## Monte Carlo Methods for Partial Differential Equations

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#### Early History of MCMs for PDEs

Probabilistic Representations of PDEs

Introduction

Probabilistic Representation of Elliptic PDEs via Feynman-Kac Probabilistic Representation of Parabolic PDEs via Feynman-Kac Probabilistic Approaches of Reaction-Diffusion Equations Monte Carlo Methods for PDEs from Fluid Mechanics

Monte Carlo Methods and Linear Algebra

Parallel Computing Overview
General Principles for Constructing Parallel Algorithms
Parallel N-body Potential Evaluation



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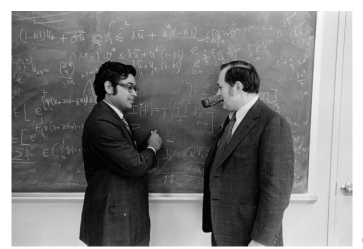
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## Dedicated to My Probability Professors at Courant: Raghu Varadhan and Monroe Donsker



- Courant, Friedrichs, and Lewy: Their pivotal 1928 paper has probabilistic interpretations and MC algorithms for linear elliptic and parabolic problems
- Fermi/Ulam/von Neumann: Atomic bomb calculations were done using Monte Carlo methods for neutron transport, their success inspired much post-War work especially in nuclear reactor design
- Kac and Donsker: Used large deviation calculations to estimate eigenvalues of a linear Schrödinger equation
- 4. Forsythe and Leibler: Derived a MCM for solving special linear systems related to discrete elliptic PDE problems





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- Curtiss: Compared Monte Carlo, direct and iterative solution methods for Ax = b
  - General conclusions of all this work (as other methods were explored) is that random walk methods do worse than conventional methods on serial computers except when modest precision and few solution values are required.
  - Much of this "conventional wisdom" needs revision due to complexity differences with parallel implementations

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### Brownian Motion and the Diffusion Equation

Cauchy problem for the diffusion equation:

$$u_t = \frac{1}{2}\Delta u \tag{1}$$

$$u(x,0) = u_0(x) \tag{2}$$

has the solution in 1-D:

$$u(x,t) = \int_{-\infty}^{\infty} \omega(x-y,t)u_0(y)dy$$
 (3)

where

$$\omega(x - y, t) = \frac{1}{\sqrt{2\pi t}} e^{-\frac{(x - y)^2}{2t}}$$
 (4)



### Brownian Motion and the Diffusion Equation

$$u(x,t) = \mathbb{E}_x[u_0(X^x(t))] \tag{5}$$

- ▶  $X^x(t)$ : a Brownian motion which has  $\omega(x y, t)$  as the transition probability of going from x to y in time t
- $\triangleright$   $\mathbb{E}_{x}[.]$ : an expectation w. r. t. Brownian motion

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A related elliptic boundary value problem (Dirichlet problem):

$$\Delta u(\mathbf{x}) = 0, \quad \mathbf{x} \in \Omega$$
  
 $u(\mathbf{x}) = f(\mathbf{x}), \quad \mathbf{x} \in \partial \Omega$  (7)

- Distribution of z is uniform on the sphere
- Mean of the values of u(z) over the sphere is u(x)
- $\triangleright$  u(x) has mean-value property and harmonic
- ightharpoonup Also, u(x) satisfies the boundary condition

$$u(\mathbf{x}) = \mathbb{E}_{x}[f(X^{\mathbf{x}}(t_{\partial\Omega}))]$$

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- Elliptic PDEs describe equilibrium, like the electrostatic field set up by a charge distribution, or the strain in a beam due to loading
- No time dependence in elliptic problems so it is natural to have the interior configuration satisfy a PDE with boundary conditions to choose a particular global solution
- Elliptic PDEs are thus part of boundary value problems (BVPs) such as the famous Dirichlet problem for Laplace's equation:

$$\frac{1}{2}\Delta u(x) = 0, \quad x \in \Omega, \quad u(x) = g(x), \ x \in \partial\Omega$$
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Probabilistic Representation of Elliptic PDEs via Fevnman-Kac

### Elliptic PDEs as Boundary Value Problems

An important equivalence for the Laplace equation is the mean value property (MVP), i.e. if u(x) is a solution to (7) then:

$$u(x) = \frac{1}{|\partial S^n(x,r)|} \int_{\partial S^n(x,r)} u(y) \, dy$$

 $\partial S^n(x,r)$  is the surface of an *n*-dimensional sphere centered at x with radius r

- Another way to express u(x) is via the Green's function:  $u(x) = \int_{\partial\Omega} G(x, y) u(y) dy$
- ▶ Showing a function has the MVP and the right boundary values establishes it as the unique solution to (7)



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Probabilistic Representation of Elliptic PDEs via Fevnman-Kac

- Early this century probabilists placed measures on different sets including sets of continuous functions
  - 1. Called Wiener measure
  - 2. Gaussian based:  $\frac{1}{\sqrt{2\pi t}}e^{-\frac{x^2}{2t}}$
  - 3. Sample paths are Brownian motion
  - 4. Related to linear PDEs
- ▶ E.g.  $u(x) = E_x[g(\beta(\tau_{\partial\Omega}))]$  is the Wiener integral representation of the solution to (7), to prove it we must check:
  - 1. u(x) = g(x) on  $\partial\Omega$ 2. u(x) has the MVP
- Interpretation via Brownian motion and/or a probabilistic Green's function
- Important:  $\tau_{\partial\Omega}=$  first passage (hitting) time of the path  $\beta(\cdot)$  started at x to  $\partial\Omega$ , statistics based on this random variable are intimately related to elliptic problems

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- ► Can generalize Wiener integrals to different BVPs via the relationship between elliptic operators, stochastic differential equations (SDEs), and the Feynman-Kac formula
- ► E.g. consider the general elliptic PDE:

$$Lu(x) - c(x)u(x) = f(x), x \in \Omega, c(x) \ge 0,$$
 (10)

$$u(x) = g(x), x \in \partial\Omega \tag{11}$$

$$L = \frac{1}{2} \sum_{i,i=1}^{s} a_{ij}(x) \frac{\partial^{2}}{\partial x_{i} \partial x_{j}} + \sum_{i=1}^{s} b_{i}(x) \frac{\partial}{\partial x_{i}},$$
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▶ The Wiener integral representation is:

$$u(x) = E_x^L \left[ \int_0^{\tau_{\partial\Omega}} \left\{ \frac{g(\beta(\tau_{\partial\Omega}))}{\tau_{\partial\Omega}} - f(\beta(t)) \right\} e^{-\int_0^t c(\beta(s)) \, ds} \, dt \right]$$
(13)

the expectation is w.r.t. paths which are solutions to the following (vector) SDE:

$$d\beta(t) = \sigma(\beta(t)) dW(t) + b(\beta(t)) dt, \ \beta(0) = x$$
 (14)

