### Monte Carlo and Simulation

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Introduction

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## Monte Carlo: principle

$$\mathbb{E}[\varphi(X)] \approx \frac{1}{N} \sum_{n=1}^{N} \varphi(X_n)$$

with  $\varphi: \mathcal{X} \to \mathbb{R}$ .

Rationale: MSE, Law of large number, central limit theorems.

Also: confidence intervals!

Need for *simulation* methods. Note that simulation has other uses beyond Monte Carlo.

## Bibliography

- Stats, MCMC: Monte Carlo statistical methods, C.P Robert, Springer
- IID simulation: non-uniform random variate generation, L. Devroye, Springer
- in French: La simulation de Monte Carlo, B. Tuffin, Lavoisier
- QMC: Monte Carlo and quasi-Monte Carlo sampling, C. Lemieux,
   Springer
- finance: *Monte Carlo methods in financial engineering*, P. Glasserman, Springer

Pseudo-random number generators

## Pseudo-random number generators

## Famous quotes, outline

Anyone who uses software to produce random numbers is in a "state of sin". John von Neumann

One should not use a random method to generate random numbers. Donald Knuth

A PRNG is a convenient fiction. Ideally, it should:

- be fast,
- be reproducible,
- look random (at least according to statistical tests, e.g. "die-hard").

### PRNGs: a few facts

- The general structure of a PRNG:  $x_t = f(x_{t-1})$ , where  $x_t \in \{0, \dots, 2^k 1\}$ ; by construction,  $x_t$  is **periodic**.
- LCG (linear congruential generators):

$$x_{t+1} = (ax_t + c) \pmod{m}$$

and take  $u_t = x_t/m$  so that the  $u_t$ 's are in [0,1].

- Take c = 0 for simplicity (then seed 0 is forbidden; and 0 is never generated, provided m is prime, and a < m).
- Assuming m is prime, the period is m-1 iff  $a^k-1$  is a multiple of m for k=m-1, but not  $k\leq m-2$ .

### Lattice structure

- Vectors of dim d lie on at most  $(d!m)^{1/d}$  hyperplanes in the d-dimensional unit cube; e.g. for  $m=2^{31}-1$ , 108 for d=3 and 39 for d=10.
- RANDU, the most ill-conceived random number generators ever designed... has  $a = 65539 = 2^{16} + 3$ ,  $c = 2^{31}$ , and is such that  $x_t = 6x_{t-1} 9x_{t-2}$ .
- See Table 2.1 p 44 of Glasserman for better choices of (a,c).
- note that if a is not small, then computing a \* x is not easy even using floating point operations. We could take  $a = 2^k$ , but then generators typically have bad properties (see RANDU).

### More modern PRNGs

- basic LCGs (even with *good* values of *a* and *c*) are now considered obsolete.
- Combine several generators to (a) increase period; and (b) reduce lattice structure: e.g. take the sum of K generators modulo one (Wichmann-Hill).
- Mersenne twister: very popular 32-bit PRNG (Python, R, Matlab, etc), has period  $2^{19937} 1$ .
- Also push for 64-bit PRNG.

### Main conclusion

- **DO NOT** use C standard implementation rand().
- DO NOT implement your own PRNG.
- **DO** resort to some **modern** implementation of a **modern** generator, such as Mersenne twister; see e.g. GSL in C.

Non-uniform simulation

## Non-uniform simulation

### Outline

### A few general recipes:

- inversion
- rejection
- chain rule

plus several specialised ones (e.g. Box-Muller).

### inversion

### inversion algorithm

If X has CDF F, take

$$X = F^{-1}(U), \quad U \sim \mathcal{U}[0, 1].$$

Applications: exponential, Laplace, Gaussian?

### Box-Muller

#### Box-Muller

$$\begin{cases} X = \sqrt{-2\log(U)} * \cos(2\pi V) \\ Y = \sqrt{-2\log(U)} * \sin(2\pi V) \end{cases}$$

Then X,  $Y \sim N(0,1)$ , independently.

## A sneaky introduction to rejection

To understand the coming slides, note that the following algorithm

### Rejection

Repeat  $X \sim \mathcal{U}(A)$ 

Until  $X \in \mathcal{B}$ .

draws from  $\mathcal{U}(\mathcal{B})$  (provided  $\mathcal{B} \subset \mathcal{A}$ ).

### Modified Box-Muller

### Box-Muller with rejection

Repeat

$$U$$
,  $V\sim \mathcal{U}[-1,1]$  Until  $S:=U^2+V^2\leq 1$ .

Return

$$\begin{cases} X = U\sqrt{-2\log(S)/S} \\ Y = V\sqrt{-2\log(S)/S} \end{cases}$$

Then X,  $Y \sim N(0,1)$ , independently.

