). Let \( H_t := (x_0^*, x_1^*, \ldots, x_t^*) \) for notational convenience. We will say that step \( t + 1 \) preserves the correct distribution if

\[
P(X_{t+1} = x_{t+1} \mid H_t) \equiv \pi_{x_t^*}(x_{t+1})
\]

(1)

\[
P(X_{t+1} = x_{t+1} \mid H_{t+1}) \equiv \pi_{x_{t+1}^*}(x_{t+1}+1).
\]

This requirement that RR preserve the correct distribution is somewhat analogous to the design requirement that a Markov chain be reversible. It gives us a straightforward approach to designing an RR.

Just as the heat bath approach gives a means for designing Markov chains that are reversible, it also gives us a method for designing RR algorithms that preserve the correct distribution. For a specified vertex \( v \) in \( V \) and coloring \( z \), let \( \pi_{v}(\cdot ; z) \) denote the conditional probability distribution of \( X(v) \) given that \( X|V\setminus\{v\} = z|V\setminus\{v\} \) when \( X \) has the stationary distribution \( \pi \). From current state \( x \), the heat bath (or Gibbs sampler) Markov chain approach is to choose \( v \) uniformly at random and then choose a new color for \( v \) distributed according to \( \pi_{v}(\cdot ; x) \).
In heat bath RR, the vertex $v$ is chosen any way the user desires from $V \setminus V_1$, and then a new color is picked according to $\pi_v (\cdot | x)$. However, this color is not always accepted. We compute the acceptance probability as follows, with the goal being to preserve the correct distribution. According to Theorem 2 below, this goal is indeed met.

Given values $x_t^*, x_t, x_t^{t+1}$, and $x_{t+1}$ that correspond to a possible acceptance step in which vertex $v$ is added to the growing vertex set, define $\rho(x_t^*, x_t, x_t^{t+1}, x_{t+1})$ to be the ratio

$$\rho(x_t^*, x_t, x_t^{t+1}, x_{t+1}) := \frac{\pi_{x_{t+1}}^*(x_{t+1})}{\pi_v(x_{t+1}(v); x_t) \pi_{x_t^*}(x_t)}.$$

Also define

$$M(x_t^*, x_t^{t+1}) := \max_{x_t, x_t^{t+1}} \rho(x_t^*, x_t, x_t^{t+1}, x_{t+1}).$$

Then the probability that we accept a possible transition from $(x_t^*, x_t)$ to $(x_t^{t+1}, x_{t+1})$ is taken to be $\rho(x_t^*, x_t, x_t^{t+1}, x_{t+1}) / M(x_t^*, x_t^{t+1}).$

We do not have to use the heat bath probabilities. It is also valid to use the same acceptance probability, with the distributions $\pi_v (\cdot | x)$ replaced by arbitrary distributions $p_v (\cdot | x)$, when the distribution $p_v (\cdot | x)$ is used to color a selected $v$ when at a configuration $x$.

While these acceptance probabilities may appear daunting, for many problems they simplify considerably. For instance, in the independent set case, suppose first that $v$ has no neighbor colored 1. Then the heat bath probabilities are $1/(1 + \lambda)$ for color 0 and $\lambda/(1 + \lambda)$ for color 1. The acceptance probability in this first case will always be 1. If instead some neighbor of $v$ is colored 1, then heat bath assigns probability 1 to the color 1. The acceptance probability, however, works out to $1/(1 + \lambda)$. Careful examination of the independent set algorithm in Section 2 shows that this is exactly how the color for $v$ is chosen, with the same acceptance probabilities.

To show that the heat bath randomness recycler approach actually works (in general), we need to show that every step preserves the correct distribution. We will first consider acceptance steps, for which the following lemma gives a sufficient condition.