- ▶ The matrix  $\sigma(\cdot)$  is the Choleski factor (matrix-like square root) of  $a_{ij}(\cdot)$  (12)
- To use these ideas to construct MCMs for elliptic BVPs one must:

Simulate sample paths via SDEs (1

Evaluate (13) on the sample paths

3. Sample until variance is acceptable





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$$d\beta(t) = \sigma(\beta(t)) dW(t) + b(\beta(t)) dt, \ \beta(0) = x$$
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- ► The matrix  $\sigma(\cdot)$  is the Choleski factor (matrix-like square root) of  $a_{ij}(\cdot)$  (12)
- To use these ideas to construct MCMs for elliptic BVPs one must:
  - 1. Simulate sample paths via SDEs (14)
  - 2. Evaluate (13) on the sample paths
  - 3. Sample until variance is acceptable





- Can generalize Wiener integrals to a wide class of IBVPs via the relationship between elliptic operators, stochastic differential equations (SDEs), and the Feynman-Kac formula
- ▶ Recall that  $t \to \infty$  parabolic  $\to$  elliptic
- ► E.g. consider the general parabolic PDE:

$$u_t = Lu(x) - c(x)u(x) - f(x), x \in \Omega, c(x) \ge 0,$$
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$$u(x) = g(x), x \in \partial\Omega$$
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$$L = \frac{1}{2} \sum_{i,j=1}^{s} a_{ij}(x) \frac{\partial^{2}}{\partial x_{i} \partial x_{j}} + \sum_{i=1}^{s} b_{i}(x) \frac{\partial}{\partial x_{i}},$$



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the expectation is w.r.t. paths which are solutions to the following (vector) SDE:

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- ► The SDE gives us a process, and the process defines L (note: a complete definition of L includes the boundary conditions)
- We have solved only the Dirichlet problem, what about other BCs?
- ▶ Neumann Boundary Conditions:  $\frac{\partial u}{\partial n} = g(x)$  on  $\partial \Omega$
- If one uses reflecting Brownian motion, can sample over these paths
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Parabolic PDEs are evolution equations: the heat equation specifies how an initial temperature profile evolves with time, a pure initial value problem (IVP):

$$\frac{\partial u}{\partial t} = \frac{1}{2} \Delta u, \quad u(x,0) = u_0(x)$$
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- ▶ As with elliptic PDEs, there are Feynman-Kac formulas for IVPs and initial-boundary value problems (IBVPs) for parabolic PDEs
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- ▶ To solve (20) with  $u_0(x)$  general, must approximate  $u_0(x)$  with spikes and "move" the spikes via their individual normal distributions
- ▶ The approximation of a smooth  $u_0(x)$  by spikes is quite poor, and so the MCM above gives a solution with large statistical fluctuations (variance)
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- This variance reduction idea is the basis of the random gradient method:
  - Set up the gradient problem
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$$\frac{\partial u}{\partial t} = \frac{1}{2}\Delta u + cu, \quad u(x,0) = u_0(x)$$
 (22)

- The first term on the r.h.s. is diffusion and its effect may be sampled via a normally distributed random number
- ► The second term on the r.h.s. is an exponential growth/shrinkage term, can also sample its effect probabilistically:
- Think of dispatching random walkers to do the sampling
  - 1. Choose  $\Delta t$  s.t.  $\Delta t |c| <$
  - 2. Move all walkers via  $N(0, \Delta t)$
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  - 1. Choose  $\Delta t$  s.t.  $\Delta t |c| < 1$
  - 2. Move all walkers via  $N(0, \Delta t)$
  - 3. Create/destroy with prob. =  $\Delta t |c|$
  - 4. c > 0: create by doubling
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$$\frac{\partial u}{\partial t} = \frac{1}{2} \Delta u + cu, \quad u(x,0) = u_0(x)$$
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Consider the IVP for a nonlinear scalar reaction diffusion equation:

$$\frac{\partial u}{\partial t} = \frac{1}{2}\Delta u + c(u), \quad u(x,0) = u_0(x)$$
 (24)

► The associated gradient equation is:

$$\frac{\partial v}{\partial t} = \frac{1}{2} \Delta v + c'(u)v, \quad v(x,0) = \frac{\partial u_0(x)}{\partial x}$$
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► The similarity of (25) to (23) make it clear how to extend the RGM method to these nonlinear scalar reaction diffusion equations



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Probabilistic Approaches of Reaction-Diffusion Equations

- ➤ Differences/advantages of the RGM and conventional (finite-difference/finite-element) methods:
  - 1. It is computationally easy to sample from  $N(0, \Delta t)$
  - Only costly operation each time step is to sort the remaining cohort of walkers by their position
  - 3. Can use ensemble averaging to reduce the variance of the solution
  - 4. Can choose to use either gradient "particles" of equal mass or allow the mass of each particle to change
  - 5. The RGM is adaptive, computational elements (pieces of the gradient) are created where c'(u) is greatest, this is where sharp fronts appear and so fewer total computational elements are needed to spatially resolve jump-like solutions





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#### The RGM for Nonlinear Parabolic IVPs

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- ► The RGM in 2D is the same as in 1D except for recovery from the gradient
- ► Write  $u(\mathbf{x}) = G(\mathbf{x}, \mathbf{y}) * \Delta u(\mathbf{y}) = \nabla_{\mathbf{n}} G(\mathbf{x}, \mathbf{y}) * \nabla_{\mathbf{n}} u(\mathbf{y})$  (integration by parts)
- ▶ If  $\nabla_{\mathbf{n}} u(\mathbf{y}) = \delta(\mathbf{y} \mathbf{y}_0)$  then  $u(\mathbf{x}) = \nabla_{\mathbf{n}} G(\mathbf{x}, \mathbf{y}_0) = \frac{-1}{2\pi} \frac{(\mathbf{x} \mathbf{y}_0) \cdot \mathbf{n}}{\|\mathbf{x} \mathbf{y}_0\|^2}$
- ► Thus gradient recovery can be done via 2 n-body evaluations with charges n₁ and n₂ or with 1 Hilbert matrix application to the complex vector with n
- ► Can (and do) use the Rokhlin-Greengard fast multipole algorithm for gradient recovery
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A model equation for fluid dynamics is Berger's equation in one-dimension, as an IVP:

$$\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} = \frac{\epsilon}{2} \frac{\partial^2 u}{\partial x^2}, \quad u(x,0) = u_0(x)$$