Note: avoid computing sin and cos.

## Rejection

Let f, g PDFs such that  $f \leq Mg$  (with  $M \geq 1$ ).

#### Accept-reject

Repeat

$$X \sim g$$
,  $U \sim \mathcal{U}[0,1]$ ,

Until  $U \leq f(X)/Mg(X)$ .

Properties:  $X \sim f$ , number of draws until acceptance is Geometric(1/M). Justification: uniform sampling under the graph, see next slide.

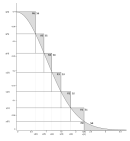
# Uniform sampling under the graph

For a function f, let  $\mathcal{G} = \{(x,y) \in \mathbb{R}^2 : 0 \le y \le Mf(x)\}$ , then

$$(X,Y) \sim \mathcal{U}(\mathcal{G}) \Leftrightarrow \begin{cases} X & \sim f \\ Y|X=x & \sim \mathcal{U}[0,Mf(x)] \end{cases}$$

Note: this construction is in fact not restricted to real-valued random variables.

# Ziggurat algorithm for N(0,1) (Marsaglia, 60?)



K Slices  $S_k = [-x_k, x_k] \times [y_k, y_{k+1}]$  constructed to have the same area.

- **1** Choose slice *k* (uniformly).
- 2 Sample (X, Y) within slice k.
- **1** If  $X \leq x_{k+1}$ , return X, else, if  $Y \leq \varphi(X)$ , return X, else go to 1.

Note: If slice 0 is selected, extra steps required (truncated Gaussian distribution).

## Multivariate simulation: chain-rule decomposition

The inverse transform method is restricted to real-valued random variables, the inverse transform *is not*.

General recipe to generate jointly (X, Y, Z), with PDF f(x, y, z):

- Generate  $X \sim f_X(x)$  (marginal). Call x the output.
- ② Generate  $Y|X=x\sim f_{Y|X}(y|x)$  (conditional given X=x). Call y the output.
- **3** Generate  $Z|X=x, Y=y\sim f_{Z|Y,X}(z|x,y)$  (full conditional). Call z the output.

### Gaussian vectors

The standard method to generate  $X \sim N_d(\mu, \Sigma)$  is:

- Generate  $Z_1, \ldots, Z_d \sim \mathcal{N}(0,1)$ .
- Compute  $C = \text{Choleksy}(\Sigma)$ . (i.e.  $\Sigma = CC^T$ , and C is lower triangular)
- Return  $X = \mu + CZ$ .

The Cholesky decomposition costs  $\mathcal{O}(d^3)$ .

Non-uniform simulation in spaces other than  $\mathbb{R}^d$ 

Non-uniform simulation in spaces other than  $\mathbb{R}^d$ 

### Outline

Some recipes to sample specific cases of

- distributions over constrained sets
- discrete distributions

## How to sample N sorted uniforms

Naive method: sample  $U_n \sim \mathcal{U}[0,1]$  for  $n=1,\ldots,N$ , return sort $(U_{1:N})$ . Cost is  $\mathcal{O}(N\log N)$  (not bad).

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- Sample  $E_1, \ldots, E_{N+1} \sim \operatorname{Exp}(1)$ .
- Compute  $V_{1:(N+1)} = \operatorname{cumsum}(E_{1:(N+1)})$ .
- Return  $(V_1/V_{N+1}, ..., V_N/V_{N+1})$ .

# How to sample uniformly on the sphere

- Sample  $X \sim N_d(0, I_d)$ .
- Return X/||X||.

## How to sample from a discrete distribution over $\mathbb N$

The inverse methods extends to the discrete case. Simply define:

$$F^{-1}(u) = \inf\{x : F(x) \ge u\}$$

In practice:

- ullet Sample  $U \sim \mathcal{U}[0,1]$
- If  $U \leq p_0$ , return 0
- If  $p_0 < U \le p_0 + p_1$ , return 1
- etc

## What if N and K are large

Suppose we want to sample N times from a distribution over  $\{0,\ldots,K-1\}$ . If we run the algorithm of the previous slide N times, we do  $\mathcal{O}(NK)$  operations (on average). Can we do better?

# What if N and K are large

Suppose we want to sample N times from a distribution over  $\{0,\ldots,K-1\}$ . If we run the algorithm of the previous slide N times, we do  $\mathcal{O}(NK)$  operations (on average). Can we do better? Solution: use as input N sorted uniforms. Then cost is  $\mathcal{O}(N+K)$ . Application: (weighted) bootstrap.