**Lemma 1** Given possible values $x_t^*$, $x_t^{t+1}$, and $x_{t+1}$ of $X_t^*$, $X_{t+1}$, and $X_t$, corresponding to an acceptance step, suppose that only one value $x_t$ of $X_t$ has positive probability. If the bivariate process $(X_t^*, X_t) \geq 2$ evolves Markovian and if for all such $x_t^*, x_t^{t+1}$, and $x_{t+1}$ and the single $x_t$ they determine we have

$$P(X_{t+1} = x_{t+1}, X_t = x_t) = \pi_{x_{t+1}}(x_{t+1}) C,$$

where $C$ does not depend on $x_t$ or $x_{t+1}$, then step $t + 1$ preserves the correct distribution.

**Proof** Let $C_t := 1/P(X_t^{t+1} = x_t^{t+1})$, and suppose that $P(X_t = x_t | H_t) = \pi_{x_t^*}(x_t)$. Let $E$ be the event that $X_{t+1} = x_{t+1} + X_t = x_t + 1$. Then

$$P(X_{t+1} = x_{t+1} | H_t) = C_t P(X_t^{t+1} = x_t^{t+1}, x_{t+1} = x_{t+1} | H_t)$$

$$= C_t P(E \cap \{ X_t = x_t \} | H_t)$$

$$= C_t P(X_t = x_t | H_t) P(E | H_t \cap \{ X_t = x_t \})$$

$$= C_t \pi_{x_t^*}(x_t) P(E | X_t^* = x_t^*, X_t = x_t)$$

$$= C_t C \pi_{x_{t+1}}(x_{t+1}),$$

where the last step is exactly our assumption.

Note that neither $C_t$ nor $C$ depends on $x_{t+1}$. Hence, summing over $x_{t+1},$

$$1 = C_t C \sum_{x_{t+1}} \pi_{x_{t+1}}(x_{t+1}) = C_t C.$$

This completes the proof. $\square$

**Theorem 2** The heat bath RR and arbitrary RR acceptance steps preserve the correct distribution.

**Proof** The acceptance probabilities were chosen precisely to match the requirements of Lemma 1. For instance, with heat bath RR, the left side of the equation in Lemma 1 equals

$$\pi_v(x_{t+1}(v); x_t) \times \frac{\pi_{x_{t+1}}^*(x_{t+1})}{\pi_v(x_{t+1}(v); x_t) \pi_{x_t^*}(x_t) M(x_t^*, x_t^{t+1})} \times \pi_{x_t^*}(x_t),$$

which reduces to the right side of the equation with $C = 1/M(x_t^*, x_t^{t+1})$. The calculation for arbitrary RR is entirely similar. $\square$

Now we turn our attention to rejection steps. In designing an RR algorithm, it is our experience that proper handling of rejection steps to ensure preservation of the correct distribution is more difficult and problem-specific to arrange than is proper handling of acceptance steps. But here are some broad guiding comments.

Determination of the acceptance probability at step $t + 1$ will reveal knowledge about the colors of some subset, call it $D_t$, of $V_t$. If we reject, we then set $V_{t+1}$ to be $V_t \setminus D_t$. This insures that when we reject, we do not bias the sample. That is, by removing $D_t$ from $V_t$, we remove all traces of our knowledge gained, and as a result the remaining sample is drawn exactly from $\pi_{V_{t+1}}$.

In the case of the independent sets, the set $D_t$ consists of precisely those vertices prescribed to be removed by the algorithm: $w$ and all its neighbors, and neighbors of $v$ with numbers lower than that of $w$. Indeed, all of these vertices
are colored 0 at time $t$, except for vertex $w$, which is colored 1.] It is not hard to check rigorously in this case that rejection steps also preserve the correct distribution, but we omit the details.

4 Applications

This section applies the randomness recycler approach to several different problems of interest. For some of these models we have theoretical bounds on the running time, while for others we have only experimental results.