The substitution  $\phi = e^{-\frac{1}{\epsilon} \int u \, dx} \iff u = -\epsilon \frac{\partial (\ln \phi)}{\partial x} = -\epsilon \frac{1}{\phi} \frac{\partial \phi}{\partial x}$  converts Berger's equation to the heat equation (Hopf, 1950)

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▶ Using the Feynman-Kac formula for the IVP for the heat equation one gets that  $\phi(x,t) = E_x[e^{-\frac{1}{\epsilon}\int_0^{\sqrt{\epsilon}\beta(t)}u_0(\xi)\,d\xi}]$ , which determines u(x,t) via the above inversion formula

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$$-i\frac{\hbar}{2\pi}u_{\tau}=\Delta u(x)-V(x)u(x)$$

- ▶ Can replace  $-i\frac{\hbar}{2\pi}\tau=t$  (imaginary time), to give a linear parabolic PDE
- ▶ Usually  $x \in \mathbb{R}^{3n}$  where there are n quantum particles, thus we are in a very high-dimensional case
- As in the above, use walks, killing, and importance sampling
- Interesting variants:
  - 1. Diffusion Monte Carlo
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The equations of two dimensional incompressible fluid dynamics are:

$$\frac{\partial \mathbf{u}}{\partial t} + (\mathbf{u} \cdot \nabla)\mathbf{u} = -\frac{1}{\rho}\nabla \rho + \gamma \Delta \mathbf{u}$$
 (26)

$$\nabla \cdot \mathbf{u} = 0, \quad \mathbf{u} = (u, v)^T$$
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- 1. Inviscid fluid:  $\gamma = 0 \implies$  Euler equations
- 2. Viscous fluid:  $\gamma \neq 0 \implies$  Navier-Stokes equations
- Since  $\nabla \cdot \mathbf{u} = 0$  there is a stream function  $\psi$  s.t.  $\mathbf{u} = (-\frac{\partial \psi}{\partial y}, \frac{\partial \psi}{\partial x})^T$ , with the vorticity  $\xi = \nabla \times \mathbf{u}$  we can rewrite as:

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- ▶ Recall the material (or total) derivative of z:  $\frac{Dz}{Dt} = \frac{\partial z}{\partial t} + (\mathbf{u} \cdot \nabla)z$ , this is the time rate of change in a quantity that is being advected in a fluid with velocity  $\mathbf{u}$
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- If we neglect boundary conditions, can represent  $\xi = \sum_{n=1}^{N} \xi_i \delta(\mathbf{x} \mathbf{x}_i)$  (spike discretization of a gradient), since in 2D the fundamental solution of the Poisson equation is  $\Delta^{-1}\delta(\mathbf{x} \mathbf{x}_i) = -\frac{1}{2\pi}\log|\mathbf{x} \mathbf{x}_i|$  we have  $\psi = \frac{1}{2\pi}\sum_{n=1}^{N} \xi_i \log|\mathbf{x} \mathbf{x}_i|$
- ▶ Can prove that if  $\xi$  is a sum of delta functions at some time  $t_0$ , it remains so for all time  $t \ge t_0$

- ► These observations on the vortex form of the Euler equations help extend ideas of a method first proposed by Chorin (Chorin, 1971):
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- This is not a MCM but only a method for converting the 2D Euler equations (PDEs) to a system of ODEs which are mathematically equivalent to the N-body problem (gravitation, particle dynamics)
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- As above, diffusion can be sampled by moving diffusing particles via the Gaussian (fundamental soln. of the diffusion eqn.)
- Thus we can extend the inviscid random vortex method (RVM) to Navier-Stokes through a fractional step approach, do the vorticity advection via the inviscid RVM and then treat the diffusion of vorticity equation by moving the vortex randomly
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Monte Carlo Methods for PDEs from Fluid Mechanics

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#### Chorin's RVM for 2D Navier-Stokes:

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- The telegrapher's equation approaches both the wave and heat equations in different limiting cases
  - 1. Wave equation:  $a \rightarrow 0$
  - 2. Heat equation:  $a, c \to \infty, 2a/c^2 \to \frac{1}{D}$
- Consider the one-dimensional telegrapher's equation, when a=0 we know the solution is given by  $F(x,t)=\frac{\phi(x+ct)+\phi(x-ct)}{2}$
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- ▶ If a particle moves with velocity c for time t it travels  $ct = \int_0^t c \, d\tau$ , if it undergoes random Poisson distributed direction reversal with probability per unit time a, the distance traveled in time t is  $\int_0^t c(-1)^{N(\tau)} d\tau$



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If we replace ct in the exact solution to the 1D wave equation by the randomized distance traveled average over all Poisson reversing paths we get:

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- In any dimension, an exact solution for the wave equation can be converted into a solution to the telegrapher's equation by replacing t in the wave equation ansatz by the randomized time  $\int_0^t (-1)^{N(\tau)} d\tau \text{ and averaging}$
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- Recall that in the Feynman-Kac formula the operator L enters in through the SDE which generates the sample paths over which expected values are taken
- If one replaces the sample paths from solutions of linear SDEs with paths derived from branching processes one can sample certain nonlinear parabolic PDEs directly (McKean, 1988)
- ► Recall that the solution to the IVP for the heat equation is represented via Feynman-Kac as:  $u(x, t) = E_x[u_0(\beta(t))]$





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Consider normal Brownian motion with exponentially distributed branching with unit branching probability per unit time, then the Feynman-Kac representation above with expectations taken over this branching process instead of normal Brownian motion solves the IVP for the Kolmogorov-Petrovskii-Piskunov equation:

$$\frac{\partial u}{\partial t} = \frac{1}{2}\Delta u + (u^2 - u), \quad u(x,0) = u_0(x)$$

has solution

$$u(x,t) = E_{x} \left[ \prod_{i=1}^{N(t)} u_{0}(x_{i}(t)) \right]$$

where the branching Brownian motion started at x at t = 0 leads to N(t) particles at time t with locations  $x_i(t)$ , i = 1, ..., N(t)

If instead of binary branching at exponentially distributed branching times there are n with probability  $p_n$ ,  $\sum_{n=2}^{\infty} p_n = 1$ , then the branching Feynman-Kac formula solves the IVP for:

$$\frac{\partial u}{\partial t} = \frac{1}{2} \Delta u + \left( \sum_{n=2}^{\infty} p_n u^n - u \right)$$

▶ Can also have  $p_n < 0$  with  $\sum_{n=2}^{\infty} |p_n| = 1$  with same result except that must have two representations for the normal and the "negative" walkers