# Inverse CDF algorithm

```
def inversecdf(su.W):
    """ Input: su[0:N] sorted uniforms
                 W[0:K] normalised weights (sum to one)
        Output: A[0:N] indexes (in \{0,\ldots,K-1\})
    11 11 11
    j=0; s=W[0]; N = su.shape[0]
    A = empty(N,'int')
    for n in xrange(N):
        while su[n]>s:
            i += 1
            s += W[j]
        A[n] = j
    return A
```

## How to sample a permutation

"Naive"  $\mathcal{O}(N \log N)$  method:  $\sigma = \operatorname{argsort}(U_{1:N})$ .

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"Naive"  $\mathcal{O}(N \log N)$  method:  $\sigma = \operatorname{argsort}(U_{1:N})$ . Smart  $\mathcal{O}(N)$  method:

- Let  $\sigma = (1, 2, ..., N)$ .
- $I \sim \mathcal{U}(1, \dots, N)$ , swap  $\sigma(1)$  and  $\sigma(I)$ .
- $I \sim \mathcal{U}(2, ..., N)$ , swap  $\sigma(2)$  and  $\sigma(I)$ .
- etc.

## Variance reduction

## Objectives, outline

Given a certain quantity

$$I = \mathbb{E}[\varphi(X)] = \int_{\mathcal{X}} f(x)\varphi(x) dx$$

find a Monte Carlo estimator with smaller variance than the standard estimator

$$\frac{1}{N}\sum_{n=1}^N \varphi(X_n).$$

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### Recipes:

- antithetic variables
- control variates

### Antithetic variables

In cases where  $\varphi(-X)$  has the same distribution as  $\varphi(X)$ , use:

$$\hat{l}_{\mathrm{anti}} = \frac{1}{2N} \sum_{n=1}^{N} \{ \varphi(X_n) + \varphi(-X_n) \}$$

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#### Lemma:

$$\operatorname{Var}(\hat{I}_{\operatorname{anti}}) \leq \operatorname{Var}(\hat{I})$$

Note: we have less variance, but twice as many evaluations of  $\varphi$ ...

## Control variates (univariate case)

Let Z a real-valued r.v. such that  $\mathbb{E}(Z) = 0$ . For any  $\beta$ ,

$$\hat{I}_{cv} = \frac{1}{N} \sum_{n=1}^{N} \{ \varphi(X_n) + \beta Z_n \}$$

is an unbaised estimator of  $I = \mathbb{E}[\varphi(X)]$ .

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The smallest variance is obtained with  $\beta_{\mathrm{opt}} = -\frac{\mathrm{Cov}(\varphi(X), Z)}{\mathrm{Var}[\varphi(X)]}$ .

## Control variates (multivariate case)

 $Z^1, \ldots, Z^K$  are mean-zero real-valued r.v. Take

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In practice, replace  $\beta_k$  by  $\hat{\beta}_k$ , the OLS estimate for regression:

$$\varphi(X_n) = \alpha + \sum_{k=1}^K \beta_k Z_n^k + \varepsilon_n.$$

#### Variance reduction and Rao-Blackwellisation

Often variance reduction techniques may be cast as particular **Rao-Blackwellisation** schemes, i.e. the idea that

$$\operatorname{Var}\left[\mathbb{E}[\varphi(X)|Z]\right] \leq \operatorname{Var}[\varphi(X)].$$

Importance sampling

# Importance sampling

#### A simple identity

$$\mathbb{E}[\varphi(X)] = \int_{\mathcal{X}} \varphi(x) f(x) dx$$

$$= \int_{\mathcal{X}} \varphi(x) \frac{f(x)}{g(x)} g(x) dx = \mathbb{E}_g \left[ \frac{f(X)}{g(X)} \varphi(X) \right]$$

assuming  $\operatorname{Supp}(f) \subset \operatorname{Supp}(g)$ .

Any expectation w.r.t. PDF f may be rewritten thusly as an expectation w.r.t. PDF g (which may be easier to simulate from):

$$\hat{I}_{IS} = \frac{1}{N} \sum_{n=1}^{N} \frac{f(X_n)}{g(X_n)} \varphi(X_n).$$

### How to choose proposal g?

• Check that variance exists,  $\Leftrightarrow \mathbb{E}_g\left[\varphi(X)^2\frac{f(X)^2}{g(X)^2}\right] < \infty$ . (Sufficient condition:  $f/g \leq M$ , and  $\mathbb{E}_f[\varphi^2] < \infty$ .)

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- Optimal (in terms of minimizing variance) is

$$g_{\rm opt}(x) \propto f(x) |\varphi(x)|$$
.

It is often not possible to simulate from  $g_{\rm opt}$ , so more generally, it is recommended to take  $g \approx f$ .

