# SLICE SAMPLING ${ }^{1}$ 

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#### Abstract

Markov chain sampling methods that adapt to characteristics of the distribution being sampled can be constructed using the principle that one can sample from a distribution by sampling uniformly from the region under the plot of its density function. A Markov chain that converges to this uniform distribution can be constructed by alternating uniform sampling in the vertical direction with uniform sampling from the horizontal "slice" defined by the current vertical position, or more generally, with some update that leaves the uniform distribution over this slice invariant. Such "slice sampling" methods are easily implemented for univariate distributions, and can be used to sample from a multivariate distribution by updating each variable in turn. This approach is often easier to implement than Gibbs sampling and more efficient than simple Metropolis updates, due to the ability of slice sampling to adaptively choose the magnitude of changes made. It is therefore attractive for routine and automated use. Slice sampling methods that update all variables simultaneously are also possible. These methods can adaptively choose the magnitudes of changes made to each variable, based on the local properties of the density function. More ambitiously, such methods could potentially adapt to the dependencies between variables by constructing local quadratic approximations. Another approach is to improve sampling efficiency by suppressing random walks. This can be done for univariate slice sampling by "overrelaxation," and for multivariate slice sampling by "reflection" from the edges of the slice.


1. Introduction. Markov chain methods such as Gibbs sampling and the Metropolis algorithm can be used to sample from many of the complex, multivariate distributions encountered in statistics. However, to implement Gibbs sampling, one may need to devise methods for sampling from nonstandard univariate distributions, and to use the Metropolis algorithm, one must find an appropriate "proposal" distribution that will lead to efficient sampling. The need for such special tailoring limits the routine use of these methods and inhibits the development of software that automatically constructs Markov chain samplers from model specifications. Furthermore, many common Markov chain samplers are inefficient, due to a combination of two flaws. First, they may try to make changes that are not well adapted to the local properties of the density function,

[^0]with the result that changes must be made in small steps. Second, these small steps take the form of a random walk, in which about $n^{2}$ such steps are needed in order to move a distance that could be traversed in only $n$ steps if these steps moved consistently in one direction.

In this paper, I describe a class of "slice sampling" methods that can be applied to a wide variety of distributions. Simple forms of univariate slice sampling are an alternative to Gibbs sampling that avoids the need to sample from nonstandard distributions. These slice sampling methods can adaptively change the scale of changes made, which makes them easier to tune than Metropolis methods and also avoids problems that arise when the appropriate scale of changes varies over the distribution. More complex slice sampling methods can adapt to the dependencies between variables, allowing larger changes than would be possible with Gibbs sampling or simple Metropolis methods. Slice sampling methods that improve sampling by suppressing random walks can also be constructed.

Slice sampling originates with the observation that to sample from a univariate distribution, we can sample points uniformly from the region under the curve of its density function and then look only at the horizontal coordinates of the sample points. A Markov chain that converges to this uniform distribution can be constructed by alternately sampling uniformly from the vertical interval defined by the density at the current point and from the union of intervals that constitutes the horizontal "slice" though the plot of the density function that this vertical position defines. If this last step is difficult, we may substitute some other update that leaves the uniform distribution over the slice invariant. To sample from a multivariate distribution, such single-variable slice sampling updates can be applied to each variable in turn. Section 4 presents these single-variable slice sampling methods.

We can also apply the slice sampling approach to a multivariate distribution directly, as described in Section 5, by sampling uniformly under the multidimensional plot of its density function. As for a univariate distribution, this can be done by alternately sampling uniformly from the vertical interval from zero up to the density at the current point and then uniformly from the slice defined by this vertical position. When the slice is high-dimensional, how to sample efficiently from it is less obvious than for single-variable slice sampling, but one gains the possibility of sampling in a way that respects the dependencies between variables. I show how, in the context of slice sampling, the way changes are proposed can be adapted to respect these dependencies, based on local information about the density function. In particular, local quadratic approximations could be constructed, as have been used very successfully for optimization problems. However, further research will be needed to fully exploit the adaptive capabilities of multivariate slice sampling.

One might instead accept that dependencies between variables will lead to the distribution being explored in small steps, but try at least to avoid exploring the distribution by an inefficient random walk, which is what happens when simple forms of the Metropolis algorithm are used. The benefits of random walk suppression are analyzed theoretically in some simple contexts by Diaconis,

Holmes and Neal (2000). Large gains in sampling efficiency can be obtained in practice when random walks are suppressed using the hybrid Monte Carlo or other dynamical methods [Duane, Kennedy, Pendleton and Roweth (1987), Horowitz (1991), Neal (1994, 1996)], or by using an overrelaxation method [Adler (1981), Barone and Frigessi (1990), Green and Han (1992), Neal (1998)]. Dynamical and overrelaxation methods are not always easy to apply, however. Use of Markov chain samplers that avoids random walks would be assisted by the development of methods that require less special programming and parameter tuning.

Two approaches to random walk suppression based on slice sampling are discussed in this paper. In Section 6, I show how one can implement an overrelaxed version of the single-variable slice sampling scheme. This may provide the benefits of Adler's (1981) Gaussian overrelaxation method for more general distributions. In Section 7, I describe slice sampling analogues of dynamical methods, which move around a multivariate slice using a stepping procedure that proceeds consistently in one direction while reflecting off the slice boundaries. These more elaborate slice sampling methods require more tuning than single-variable slice sampling, but they may still be easier to apply than alternative methods that avoid random walks.

I illustrate the benefits of the adaptive nature of slice sampling in Section 8, showing that it avoids disaster when sampling from a distribution typical of priors for hierarchical Bayesian models. Simple Metropolis methods can give the wrong answer for this problem, while providing little indication that anything is amiss.

I conclude with a discussion of the merits of the various slice sampling methods in comparison with other Markov chain methods and of their suitability for routine and automated use. Below, I set the stage by discussing general-purpose Markov chain methods that are currently in wide use.
2. General-purpose Markov chain sampling methods. Applications of Markov chain sampling in statistics often involve sampling from many distributions, such as posterior distributions for parameters of various different models, given various different datasets. For routine use of Markov chain methods, it is important to minimize the amount of effort that the data analyst must spend in order to sample from all these distributions. Ideally, a Markov chain sampler would be constructed automatically for each model and dataset.

The Markov chain method most commonly used in statistics is Gibbs sampling, popularized by Gelfand and Smith (1990). Suppose that we wish to sample from a distribution over $n$ state variables (e.g., model parameters), written as $x=\left(x_{1}, \ldots, x_{n}\right)$, with probability density $p(x)$. Gibbs sampling proceeds by sampling in succession from the conditional distributions for each $x_{i}$ given the current values of the other $x_{j}$ for $j \neq i$, with conditional densities written as $p\left(x_{i} \mid\left\{x_{j}\right\}_{j \neq i}\right)$. Repetition of this procedure defines a Markov chain which leaves the desired distribution invariant, and which in many circumstances is ergodic [e.g., when $p(x)>0$ for all $x$ ]. Running the Gibbs sampler for a sufficiently
long time will then produce a sample of values for $x$ from close to the desired distribution, from which we can estimate the expectations of those functions of state that are of interest.

Gibbs sampling can be done only if we know how to sample from all the required conditional distributions. These sometimes have standard forms for which efficient sampling methods have been developed, but there are many models for which sampling from these conditional distributions requires the development of custom algorithms, or is infeasible in practice (e.g., for multilayer perceptron networks [Neal (1996)]). Note, however, that once methods for sampling from these conditional distributions have been found, no further tuning parameters need be set in order to produce the final Markov chain sampler.

The routine use of Gibbs sampling has been assisted by the development of adaptive rejection sampling (ARS) [Gilks and Wild (1992), Gilks (1992)], which can be used to sample efficiently from any conditional distribution whose density function is log concave, given only the ability to compute some function, $f_{i}\left(x_{i}\right)$, that is proportional to the conditional density, $p\left(x_{i} \mid\left\{x_{j}\right\}_{j \neq i}\right)$ [the ability to also compute the derivative, $f_{i}^{\prime}\left(x_{i}\right)$, is helpful, but not essential]. This method has been used for some time by the BUGS software [Thomas, Spiegelhalter and Gilks (1992)] to automatically generate Markov chain samplers from model specifications. The first step in applying ARS is to find points on each side of the mode of the conditional distribution. This will in general require a search, which will in turn require the choice of some length scale for an initial step. However, the burden of setting this scale parameter is lessened by the fact that a good value for it can be chosen "retrospectively," based on past iterations of the Markov chain, without invalidating the results, since the value chosen affects only the computation time, not the distribution sampled from.

The adaptive rejection Metropolis sampling (ARMS) method [Gilks, Best and Tan (1995)] generalizes ARS to conditional distributions whose density functions may not be log-concave. However, when the density is not log-concave, ARMS does not produce a new point drawn independently from the conditional distribution, but merely updates the current point in a fashion that leaves this distribution invariant. Also, when a conditional distribution is not log-concave, the points used to set up the initial approximation to it must not be chosen with reference to past iterations, as this could result in the wrong distribution being sampled [Gilks, Neal, Best and Tan (1997)]. The initial approximation must be chosen based only on prior knowledge (including any preliminary Markov chain sampling runs), and on the current values of the other variables. Unlike ARS, neither the current value of the variable being updated, nor any statistics collected from previous updates (e.g., the typical scale of changes) can be used. This hinders routine use of the method.

Another general way of constructing a Markov chain sampler is to perform Metropolis updates [Metropolis, Rosenbluth, Rosenbluth, Teller and Teller (1953),

Hastings (1970)], either for each variable in turn, as with Gibbs sampling, or for all variables simultaneously. A Metropolis update starts with the random selection of a "candidate" state, drawn from a "proposal" distribution. The candidate state is then accepted or rejected as the new state of the Markov chain, based on the ratio of the probability densities of the candidate state and the current state. If the candidate state is rejected, the new state is the same as the old state.

A simple "random-walk" Metropolis scheme can be constructed based on a symmetric proposal distribution (e.g., Gaussian) that is centred on the current state. All variables could be updated simultaneously in such a scheme, or alternatively, one variable could be updated at a time. In either case, a scale parameter is required for each variable, in order to fix the width of the proposal distribution in that dimension. For the method to be valid, these scale parameters must not be set on the basis of past iterations, but rather only on the basis of prior knowledge (including preliminary runs), and the current values of too large a scale for the proposal distribution will result in a high rejection rate, while choosing too small a scale will result in inefficient exploration via a random walk with unnecessarily small steps. Furthermore, the appropriate scale for Metropolis proposals may vary from one part of the distribution to another, in which case no single value will produce acceptable results. Selecting a scale at random from some range can sometimes alleviate these problems, but at a large cost in wasted effort whenever the scale selected is inappropriate.

It is tempting to tune the Metropolis proposal distribution based on the rejection rate in past iterations of the Markov chain, but such "retrospective tuning" is not valid, since it can disturb the stationary distribution to which the process converges. Fixing the proposal distribution based on a preliminary run is allowed, but if the original proposal distribution was not good, such a preliminary run may not have sampled from the whole distribution, and hence may be a bad guide for tuning.