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If  $\lambda =$  the probability per unit time of branching then we can solve the IVP for:

$$\frac{\partial u}{\partial t} = \frac{1}{2} \Delta u + \lambda \left( \sum_{n=2}^{\infty} p_n u^n - u \right)$$

If we have k(x, t) as inhomogeneous branching probability per unit time, then we solve the IVP for:

$$\frac{\partial u}{\partial t} = \frac{1}{2} \Delta u + \int_0^t k(x, \tau) \, d\tau \left( \sum_{n=2}^\infty p_n u^n - u \right)$$

▶ If  $\lambda$  = the probability per unit time of branching then we can solve the IVP for:

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- Monte Carlo has been, and continues to be used in linear algebra
- ▶ Consider the linear system: x = Hx + b, if  $||H|| = \mathbb{H} < 1$ , then

$$x^{n+1} := Hx^n + b, \quad x^0 = 0,$$

$$N = \sum_{i=0}^{\infty} H^i = (I - H)^{-1}, \quad ||N|| = \sum_{i=0}^{\infty} ||H^i|| \le \sum_{i=0}^{\infty} \mathbb{H}^i = \frac{1}{1 - \mathbb{H}}$$

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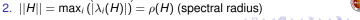
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  - 3.  $Var[a_i/p_i] \propto \sum_{i=1}^{M} \left(\frac{a_i^2}{p_i}\right)$ , so optimal choice is  $p_i \propto a_i^2$
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- ▶ We first construct a Markov chain based on H and b to sample a ℓ-fold product matrices as follows
- ▶ Define the transition probability matrix, P1.  $p_{ij} \ge 0$ , and  $p_{ij} > 0$  when  $h_{ij} \ne 0$ 2. Define  $p_{ij} = 1 \sum_{i} p_{ij}$
- Now define a Markov chain on states  $\{0, 1, ..., n\}$  with transition probability,  $p_{ii}$ , and termination probability from state i,  $p_i$
- Also define

$$V_{ij} = egin{cases} rac{h_{ij}}{p_{ij}}, & h_{ij} 
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- Given the desire to sample x<sub>i</sub> we create the following estimator based on
  - 1. Generating a Markov chain  $\gamma(i_0, i_1, \dots, i_k)$  using  $p_{ij}$  and  $p_i$ , where state  $i_k$  is the penultimate before absorbtion
  - 2 Form the estimator

$$\mathbb{X}(\gamma) = \mathbb{V}_m(\gamma) \frac{b_{i_k}}{\rho_{i_k}}, \text{ where } \mathbb{V}_m(\gamma) = \sum_{j=1}^m v_{i_{j-1}i_j}, \ m \leq k$$

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We have

$$E\left[\mathbb{X}(\gamma)|i_{0}=i\right] = \sum_{k=0}^{\infty} \left(\sum_{i_{1}} \cdots \sum_{i_{k}} p_{ii_{1}} \dots p_{i_{k-1}i_{k}} v_{ii_{1}} \dots v_{i_{k-1}i_{k}} \frac{b_{i_{k}}}{p_{i_{k}}}\right) =$$

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$$\sum_{k=0}^{\infty} \left(\sum_{i_{1}} \cdots \sum_{i_{k}} h_{ii_{1}} h_{i_{k-1}i_{k}} b_{i_{k}}\right) = \text{the } i \text{th component of } \sum_{k=0}^{\infty} H^{k} b$$

- Backward walks (Adjoint sampling)
- ▶ The lower variance Wasow estimator

$$\mathbb{X}^*(\gamma) = \sum_{m=0}^k \mathbb{V}_m(\gamma) b_{i_k}$$

- Again with only matrix-vector multiplication can do the power method and shifted variants for sampling eigenvalues
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- Now that we know all these MCMs, we must discuss how to implement these methods on parallel (and vector) computers
- There are two major classes of parallel computer being commercially produced
  - Single Instruction Multiple Data machines:
- Only one data stream so the same instruction is broadcast to all processors
- Usually these machines have many simple processors, often they are bit serial (fine grained)
- Usually these machines have distributed memory
- Connection Machine and vector-processing units are "classical" examples
- ▶ Modern examples include GPGPUs





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#### 1. Multiple Instruction Multiple Data machines:

- These machines are a collection of conventional computers with an interconnection network
- Each processor can run its own program
- Processors are usually large grained
- Can have shared or distributed memory
- Shared memory limits the number of processors though bus technology
- Distributed memory can be implemented with many different interconnection topologies
- Modern examples:
  - 1.1 Multicore machines
  - 1.2 Clustered machines
  - 1.3 Shared-memory machines





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- Use widely replicated aspects of a given problem as the basis for parallelism
  - In MCMs can map each independent statistical sample to an independent process with essentially no interprocessor communication
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- ► As a simple example, consider the Dirichlet problem for the Laplace equation (7)
- ▶ The Wiener integral representation for the solution to (7) is  $u(x) = E_x[g(\beta(\tau_{\partial\Omega}))]$
- General approaches are to use (i) "exact" samples of the Wiener integral, (ii) sample from a discretization of the Wiener integral
- ▶ Can sample with spherical processes, use the MVP and spheres to walk until  $\epsilon$  from the boundary, other elliptic L's lead to uniformity on ellipses instead of spheres (i)



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- ▶ Random Fourier series relationship to Brownian motion can be used to generate walks via a truncated series, in some cases this gives an exact random series solution which is then sampled, with complicated  $\Omega$  this approach is not practical (ii)
- Can use a high dimensional integral approximation of the infinite-dimensional (Wiener) integral, the finite-dimensional integral is then evaluated via analytic or MCMs (ii)
- ► Spatially discretize the region and sample from the discrete Wiener integral (ii)
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- All of the theory for continuous sample path Wiener integrals mentioned above carries over to the discrete cases
  - 1. Can replace the continuous region  $\Omega$  with a discretization  $\Omega_h$  where h the characteristic discretization parameter
  - 2. Replace  $\beta(\cdot)$  with random walks on  $\Omega_h$ ,  $\beta_h(\cdot)$ , this requires a transition probability matrix for the walks on the grid  $[\mathbf{P}]_{ij} = p_{ij}$
  - 3. E.g. the discrete Wiener integral solution to equation (1):  $u_h(x) = E_h^k[g(\beta_h(\tau_{\partial\Omega_h}))]$
  - 4. In this case if one has elliptic regularity of u(x) and a nonsingular discretization,  $\Omega_h$  then  $u_h(x) = u(x) + O(h^2)$