#### Auto-normalised IS

Sometimes either f or g are known only up to a constant:  $f = f_u/Z_f$ ,  $g = g_u/Z_g$ , and  $Z_f$ ,  $Z_g$  are intractable. In that case, we use the auto-normalised IS estimator:

$$\hat{I}_{\text{AIS}} = \frac{\sum_{n=1}^{N} w_n \varphi(X_n)}{\sum_{n=1}^{N} w_n}, \quad w_n = \frac{f_u(X_n)}{g_u(X_n)}.$$

This estimator is biased, and asymptotically Gaussian:

$$\sqrt{N}\left(\hat{I}_{\mathrm{AIS}}-I\right)\Rightarrow N(0,v_{f/g})$$

with  $v_{f/g} = \mathbb{E}_g[\left(\frac{f}{g}\right)^2(\varphi - I)^2]$  (assuming this quantity is  $< \infty$ ).

## How to choose g (bis repetita)

#### Same points as for standard IS:

- Check that at least  $v_{f/g} < \infty$ ; sufficient condition is (a) f/g < M and (b)  $\operatorname{Var}_f(\varphi) < \infty$ .
- $\bigcirc$  Optimal g is

$$g_{\mathrm{opt}}(x) \propto f(x) |\varphi(x) - I|$$

which depends on I... In practice, take  $g \approx f.$ 

### Estimating the Z's, effective sample size

Note that AIS also provides an estimate of  $Z_f/Z_g$ :

$$\mathbb{E}\left[\frac{1}{N}\sum_{n=1}^{N}w_{n}\right]=\frac{Z_{f}}{Z_{g}}$$

and of  $v_{f/g}$ :

$$\frac{N\sum_{n=1}^{N}w_n^2\left\{\varphi(X_n)-\hat{I}\right\}^2}{\left(\sum_{n=1}^{N}w_n\right)^2}.$$

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Similarly, the effective sample size

$$\frac{\left(\sum_{n=1}^{N} w_{n}\right)^{2}}{\sum_{n=1}^{N} \left(w_{n}\right)^{2}} \in [1, N]$$

is a good indicator of AIS efficiency.

#### Curse of dimensionality

For 
$$\mathcal{X} = \mathbb{R}^d$$
,  $f(x) = \prod_{i=1}^d f_1(x_i)$ ,  $g(x) = \prod_{i=1}^d g_1(x_i)$ , one has: 
$$\mathbb{E}_{\sigma}[f^2/g^2] = C^d, \quad C > 1.$$

We expect the variance of IS to grow exponentially with the dimension.

#### Resampling

How to transform weighted sample  $(w_n, X_n)$  into an **unweighted** sample?

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How to transform weighted sample  $(w_n, X_n)$  into an **unweighted** sample? Simply draw randomly  $\tilde{X}_n$  from

$$\sum_{n=1}^{N} W_n \delta_{X_n}, \quad W_n = \frac{w_n}{\sum_{m=1}^{N} w_m}$$

(as in the bootstrap).

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See previous chapter on multinomial sampling.

## Quasi-Monte Carlo

#### Principle

Often one may rewrite quantity of interest as:

$$I = \mathbb{E}[\varphi(U)], \quad U \sim \mathcal{U}[0,1]^d$$

and then use

$$\hat{I} = \frac{1}{N} \sum_{n=1}^{N} \varphi(U_n).$$

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Can we construct (deterministic or random) points  $U_1, \ldots, U_N$  in  $[0,1]^d$  so that the approximation error is smaller than with standard Monte Carlo (i.e.  $U_n$  are IID uniforms)?

### Stratification (d = 1)

• Generate N/K uniforms in each interval [(k-1)/K, k/K],  $k=1,\ldots,K$ . (Note the connection with antithetic variables.)

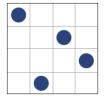
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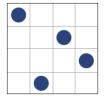
- Generate N/K uniforms in each interval [(k-1)/K, k/K],  $k=1,\ldots,K$ . (Note the connection with antithetic variables.)
- Or even take K = N, i.e. generate  $U_n \sim \mathcal{U}[(n-1)/N, n/N]$ .
- or even take  $u_n = (2n-1)/2N$ , the (deterministic) centre of interval [(n-1)/N, n/N]).

## Stratification for d > 1: Latin hypercube sampling



Generate the  $U_n$ 's so that exactly one point falls in each horizontal or vertical strip (of area 1/N).

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**Tip**: use random permutations.