We therefore see that although Gibbs sampling and Metropolis methods have been used to do much useful work, there is a need for better methods that can be routinely applied in a wider variety of situations. One aim of this paper is to find variations on slice sampling that can be used to sample from any continuous distribution, given only the ability to evaluate a "black-box" function that is proportional to its density, and in some cases, to also evaluate the gradient of this function. For many distributions, these new methods will not sample more efficiently than Gibbs sampling or a well-designed Metropolis scheme, but the slice sampling methods will often require less effort to implement and tune. For some distributions, however, slice sampling can be much more efficient, because it can adaptively choose a scale for changes appropriate to the region of the distribution currently being sampled. Slice samplers that adapt in more elaborate ways, or that suppress random walks, can potentially be much faster than simple Metropolis methods or Gibbs sampling.
3. The idea of slice sampling. Suppose we wish to sample from a distribution for a variable, $x$, taking values in some subset of $\mathfrak{R}^{n}$, whose density is proportional to some function $f(x)$. We can do this by sampling uniformly from the $(n+1)$-dimensional region that lies under the plot of $f(x)$. This idea can be formalized by introducing an auxiliary real variable, $y$, and defining a joint distribution over $x$ and $y$ that is uniform over the region $U=\{(x, y): 0<y<f(x)\}$ below the curve or surface defined by $f(x)$. That is, the joint density for $(x, y)$ is

$$
p(x, y)= \begin{cases}1 / Z, & \text { if } 0<y<f(x)  \tag{1}\\ 0, & \text { otherwise }\end{cases}
$$

where $Z=\int f(x) d x$. The marginal density for $x$ is then

$$
\begin{equation*}
p(x)=\int_{0}^{f(x)}(1 / Z) d y=f(x) / Z \tag{2}
\end{equation*}
$$

as desired. To sample for $x$, we can sample jointly for $(x, y)$, and then ignore $y$.
Generating independent points drawn uniformly from $U$ may not be easy, so we might instead define a Markov chain that will converge to this uniform distribution. Gibbs sampling is one possibility: We sample alternately from the conditional distribution for $y$ given the current $x$, which is uniform over the interval $(0, f(x))$, and from the conditional distribution for $x$ given the current $y$, which is uniform over the region $S=\{x: y<f(x)\}$, which I call the "slice" defined by $y$. Generating an independent point drawn uniformly from $S$ may still be difficult, in which case we can substitute some update for $x$ that leaves the uniform distribution over $S$ invariant. Higdon (1996) has interpreted the standard Metropolis algorithm in these terms. Beyond this, however, reducing the problem to that of updating $x$ so as to leave a uniform distribution invariant allows us to use various tricks that would not be valid for a nonuniform distribution.

Related methods have been used in the past. Chen and Schmeiser (1998) describe a method that samples from the distribution over the region $U$ by moving in random directions. Their work shares with this paper the aim of finding a method that requires little or no tuning, and hence is suitable for use as a "black-box" sampler, when little is known of the distribution being sampled from. Unfortunately, Chen and Schmeiser do not achieve this goal: though they intend that causal users fix the $a_{r}$ and $b_{r}$ parameters of their sampler at default values, these parameters in fact play essentially the same role as the proposal width in a simple random-walk Metropolis algorithm, and will sometimes have to be tuned if reasonable performance is to be achieved. This tuning is harder than for simple Metropolis updates, since the optimal setting of these parameters depends not just on the properties of the distribution but also on the normalizing constant, $Z$. In their main example, this problem is lessened by the way they periodically adjust the normalization constant based on the current point, but this state-
dependent adjustment destroys the reversibility of the transitions, undermining the correctness of the algorithm.

The highly successful Swendsen-Wang algorithm for the Ising model can also be seen as an auxiliary variable method, which led to its generalization by Edwards and Sokal (1988). In their scheme, the density (or probability mass) function is proportional to a product of $k$ functions: $p(x) \propto f_{1}(x) \cdots f_{k}(x)$. They introduce $k$ auxiliary variables, $y_{1}, \ldots, y_{k}$, and define a joint distribution for $\left(x, y_{1}, \ldots, y_{k}\right)$ which is uniform over the region in which $0<y_{i}<f_{i}(x)$ for $i=1, \ldots, k$. Gibbs sampling, or some other Markov chain procedure, can then be used to sample for $\left(x, y_{1}, \ldots, y_{k}\right)$. The slice sampling procedure described above is a special case of this procedure, when there is a single auxiliary variable (i.e., $k=1$ ). Besag and Green (1993) and Higdon (1996) have discussed applications in image analysis of these methods with $k>1$.

Mira and Tierney (2002) have shown that these auxiliary variable methods, with one or with many auxiliary variables, are uniformly ergodic under certain conditions. Roberts and Rosenthal (1999) have shown that these methods are geometrically ergodic under weaker conditions, and have also found some quantitative convergence bounds. These results all assume that the sampler generates a new value for $x$ that is uniformly drawn from $S$, independently of the old value, which is often difficult in practice.

Concurrently with the work reported here, Damien, Wakefield and Walker (1999) have viewed methods based on multiple auxiliary variables as a general approach to constructing Markov chain samplers for Bayesian inference problems. They illustrate how one can often decompose $f(x)$ into a product of $k$ factors for which the intersection of the sets $\left\{x: y_{i}<f_{i}(x)\right\}$ is easy to compute. This leads to an easily implemented sampler, but convergence is slowed by the presence of many auxiliary variables. For example, for a model of $k$ i.i.d. data points, one simple approach (similar to some examples of Damien, Wakefield and Walker) is to have one factor for each data point, whose product is the likelihood. (Suppose that prior is uniform, and so need not be represented in the posterior density.) For many models, $\left\{x: y_{i}<f_{i}(x)\right\}$ will be easy to compute when $f_{i}$ is the likelihood from one data point. However, if this approach is applied to $n$ data points that are modeled as coming from a Gaussian distribution with mean $\mu$ and variance 1 , one can show that after the $y_{i}$ are chosen, the allowable range for $\mu$ will have width of order $1 / n$. Since the width of the posterior distribution for $\mu$ will be of order $1 / \sqrt{n}$, and since the posterior will be explored by a random walk, the convergence time will be of order $n$. Gibbs sampling would, of course, converge in a single iteration when there is only one parameter, and the slice sampling methods of this paper would also converge very rapidly, for any $n$. Using a large number of auxiliary variables is a costly way to avoid difficult computations.

I therefore am concerned in this paper with methods based on slice sampling with a single auxiliary variable. So that these methods will be practical for a wide range of problems, they often use updates for $x$ that do not produce a point drawn independently from the slice, $S$, but merely change $x$ in some fashion that leaves the uniform distribution over $S$ invariant. This allows the methods to be used for any continuous distribution, provided only that we can compute some function, $f(x)$, that is proportional to the density.
4. Single-variable slice sampling methods. Slice sampling is simplest when only one (real-valued) variable is being updated. This will of course be the case when the distribution of interest is univariate, but more typically, the singlevariable slice sampling methods of this section will be used to sample from a multivariate distribution for $x=\left(x_{1}, \ldots, x_{n}\right)$ by sampling repeatedly for each variable in turn. To update $x_{i}$, we must be able to compute a function, $f_{i}\left(x_{i}\right)$, that is proportional to $p\left(x_{i} \mid\left\{x_{j}\right\}_{j \neq i}\right)$, where $\left\{x_{j}\right\}_{j \neq i}$ are the values of the other variables.

To simplify notation, I will here write the single real variable being updated as $x$ (with subscripts denoting different such points, not components of $x$ ). I will write $f(x)$ for the function proportional to the probability density of $x$. The singlevariable slice sampling methods discussed here replace the current value, $x_{0}$, with a new value, $x_{1}$, found by a three-step procedure:
(a) Draw a real value, $y$, uniformly from $\left(0, f\left(x_{0}\right)\right.$ ), thereby defining a horizontal "slice": $S=\{x: y<f(x)\}$. Note that $x_{0}$ is always within $S$.
(b) Find an interval, $I=(L, R)$, around $x_{0}$ that contains all, or much, of the slice.
(c) Draw the new point, $x_{1}$, from the part of the slice within this interval.

Step (a) picks a value for the auxiliary variable that is characteristic of slice sampling. Note that there is no need to retain this auxiliary variable from one iteration of the Markov chain to the next, since its old value is forgotten at this point anyway. In practice, it is often safer to compute $g(x)=\log (f(x))$ rather than $f(x)$ itself, in order to avoid possible problems with floating-point underflow. One can then use the auxiliary variable $z=\log (y)=g\left(x_{0}\right)-e$, where $e$ is exponentially distributed with mean one, and define the slice by $S=\{x: z<g(x)\}$.

Steps (b) and (c) can potentially be implemented in several ways, which must of course be such that the resulting Markov chain leaves the distribution defined by $f(x)$ invariant. Figure 1 illustrates one generally applicable method, in which the interval is found by "stepping out," and the new point is drawn with a "shrinkage" procedure. Figure 2 illustrates an alternative "doubling" procedure for finding the interval. These and some other variations are described in detail next, followed by a proof that the resulting transitions leave the correct distribution invariant.


FIG. 1. A single-variable slice sampling update using the stepping-out and shrinkage procedures. A new point, $x_{1}$, is selected to follow the current point, $x_{0}$, in three steps. (a) A vertical level, $y$, is drawn uniformly from $\left(0, f\left(x_{0}\right)\right)$, and used to define a horizontal "slice," indicated in bold. (b) An interval of width $w$ is randomly positioned around $x_{0}$, and then expanded in steps of size $w$ until both ends are outside the slice. (c) A new point, $x_{1}$, is found by picking uniformly from the interval until a point inside the slice is found. Points picked that are outside the slice are used to shrink the interval.


FIG. 2. The doubling procedure. In (a), the initial interval is doubled twice, until both ends are outside the slice. In (b), where the start state is different, and the initial interval's ends are already outside the slice, no doubling is done.
4.1. Finding an appropriate interval. After a value for the auxiliary variable has been drawn, defining the slice $S$, the next task is to find an interval $I=(L, R)$, containing the current point, $x_{0}$, from which the new point, $x_{1}$, will be drawn. We would like this interval to contain as much of the slice as is feasible, so as to allow the new point to differ as much as possible from the old point, but we would also like to avoid intervals that are much larger than the slice, as this will make the subsequent sampling step less efficient.

Several schemes for finding an interval are possible.

1. Ideally, we would set $L=\inf (S)$ and $R=\sup (S)$. That is, we would set $I$ to the smallest interval that contains all of $S$. This may not be feasible, however.
2. If the range of possible values of $x$ is bounded, we might let $I$ be that range. However, this may not be good if the slice is typically much smaller than this.
3. Given an estimate, $w$, for the scale of $S$, we can randomly pick an initial interval of size $w$, containing $x_{0}$, and then perhaps expand it by a "stepping out" procedure.
4. Similarly, we can randomly pick an initial interval of size $w$, and then expand it by a "doubling" procedure.

For each scheme, we must also be able to find the set $A$ of acceptable successor states, defined as follows:
(3) $A=\left\{x: x \in S \cap I\right.$ and $P($ Select $I \mid$ At state $x)=P\left(\right.$ Select $I \mid$ At state $\left.\left.x_{0}\right)\right\}$.

That is, $A$ is the set of states from which we would be as likely to choose the interval $I$ as we were to choose this $I$ from the current state. When we subsequently sample from within $I$ (see Section 4.2), we will ensure that the state chosen is in $A$, a fact that will be used in the proof of correctness in Section 4.3. Clearly, for schemes (1) and (2), $A=S$. For scheme (3), we will arrange that $A=S \cap I$. For scheme (4), a special test of whether a state is in $A$ may be necessary.