The Ising and Potts models In the Ising model, vertices in a graph $(V, E)$ are colored from the set $\{-1, 1\}$. The distribution $\pi$ from which we wish to sample is defined by

$$\pi(x) := \frac{\exp(-\beta J H(x))}{Z_\beta},$$

where

$$H(x) := -\sum_{\{v_1, v_2\} \in E} x(v_1) x(v_2)$$

is known as the energy of the coloring, $\beta$ is (proportional to) a positive parameter known as inverse temperature, and $J$ is 1 in the ferromagnetic model and $-1$ in the antiferromagnetic model. Generating approximate samples may be done in (nonlinear) polynomial time in the ferromagnetic case using Markov chain techniques of Jerrum and Sinclair [10] [16].

The RR approach has provably linear expected running time for both the ferromagnetic and antiferromagnetic models when $\beta$ is small (i.e., the temperature is high). The set $D_t$ to be removed from $V_t$ in case of rejection is just the set of neighbors of the vertex $v$ that we tried to add. Omitting details and proofs, we simply state the running time bound in the following theorem.

Theorem 3 Let $\Delta$ be the maximum degree of the graph. If $e^\beta < \left(1 + \frac{1}{\Delta}\right)^{\frac{1}{2}}$, then the expected running time of the heat bath RR procedure for the Ising model is $O(n)$.

Comments like those following the proof of Theorem 1 apply here, where now the expected increase $\frac{1}{1 + \beta} \left[1 - (2\Delta - 1) e^{-\beta\Delta}\right]$ in $|V_t|$ becomes $(\Delta + 1) e^{-\beta\Delta} - \Delta$.

Although not needed for the theorem, in practice it helps to introduce a third color 0 to supplement $\{-1, 1\}$. Notice that no edge with an endpoint colored 0 contributes to $H$. At the completion of step $t$, every vertex in $V \setminus V_t$ which is surrounded entirely by vertices in $V \setminus V_t$ may be recolored 0 since this action does not affect the vertices in $V_t$ at all.

The Potts model differs from the Ising model in that more than two colors are used, but the energy depends (in a natural way) only on whether edges are colored concordantly or discordantly, and the running time Theorem 3 remains valid verbatim.

The Random Cluster Model The random cluster model is an extension of the Potts model to noninteger numbers of colors [4]; this is discussed further below. Unlike our previous examples, which colored vertices, the random cluster model colors edges of a given graph $G = (V, E)$ with colors from $\{0, 1\}$. If $A$ is the set of edges colored 1, then the distribution is

$$\pi(A) := p^{\vert A \vert} (1 - p)^{|E \setminus A|} q^{e(A)}/Z_{p,q}, \quad A \subseteq E,$$

where $p \in [0, 1]; q > 0$ is not necessarily an integer, and we shall assume $q > 1; c(A)$ is the number of connected components in the graph $(V, A)$; and $Z_{p,q}$ is a normalizing constant.

The RR approach is as follows. We represent a set $A \subseteq E$ by a binary vector $x$, by setting $x(e) = 1$ for $e \in A$, and $x(e) = 0$ otherwise. At each step, we keep track of such a vector $x_t$ and a set $E_t$ of edges, namely, the edges on which $x_t$ is random; all other edges will be colored 0. We choose an oriented edge $e = (v, w) \in E \setminus E_t$, until such an edge $e$ no longer exists. We set $x_{t+1}(e) = 1$ with probability $p$, and $x_{t+1}(e) = 0$ with probability $1 - p$. If $v$ and $w$ are already connected in $x_t$ (i.e., in the graph $(V, A_t)$ where $A_t = \{e' : x_t(e') = 1 \} \subseteq E_t$), then we accept the edge and set $E_{t+1} = E_t \cup \{e\}$. If $v$ and $w$ are not already connected, then we always accept $x_{t+1}(e) = 0$, but we accept $x_{t+1}(e) = 1$ only with probability $1/q$ (since by adding this edge we reduce by 1 the number of connected components).