### Koksma-Hlawka inequality

$$|\hat{I} - I| \leq V(\varphi) D^*(u_{1:N})$$

where  $V(\varphi)$  is the variation of  $\varphi$  (in the sense of Hardy and Krause), and

$$D^{\star}(u_{1:N}) = \sup_{[0,b] \subset [0,1]^d} \left| N^{-1} \sum_{n=1}^{N} \mathbb{I}_{[0,b]}(u_n) - \prod_{i=1}^d b_i \right|$$

is the **star discrepancy**.

#### Proof for d = 1

$$N^{-1}\sum_{n=1}^N \varphi(u_n) - \int_0^1 \varphi(u)\,du = \int_0^1 \delta(u)\varphi'(u)\,du$$

where  $\delta(u) = u - N^{-1} \sum_{n=1}^{N} \mathbf{1}(u_n \leq u)$ .

### Why is the previous inequality so important?

Because we know how to construct:

- ullet point-sets such that  $D^\star(u_{1:N}) = \mathcal{O}\left(rac{(\log N)^{d-1}}{N}
  ight)$
- ullet sequences such that  $D^\star(u_{1:N}) = \mathcal{O}\left(\frac{(\log N)^d}{N}\right)$

hence we can do **better** than Monte Carlo, i.e.  $\mathcal{O}_P(\frac{1}{\sqrt{N}})$ .

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Side note: there are good reasons to believe that these rates are optimal.

$$d = 1$$

Take 
$$u_n=(2n-1)/2N,\; n=1,\ldots,N.$$
 Then 
$$D^\star(u_{1:N})=\frac{1}{2N}.$$

## Van der Corput (sequence for d = 1)

In base b, for  $n = \sum_{j=0}^{k} a_j(n)b^j$ , take

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e.g. for b = 2: 1/2, 1/4, 3/4, 1/8, . . .

For b = 3: 1/3, 2/3, 1/9, ...

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Then  $D^*(u_{1:N}) = \mathcal{O}(\log N/N)$ .

#### d > 1: Halton & Hammersley

**Halton sequence**: component j is a van der Corput sequence in base  $b_j$ , where the  $b_j$  are the first d prime numbers. Discrepancy is  $\mathcal{O}((\log N)^d/N)$ .

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# Other low-discrepancy sequences and point sets

- Niederreiter
- Faure
- Sobol'
- .

# RQMC (randomised QMC)

QMC is purely deterministic. It lacks a simple way to evaluate the numerical error. Imagine we are able to randomise  $U_{1:N}$  so that

- $U_n \sim \mathcal{U}[0,1]^d$  (marginally).
- $Oldsymbol{0}$   $U_{1:N}$  is still a low-discrepancy point-set (or sequence).

Then

$$\mathbb{E}\left[\frac{1}{N}\sum_{n=1}^{N}\varphi(U_n)\right]=\mathbb{E}[\varphi(U)]$$

and we can evaluate the numerical error through the empirical variance (over repeated runs).

### RQMC: random shift

The simplest RQMC strategy is to generate a low-discrepancy point set  $v_{1:N}$ ,  $W \sim \mathcal{U}[0,1]^d$ , then take:

$$U_n = v_n + W \pmod{1}$$
 (componentwise)

# RQMC: a surprising result

Owen (1998) showed that for smooth functions  $\varphi$ 

$$\operatorname{Var}[\hat{I}] = \mathcal{O}\left(\frac{(\log N)^{(d-1)/2}}{N^3}\right)$$

when scrambling (a particular RQMC technique) is used.

### conclusion: QMC vs MC

- QMC has a better convergence rate.
- But for large d, QMC might need a very large N to beat MC.
- With MC, the (square) error is simple to estimate, whereas for QMC, we have only a deterministic bound, which is hard to evaluate, and is often pessimistic. See RQMC however.
- Variance reduction: may be used in conjunction with (R)QMC. (Recommendation is to do variance reduction, then replace MC with QMC).
- Practical recommendation: scrambled Sobol' seems like a good default choice (or Latin Hypercube sampling for very high dimensions).

#### Markov chain Monte Carlo

#### Outline

In some settings, simulating independently  $X \sim \pi(dx)$  is difficult, but it is possible to simulate a Markov chain  $(X_n)$  that leaves  $\pi(dx)$  invariant. Then, we still have

$$\frac{1}{N}\sum_{n=1}^{N}\varphi(X_n)\approx \mathbb{E}_{\pi}[\varphi(X)]$$

in some sense.

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This is the case in particular when density  $\pi$  is known only up to a constant.

#### **Definitions**

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- A Markov kernel K leaves distribution  $\pi$  invariant iff

$$\int_{\mathcal{X}} \pi(dx) K(x, dy) = \pi(dy).$$

• A Markov kernel is **reversible** w.r.t.  $\pi$  iff

$$\pi(dx)K(x,dy)=\pi(dy)K(y,dx).$$

This implies that  $\pi$  is invariant.