Scheme (1), in which $I$ is set to the smallest interval containing $S$, will be feasible when all solutions of $f(x)=y$ can be found analytically, or by an efficient and robust numerical method, but one cannot expect this in general. Often, even the number of disjoint intervals making up $S$ will be hard to determine.

Scheme (2) is certainly easy to implement when the range of $x$ is bounded, and we can always arrange this by applying a suitable transformation. However, if the slice is usually much smaller than the full range, the subsequent sampling (see Section 4.2) will be inefficient. This scheme has been used by Frey (1997).

The "stepping out" procedure [scheme (3) above] is appropriate for any distribution, provided that some rough estimate, $w$, for the typical width of the slice is available. The manner in which an interval is found by stepping out is illustrated in Figure 1(b) and the procedure is given in detail in Figure 3. The size of the interval found can be limited to $m w$, for some specified integer $m$, or the interval can be allowed to grow to any size (i.e., $m$ can be set to infinity), in which case the procedure can be simplified in an obvious way. Simplification is

```
Input: \(\quad f=\) function proportional
        to the density
    \(x_{0}=\) the current point
    \(y=\) the vertical level defining
        the slice
    \(w=\) estimate of the typical
        size of a slice
    \(m=\) integer limiting the size
        of a slice to \(m w\)
```

Output: $(L, R)=$ the interval found

$$
\begin{aligned}
& U \sim \operatorname{Uniform}(0,1) \\
& L \leftarrow x_{0}-w * U \\
& R \leftarrow L+w \\
& V \sim \operatorname{Uniform}(0,1) \\
& J \leftarrow \text { Floor }(m * V) \\
& K \leftarrow(m-1)-J
\end{aligned}
$$

repeat while $J>0$ and $y<f(L)$ :

$$
\begin{aligned}
& L \leftarrow L-w \\
& J
\end{aligned} \leftarrow \leftarrow=1
$$

repeat while $K>0$ and $y<f(R)$ :

$$
\begin{aligned}
& R \leftarrow R+w \\
& K \leftarrow K-1
\end{aligned}
$$

FIG. 3. The "stepping out" procedure for finding an interval around the current point. The notation $U \sim \operatorname{Uniform}(0,1)$ indicates that $U$ is set to a number randomly drawn from the uniform distribution on $(0,1)$.
also possible when $m$ is 1 , in which case the interval will always be of size $w$, and there will be no need to evaluate $f$ at its endpoints. Note that the random positioning of the initial interval and the random apportioning of the maximum number of steps $m$ into a limit on going to the left and a limit on going to the right are essential for correctness, as they ensure that the final interval could equally well have been produced from any point within $S \cap I$.

The "doubling" procedure [scheme (4)] can expand the interval faster than the stepping out procedure, and hence may be more efficient when the estimated size of the slice $(w)$ turns out to be too small. This procedure is illustrated in Figure 2, and given in detail in Figure 4. Doubling produces a sequence of intervals, each

$$
\text { Input: } \begin{aligned}
f= & \text { function proportional } \\
& \text { to the density } \\
x_{0}= & \text { the current point } \\
y= & \text { the vertical level defining } \\
& \text { the slice } \\
w= & \text { estimate of the typical } \\
& \text { size of a slice } \\
p= & \text { integer limiting the size } \\
& \text { of a slice to } 2^{p} w
\end{aligned}
$$

Output: $(L, R)=$ the interval found

```
\(U \sim \operatorname{Uniform}(0,1)\)
\(L \leftarrow x_{0}-w * U\)
\(R \leftarrow L+w\)
\(K \leftarrow p\)
repeat while \(K>0\)
    and \(\{y<f(L)\) or \(y<f(R)\}\) :
    \(V \sim \operatorname{Uniform}(0,1)\)
    if \(V<1 / 2\) then \(L \leftarrow L-(R-L)\)
        else \(R \leftarrow R+(R-L)\)
    \(K \leftarrow K-1\)
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FIG. 4. The "doubling" procedure for finding an interval around the current point. Note that it is possible to save some computation in second and later iterations of the loop, since only one of $f(L)$ and $f(R)$ will have changed from the previous iteration.
twice the size of the previous one, until an interval is found with both ends outside the slice, or a predetermined limit is reached. Note that when the interval is doubled the two sides are not expanded equally. Instead just one side is expanded, chosen at random (irrespective of whether that side is already outside the slice). This is essential to the correctness of the method, since it produces a final interval that could have been obtained from points other than the current one. The set $A$ of acceptable next states is restricted to those for which the same interval could have been produced, and is in general not all of $S \cap I$. This complicates the subsequent sampling somewhat, as described below.
4.2. Sampling from the part of the slice within the interval. Once an interval, $I=(L, R)$, has been found containing the current point, $x_{0}$, the final step of the single-variable slice sampling procedure is to randomly draw a new point, $x_{1}$, from within this interval. This point must lie within the set $A$ of points acceptable as the next state of the Markov chain, defined in (3).

Two methods could be used to sample from $I$.
(i) Repeatedly sample uniformly from $I$ until a point is drawn that lies within $A$.
(ii) Repeatedly sample uniformly from an interval that is initially equal to $I$, and which shrinks each time a point is drawn that is not in $A$, until a point within $A$ is found.

Method (i) could be very inefficient, if $A$ turns out to be a tiny portion of $I$. The shrinkage of the interval in method (ii) ensures that the expected number of points drawn will not be too large, making it a more appropriate method for general use.

The shrinkage procedure is shown in detail in Figure 5. Note that each rejected point is used to shrink the interval in such a way that the current point remains
Input: $\quad f=$ function proportional
Input: $\quad f=$ function proportional
to the density
to the density
$x_{0}=$ the current point
$x_{0}=$ the current point
$y=$ the vertical level defining
$y=$ the vertical level defining
the slice
the slice
$(L, R)=$ the interval to sample from
$(L, R)=$ the interval to sample from
Output: $x_{1}=$ the new point
Output: $x_{1}=$ the new point

$$
\bar{L} \leftarrow L, \quad \bar{R} \leftarrow R
$$

Repeat:
$U \sim \operatorname{Uniform}(0,1)$
$x_{1} \leftarrow \bar{L}+U *(\bar{R}-\bar{L})$
if $y<f\left(x_{1}\right)$ and $\operatorname{Accept}\left(x_{1}\right)$ then exit loop
if $x_{1}<x_{0}$ then $\bar{L} \leftarrow x_{1}$

$$
\text { else } \bar{R} \leftarrow x_{1}
$$

FIG. 5. The "shrinkage" procedure for sampling from the interval. Accept ( $x_{1}$ ) is notation for a test of whether a point, $x_{1}$, that is, within $S \cap I$ is an acceptable next state. If scheme (1), (2) or (3) was used for constructing the interval, all points within $S \cap I$ are acceptable. If the doubling procedure [scheme (4)] was used, the point must pass the test of Figure 6.