When we reject, we know that $v$ and $w$ lie in separate components in $(V, A_t)$. To counteract this knowledge, to form $E_{t+1}$ we remove from $E_t$ all the edges in the component of $(V, A_t)$ that contains $w$, together with all edges of $E_t$ that lead out of this component (and which therefore do not belong to $A_t$).

We could cease our handling of a rejection step at this point and prove that (a) the algorithm works correctly and (b) Theorem 4 below holds (and the proof simplifies somewhat) with the bound on $p$ decreased to

$$p < 1/((\Delta - (1/q))).$$

However, we shall omit the formal proof of correctness and instead discuss a small (provably valid) trick which gains us some efficiency.

Suppose that there are $M$ vertices in the removed component. Consider the (connected!) graph consisting of the
vertices and edges in this component, together with the vertex \( v \) and the edge \( \{v, w\} \). Choose (in any fashion) a spanning tree \( T \) of this graph; \( T \) will comprise \( M + 1 \) vertices and therefore \( M \) edges. Add back all these \( M \) edges to get \( E_{t+1} \). Sample from the random cluster model on \( T \), and add back in the edges thereby colored \( 1 \) to get \( A_{t+1} \).

The key observation here is that it is elementary to sample from the random cluster model when the graph is a tree. Indeed, then each edge independently is colored \( 1 \) with probability \( \rho/(1 - p + \rho) \) and \( 0 \) with probability \((1 - p)/(1 - p + \rho)\), where

\[
p := \frac{p}{q}.
\]

The random cluster model is an extension of the ferromagnetic Ising and Potts models. When \( q > 1 \) is an integer, and \( p = 1 - \exp(-\beta) \), then samples from the random cluster model may be used to generate samples from the ferromagnetic Potts model with \( q \) colors by independently taking each connected component of \( (V, A) \), uniformly choosing one of the \( q \) colors, and assigning to every vertex in the component that color. For certain instances of the random cluster model, the heat bath Markov chain approach is believed from experimental evidence to be rapidly mixing [15], but no theoretical rapid mixing results in the positive direction are known for any nontrivial instances of the problem. For some instances, the Markov chain approach is known not to be rapidly mixing [5]. For the RR approach, we know that when \( p \) is small (corresponding to small \( \beta \)), the approach takes an expected number of steps which is linear in the number of edges:

**Theorem 4** Suppose that

\[
p < \frac{\Delta - (1/q) - \sqrt{[\Delta - (1/q)]^2 - 4[1 - (1/q)][\Delta - 1]}}{2[1 - (1/q)][\Delta - 1]}.
\]

Then the expected number of steps required by the RR algorithm is \( O(|E|) \).

For example, if \( \Delta = 4 \) (as on a 2-dimensional rectangular grid) and \( q = 2 \) (corresponding to the Ising model), then our restriction is that \( p < 1/3 \); this improves on the restriction \( p < 1/2(\Delta - 1/q) \) \( = 2/7 \) obtained when the “add a tree” trick is not employed.

Comments analogous to those following the proof of Theorem 1 again apply.

**Proof** We use a potential function that rewards us for adding edges and penalizes us for connecting components. Let

\[
\phi(E_t, A_t) := |E_t| - \alpha c(A_t),
\]

where \( \alpha \) will be determined later.

When the edge \( \{v, w\} \) we attempt to add to \( E_t \) is between two vertices already connected in \( A_t \), then \( \phi \) always goes up by 1, making this case uninteresting. It is when \( \{v, w\} \) would connect two previously unconnected components of \( A_t \) that the calculation becomes interesting.