## Metropolis-Hastings

Let Q(x, dy) a Markov kernel, such that Q(x, dy) = q(x, y)dy.

#### Hastings-Metropolis step

Input:  $X_{n-1}$ 

- Generate  $Y \sim Q(X_{n-1}, dy)$
- ② With probability  $1 \wedge r(X_{n-1}, Y)$ , where

$$r(x,y) = \frac{\pi(y)q(y,x)}{\pi(x)q(x,y)}$$

accept Y, i.e.  $X_n = Y$ ; otherwise  $X_n = X_{n-1}$ .

**Property:** This kernel is reversible (w.r.t.  $\pi$ ).

#### An important practical point

Note that Hastings-Metropolis may be implemented even if  $\pi$  is known only up to a constant:  $\pi(x) = \pi_u(x)/Z$ , Z is intractable. Then

$$r(x,y) = \frac{\pi(y)q(y,x)}{\pi(x)q(x,y)} = \frac{\pi_u(y)q(y,x)}{\pi_u(x)q(x,y)}$$

# Examples of MH algorithms

• q(x,y) = q(y,x), for instance  $q(x,y) = N(x;y,\Sigma)$  (Gaussian random walk); then

$$r = \frac{\pi(y)}{\pi(x)}$$

**2** q(x,y) = q(y): **independent Metropolis**; then

$$r = \frac{\pi(y)q(x)}{\pi(x)q(y)}$$

Langevin proposal:

$$Y \sim N(x + \frac{1}{2}\Sigma\nabla\log\pi(X_n), \Sigma)$$

# (two-block) Gibbs sampling

Assume  $X = (X_1, X_2)$ ,  $\pi(x) = \pi(x_1, x_2)$ , with conditional distributions  $\pi_{1|2}(x_1|x_2)$ ,  $\pi_{2|1}(x_2|x_1)$ .

#### Gibbs sampling step

Input:  $X_{n-1} = (X_{n-1,1}, X_{n-1,2})$ 

- **①** Generate  $X_{n,1} \sim f_{1|2}(\bullet|X_{n-1,2})$ .
- ② Generate  $X_{n,2} \sim f_{2|1}(\bullet|X_{n,1})$ .

Again, this leaves invariant  $\pi$ . Gibbs can be generalised to k > 2 blocks.

#### Combining chains

- If  $K_1$ ,  $K_2$  leave  $\pi$  invariant, then so does  $K_1K_2$ .
- **② Within** Gibbs, we can replace the exact simulation of  $X_1|X_2$  (say) by a Metropolis step that leaves invariant  $\pi_{1|2}$ .

# MCMC in practice

- Assess how long it takes for the chain to reach stationarity;
- When chain seems stationary, check for intra-correlations, i.e. look at ACF (Auto-Correlation Function).

Then we compute averages

$$\frac{1}{N-N_0}\sum_{n=N_0+1}^N\varphi(X_n)$$

where  $N_0$  is burn-in time, and  $N-N_0$  is sufficiently large relative to the **auto-correlation time** (i.e. time k so that  $X_n$  and  $X_{n+k}$  are nearly uncorelated).

### scaling random walk Metropolis

One big hurdle with random walk Metropolis is the choice of  $\Sigma$ , in the proposal  $N(x, \Sigma)$ . If too small, chain moves slowly, if too large, proposals always get rejected.

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One big hurdle with random walk Metropolis is the choice of  $\Sigma$ , in the proposal  $N(x,\Sigma)$ . If too small, chain moves slowly, if too large, proposals always get rejected.

Theory (e.g. Roberts and Rosenthal, 2004) indicates that one should take

$$\Sigma = c\Sigma_{\pi}$$

where  $\Sigma_{\pi}$  is the covariance matrix of target  $\pi$ , and c calibrated so that acceptance rate is  $\approx 0.25$ .

## a tiny bit of MCMC theory

**1** From an arbitrary starting point  $X_0 = x_0$ , and any  $\varepsilon > 0$ , we have

$$\|K^n(x_0, dx_n) - \pi(dx_n)\|_{\mathrm{TV}} \le \varepsilon$$

for *n* large enough.

CLT:

$$\sqrt{N}\left(\frac{1}{N}\sum_{n=1}^N\varphi(X_n)-I\right)\Rightarrow N(0,V(\varphi))$$

with

$$V(\varphi) = \operatorname{Var}_{\pi}(\varphi) + 2 \sum_{k=1}^{\infty} \gamma_k(\varphi)$$

and  $\gamma_k(\varphi) = \text{Cov}[\varphi(X_n), \varphi(X_{n+k})].$ 

#### Adaptive MCMC?

Can we use past samples to automatically calibrate Metropolis-Hastings? e.g. at time t, do a random walk Metropolis step, of size  $\Sigma = c \hat{\Sigma}_t$ , where  $\hat{\Sigma}_t$  is the empirical covariance matrix computed from  $X_0, \ldots, X_{t-1}$ .