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- N-body potential problems are common in biochemistry, stellar dynamics, fluid dynamics, materials simulation, the boundary element method for elliptic PDEs, etc.
- The solution of N-body potential problems requires evaluation of function of the form:  $\Phi(\mathbf{x}) = \sum_{i=1}^{N} \phi(\mathbf{x} \mathbf{x}_i)$  for all values of  $\mathbf{x} = \mathbf{x}_i, j = 1, \dots, N$
- One heuristic solution is to replace  $\phi(\mathbf{x})$  with a cutoff version  $\phi_{co}(\mathbf{x}) = \phi(\mathbf{x})$  for  $|\mathbf{x}| \le r$  and  $\phi_{co}(\mathbf{x}) = 0$  otherwise, this reduces the problem to only keeping track of r-neighborhood points
- Can use the  $\mathbf{x}_i$ ,  $\mathbf{x}_j$  interaction as the basis for parallelism and use  $N^2$  processors to calculate the N(N-1)/2 terms in parallel, initialization of the coordinates and accumulation the results requires  $O(\log_2 N)$  operations





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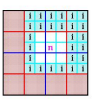
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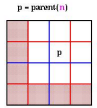


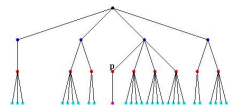


### Parallel N-body Potential Evaluation

Interaction Set(n) for the Fast Multipole Method







► The fast multipole method is an efficiency improvement over direct methods





- Algorithm is based on multipole expansion and some theory from complex variable series, consider the electrostatic description
- ▶ If  $z \in \mathbb{C}$  then a charge of intensity q at  $z_0$  results in a complex potential via Laurent series for  $|z| > |z_0|$ :

$$\phi_{z_0}(z) = q \ln(z - z_0) = q \left[ \ln(z) - \sum_{k=1}^{\infty} \frac{1}{k} \left( \frac{z_0}{z} \right)^k \right]$$

1. If we have m charges  $q_i$  at locations  $z_i$  then the potential induced by them is given by the multipole expansion for  $|z| > r = \max_i |z_i|$ :

$$\phi(z) = Q \ln(z) + \sum_{k=1}^{\infty} \frac{a_k}{z^k},$$

where 
$$Q = \sum_{i=1}^m q_i$$
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- ▶ Given an accuracy,  $\epsilon$ , one can truncate the multipole expansion to a fixed number,  $p = \lceil -\log_c \epsilon \rceil$ , of terms, where  $c = \lfloor \frac{z}{r} \rfloor$
- ▶ With p determined one can store a multipole expansion as  $\{a_1, a_2 \dots, a_p\}$
- We can move a multipole's center from z<sub>0</sub> to the origin with new coefficients:

$$b_l = -\frac{Qz_0^l}{l} + \sum_{k=1}^l a_k z_0^{l-k} {l-1 \choose k-1}$$

Note that  $\{a_1, a_2 \dots, a_p\}$  can be used to exactly compute  $\{b_1, b_2 \dots, b_p\}$ 



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1. Can also convert a multipole Laurent expansion into a local Taylor expansion:

$$\phi(z) = \sum_{l=0}^{\infty} c_l z^l, \text{ where}$$

$$c_0 = Q \ln(-z_0) + \sum_{k=0}^{\infty} \frac{a_k}{z_0^k} (-1)^k, \text{ and}$$

$$c_l = -\frac{Q}{l \dot{z}_0^l} + \frac{1}{z_0^l} \sum_{k=1}^{\infty} \frac{a_k}{z_0^k} \binom{l+k-1}{k-1} (-1)^k$$

2. And translate local expansions:

$$\sum_{k=0}^{n} a_k (z - z_0)^k = \sum_{l=0}^{n} \left( \sum_{k=l}^{n} a_k \binom{k}{l} (-z_0)^{k-l} \right) z^l$$



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- Items (1)-(4) above are the machinery required to allow the construction and use of multipole expansions, and given a multipole expansion, it requires O(N) operations to evaluate it at N points, thus an algorithm for the construction of a multipole expansion from N point charges that requires O(N) operations reduces the complexity of N-body problems to O(N) complexity
- ► The Rokhlin-Greengard algorithm achieves this by using a multiscale approach and (1)-(4)
- ▶ Consider a box enclosing  $z_1, z_2, ..., z_N$ , and  $n \approx \lceil \log_4 N \rceil$  refinements of the box, in 2D one parent box becomes four children boxes





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- Goal is to construct all p-term multipole expansions due to the particles in each box and level (upward pass) and then use these to construct local expansions in each box and level (downward pass)
  - 1. The upward pass:
    - At the finest level construct box-centered p-term multipole expansions due to the particles in each box using (1)
       At each coarser level shift child p-term multipole expansions to build box-centered p-term multipole expansions due to the particles in the particles in the particles in the particles.
  - 2. The downward pass
    - Construct local Taylor expansion in each box at the finest level by converting the p-term multipole expansions of boxes in the "interaction list" via (3)





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- ► Thus the parallel complexity is proportional to the number of levels, i.e.  $O(\log_4 N) = O(n)$
- For the nonadaptive version there a serious load balancing problem due to whether at the each level, I, each box contains exactly  $N/n^I$  particles
- This is a multigrid algorithm and so there will be the problem of idle processors at coarse levels on parallel machines with fine grainsize
- One can also implement the adaptive version of the multipole method in parallel
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- Booth, T. E. (1981) "Exact Monte Carlo solutions of elliptic partial differential equations," *J. Comp. Phys.*, **39**: 396–404.
- Brandt A. (1977) "Multi-level adaptive solutions to boundary value problems," *Math. Comp.*, **31**: 333–390.
- Chorin, A. J. (1973) "Numerical study of slightly viscous flow," *J. Fluid Mech.*, **57**: 785–796.
- Chorin, Alexandre J. and Hald, Ole H. (2006) Stochastic Tools in Mathematics and Science, Surveys and Tutorials in the Applied Mathematical Sciences, Vol. 1, viii+147, Springer, New York.
- Courant, R., K. O. Friedrichs, and H. Lewy (1928) "Über die partiellen Differenzengleichungen der mathematischen Physik," *Math. Ann.*, **100**: 32–74 (In German), Reprinted in *I.B.M. J. Res. and Dev.*, **11**: 215–234 (In English).
- Curtiss, J. H. (1953) "Monte Carlo methods for the iteration of linear operators," *J. Math. and Phys.*, **32**: 209–232.