If the edge is chosen to be excluded from \( A_{t+1} \), then \( \phi \) increases by 1. If the edge is proposed to be included in \( A_{t+1} \), then \( \phi \) changes by \( 1 - \alpha \) if we accept. If we reject, we remove from \( A_t \) (and also from \( E_t \)) a component of size \( M \) and (from \( E_t \)) all of its adjacent edges. Not counting the edge \( \{v, w\} \) and making sure that we do not double-count, this totals at most \( M(\Delta - 1) \) edges removed from \( E_t \). However, we add exactly \( M - 1 \) new components to \( A_t \) by removing these edges. When we add the tree \( T \) back in, this produces \( M \) new edges for \( E_{t+1} \), but for each such edge there is a \( p/(1 - p + \rho) \) chance of including the edge in \( A_{t+1} \) and thereby reducing the number of components by 1. Therefore, when we attempt to add \( \{v, w\} \) to \( A_{t+1} \), but reject instead, the expected contribution to the change in \( \phi \) is at least

\[
-M(\Delta - 1) + M + \alpha \left( M - 1 - M \frac{\rho}{1 - p + \rho} \right).
\]

Now \( M \) may be very large (nearly as large as \( n \)), so we choose \( \alpha \) in such a way that the coefficient of \( M \) in this expression vanishes. That is, we set

\[
\alpha := (\Delta - 2)\frac{1 - p + \rho}{1 - p}
\]

and so the contribution in this case is bounded below by \(-\alpha\).

We try to put the edge in with probability \( p \) and to leave it out with probability \( 1 - p \). We accept an inclusion with probability \( 1/q \). Putting everything together, we find that the expected change in \( \phi \) at any time step when \( u \) and \( w \) are not already connected in \( A_t \) is at least

\[
(1 - p) + p \left[ \frac{1}{q} (1 - \alpha) + \left(1 - \frac{1}{q}\right) (-\alpha) \right],
\]

which is positive exactly when

\[
p < \frac{\Delta - (1/q) - \sqrt{[\Delta - (1/q)]^2 - 4[1 - (1/q)][\Delta - 1]}}{2[1 - (1/q)][\Delta - 1]}.
\]

\( \Box \)

In this case, the Markov chain approach does not have theoretical guarantees on the running time for any nontrivial value of \( p \). While coupling from the past may also be used to generate perfect samples, there is no a priori bound on its running time.

As with CFTP, we may still use the RR approach for values of \( p \) for which no theoretical bound exists. We simply do not know beforehand how long the algorithm will take. Unlike CFTP, the RR approach is interruptible, so we may abort the procedure if it needs too many steps, without introducing bias into the sample.
Proper colorings of a graph  Finding the number of proper colorings of a graph is a \#P-complete problem \[9\].
Recall that a proper coloring of a graph assigns each vertex a color such that no edge has both endpoints colored the same color. The ability to sample from the set of proper colorings leads to an approximation algorithm for counting the number of such colorings.

Markov chain approaches require that \(k\), the number of colors, be at least \((11/6)\Delta\) (where \(\Delta\) is again the maximum degree of the graph) \[19\] in order to guarantee rapid mixing for the chain. Perfect sampling using bounding chains \[7, 6\] is only guaranteed to run in polynomial time when the number of colors is \(\Omega(\Delta^2)\). Unfortunately, the straightforward RR approach does not match these bounds. Somewhat roughly stated,

**Theorem 5** The heat bath RR approach to generating perfect colorings requires only a linear expected number of steps when \(k\) is \(\Omega(\Delta^4)\).

As with the bounding chain procedure, however, this algorithm may be run even when \(k\) is much smaller; we simply have no reasonable a priori bound on the running time in such cases.

The Move Ahead 1 chain  Finally, we present a problem where an RR-based algorithm seems experimentally to run fast although we cannot give any theoretical bounds. In the list update problem, a set of items is kept in a list. To access an item, a user starts at the beginning of the list and steps through the items until the desired item is located. The located item may be replaced in the list anywhere between its current position and the front of the list, at fixed cost. The goal is to use a replacement strategy that keeps small the access times (i.e., item depths in the list) needed for items.