```
Input: \(\quad f=\) function proportional to the density
\(x_{0}=\) the current point
\(x_{1}=\) the possible next point
\(y=\) the vertical level defining the slice
\(w=\) estimate of the typical size of a slice
\((L, R)=\) the interval found by the doubling procedure, using \(w\)
```

Output: whether or not $x_{1}$ is acceptable as the next state
$\hat{L} \leftarrow L, \quad \hat{R} \leftarrow R$
$D \leftarrow$ false
repeat while $\hat{R}-\hat{L}>1.1 * w$ :
$M \leftarrow(\hat{L}+\hat{R}) / 2$
if $\left\{x_{0}<M\right.$ and $\left.x_{1} \geq M\right\}$ or
$\left\{x_{0} \geq M\right.$ and $\left.x_{1}<M\right\}$ then
$D \leftarrow$ true
if $x_{1}<M$ then $\hat{R} \leftarrow M$ else $\hat{L} \leftarrow M$
if $D$ and $y \geq f(\hat{L})$ and $y \geq f(\hat{R})$ then the new point is not acceptable
The new point is acceptable if it is not rejected in the loop above

Fig. 6. The test for whether a new point, $x_{1}$, that is, within $S \cap I$ is an acceptable next state, when the interval was found by the "doubling" procedure. The multiplication by 1.1 in the "while" condition guards against possible round-off error. The variable D tracks whether the intervals that would be generated from the new point differ from those leading to the current point. When they don't, time can be saved by omitting a check.
within it. Since the current point is always within $A$, the interval used always contains acceptable points, ensuring that the procedure will terminate.

If the interval was found by scheme (1), (2), or (3), the set $A$ is simply $S \cap I$. However, if the doubling procedure [scheme (4)] was used, $A$ may be a smaller subset of $S \cap I$. This is illustrated in Figure 2. In 2(a), an interval is found by doubling an initial interval until both ends are outside the slice. A different starting point is considered in 2(b), one which might have been drawn from the interval found in 2(a). The doubling procedure terminates earlier starting from here, so this point is not in $A$. (Note that $A$ is here defined conditional on the alignment of the initial interval.)

The Accept $\left(x_{1}\right)$ predicate in Figure 6 tests whether a point in $S \cap I$ is in $A$ when the doubling procedure [scheme (4)] was used. This procedure works backward through the intervals that the doubling procedure would pass through to arrive at $I$ when starting from the new point, checking that none of them has both ends outside the slice, which would lead to earlier termination of the doubling procedure.
4.3. Correctness of single-variable slice sampling. To show that singlevariable slice sampling is a correct procedure, we must show that each update leaves the desired distribution invariant. To guarantee convergence to this distribution, the resulting Markov chain must also be ergodic. This is not always true, but it is in those situations [such as when $f(x)>0$ for all $x$ ] for which one can easily
show that Gibbs sampling is ergodic. I will not discuss the more difficult situations here.

To show invariance, we suppose that the initial state, $x_{0}$, is distributed according to $f(x)$. In step (a) of single-variable slice sampling, a value for $y$ is drawn uniformly from $(0, f(x))$. The joint distribution for $x_{0}$ and $y$ will therefore be as in equation (1). If the subsequent steps update $x_{0}$ to $x_{1}$ in a manner that leaves this joint distribution invariant, then when we subsequently discard $y$, the resulting distribution for $x_{1}$ will be the marginal of this joint distribution, which is the same as that defined by $f(x)$, as desired.

We therefore need only show that the selection of $x_{1}$ to follow $x_{0}$ in steps (b) and (c) of the single-variable slice sampling procedure leaves the joint distribution of $x$ and $y$ invariant. Since these steps do not change $y$, this is the same as leaving the conditional distribution for $x$ given $y$ invariant, and this conditional distribution is uniform over $S=\{x: y<f(x)\}$, the slice defined by $y$. We can show invariance of this distribution by showing that the updates satisfy detailed balance, which for a uniform distribution reduces to showing that the probability density for $x_{1}$ to be selected as the next state, given that the current state is $x_{0}$, is the same as the probability density for $x_{0}$ to be the next state, given that $x_{1}$ is the current state, for any states $x_{0}$ and $x_{1}$ within $S$.

In the process of picking a new state, various intermediate choices are made randomly. When the interval is found by the stepping out procedure of Figure 3, the alignment of the initial interval is randomly chosen, as is the division of the maximum number of intervals into those used to extend to the left and those used to extend to the right. For the doubling procedure of Figure 4, the alignment of the initial interval is random and the decisions whether to extend to the right or to the left are also made randomly. When sampling is done using the shrinkage procedure of Figure 5, zero or more rejected points will be chosen before the final point. Let $r$ denote these intermediate random choices. I will prove that detailed balance holds for the entire procedure by showing the following stronger result:

$$
\begin{align*}
& P\left(\text { next state }=x_{1}, \text { intermediate choices }=r \mid \text { current state }=x_{0}\right) \\
& \quad=P\left(\text { next state }=x_{0}, \text { intermediate choices }=\pi(r) \mid \text { current state }=x_{1}\right) \tag{4}
\end{align*}
$$

where $\pi(r)$ is some one-to-one mapping that has Jacobian one (with regard to the real-valued variables), which may depend on $x_{0}$ and $x_{1}$. Integrating over all possible values for $r$ then gives the desired result.

The mapping $\pi$ used is as follows. If the interval $I$ is found by the stepping out or doubling procedure, an intermediate value, $U$, will be generated by the procedure of Figure 3 or 4 , and used to define the initial interval. We define $\pi$ so that it maps the value $U_{0}$ chosen when the state is $x_{0}$ to the following $U_{1}$ when the state is $x_{1}$ :

$$
\begin{equation*}
U_{1}=\operatorname{Frac}\left(U_{0}+\left(x_{1}-x_{0}\right) / w\right) \tag{5}
\end{equation*}
$$

where Frac $(x)=x$ - Floor $(x)$ is the fractional part of $x$. This mapping associates values that produce the same alignment of the initial interval. Note also that it has Jacobian one. If the stepping out procedure is used, a value for $J$ is also generated, uniformly from the set $\{0, \ldots, m-1\}$. The mapping $\pi$ associates the $J_{0}$ found when the state is $x_{0}$ with the following $J_{1}$ when the state is $x_{1}$ :

$$
\begin{equation*}
J_{1}=J_{0}+\left(x_{1} / w-U_{1}\right)-\left(x_{0} / w-U_{0}\right) \tag{6}
\end{equation*}
$$

Here, $\left(x_{1} / w-U_{1}\right)-\left(x_{0} / w-U_{0}\right)$ is an integer giving the number of steps (of size $w$ ) from the left end of the interval containing $x_{0}$ to the left end of the interval containing $x_{1}$. This is the amount by which we must adjust $J_{0}$ in order to ensure that if the interval found starting from $x_{0}$ grows to its maximum size, the associated interval found starting from $x_{1}$ will be identical. Similarly, if the doubling procedure of Figure 4 is used, the sequence of random decisions as to which side of the interval to expand is mapped by $\pi$ to the sequence of decisions that would cause the interval expanding from $x_{1}$ to become identical to the interval expanding from $x_{0}$ when the latter first includes $x_{1}$, and to remain identical through further expansions. Note here that there is at most one way to obtain a given final interval by successive doublings from a given initial interval, and that the alignment of the initial intervals by the association of $U_{0}$ with $U_{1}$ ensures that doubling starting from $x_{1}$ can indeed lead to the same interval as found from $x_{0}$. Finally, to complete the definition, $\pi$ maps the sequence of rejected points used to shrink the interval found from $x_{0}$ (see Figure 5) to the same sequence of points when $x_{1}$ is the start state.

It remains to show that with this definition of $\pi$, (4) does indeed hold, for all points $x_{0}$ and $x_{1}$, and all possible intermediate values $r$. The equation certainly holds when both sides are zero, so we can ignore situations where movement between $x_{0}$ and $x_{1}$ is impossible (in conjunction with the given intermediate values).

Consider first the probability (density) for producing the intermediate values that define the interval $I$. For the stepping out and doubling procedures, the values $U_{0}$ and $U_{1}$ (related by $\pi$ ) that are generated from $x_{0}$ and $x_{1}$ will certainly have the same probability density, since $U$ is drawn from a uniform distribution. Similarly, for the stepping out procedure, the values $J_{0}$ and $J_{1}$ are drawn from a uniform distribution over $\{0, \ldots, m-1\}$, and hence have the same probability as long as $J_{0}$ and $J_{1}$ are both in this set, which will be true whenever movement between $x_{0}$ and $x_{1}$ is possible. For the doubling procedure, a sequence of decisions as to which side to extend is made, with all sequences of a given length having the same probability. Here also, the sequences associated by $\pi$ will have the same probability, provided the same number of doublings are done starting from $x_{0}$ as from $x_{1}$. This need not be true in general, but if the sequence from $x_{1}$ is shorter, the test of Figure 6 will eliminate $x_{1}$ as a possible successor to $x_{0}$, and if the sequence from $x_{0}$ is shorter, $x_{1}$ will not be a possible successor because it will be outside the interval $I$ found from $x_{0}$. Both sides of (4) will therefore be zero in this situation.

Note next that the intervals found by any of the schemes of Section 4.1 will be the same for $x_{0}$ as for $x_{1}$, when the intermediate values chosen are related by $\pi$, assuming a transition from $x_{0}$ to $x_{1}$ is possible. For the stepping out procedure, the maximum extent of the intervals will be the same because of the relationships between $U_{0}$ and $U_{1}$ and between $J_{0}$ and $J_{1}$. Furthermore, the actual intervals found by stepping out (limited by the maximum) must also be the same whenever a transition between $x_{0}$ and $x_{1}$ is possible, since if the interval starting from $x_{0}$ has reached $x_{1}$, expansion of both intervals will continue in the same direction until the outside of the slice or the maximum is reached, and likewise in the other direction. Similarly, the mapping $\pi$ is defined to be such that if the interval found by the doubling procedure starting from $x_{0}$ includes $x_{1}$, the same interval would be found from $x_{1}$, provided the process was not terminated earlier (by both ends being outside the slice), in which case $x_{1}$ is not a possible successor (as it would be rejected by the procedure of Figure 6). Note also that since the set $A$ is determined by $I$ (for any start state), it too will be the same for $x_{0}$ as for $x_{1}$.