- Booth, T. E. (1981) "Exact Monte Carlo solutions of elliptic partial differential equations," *J. Comp. Phys.*, **39**: 396–404.
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- Chorin, A. J. (1973) "Numerical study of slightly viscous flow," *J. Fluid Mech.*, **57**: 785–796.
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  - Curtiss, J. H. (1953) "Monte Carlo methods for the iteration of linear operators," *J. Math. and Phys.*, **32**: 209–232.

- Booth, T. E. (1981) "Exact Monte Carlo solutions of elliptic partial differential equations," *J. Comp. Phys.*, **39**: 396–404.
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- Chorin, Alexandre J. and Hald, Ole H. (2006) Stochastic Tools in Mathematics and Science, Surveys and Tutorials in the Applied Mathematical Sciences, Vol. 1, viii+147, Springer, New York.
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  - Curtiss, J. H. (1953) "Monte Carlo methods for the iteration of linear operators," *J. Math. and Phys.*, **32**: 209–232.

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Curtiss, J. H. (1956) "A theoretical comparison of the efficiencies of two classical methods and a Monte Carlo method for computing one component of the solution of a set of linear algebraic equations," in Symp. Monte Carlo Methods, H. A. Meyer, Ed., Wiley: New York, pp. 191–233.



Donsker, M. D., and M. Kac (1951) "A sampling method for

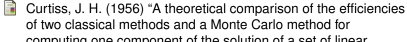


Ermakov, S. M. and G. A. Mikhailov (1982) Statistical Modeling,





Forsythe, G. E., and R. A. Leibler (1950) "Matrix inversion by a Monte Carlo method," Math. Tab. Aids Comput., 4: 127-129.



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Donsker, M. D., and M. Kac (1951) "A sampling method for determining the lowest eigenvalue and the principle eigenfunction of Schrödinger's equation,"

J. Res. Nat. Bur. Standards. 44: 551-557.

Ermakov, S. M. and G. A. Mikhailov (1982) Statistical Modeling,

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  J. Res. Nat. Bur. Standards, 44: 551–557.
- Ermakov, S. M. and G. A. Mikhailov (1982) Statistical Modeling,
- Nauka, Moscow, (in Russian).

  Frmakov, S. M. V. V. Nekrutkin and A. S. Sipin (1989) *Bandom*
- Ermakov, S. M., V. V. Nekrutkin and A. S. Sipin (1989) Random Processes for Classical Equations of Mathematical Physics, Kluwer Academic Publishers: Dordrecht.
- Forsythe, G. E., and R. A. Leibler (1950) "Matrix inversion by a Monte Carlo method," *Math. Tab. Aids Comput.*, **4**: 127–129.

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- Forsythe, G. E., and R. A. Leibler (1950) "Matrix inversion by a Monte Carlo method," *Math. Tab. Aids Comput.*, **4**: 127–129.

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- Donsker, M. D., and M. Kac (1951) "A sampling method for determining the lowest eigenvalue and the principle eigenfunction of Schrödinger's equation,"
  - J. Res. Nat. Bur. Standards. 44: 551-557.
- Ermakov, S. M. and G. A. Mikhailov (1982) Statistical Modeling, Nauka, Moscow, (in Russian).
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- Freidlin, M. (1985) Functional Integration and Partial Differential Equations, Princeton University Press: Princeton.
- Ghoniem, A. F. and F. S. Sherman (1985) "Grid-free simulation of diffusion using random walk methods," *J. Comp. Phys.*, **61**: 1-37.
- Greengard, L. F. (1988) *The Rapid Evaluation of Potential Fields in Particle Systems*, MIT Press: Cambridge, MA.
- Greengard, L. F. and W. Gropp (1990) "A parallel version of the fast multipole method," *Computers Math. Applic.*, **20**: 63-71.
- Hall, A. (1873) "On an experimental determination of  $\pi$ ," *Messeng. Math.*, **2**: 113–114.
- Halton, J. H. (1970), "A Retrospective and Prospective Survey of the Monte Carlo Method," *SIAM Review*, **12(1)**: 1–63,.
- Hammersley, J. M., and D. C. Handscomb (1964) *Monte Carlo Methods*, Chapman and Hall, London.

- Freidlin, M. (1985) Functional Integration and Partial Differential Equations, Princeton University Press: Princeton.
- Ghoniem, A. F. and F. S. Sherman (1985) "Grid-free simulation of diffusion using random walk methods," *J. Comp. Phys.*, **61**: 1-37.
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- Hammersley, J. M., and D. C. Handscomb (1964) *Monte Carlo Methods*, Chapman and Hall, London.

- Freidlin, M. (1985) Functional Integration and Partial Differential Equations, Princeton University Press: Princeton.
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- Hammersley, J. M., and D. C. Handscomb (1964) *Monte Carlo Methods*, Chapman and Hall, London.

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- Halton, J. H. (1970), "A Retrospective and Prospective Survey of the Monte Carlo Method," *SIAM Review*, **12(1)**: 1–63,.
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- Halton, J. H. (1970), "A Retrospective and Prospective Survey of the Monte Carlo Method," *SIAM Review*, **12(1)**: 1–63,.
- Hammersley, J. M., and D. C. Handscomb (1964) *Monte Carlo Methods*, Chapman and Hall, London.



Halton, J. H. (1989) "Pseudo-random trees: multiple independent sequence generators for parallel and branching computations," *J. Comp. Phys.*, **84**: 1–56.



Hillis, D. (1985) *The Connection Machine*, M.I.T. University Press:



Hopf, E. (1950) "The partial differential equation  $u_t + uu_x = \mu_{xx}$ ," *Comm. Pure Applied Math.*, **3**: 201–230.



Itô, K. and H. P. McKean, Jr. (1965) *Diffusion Processes and Their Sample Paths*, Springer-Verlag: Berlin, New York.



Kac, M. (1947) "Random Walk and the Theory of Brownian Motion," *The American Mathematical Monthly*, **54(7)**: 369–391



Kac, M. (1956) *Some Stochastic Problems in Physics and Mathematics*, Colloquium Lectures in the Pure and Applied Sciences, No. 2, Magnolia Petroleum Co., Hectographed.