Call the strategy which moves the accessed item to the front of the list the **Move to Front** (MTF) rule. A worst-case analysis shows that the MTF rule yields a 2-approximation for the optimal total access time for any sequence of item requests \[18\]. Alternatively, it is useful to employ probabilistic models to describe how list items are chosen to be accessed. Commonly, such an access model will induce a Markov chain model on the evolution of the order of the list. Characteristics such as the limiting distribution as \(t \to \infty\) of \(A_t\), where \(A_t\) is the access time for the item accessed at time \(t\), can then be estimated by drawing from the stationary distribution of the chain.

To be specific, label the items with identification numbers \(1, \ldots, n\); suppose that at each time step, independently of previous time steps, any particular item \(i\) is accessed with probability \(p_i > 0\) (independently of the order of the list); and suppose that after each selection, the accessed item is moved forward one rank in the list, i.e., is transposed with its predecessor in the list. (If the accessed item is already at the front of the list, the order of the list is left unchanged.) The self-organization rule we have described is called the **Move Ahead 1** (MA1) rule. The limiting expected access time for MA1 is known to be, for any access probability vector \(p\), no more than that for MTF \[17\]. Further Monte Carlo study of the limiting access time distribution is complicated by the fact that sampling from the limiting list-order distribution \(\pi\) (for which a formula is known, but only up to a normalizing constant) seems to be quite difficult in general.

Coupling from the past approaches to sampling from \(\pi\) exist \[8\], but experimental evidence suggests that use of RR gives a faster algorithm. Suppose that \(p_i \propto r^i\) for some ratio \(0 < r < 1\). Then experimental evidence suggests that for each fixed value of \(r \in (0, 1]\) the expected running time is linear in \(n\), although the constant of linearity does vary with \(r\). The Markov chain approach to this problem is only known to be rapidly mixing when \(r < 0.2\) \[8\].

5 Conclusion

The RR approach to perfect sampling gives exact samples from difficult distributions without using the traditional Markov chain. It is quite different from other recent approaches to perfect sampling such as coupling from the past.

Because it dispenses with the Markov chain, the RR approach yields, for restricted versions of some of these problems, the first expected linear time algorithms for these problems. Even when the running time of RR is unknown, the algorithm may be run and the output will be guaranteed to come from the correct distribution.

Unlike coupling from the past, RR is interruptible, so the user may set a time limit on the algorithm’s running time (if measured in number of iterations of the basic Repeat loop) without introducing bias into the sample. Like read-once coupling from the past \[20\], this algorithm does not require storage of any random bits. (Another perfect sampling approach, that of Fill, Machida, Murdoch, and Rosenthal \[3\] is also interruptible but not read-once, and so does requires storage of random bits. We wish to stress that these existing means for perfect sampling rely on finding a “good” Markov chain for the problem at hand. RR does away with the chain, and in doing so breaks the \(O(n \ln n)\) barrier that has characterized so many of these problems.

For independent sets and for proper colorings, the theoretical bounds obtained apply only for a more restricted set of parameters than do those based on Markov chain approaches. However, when the appropriate restriction is met, our RR methods are faster, yielding samples in a linear (expected) number of steps. Moreover, much work has gone into analyses of Markov chains, while our work is still rather new, and we might hope with time and further effort eventually to match or even to relax the restrictions needed...
for the Markov chain approaches. For the Move Ahead 1 chain we do not know any theoretical bounds on the running time of our method. However, computer experiments show that for this problem the RR method works much better in practice than does the CFTP method.

For the random cluster model, our RR technique is guaranteed to run in a linear (expected) number of steps for a range of values of $p$. This is in sharp constrast to the Markov chain approach, where no polynomial running time bounds are known except in trivial cases.

In summary, the randomness recycler is not applicable in all situations where Markov chain approaches are used, but RR often gives a fast read-once interruptible means for generating perfect samples that in restricted cases gives the first linear time algorithms for some difficult and important problems.

References