If we sample from this $I$ by simple rejection [scheme (i) in Section 4.2], the state chosen will be uniformly distributed over $A$, so the probability of picking $x_{0}$ will be the same as that of picking $x_{1}$. If we instead use the shrinkage procedure [scheme (ii) in Section 4.2, detailed in Figure 5], we need to consider as intermediate values the sequence of rejected points that was used to narrow the interval (recall that under $\pi$ this sequence is the same for $x_{0}$ as for $x_{1}$ ). The probability density for the first of these is clearly the same for both starting points, since $I$ is the same. As the interval shrinks, it remains the same for both $x_{0}$ and $x_{1}$, since the rejection decisions (based on $A$ ) are the same, and since we need consider only the case where the same end of the interval is moved to the rejected point (as otherwise a transition between $x_{0}$ and $x_{1}$ in conjunction with these intermediate values would be impossible). The probability densities for later rejected points, and for the final accepted state, are therefore also the same.

This completes the proof. Various seemingly reasonable modifications, such as changing the doubling procedure of Figure 4 so as not to expand the interval on a side that is already outside the slice, would undermine the argument of the proof and hence cannot be used.
4.4. Shortcuts for unimodal distributions. Some shortcuts are, however, allowable when the distribution is unimodal, because the slice, $S$, is then guaranteed to consist of a single interval. The acceptance test in Figure 6 can be omitted, since one can show that it will always indicate that the new point is acceptable. The interval found by the doubling procedure can also be shrunk at the outset by setting its endpoints to the first point in each direction that was found to lie outside the slice, since for a unimodal distribution, this shrinkage cannot eliminate any points within the slice and hence will not change the distribution of the point selected.

If the distribution is known to be unimodal and no limit is imposed on the size of the interval found (i.e., $m$ and $p$ in Figures 3 and 4 are infinite), the estimate, $w$, for the typical size of a slice can be set on the basis of past iterations. One could, for example, set $w$ to the average distance between the old and new points in past iterations. This is valid because the distribution of the new point does not depend on $w$ in this situation, even though $w$ influences how efficiently this new point is found. Indeed, when the distribution is known to be unimodal, one can use any method at all for finding an interval that contains the current point and has both ends outside the slice, as any such interval will lead to the new point finally chosen being drawn uniformly from the slice.
5. Multivariate slice sampling methods. Rather than sample from a distribution for $x=\left(x_{1}, \ldots, x_{n}\right)$ by applying one of the single-variable slice sampling procedures described above to each $x_{i}$ in turn, we might try instead to apply the idea of slice sampling directly to the multivariate distribution. I will start by describing a straightforward generalization of the single-variable methods to multivariate distributions, and then describe a more sophisticated method, which can potentially allow for adaptation to the local dependencies between variables.
5.1. Multivariate slice sampling with hyperrectangles. We can generalize the single-variable slice sampling methods of Section 4 to methods for performing multivariate updates by replacing the interval $I=(L, R)$ by an axis-aligned hyperrectangle $H=\left\{x: L_{i}<x_{i}<R_{i}\right.$ for all $\left.i=1, \ldots, n\right\}$. Here, $L_{i}$ and $R_{i}$ define the extent of the hyperrectangle along the axis for variable $x_{i}$.

The procedure for finding the next state, $x_{1}=\left(x_{1,1}, \ldots, x_{1, n}\right)$, from the current state, $x_{0}=\left(x_{0,1}, \ldots, x_{0, n}\right)$, parallels the single-variable procedure:
(a) Draw a real value, $y$, uniformly from $\left(0, f\left(x_{0}\right)\right)$, thereby defining the slice $S=\{x: y<f(x)\}$.
(b) Find a hyperrectangle, $H=\left(L_{1}, R_{1}\right) \times \cdots \times\left(L_{n}, R_{n}\right)$, around $x_{0}$, which preferably contains at least a big part of the slice.
(c) Draw the new point, $x_{1}$, from the part of the slice within this hyperrectangle.

It would perhaps be ideal for step (b) to set $H$ to the smallest hyperrectangle containing $S$, but this is unlikely to be feasible. When all the variables have bounded ranges we might set $H$ to the entire space, but this may be inefficient, since $S$ is likely to be much smaller. We may therefore have to be content with finding an $H$ that contains the current point, $x_{0}$, but probably not all of $S$. We will need estimates, $w_{i}$, for the appropriate dimensions of $H$ along each axis, which we might set to a common value, $w$, if we know nothing about the relative scales of the variables. The simplest way of finding $H$ is then to randomly position a hyperrectangle with these dimensions, uniformly over positions that lead to $H$ containing $x_{0}$. This generalizes the random positioning of the initial interval $I$ for
single-variable slice sampling. The stepping out and doubling procedures do not generalize so easily, however. The goal of finding an interval whose endpoints are outside the slice would generalize to finding a hyperrectangle all of whose vertices are outside the slice, but since an $n$-dimensional hyperrectangle has $2^{n}$ vertices, we would certainly not want to test for this when $n$ is large. The stepping out procedure seems to be too time consuming in any case, since one would need to step out in each of the $n$ directions. The doubling procedure does generalize appropriately, and one could decide to stop doubling when a randomly drawn point picked uniformly from the current hyperrectangle is outside the slice. Here, however, I will consider only the simplest scheme, which is to use the randomly positioned hyperrectangle without any expansion, though it is then crucial that the $w_{i}$ not be much smaller than they should be.

The shrinkage procedure of Figure 5 generalizes easily to multiple dimensions: the hyperrectangle is simply shrunk independently along each axis. Combining this with random positioning of $H$ gives the multivariate slice sampling method shown in Figure 7(a), and given in detail in Figure 8. The validity of this method can be proved in the same way as was done for single-variable slice sampling in Section 4.3.

Although this simple multivariate slice sampling method is easily implemented, in one respect it works less well than applying single-variable slice sampling to each variable in turn. When each variable is updated separately, the interval for that variable will be shrunk only as far as needed to obtain a new value within the slice. The amount of shrinkage can be different for different variables. In contrast, the procedure of Figure 8 shrinks all dimensions of the hyperrectangle until a point inside the slice is found, even though the probability density may not vary rapidly in some of these dimensions, making shrinkage in these directions unnecessary.

One way to try to avoid this problem is illustrated in Figure 7(b). Rather than shrink all dimensions of the hyperrectangle when the last point chosen was outside the slice, we can instead shrink along only one axis, basing the choice on the gradient of $\log f(x)$, evaluated at the last point. Specifically, only the axis corresponding to variable $x_{i}$ is shrunk, where $i$ maximizes the following product:

$$
\begin{equation*}
\left(R_{i}-L_{i}\right)\left|G_{i}\right|, \tag{7}
\end{equation*}
$$

where $G$ is the gradient of $\log f(x)$ at the last point chosen. By multiplying the magnitude of component $i$ of the gradient by the width of the hyperrectangle in this direction, we get an estimate of the amount by which $\log f(x)$ changes along axis $i$. The axis for which this change is thought to be largest is likely to be the best one to shrink in order to eliminate points outside the slice. Unfortunately, if this decision were based as well on whether the sign of the gradient indicates that $\log f(x)$ is increasing or decreasing as we move toward the current point, $x_{0}$, the shrinkage decision might be different if we were to shrink from the final accepted point, $x_{1}$, which would invalidate the method.
(a)

(b)


Fig. 7. Multivariate slice sampling with hyperrectangles. The heavy line outlines the slice, containing the current point, $x_{0}$. The large square is the initial hyperrectangle. In (a), the hyperrectangle is shrunk in all directions when the point drawn is outside the slice, until a new point, $x_{1}$, inside the slice is found. In (b), the hyperrectangle is shrunk along only one axis, determined from the gradient and the current dimensions of the hyperrectangle. The dashed lines are contours of the density function, indicating the direction of the gradient.

Many more elaborate schemes along these lines are possible. For instance, we might shrink along all axes for which the product (7) is greater than some

Input: $\quad f=$ function proportional to the density
$x_{0}=$ the current point, of dimension $n$
$w_{i}=$ scale estimates for each variable, $i=1, \ldots, n$

Output: $x_{1}=$ the new point

Step (a): Find value of $y$ that defines the slice

$$
y \sim \operatorname{Uniform}\left(0, f\left(x_{0}\right)\right)
$$

Step (b): Randomly position the hyperrectangle $H=\left(L_{1}, R_{1}\right) \times \cdots \times\left(L_{n}, R_{n}\right)$

For $i=1$ to $n$ :

$$
\begin{aligned}
& U_{i} \sim \operatorname{Uniform}(0,1) \\
& L_{i} \leftarrow x_{0, i}-w_{i} * U_{i} \\
& R_{i} \leftarrow L_{i}+w_{i}
\end{aligned}
$$

Step (c): Sample from $H$, shrinking when points are rejected.
Repeat:
For $i=1$ to $n$ :

$$
\begin{aligned}
& U_{i} \sim \operatorname{Uniform}(0,1) \\
& x_{1, i} \leftarrow L_{i}+U_{i} *\left(R_{i}-L_{i}\right)
\end{aligned}
$$

if $y<f\left(x_{1}\right)$ then exit loop
For $i=1$ to $n$ :

$$
\begin{array}{r}
\text { if } x_{1, i}<x_{0, i} \text { then } L_{i} \leftarrow x_{1, i} \\
\text { else } R_{i} \leftarrow x_{1, i}
\end{array}
$$

FIG. 8. A simple multivariate slice sampling procedure, with randomly positioned hyperrectangle and shrinkage in all directions, as in Figure 7(a).
threshold. A good scheme might preserve the ability of single-variable slice sampling to adapt differently for different variables, while keeping the advantages that simultaneous updates may sometimes have (e.g., in producing an ergodic chain when there are tight dependencies between variables).

More ambitiously, we might hope that a multivariate slice sampler could adapt to the dependencies between variables, not just to their different scales. This requires that we go beyond axis-aligned hyperrectangles, as is done in the next section.
5.2. A framework for adaptive multivariate slice sampling. We would like a more general framework by which trial points outside the slice that were previously rejected can be used to guide the selection of future trial points. In contrast to schemes based on hyperrectangles, we would like future trial points to potentially come from distributions that take account of the dependencies between variables. The scheme I present here achieves this by laying down a trail of "crumbs" that guides the selection of future trial points, leading eventually to a point inside the slice. A crumb can be anything, for example, a discrete value, a real number, a vector, a hyperrectangle, or a point in the state space being sampled from.