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Interlude: Bayesian classification

Interlude: Bayesian classification

#### Outline

Consider model with responses  $y_i \in \{-1, 1\}$ , covariates  $x_i \in \mathbb{R}^p$ , likelihood

$$L(x, y; \beta) = \prod_{i=1}^{n_d} F(y_i \beta^T x_i)$$

with  $F = \Phi$  (probit), or F = L (logit), and prior

$$\pi(\beta) = 1$$

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with  $F = \Phi$  (probit), or F = L (logit), and prior

$$\pi(\beta) = 1$$

The posterior is

$$\pi(\beta|x,y) \propto \prod_{i=1}^{n_d} F(y_i \beta^T x_i)$$

We will use this example to discuss many of the approaches seen so far.

## Monte Carlo optimisation

### Objectives

Numerical maximisation:

$$\max_{\theta \in \Theta} \psi(\theta)$$

#### when

- $\mathbf{2}$   $\psi$  is an (intractable) expectation:

$$\psi(\theta) = \mathbb{E}_{\theta}[h(X, \theta)]$$

#### Stochastic approximation

Statistical applications: MLE

#### Exploration

When  $\psi$  can be evaluated point-wise, one may sample N times from some distribution  $\pi(d\theta)$ , and return  $\max_{n=1,\dots,N} \psi(\theta_n)$ ; for instance if  $\Theta$  is compact, take  $\pi(d\theta)$  to be the Uniform dist. over  $\Theta$ .

#### **Exploration**

When  $\psi$  can be evaluated point-wise, one may sample N times from some distribution  $\pi(d\theta)$ , and return  $\max_{n=1,\dots,N} \psi(\theta_n)$ ; for instance if  $\Theta$  is compact, take  $\pi(d\theta)$  to be the Uniform dist. over  $\Theta$ . In particular, consider

$$\pi_{\lambda}(\theta) \propto \exp\{\lambda \psi(\theta)\}.$$

When  $\lambda$  (inverse temperature) increases, support of  $\pi_{\lambda}$  gets more concentrated around modal regions, but in return it may be more difficult to sample from  $\pi_{\lambda}$ .

#### Simulated annealing

Simulate a (inhomogeneous) Markov chain as follows: at iteration t, do a Metropolis step w.r.t.  $\pi_{\lambda_t}$ , and make  $\lambda_t$  increase at a logarithmic rate.

# The Cross-Entropy method

For some parametric family  $\{f_{\xi}, \xi \in \Xi\}$ , choose initial  $\xi_0$ , then iteratively:

- Sample  $\theta_1, \ldots, \theta_n \sim f_{\xi_t}$ .
- ② Estimate (using e.g. MLE)  $\xi_{t+1}$  from the 10% best of the  $\theta_i$  (in terms of  $\psi(\theta_i)$ ).

## Other heuristic optimisation procedures

- genetic algorithms
- tabu search
- ant colony algorithm

and also more specialised ones.

### Stochastic approximation

One has:  $\psi(\theta) = \mathbb{E}_{\theta}[h(X, \theta)]$  (double dependence on  $\theta$ ). Possible approaches:

- If Expectation is w.r.t. a fixed dist' f,  $\psi(\theta) = \mathbb{E}[h(X, \theta)]$ , generate  $X_1, \dots, X_n \sim f$ , maximise  $\theta \to N^{-1} \sum_{n=1}^N h(X_n, \theta)$ .
- @ Gradient-based approach, e.g.

$$\theta_{t+1} = \theta_t + \alpha_t \hat{\nabla} \psi(\theta_t)$$

where  $\hat{\nabla}\psi(\theta_t)$  is some MC estimate of the gradient of  $\psi$ .

#### Robins-Monroe

Take  $\alpha_t$  such that  $\alpha_t \to 0$ , and  $\sum_t \alpha_t = \infty$ ; e.g.  $\alpha_t = Ct^{-b}$ ,  $1/2 < b \le 1$ .