Kac, M. (1980) Integration in Function Spaces and Some of Its
Applications, Lezioni Fermiane, Accademia Nazionale Dei Lir

- - Halton, J. H. (1989) "Pseudo-random trees: multiple independent sequence generators for parallel and branching computations," *J. Comp. Phys.*, **84**: 1–56.
- Hillis, D. (1985) *The Connection Machine*, M.I.T. University Press: Cambridge, MA.
- Hopf, E. (1950) "The partial differential equation  $u_t + uu_x = \mu_{xx}$ ," *Comm. Pure Applied Math.*, **3**: 201–230.
- Itô, K. and H. P. McKean, Jr. (1965) *Diffusion Processes and Their Sample Paths*, Springer-Verlag: Berlin, New York.
- Kac, M. (1947) "Random Walk and the Theory of Brownian Motion," *The American Mathematical Monthly*, **54(7)**: 369–391.
- Kac, M. (1956) Some Stochastic Problems in Physics and Mathematics, Colloquium Lectures in the Pure and Applied Sciences, No. 2, Magnolia Petroleum Co., Hectographed.
- Kac, M. (1980) Integration in Function Spaces and Some of Its Applications, Lezioni Fermiane, Accademia Nazionale Dei Lire Scuola Normale Superiore. Pisa.

- Halton, J. H. (1989) "Pseudo-random trees: multiple independent sequence generators for parallel and branching computations," *J. Comp. Phys.*, **84**: 1–56.
- Hillis, D. (1985) *The Connection Machine*, M.I.T. University Press: Cambridge, MA.
- Hopf, E. (1950) "The partial differential equation  $u_t + uu_x = \mu_{xx}$ ," *Comm. Pure Applied Math.*, **3**: 201–230.
- Itô, K. and H. P. McKean, Jr. (1965) *Diffusion Processes and Their Sample Paths*, Springer-Verlag: Berlin, New York.
- Kac, M. (1947) "Random Walk and the Theory of Brownian Motion," The American Mathematical Monthly, **54(7)**: 369–391.
- Kac, M. (1956) Some Stochastic Problems in Physics and Mathematics, Colloquium Lectures in the Pure and Applied Sciences, No. 2, Magnolia Petroleum Co., Hectographed.
- Kac, M. (1980) Integration in Function Spaces and Some of Its Applications, Lezioni Fermiane, Accademia Nazionale Dei Lir

- Halton, J. H. (1989) "Pseudo-random trees: multiple independent sequence generators for parallel and branching computations," *J. Comp. Phys.*, **84**: 1–56.
- Hillis, D. (1985) *The Connection Machine*, M.I.T. University Press: Cambridge, MA.
- Hopf, E. (1950) "The partial differential equation  $u_t + uu_x = \mu_{xx}$ ," *Comm. Pure Applied Math.*, **3**: 201–230.
- Itô, K. and H. P. McKean, Jr. (1965) *Diffusion Processes and Their Sample Paths*, Springer-Verlag: Berlin, New York.
- Kac, M. (1947) "Random Walk and the Theory of Brownian Motion," *The American Mathematical Monthly*, **54(7)**: 369–391
- Kac, M. (1956) Some Stochastic Problems in Physics and Mathematics, Colloquium Lectures in the Pure and Applied Sciences, No. 2, Magnolia Petroleum Co., Hectographed.
- Kac, M. (1980) Integration in Function Spaces and Some of Its Applications, Lezioni Fermiane, Accademia Nazionale Dei Lir Scuola Normale Superiore, Pisa.

- Halton, J. H. (1989) "Pseudo-random trees: multiple independent sequence generators for parallel and branching computations," *J. Comp. Phys.*, **84**: 1–56.
- Hillis, D. (1985) *The Connection Machine*, M.I.T. University Press: Cambridge, MA.
- Hopf, E. (1950) "The partial differential equation  $u_t + uu_x = \mu_{xx}$ ," Comm. Pure Applied Math., **3**: 201–230.
- Itô, K. and H. P. McKean, Jr. (1965) *Diffusion Processes and Their Sample Paths*, Springer-Verlag: Berlin, New York.
- Kac, M. (1947) "Random Walk and the Theory of Brownian Motion," *The American Mathematical Monthly*, **54(7)**: 369–391.
- Kac, M. (1956) Some Stochastic Problems in Physics and Mathematics, Colloquium Lectures in the Pure and Applied Sciences, No. 2, Magnolia Petroleum Co., Hectographed.
- Kac, M. (1980) Integration in Function Spaces and Some of Its Applications, Lezioni Fermiane, Accademia Nazionale Dei Lir Scuola Normale Superiore, Pisa.

- Halton, J. H. (1989) "Pseudo-random trees: multiple independent sequence generators for parallel and branching computations," *J. Comp. Phys.*, **84**: 1–56.
- Hillis, D. (1985) *The Connection Machine*, M.I.T. University Press: Cambridge, MA.
- Hopf, E. (1950) "The partial differential equation  $u_t + uu_x = \mu_{xx}$ ," Comm. Pure Applied Math., **3**: 201–230.
- ltô, K. and H. P. McKean, Jr. (1965) *Diffusion Processes and Their Sample Paths*, Springer-Verlag: Berlin, New York.
- Kac, M. (1947) "Random Walk and the Theory of Brownian Motion," *The American Mathematical Monthly*, **54(7)**: 369–391.
- Kac, M. (1956) Some Stochastic Problems in Physics and Mathematics, Colloquium Lectures in the Pure and Applied Sciences, No. 2, Magnolia Petroleum Co., Hectographed.
- Kac, M. (1980) Integration in Function Spaces and Some of Its Applications, Lezioni Fermiane, Accademia Nazionale Dei Lir Scuola Normale Superiore, Pisa.

- Halton, J. H. (1989) "Pseudo-random trees: multiple independent sequence generators for parallel and branching computations," *J. Comp. Phys.*, **84**: 1–56.
- Hillis, D. (1985) *The Connection Machine*, M.I.T. University Press: Cambridge, MA.
- Hopf, E. (1950) "The partial differential equation  $u_t + uu_x = \mu_{xx}$ ," *Comm. Pure Applied Math.*, **3**: 201–230.
- Itô, K. and H. P. McKean, Jr. (1965) *Diffusion Processes and Their Sample Paths*, Springer-Verlag: Berlin, New York.
- Kac, M. (1947) "Random Walk and the Theory of Brownian Motion," *The American Mathematical Monthly*, **54(7)**: 369–391.
- Kac, M. (1956) Some Stochastic Problems in Physics and Mathematics, Colloquium Lectures in the Pure and Applied Sciences, No. 2, Magnolia Petroleum Co., Hectographed.
- Kac, M. (1980) Integration in Function Spaces and Some of Its Applications, Lezioni Fermiane, Accademia Nazionale Dei Ling Scuola Normale Superiore, Pisa.