As with the previous slice sampling schemes, we start by choosing a value $y$ uniformly between zero and $f\left(x_{0}\right)$, where $x_{0}$ is the current point. A crumb, $c_{1}$, is then drawn at random from some distribution with density (or probability mass) function $g_{1}\left(c ; x_{0}, y\right)$. Note that this distribution may depend on both the current point, $x_{0}$, and on the value of $y$ that defines the slice. A first trial point, $x_{1}^{*}$, is then drawn from the distribution with density $h_{1}\left(x^{*} ; y, c_{1}\right)=g_{1}\left(c_{1} ; x^{*}, y\right) / Z_{1}\left(y, c_{1}\right)$, where $Z_{1}\left(y, c_{1}\right)=\int g_{1}\left(c_{1} ; x^{*}, y\right) d x^{*}$ is the appropriate normalizing constant. One can view $x_{1}^{*}$ as being drawn from a pseudo-posterior distribution, based on a uniform prior, and the "data" that the first crumb was $c_{1}$. If $x_{1}^{*}$ is inside the slice, it becomes the new state, and we are finished. Otherwise, a second crumb $c_{2}$, is drawn from some distribution $g_{2}\left(c ; x_{0}, y, c_{1}, x_{1}^{*}\right)$, which may depend on the previous crumb and the previous trial point, as well as $x_{0}$ and $y$. The second trial point is then drawn from the pseudo-posterior distribution based on the "data" $c_{1}$ and $c_{2}$; that is, $x_{2}^{*}$ is drawn from

$$
\begin{equation*}
h_{2}\left(x^{*} ; y, c_{1}, x_{1}^{*}, c_{2}\right)=g_{1}\left(c_{1} ; x^{*}, y\right) g_{2}\left(c_{2} ; x^{*}, y, c_{1}, x_{1}^{*}\right) / Z_{2}\left(y, c_{1}, x_{1}^{*}, c_{2}\right), \tag{8}
\end{equation*}
$$

where $Z_{2}\left(y, c_{1}, x_{1}^{*}, c_{2}\right)=\int g_{1}\left(c_{1} ; x^{*}, y\right) g_{2}\left(c_{2} ; x^{*}, y, c_{1}, x_{1}^{*}\right) d x^{*}$. If $x_{2}^{*}$ is inside the slice, it becomes the new state. Otherwise, we draw a third crumb, from a distribution that may depend on the current state, the value defining the slice, the previous crumbs and the previous trial points, generate a third trial point using this and the previous crumbs, and so forth until a trial point lying within the slice is found.

The distributions of trial points in this sequence will become more and more concentrated, since they are pseudo-posterior distributions based on more and more pseudo-data (the crumbs). Since this pseudo-data is generated from the
current point, the concentration will be around this point, which is of course within the slice. The probability of a trial point being within the slice therefore increases towards one as the procedure progresses.

To show that this procedure leaves the distribution with density $f(x) / Z$ invariant, it suffices to show that it separately satisfies detailed balance with respect to transitions that occur in conjunction with any given number of crumbs being drawn. In the case, for instance, of transitions involving two crumbs, we can show this by showing the stronger property that for any $x_{1}^{*}$ that is not in the slice defined by $y$ and any $x_{2}^{*}$ that is in this slice, the following will hold:

$$
\begin{equation*}
P\left(x_{0}\right) P\left(y, c_{1}, x_{1}^{*}, c_{2}, x_{2}^{*} \mid x_{0}\right)=P\left(x_{2}^{*}\right) P\left(y, c_{1}, x_{1}^{*}, c_{2}, x_{0} \mid x_{2}^{*}\right) . \tag{9}
\end{equation*}
$$

Here, $P\left(x_{0}\right)$ and $P\left(x_{2}^{*}\right)$ are the probability densities for the current point and the point that will become the new point [which are proportional to $f(x)$ ]. The conditional probabilities above are the densities for the given sequence of values being chosen during the procedure, given that the current point is the one conditioned on. The left-hand side of (9) can be written as follows:

$$
\begin{aligned}
& P\left(x_{0}\right) P\left(y \mid x_{0}\right) P\left(c_{1} \mid x_{0}, y\right) P\left(x_{1}^{*} \mid y, c_{1}\right) P\left(c_{2} \mid x_{0}, y, c_{1}, x_{1}^{*}\right) P\left(x_{2}^{*} \mid y, c_{1}, x_{1}^{*}, c_{2}\right) \\
&= {\left[f\left(x_{0}\right) / Z\right]\left[1 / f\left(x_{0}\right)\right] g_{1}\left(c_{1} ; x_{0}, y\right)\left[g_{1}\left(c_{1} ; x_{1}^{*}, y\right) / Z_{1}\left(y, c_{1}\right)\right] } \\
& \quad \times g_{2}\left(c_{2} ; x_{0}, y, c_{1}, x_{1}^{*}\right)\left[g_{1}\left(c_{1} ; x_{2}^{*}, y\right) g_{2}\left(c_{2} ; x_{2}^{*}, y, c_{1}, x_{1}^{*}\right) / Z_{2}\left(y, c_{1}, x_{1}^{*}, c_{2}\right)\right] .
\end{aligned}
$$

The right-hand side is

$$
\begin{aligned}
& P\left(x_{2}^{*}\right) P\left(y \mid x_{2}^{*}\right) P\left(c_{1} \mid x_{2}^{*}, y\right) P\left(x_{1}^{*} \mid y, c_{1}\right) P\left(c_{2} \mid x_{2}^{*}, y, c_{1}, x_{1}^{*}\right) P\left(x_{0} \mid y, c_{1}, x_{1}^{*}, c_{2}\right) \\
&= {\left[f\left(x_{2}^{*}\right) / Z\right]\left[1 / f\left(x_{2}^{*}\right)\right] g_{1}\left(c_{1} ; x_{2}^{*}, y\right)\left[g_{1}\left(c_{1} ; x_{1}^{*}, y\right) / Z_{1}\left(y, c_{1}\right)\right] } \\
& \quad \times g_{2}\left(c_{2} ; x_{2}^{*}, y, c_{1}, x_{1}^{*}\right)\left[g_{1}\left(c_{1} ; x_{0}, y\right) g_{2}\left(c_{2} ; x_{0}, y, c_{1}, x_{1}^{*}\right) / Z_{2}\left(y, c_{1}, x_{1}^{*}, c_{2}\right)\right] .
\end{aligned}
$$

These are equal, as is true in general for transitions involving any number of crumbs.

The hyperrectangle methods of Section 5.1 can be viewed in this framework. The randomly placed initial hyperrectangle is the first crumb. The first trial point is chosen from those points that could produce this initial hyperrectangle, which is simply the set of points within the hyperrectangle. The second and later crumbs are the shrunken hyperrectangles. Conditional on the current point, the previous crumb (i.e., the previous hyperrectangle), and the previous trial point, these later crumbs have degenerate distributions, concentrated on a single hyperrectangle. The possible corresponding trial points are the points within the shrunken hyperrectangle.

By using different sorts of crumbs, and different distributions for them, a huge variety of methods could be constructed within this framework. I discuss here only methods in which the crumbs are points in the state space, and have multivariate Gaussian distributions. The distributions of the trial points given the crumbs will then also be multivariate Gaussians.

In the simplest method of this sort, every $g_{i}$ is Gaussian with mean $x_{0}$ and covariance matrix $\sigma^{2} I$, for some fixed $\sigma^{2}$. The distribution, $h_{i}$, for $x_{i}^{*}$ is then Gaussian with mean $\bar{c}_{i}=\left(c_{1}+\cdots+c_{i}\right) / i$ and covariance matrix $\left(\sigma^{2} / i\right) I$. As more and more trial points are generated, they will come from narrower and narrower distributions, which will be concentrated closer and closer to the current point (since $\bar{c}_{i}$ will approach $x_{0}$ ). This is analogous to shrinkage in the hyperrectangle method. In practice, it is probably desirable to let $\sigma_{i}^{2}$ decrease with $i$ (perhaps exponentially), so that the trial points would be forced closer to $x_{0}$ more quickly. Alternatively, we might look at $f\left(x_{i-1}^{*}\right) / y$ in order to estimate what value for $\sigma_{i}$ would produce a distribution for the next trial point, $x_{i}^{*}$, that is likely to lie within the slice.

More generally, $g_{i}$ could be a multivariate Gaussian with mean $x_{0}$ and some covariance matrix $\Sigma_{i}$, which may depend on the value of $y$, the previous crumbs, and the previous trial points. In particular, $\Sigma_{i}$ could depend on the gradients of $f\left(x_{j}^{*}\right)$ for $j<i$, which provide information on what Gaussian distribution would be a good local approximation to $f(x)$. The distribution, $h_{i}$, for trial point $x_{i}^{*}$ will then have covariance $\Sigma_{i}^{*}=\left[\Sigma_{1}^{-1}+\cdots+\Sigma_{i}^{-1}\right]^{-1}$ and mean $\bar{c}_{i}=$ $\Sigma_{i}^{*}\left[\Sigma_{1}^{-1} c_{1}+\cdots+\Sigma_{i}^{-1} c_{i}\right]$.

When $x$ is of only moderate dimensionality, operations on covariance matrices would be fairly fast, and a wide variety of ways for producing $\Sigma_{i}$ would be feasible. For higher-dimensional problems, such operations would need to be avoided, as is done in an optimization context with the conjugate gradient and related methods. Further research is therefore needed in order to fully exploit the potential of this promising framework for adaptation, and to compare it with methods based on the "delayed rejection" (also called "splitting rejection") framework of Tierney and Mira [Mira (1998), Chapter 5; Tierney and Mira (1999); Green and Mira (2001)].
6. Overrelaxed slice sampling. When variables are updated in ways that do not take account of their dependencies, changes must be small, and many updates will be needed to move from one part of the distribution to another. Sampling efficiency can be improved in this context by suppressing the random walk behavior characteristic of simple schemes such as Gibbs sampling. One way of achieving this is by using "overrelaxed" updates. Like Gibbs sampling, overrelaxation methods update each variable in turn, but rather than drawing a new value for a variable from its conditional distribution independently of the current value, the new value is instead chosen to be on the opposite side of the mode from the current value. In Adler's (1981) scheme, applicable when the conditional distributions are Gaussian, the new value for variable $i$ is

$$
\begin{equation*}
x_{i}^{\prime}=\mu_{i}+\alpha\left(x_{i}-\mu_{i}\right)+\sigma_{i}\left(1-\alpha^{2}\right)^{1 / 2} n \tag{10}
\end{equation*}
$$

where $\mu_{i}$ and $\sigma_{i}$ are the conditional mean and standard deviation of variable $i$, $n$ is a Gaussian with mean zero and variance one, and $\alpha$ is a parameter slightly
greater than -1 . This method is analyzed and discussed by Barone and Frigessi (1990) and by Green and Han (1992), though these discussions fail in some respects to elucidate the true benefits and limitations of overrelaxation: the crucial advantage being that it sometimes (though not always) suppresses random walks [Neal (1998)].

Various attempts have been made to produce overrelaxation schemes that can be used when the conditional distributions are not Gaussian. I have reviewed several such schemes and introduced one of my own [Neal (1998)]. The concept of overrelaxation seems to apply only when the conditional distributions are unimodal, so we may assume that this is usually the case, though we would like the method to at least remain valid (i.e., leave the desired distribution invariant) even if this assumption turns out to be false. To obtain the full benefits of overrelaxation, it is important that almost every update be overrelaxed, with few or no "rejections" that leave the state unchanged, as such rejections reintroduce an undesirable random walk aspect to the motion through state space [Neal (1998)].

In this section, I will show how overrelaxation can be done using slice sampling. Of the many possible schemes, I will describe only one in detail, based on stepping out and bisection, which is illustrated in Figure 9, and detailed in Figure 10.

To begin, we apply the stepping out procedure of Figure 3 to find an interval around the current point. Normally, we would set the maximum size of the interval ( $m$ ) to infinity, since a proper overrelaxation operation requires that the entire slice be found, but the scheme remains valid for any $m$.

If the stepping out procedure finds an interval around the slice that is bigger than the initial interval, the two outermost steps will serve to locate the endpoints


Fig. 9. Overrelaxation using the stepping out procedure and bisection. In (a), an interval, ( $L, R$ ), with both ends outside the slice is found by stepping out from the current point. In (b), the endpoints of the slice are located more accurately using bisection. In (c), a candidate point is found by flipping through the point half-way between the approximations to the endpoints. In this case, the candidate point is accepted, since it is within the slice, and within the interval prior to bisection.