#### Robins-Monroe

Take  $\alpha_t$  such that  $\alpha_t \to 0$ , and  $\sum_t \alpha_t = \infty$ ; e.g.  $\alpha_t = Ct^{-b}$ ,  $1/2 < b \le 1$ . To estimate the gradient, if  $\psi(\theta) = \mathbb{E}_{\theta}[h(X)]$ , one has

$$abla \psi(\theta) = \mathbb{E}_{\theta}[h(X)s_{\theta}(X)], \quad s_{\theta}(x) = \frac{\partial}{\partial \theta}\log f_{\theta}(x)$$

and thus a possible choice is:

$$\hat{\nabla}\psi(\theta) = \frac{1}{N} \sum_{n=1}^{N} h(X_n) s_{\theta}(X_n)$$

## Selected applications of Monte Carlo

#### Outline

- Oerivative pricing
- Statistical applications: MCEM, Bayesian inference, ABC
- Enumeration
- Go playing...

### Derivative pricing: statement

There, X is continous-time process on [0, T], and  $\varphi$  could be:

- $\varphi(X) = (K X_T)^+$
- $\varphi(X) = (K \frac{1}{k} \sum_{i} X_{t_i})^+$
- $\varphi(X) = (K \int X_t dt)^+$
- $\varphi(X) = \mathbb{I}\{\tau_b > T\}(X_T K)^+$ , with  $\tau_b = \inf\{t : X_t \le b\}$
- etc.

## Simulating Brownian paths

For a Brownian motion  $\{W_t\}$ , several ways to simulate *exactly*  $(W_{t_1}, \ldots, W_{t_k})$ :

- random walk:  $W_{t_i}|W_{t_{i-1}} \sim N(W_{t_{i-1}}, t_i t_{i-1})$
- Brownian bridge:  $W_{t_i}|W_{t_{i-1}},W_{t_{i+1}}\sim$

$$N\left(\frac{(t_{i+1}-t_i)W_{t_{i-1}}+(t_i-t_{i-1})W_{t_{i+1}}}{t_{i+1}-t_{i-1}},\frac{(t_{i+1}-t_i)(t_i-t_{i-1})}{t_{i+1}-t_{i-1}}\right)$$

principal components

Try to think about the implications for QMC...

## QMC and Brownian paths

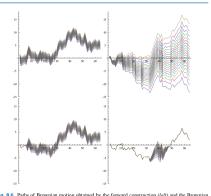


Fig. 8.6 Paths of Brownian motion obtained by the forward construction (left) and the Brownian bridge construction (right). All but one parameters are fixed

Top: all but first component fixed; bottom, all but seventh component fixed

#### Discretization

In general, diffusion processes need to be discretized:

$$dX_t = \mu(X_t)dt + \sigma(X_t)dW_t$$

becomes

$$X_{t+1} - X_t = \delta \mu(X_t) + \sigma(X_t)\epsilon_t, \quad \epsilon_t \quad N(0, \delta)$$

where  $\delta$  is the discretization step.

Choice of  $\delta$ : trade-off between discretization bias and CPU time.

#### Multi-level Monte Carlo

Consider a sequence of decreasing steps:  $\delta_0 > \ldots > \delta_L$ ; say  $\delta_l = 2^{-l}$ .

$$\mathbb{E}_{\delta_L}(arphi) = \mathbb{E}_{\delta_0}(arphi) + \sum_{l=1}^L \left\{ \mathbb{E}_{\delta_l}(arphi) - \mathbb{E}_{\delta_{l-1}}(arphi) 
ight\}$$

To get a low-variance estimate for each level, use **coupling**: e.g. use Brownian bridge construction to obtain the finer level from the coarser level.

To minimise variance, choose  $N_l$  (number of samples for level l) to be:

$$N_I \propto \sqrt{V_I/C_I}$$

where  $V_l$  (resp.  $C_l$ ) is variance (resp. CPU cost per sample) of estimate for level l.

## Other worthy points

- control variates: simulation invoves many Gaussian variables, with known mean and variance
- antithetic variables (Gaussians variables are symetric)
- QMC very popular nowadays in option pricing

### Statistical applications

- Bayesian estimation: already covered, see MCMC
- Frequentist estimation: MC for the E part of EM: MCEM
- Likelihood-free inference: see next slide

# ABC (likelihood-free inference)

Data  $y^*$ , model  $p(y|\theta)$  such that (a) one can simulate from  $p(y|\theta)$ ; (b) one cannot compute the likelihood  $p(y|\theta)$ . (Many scientific models fall in this category.)

# ABC (likelihood-free inference)

Data  $y^*$ , model  $p(y|\theta)$  such that (a) one can simulate from  $p(y|\theta)$ ; (b) one cannot compute the likelihood  $p(y|\theta)$ . (Many scientific models fall in this category.)

ABC (Approximate Bayesian inference) samples from:

$$p_{\varepsilon}(\theta, y|y^{\star}) \propto p(\theta)p(y|\theta)\mathbb{I}(\|s(y) - s(y^{\star})\| \leq \varepsilon).$$