```
Input: \(\quad f=\) function proportional to the density
\(x_{0}=\) the current point
\(y=\) the vertical level defining the slice
\(w=\) estimate of the typical size of a slice
\(a=\) integer limiting endpoint accuracy to \(2^{-a} w\)
\((L, R)=\) interval found by the stepping out procedure using stepsize \(w\)
```

Output: $x_{1}=$ the new point

$$
\begin{aligned}
& \bar{L} \leftarrow L, \quad \bar{R} \leftarrow R \\
& \bar{w} \leftarrow w, \quad \bar{a} \leftarrow a
\end{aligned}
$$

When the interval is only of size $w$, narrow it until the mid-point is inside the slice (or accuracy limit is reached).
if $R-L<1.1 * w$ then
repeat:

$$
\begin{aligned}
& M \leftarrow(\bar{L}+\bar{R}) / 2 \\
& \text { if } \bar{a}=0 \text { or } y<f(M) \text { then } \\
& \quad \text { exit loop } \\
& \text { if } x_{0}>M \text { then } \bar{L} \leftarrow M \\
& \quad \text { else } \bar{R} \leftarrow M \\
& \bar{a} \leftarrow \bar{a}-1 \\
& \bar{w} \leftarrow \bar{w} / 2
\end{aligned}
$$

Refine endpoint locations by bisection, to the specified accuracy.
$\hat{L} \leftarrow \bar{L}, \quad \hat{R} \leftarrow \bar{R}$
repeat while $\bar{a}>0$ :

$$
\begin{aligned}
& \bar{a} \quad \leftarrow \bar{a}-1 \\
& \bar{w} \leftarrow \bar{w} / 2 \\
& \text { if } y \geq f(\hat{L}+\bar{w}) \text { then } \hat{L} \leftarrow \hat{L}+\bar{w} \\
& \text { if } y \geq f(\hat{R}-\bar{w}) \text { then } \hat{R} \leftarrow \hat{R}-\bar{w}
\end{aligned}
$$

Find a candidate point by flipping from the current point to the opposite side of $(\hat{L}, \hat{R})$, then test it for acceptability.
$x_{1} \leftarrow \hat{L}+\hat{R}-x_{0}$
if $x_{1}<\bar{L}$ or $x_{1}>\bar{R}$ or $y \geq f\left(x_{1}\right)$ then
$x_{1} \leftarrow x_{0}$

Fig. 10. The overrelaxation procedure using bisection.
of the slice to within an interval of size $w$ (assuming the slice consists of a single interval, as it will if the distribution is unimodal). We then locate the endpoints more precisely using a bisection procedure. For each endpoint, we test whether the midpoint of the interval in which it is located is inside or outside the slice and shrink this interval appropriately to narrow the location of the endpoint. After this is done $a$ times, each endpoint will be known to lie in an interval of size $2^{-a} w$.

If the stepping out procedure finds that the initial interval (of size $w$ ) already has both ends outside the slice, then before doing any bisection, we narrow this interval, by shrinking it in half repeatedly until its midpoint is within the slice. We then use bisection as above to locate the endpoints to within an interval of size $2^{-a} w$.

After the locations of the endpoints have been narrowed down, we approximate the entire slice by the interval ( $\hat{L}, \hat{R}$ ), formed from the outer bounds on the endpoint locations. To do an overrelaxed update, we flip from the current point, $x_{0}$, to a new point, $x_{1}$, that is the same distance as the current point from the middle of
this interval, but on the opposite side. That is, we let

$$
\begin{equation*}
x_{1}=\frac{\hat{L}+\hat{R}}{2}-\left(x_{0}-\frac{\hat{L}+\hat{R}}{2}\right)=\hat{L}+\hat{R}-x_{0} . \tag{11}
\end{equation*}
$$

We must sometimes reject this candidate point, in which case the new point is the same as the current point. First, we must reject $x_{1}$ if it lies outside the interval, $(\bar{L}, \bar{R})$, that was found prior to bisection, since the interval found from $x_{1}$ would then be different, and detailed balance would not hold. However, this cannot happen when the distribution is unimodal. Secondly, we must reject $x_{1}$ if it lies outside the slice. This can happen even for a unimodal distribution, when the endpoints of the slice have not been located exactly. However, the probability of rejection for a unimodal distribution can be reduced to as low a level as desired, at moderate cost, by locating the endpoints more precisely using more iterations of bisection.

The correctness of this procedure can be seen using arguments similar to those of Section 4.3. The interval before bisection can be found by the doubling procedure instead of stepping out, provided the point found is rejected if it fails the acceptance test of Figure 6. However, rejection for this reason will not occur in the case of a unimodal distribution, which is presumably typical, since overrelaxation is likely inappropriate for a multimodal distribution.

Many methods other than bisection could be used to locate the endpoints before overrelaxing. If the derivative of $f(x)$ can easily be calculated, we could use Newton iteration, whose rapid convergence would often allow the endpoints to be calculated to machine precision in a few iterations. For unimodal distributions, such exact calculations would eliminate the possibility of rejection, and make the final result be independent of the way the interval containing the slice was found, thereby allowing use of retrospective methods for tuning the procedure for finding this interval.

To obtain a full sampling scheme, overrelaxed updates of this sort would be applied to each variable in turn, in a fixed order, for a number of cycles, after which a cycle of normal single-variable slice sampling updates would be done. Alternatively, each update could be done normally with some small probability. A Markov chain consisting solely of overrelaxed updates might not be ergodic (perhaps staying on one contour of the probability density), and might in any case suppress random walks for too long. The frequency of normal updates is a tuning parameter, analogous to the choice of $\alpha$ in Adler's overrelaxation method, and would ideally be set so that the Markov chain moves systematically, rather than in a random walk, for long enough that it traverses a distance comparable to the largest dimension of the multivariate distribution, but for no longer than this. To keep from doing a random walk for around $k$ steps, one would do every $k$ th update normally and also arrange for the rejection rate for the overrelaxed updates to be less than $1 / k$.
7. Reflective slice sampling. Multivariate slice sampling methods can also be designed to suppress random walks. In this section I describe methods that "reflect" off the boundaries of the slice. Such movement with reflection can be seen as a specialization to uniform distributions of the Hamiltonian dynamics on which the hybrid Monte Carlo method of Duane, Kennedy, Pendleton and Roweth (1987) is based.

As before, suppose we wish to sample from a distribution over $\mathfrak{R}^{n}$, defined by a function $f(x)$ that is proportional to the probability density. We assume here that we can compute both $f(x)$ and its gradient. In each iteration of the Markov chain, we will draw a value for an auxiliary variable, $y$, uniformly from $(0, f(x))$, thereby defining an $n$-dimensional slice $S=\{x: y<f(x)\}$. We also introduce $n$ additional "momentum" variables, written as a vector, $p$, which serve to indicate the current direction and speed of motion through state space. At the start of each iteration, we pick a value for $p$, independently of $x$, from some rotationally symmetric distribution, typically Gaussian with mean zero and identity covariance.

Once $y$ and $p$ have been drawn, we repeatedly update $x$ by stepping in the direction of $p$. After some predetermined number of steps, we take the final value of $x$ as our new state (provided it is acceptable). In each step, we try to set $x^{\prime}=x+w p$, for some scale parameter $w$ that determines the average step size. However, if the resulting $x^{\prime}$ is outside the slice $S$ [i.e., $y \geq f\left(x^{\prime}\right)$ ], we must somehow try to bring it back inside. The schemes considered here all do this by some form of reflection, but differ in the exact procedure used.

Ideally, we would reflect from the exact point at which movement in the direction of $p$ first takes us outside the slice. This reflection operation modifies $p$, after which motion continues in the new direction, until we again encounter the boundary of the slice. When we hit the boundary at a point where the gradient of $f(x)$ is $h$, reflection will change $p$ as follows:

$$
\begin{equation*}
p^{\prime}=p-2 h \frac{p \cdot h}{|h|^{2}} \tag{12}
\end{equation*}
$$

This ideal reflection scheme is illustrated for a two-dimensional slice in Figure 11. Using the fact that the reflection transformation above has Jacobian one and is its own inverse, one can show that movement with reflection for some predetermined duration leaves invariant the joint distribution of $x$ (uniform within the slice) and $p$ (rotationally symmetric, independent of $x$ ), so this way of sampling is valid, with no need for an acceptance test. One can also see from the figure how such motion can proceed consistently in one direction (until the end of the slice is reached), rather than in a random walk.

Ideal reflection may be difficult to implement, however, as it requires precise calculation of where the current path intersects the boundary of the slice. Finding this point analytically is sometimes possible, as in the application of reflective slice sampling by Downs, MacKay and Lee (2000). We might instead try to solve for the intersection point numerically, but if the slice is not known to be convex, it may be


FIG. 11. Moving around a two-dimensional slice by reflection from the exact boundaries.
difficult even to determine with certainty that any intersection point that has been found is in fact the first one that would be encountered. Rather than attempt such exact calculations, we can instead employ one of two approximate schemes, based on "inside" or "outside" reflection, although the trajectories these schemes produce must sometimes be rejected in order to produce the exactly correct equilibrium distribution (in which case, the chain remains in the state from which the trajectory began).

When stepping from $x$ to $x^{\prime}=x+w p$ takes us outside the slice, we can try to reflect from the last inside point, $x$, instead of from the exact point where the path intersects the boundary, using the gradient of $f(x)$ at this inside point. The process is illustrated in Figure 12. However, for this method to be valid, we must check that the reverse trajectory would also reflect at this point, by verifying that a step in the direction opposite to our new heading would take us outside the slice. If this is not so, we must either reject the entire trajectory of which this reflection step forms a part, or alternatively, set $p$ and $x$ so that we retrace the path taken to this point (starting at the inside point where the reflection failed).

Alternatively, when we step outside the slice, we can try to reflect from the outside point, $x^{\prime}$, based on the gradient at that point. A trajectory with several such reflections is shown in Figure 13. After performing a predetermined number of


FIG. 12. Reflection from an inside point. The trajectories here go in steps of size $w|p|$, starting from the top right, until a point outside the slice is reached, when a reflection is attempted based on the inner contour shown. In (a), the reflection is successful; in (b), it must be rejected, since the reverse trajectory would not reflect at this point.


Fig. 13. Reflection from outside points. Starting from the left, two reflections based on outside contours lead back inside the slice after the next step. The step after the third reflection is still outside the slice, so further reflections must be done. In this case, the trajectory eventually returns to the slice, and its endpoint would therefore be accepted.
steps, we accept the trajectory if the final point is inside the slice. Note that for this method to be valid, one must reflect whenever the current point is outside the slice, even if this leads one away from the slice rather than toward it. This will sometimes result in the trajectory never returning to the slice, and hence being rejected, but other times, as in the figure, it does return eventually.

Many variations on these procedures are possible. Previously, it was assumed that values for $y$ and $p$ are randomly drawn at the beginning of a trajectory, and then kept the same for many steps (apart from changes to $p$ due to reflections). When using inside reflection, we might instead pick a new value for $y$ more often, perhaps before every step, and we might also partially update $p$, as is done in Horowitz's (1991) variation on hybrid Monte Carlo. When using outside reflection, the acceptance rate can be increased by terminating the trajectory when either some preset maximum number of steps have been taken, or some preset number of steps have ended inside the slice. When termination occurs for the latter reason, the final point will necessarily be inside the slice, and the trajectory will therefore be accepted.
8. A demonstration. To illustrate the benefits stemming from the adaptive nature of slice sampling, I show here how it can help avoid a disastrous scenario, in which a seriously wrong answer is obtained without any obvious indication that something is amiss.

The task is to sample from a distribution for ten real-valued variables, $v$ and $x_{1}$ to $x_{9}$. The marginal distribution of $v$ is Gaussian with mean zero and standard deviation 3. Conditional on a given value of $v$, the variables $x_{1}$ to $x_{9}$ are independent, with the conditional distribution for each being Gaussian with mean zero and variance $e^{v}$. The resulting shape resembles a ten-dimensional funnel, with small values for $v$ at its narrow end, and large values for $v$ at its wide end. Such
a distribution is typical of priors for components of Bayesian hierarchical models: $x_{1}$ to $x_{9}$ might, for example, be random effects for nine subjects, with $v$ being the $\log$ of the variance of these random effects. If the data happens to be largely uninformative, the problem of sampling from the posterior will be similar to that of sampling from the prior, so this test is relevant to actual Bayesian inference problems.

It is of course possible to sample from this distribution directly, by simply sampling for $v$, and then sampling for each of $x_{1}$ to $x_{9}$ given this value for $v$, thereby obtaining independent points from exactly the correct distribution. And in any case, we already know the correct marginal distribution for $v$, which will be the main focus of this test. We will pretend, however, that we don't already know the answer and compare what we would then conclude using various Markov chain methods to what we know is actually correct.

Figure 14 shows the results of trying to sample from this distribution using Metropolis methods and single-variable slice sampling. The upper plot shows 2,000 iterations of a run in which each iteration consists of 10,000 multivariate Metropolis updates (i.e., 20 million Metropolis updates altogether). The proposal distribution was a spherical Gaussian centered on the current state, with standard deviation one for each of the ten variables. The initial state had $v=0$ and all $x_{i}=1$. The points plotted are the value of $v$ at each iteration, with dotted lines shown at $v= \pm 7.5$.

The results of this run are grossly incorrect. We know that the marginal distribution for $v$ is Gaussian with mean zero and standard deviation 3. One would expect that out of 2,000 points from this distribution, on average 95.6 $(4.8 \%)$ should be less than -5 , but none of the points sampled by the multivariate Metropolis method is in this region. Moreover, there is little in the plot to indicate that anything is wrong. In an actual application, the results of a run such as this could easily be accepted as being correct, with serious consequences.

The source of the problem is the low probability of accepting a proposal when in a state where $v$ is small. When $v$ is -4 , for example, the standard deviation of the $x_{i}$ conditional on this value for $v$ is 0.135 . The chances that a multivariate Metropolis proposal in which each $x_{i}$ has standard deviation one will produce values for all the $x_{i}$ that are within this range of zero are about $0.135^{9} \approx 1.5 \times 10^{-8}$. The proposal will include a change to $v$ as well as the $x_{i}$, so this calculation does not give the exact acceptance probability, but it does indicate that when $v$ is small, the acceptance probability can become very small, and the chain will remain in the same state for a very long time. Since the Markov chain leaves the correct distribution invariant, it follows that the chain will only very rarely move from a large value of $v$ (which happens to be where this run was started) to a small value for $v$; indeed, this never occurred in the actual run.

Once one suspects a problem of this sort, signs of it can be seen in the plot. In particular, starting at iteration 1,198 , the value of $v$ stays at around -3.3 for 25 iterations (i.e., for 250,000 Metropolis updates). However, there are no obvious


FIG. 14. Sampling from the funnel distribution using Metropolis and slice-sampling methods.
occurrences of this sort in the first 1,000 iterations, so the problem would not be apparent even to a suspicious user if only half as many iterations had been done. Running several chains from different starting states might have revealed the problem, but when sampling from more complex distributions, it is difficult to be sure that an appropriate variety of starting states has been tried.

The second plot in Figure 14 shows the results of sampling from the funnel distribution using single-variable Metropolis updates, applied to each variable in sequence. The proposal distribution was a Gaussian centered on the current value, with standard deviation one. Each iteration for this run consisted of 1,300 updates for each variable in turn, which take approximately as long as 10,000 multivariate Metropolis updates (with the program and machine used). As before, the plot shows the value of $v$ after each of 2,000 such iterations.

The results using single-variable Metropolis updates are not as grossly wrong as those obtained using multivariate Metropolis updates. Small values for $v$ are obtained in the expected proportion. The previous problem of very low acceptance rates when $v$ is small is avoided because even when the standard deviation for one of the $x_{i}$ given $v$ is much smaller than the proposal standard deviation, proposals to change a single $x_{i}$ are still accepted occasionally (e.g., when $v=-9$, the standard deviation of the $x_{i}$ is 0.011 , and about one proposal in 100 is accepted).

However, large values for $v$ are sampled poorly in this run. About $0.6 \%$ of the values should be greater than 7.5 (which is marked by a dotted line), but no such values are seen in the first half of the run ( 1,000 iterations, 1.3 million updates for each variable). Around iteration 1,200, the chain moves to large values of $v$ and stays there for 17 iterations ( 22,100 updates for each variable). This number of points above 7.5 is not too far from the expected number in 2,000 iterations, which is 12.4 , so in this sense the run produced approximately the right answer. However, it is clear that this was largely a matter of luck. Movement to large values of $v$ is rare, because once such a value for $v$ is reached, the chain is likely to stay at a large value for $v$ for a long time. In this case, the problem is not a high rejection rate, but rather slow exploration of the space in small steps. For example, the standard deviation of the $x_{i}$ when $v$ is 7.5 is 42.5 . Exploring a range of plus or minus twice this by a random walk with steps of size around 1 takes about $(4 \times 42.5)^{2}=28,900$ updates of each variable. While exploring this range, substantial amounts of time will be spent with values for the $x_{i}$ that are not compatible with smaller values of $v$. (This problem is not as severe in the previous run, because the multivariate proposals take larger steps, since they change all variables at once.)

We might try to avoid the problems with sampling for both large and small values of $v$ by picking the proposal standard deviation at random, from a wide range. The third plot in Figure 14 shows the results when using multivariate Metropolis proposals in which the log base 10 of the proposal standard deviation is chosen uniformly from the interval $(-3,3)$. Large values for $v$ are sampled fairly well, but small values for $v$ are still a problem, though the results are not as bad as for multivariate Metropolis with the proposal standard deviation fixed
at 1. Increasing the range of proposal standard deviations to even more than six orders of magnitude might fix the problem, but at an even greater cost in wasted computation when the random choice is inappropriate.

The bottom plot in Figure 14 shows the results of trying to sample from the funnel distribution using single-variable slice sampling. The initial interval was of size 1 , and was expanded by the stepping-out procedure (Figure 3) until both ends are outside the slice, and then sampled from with the shrinkage procedure (Figure 5). Each of the 2,000 iterations done consisted of 120 such updates for each variable in turn, which takes approximately the same amount of time as the Metropolis methods. The average number of evaluations of $f$ for these slice sampling updates was 12.7 , but a few updates required more than a hundred evaluations.

The results with single-variable slice sampling are quite good. Small values of $v$ are perhaps sampled slightly less well than with single-variable Metropolis updates, but the difference is not large. Large values of $v$ are sampled better than with any of the Metropolis methods. This good performance is due to the way the stepping out and shrinkage procedures can adapt to the local characteristics of the distribution.
9. Discussion. The table in Figure 15 summarizes the characteristics of the slice sampling methods discussed in this paper and of some competing approaches for sampling from general distributions on continuous state spaces. I list methods that update a single variable at a time separately from multivariate methods. Single-variable methods may be preferred when the coordinate system used is such that one expects many of the variables to be almost independent. Furthermore, for some distributions, recomputing the probability density after a change to one variable may be much faster than recomputing it after a change to all variables. When there are strong dependencies between variables, however, single-variable updates may converge slowly, or even be nonergodic, though simple-minded multivariate methods will not necessarily be better in such a situation.

The first column in the table indicates whether the method requires that derivatives of the log probability density be computable. Derivatives are needed by dynamical methods and reflective slice sampling, which limits their applicability. Adaptive rejection sampling [Gilks and Wild (1992), Gilks (1992)] and overrelaxed slice sampling can take advantage of derivatives, but can operate without them with only a moderate loss of efficiency; for example, with no derivatives available, overrelaxed slice sampling can find endpoints using bisection rather than Newton iteration.

The second and third columns indicate how critical it is that tuning parameters be set to good values, and whether or under what conditions "retrospective tuning" is allowed; that is, whether parameters of the method can be set based on information from past iterations. Adaptive rejection sampling (ARS) for log concave distributions is very good in these respects; one must specify a stepsize

|  | Derivatives needed? | How critical is tuning? | Retrospective tuning allowed? | Can suppress random walks? |
| :---: | :---: | :---: | :---: | :---: |
| Single-variable methods |  |  |  |  |
| ARS/ARMS | No (but helpful) | Low-Medium | If $\log$ concave | No |
| Single-variable Metropolis | No | Medium | No | No |
| Single-variable slice sampling | No | Low | If unimodal | No |
| Overrelaxed slice sampling | No (but helpful) | Low | If unimodal and endpoints exact | Yes |
| Multivariate methods |  |  |  |  |
| Multivariate Metropolis | No | Medium-High | No | No |
| Dynamical methods | s Yes | High | No | Yes |
| Slice sampling with hyperrectangles | No | Low-Medium | No | No |
| Slice sampling with Gaussian crumbs | Possibly helpful | Low-Medium | No | No |
| Reflective slice sampling | Yes | Medium-High | No | Yes |

FIG. 15. Characteristics of some general-purpose Markov chain sampling methods.
to use in searching for a point on the other side of the mode, but this parameter can be tuned retrospectively, and if it is too small, it can be rapidly increased by doubling. Parameter tuning is more of a problem when ARMS [Gilks, Best and Tan (1995)] is used for distributions not known to be log concave. A poor choice of parameters may have worse effects, and retrospective tuning is not allowed [Gilks, Neal, Best and Tan (1997)]. Tuning is also a problem for single-variable and multivariate Metropolis methods: proposing changes that are too small leads to an inefficient random walk, while proposing changes that are too large leads to frequent rejections. Metropolis methods must not be tuned retrospectively.

Single-variable slice sampling and overrelaxed slice sampling offer advantages over other methods in these respects. Whereas ARS/ARMS allows retrospective tuning only for $\log$ concave distributions, it is allowed for these slice sampling methods when they are applied to any unimodal distribution (provided the interval is expanded to the whole slice, and endpoints for overrelaxation are computed exactly). Furthermore, the tuning is less critical for slice sampling than for the other methods (apart from ARS), as discussed further below.

The final column indicates whether the method can potentially suppress random walk behavior. This is important when sampling from a distribution with high dependencies between variables, which may have to be explored in small steps, since the difference in efficiency between diffusive and systematic exploration of the distribution can then be very large. Hybrid Monte Carlo works very well as a way of suppressing random walks, provided it is tuned properly. However, using a stepsize for hybrid Monte Carlo that is too large is disastrous, since the dynamical simulation becomes unstable, and very few changes are accepted. Reflective slice sampling offers an alternative that may sometimes be easier to tune.

We can also explore the differences between these methods by seeing how well they work in various circumstances. The most favourable situation is when our prior knowledge lets us choose good tuning parameters for all methods. A Metropolis algorithm with a simple proposal distribution will then explore the distribution fairly efficiently (although in a random walk), and will have low overhead, since it requires evaluation of $f(x)$ at only a single new point in each iteration. Single-variable slice sampling will be comparably efficient, however, provided we stick with the interval chosen initially (i.e., set $m=1$ ). There will then be no need to evaluate $f(x)$ at the boundaries of the interval, and if the first point chosen from this interval is within the slice, only a single evaluation of $f(x)$ will be done. If this point is outside the slice, further evaluations will be required, but this inefficiency corresponds to the possibility of rejection with the Metropolis algorithm. Multivariate slice sampling with an initial hyperrectangle that is not expanded behaves analogously. Metropolis and slice sampling methods should therefore have similar performance when both are tuned well. However, slice sampling will work better if it turns out that we mistakenly chose too large a width for the Metropolis proposal distribution and initial slice sampling interval. This will lead to a high rejection rate for the Metropolis algorithm, but the sampling procedures of Figures 5 and 8 efficiently use rejected points to shrink the interval, lessening the impact of such a bad choice.

As seen in the demonstration of Section 8, the advantage of slice sampling over Metropolis methods can be quite dramatic if we do not know enough to choose a good tuning parameter, or if no single value of the tuning parameter is appropriate.

Another possibility is that we know that the conditional distributions are log concave, but we do not know how wide they are. Adaptive rejection sampling (ARS) with retrospective tuning will then work quite well. Single-variable slice sampling will also work well, since in this situation it too can be tuned retrospectively (provided no limit is set on the size of the interval). However, ARS does true Gibbs sampling, whereas the slice sampling updates do not produce points that are independent of the previous point. This dependency probably slows convergence, so ARS may be better than single-variable slice sampling in this context (though this will depend also on how many function evaluations each method requires).

Suppose, however, that we know only that the conditional distributions are unimodal, but not necessarily log concave. We would then need to use ARMS rather than ARS, and would not be able to tune it retrospectively, whereas we can still use single-variable slice sampling with retrospective tuning. This will likely not be as good as true Gibbs sampling, however, which we should prefer if the conditional distribution happens to be one that can be efficiently sampled from. In particular, if slice sampling is used to sample from a heavy-tailed distribution, it may move only infrequently between the tails and the central region, since this transition can occur only when we move to a point under the curve of $f(x)$ that is as low as the region under the tails. However, there appears to be no general purpose scheme that avoids problems in this situation.

Finally, consider a situation where we do not know that the conditional distributions are unimodal, and have only a rough idea of an appropriate width for a proposal distribution or initial slice sampling interval. Single-variable slice sampling copes fairly well with this uncertainty. If the initial interval is too small it can be expanded as needed, either by stepping out or by doubling-which is better depends on whether the faster expansion of doubling is worth the extra overhead from the acceptance test of Figure 6. If instead the initial interval is too big, it will be shrunk efficiently by the procedure of Figure 5 . We might try to achieve similar robustness with the Metropolis algorithm by doing several updates for each variable, using proposal distributions with a range of widths. For example, if $w$ is our best guess at an appropriate width, we might do updates with widths of $w / 4$, $w / 2, w, 2 w$ and $4 w$. This may ensure that an appropriate proposal distribution is used some of the time, but it is unattractive for two reasons. First, the limits of the range (e.g., from $w / 4$ to $4 w$ ) must be set a priori. Second, for this approach to be valid, we must continue through the original sequence of widths even after it is clear that we have gone past the appropriate one. These problems are not present with slice sampling.

Multivariate slice sampling using hyperrectangles will usually not offer much advantage over single-variable slice sampling (as is also the case with multivariate versus single-variable Metropolis methods). However, the more general framework for multivariate slice sampling based on "crumbs" that was outlined in Section 5.2 offers the possibility of adapting not just to the scales of the variables, but also to the dependencies between them. The benefits of such methods can only be determined after further research, but huge increases in efficiency would seem conceivable, if one is to judge from the analogous comparison of minimization by simple steepest descent versus more sophisticated quasi-Newton or conjugate gradient methods.

The practical utility of the slice sampling methods described here will ultimately be determined by experience in a variety of applications. Some applications will involve tailor-made sampling schemes for particular models-for instance, Frey (1997) used single-variable slice sampling to sample for latent variables
in a slice sampling for another neural network model. The adaptivity of slice sampling should prove particularly useful when using tempering or annealing methods [Geyer and Thompson (1995), Neal (2001)] in order to avoid problems with multimodality, since these methods require sampling from a whole sequence of distributions, and we would rather not have to manually tune a sampler for each one. Slice sampling is also particularly suitable for use in automatically generated samplers, and is now used in some situations by the WinBUGS system [Lunn, Thomas, Best and Spiegelhalter (2000)]. Readers can try out slice sampling methods for themselves, on a variety of Bayesian models, using the "software for flexible Bayesian modeling" that is available from my web page. This software (version of 2000-08-21) implements most of the methods discussed in this paper.

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