- Knuth, D. E. (1981) *The Art of Computer Programming, Vol. 2:* Seminumerical Algorithms, Second edition, Addison-Wesley: Reading, MA.
- Marsaglia, G. and A. Zaman "A new class of random number generators," submitted to SIAM J. Sci. Stat. Comput.
- Mascagni, M. (1991) "High dimensional numerical integration and massively parallel computing," *Contemp. Math.*, , **115**: 53–73.
- Mascagni, M. (1990) "A tale of two architectures: parallel Wiener integral methods for elliptic boundary value problems," *SIAM News*. **23**: 8.12.
- McKean, H. P. (1975) "Application of Brownian Motion to the Equation of Kolmogorov-Petrovskii-Piskunov" *Communications on Pure and Applied Mathematics*, **XXVIII**: 323–331.



- Knuth, D. E. (1981) *The Art of Computer Programming, Vol. 2:* Seminumerical Algorithms, Second edition, Addison-Wesley: Reading, MA.
- Marsaglia, G. and A. Zaman "A new class of random number generators," submitted to SIAM J. Sci. Stat. Comput.
- Mascagni, M. (1991) "High dimensional numerical integration and massively parallel computing," *Contemp. Math.*, , **115**: 53–73.
- Mascagni, M. (1990) "A tale of two architectures: parallel Wiener integral methods for elliptic boundary value problems," *SIAM News*, **23**: 8.12.
- McKean, H. P. (1975) "Application of Brownian Motion to the Equation of Kolmogorov-Petrovskii-Piskunov" *Communications on Pure and Applied Mathematics*, **XXVIII**: 323–331.



- Knuth, D. E. (1981) *The Art of Computer Programming, Vol. 2: Seminumerical Algorithms*, Second edition, Addison-Wesley: Reading, MA.
- Marsaglia, G. and A. Zaman "A new class of random number generators," submitted to SIAM J. Sci. Stat. Comput.
- Mascagni, M. (1991) "High dimensional numerical integration and massively parallel computing," *Contemp. Math.*, , **115**: 53–73.
- Mascagni, M. (1990) "A tale of two architectures: parallel Wiener integral methods for elliptic boundary value problems," *SIAM News.* **23**: 8.12.
- McKean, H. P. (1975) "Application of Brownian Motion to the Equation of Kolmogorov-Petrovskii-Piskunov" *Communications on Pure and Applied Mathematics*, XXVIII: 323–331.



- Knuth, D. E. (1981) *The Art of Computer Programming, Vol. 2:* Seminumerical Algorithms, Second edition, Addison-Wesley: Reading, MA.
- Marsaglia, G. and A. Zaman "A new class of random number generators," submitted to SIAM J. Sci. Stat. Comput.
- Mascagni, M. (1991) "High dimensional numerical integration and massively parallel computing," *Contemp. Math.*, , **115**: 53–73.
- Mascagni, M. (1990) "A tale of two architectures: parallel Wiener integral methods for elliptic boundary value problems," *SIAM News*, **23**: 8,12.
- McKean, H. P. (1975) "Application of Brownian Motion to the Equation of Kolmogorov-Petrovskii-Piskunov" Communications on Pure and Applied Mathematics, XXVIII: 323–331.



- Knuth, D. E. (1981) *The Art of Computer Programming, Vol. 2:* Seminumerical Algorithms, Second edition, Addison-Wesley: Reading, MA.
- Marsaglia, G. and A. Zaman "A new class of random number generators," submitted to SIAM J. Sci. Stat. Comput.
- Mascagni, M. (1991) "High dimensional numerical integration and massively parallel computing," *Contemp. Math.*, , **115**: 53–73.
- Mascagni, M. (1990) "A tale of two architectures: parallel Wiener integral methods for elliptic boundary value problems," *SIAM News*, **23**: 8.12.
- McKean, H. P. (1975) "Application of Brownian Motion to the Equation of Kolmogorov-Petrovskii-Piskunov" *Communications on Pure and Applied Mathematics*, **XXVIII**: 323–331.





McKean, H. P. (1988) Private communication.







pseudo-random numbers," Bull. Amer. Math. Soc., 84: 957-1041.







and a new version of the Random Walk on Boundary method." Monte Carlo Methods and Applications, 15(3): 257-284. 4 D > 4 P > 4 E > 4 E >



M. E. Muller, "Some Continuous Monte Carlo Methods for the Dirichlet Problem," The Annals of Mathematical Statistics, 27(3): 569-589, 1956.

Mikhailov, G. A. (1995) New Monte Carlo Methods With Estimating Derivatives, V. S. P. Publishers.

Niederreiter, H. (1978) "Quasi-Monte Carlo methods and pseudo-random numbers," *Bull. Amer. Math. Soc.*, **84**: 957–1041.

Rubenstein, M. (1981) Simulation and the Monte Carlo Method, Wiley-Interscience: New York.

Sabelfeld, K. K. (1991), *Monte Carlo Methods in Boundary Value Problems*, Springer-Verlag, Berlin, Heidelberg, New York.

Sabelfeld, K. and N. Mozartova (2009) "Sparsified Randomization Algorithms for large systems of linear equations and a new version of the Random Walk on Boundary method "Monte Carlo Methods and Applications, 15(3): 257–284.

- McKean, H. P. (1988) Private communication.
- M. E. Muller, "Some Continuous Monte Carlo Methods for the Dirichlet Problem," The Annals of Mathematical Statistics, 27(3): 569-589, 1956.
- Mikhailov, G. A. (1995) New Monte Carlo Methods With Estimating Derivatives, V. S. P. Publishers.
- Niederreiter, H. (1978) "Quasi-Monte Carlo methods and pseudo-random numbers," Bull. Amer. Math. Soc., 84: 957–1041.
- Rubenstein, M. (1981) Simulation and the Monte Carlo Method, Wiley-Interscience: New York.
- Sabelfeld, K. K. (1991), *Monte Carlo Methods in Boundary Value Problems*, Springer-Verlag, Berlin, Heidelberg, New York.
- Sabelfeld, K. and N. Mozartova (2009) "Sparsified Randomization Algorithms for large systems of linear equations and a new version of the Random Walk on Boundary method "Monte Carlo Methods and Applications, 15(3): 257–284.



M. E. Muller, "Some Continuous Monte Carlo Methods for the Dirichlet Problem," The Annals of Mathematical Statistics, 27(3): 569-589, 1956.

Mikhailov, G. A. (1995) New Monte Carlo Methods With Estimating Derivatives, V. S. P. Publishers.

Niederreiter, H. (1978) "Quasi-Monte Carlo methods and pseudo-random numbers," *Bull. Amer. Math. Soc.*, **84**: 957–1041.

Rubenstein, M. (1981) *Simulation and the Monte Carlo Method*, Wiley-Interscience: New York.

Sabelfeld, K. K. (1991), *Monte Carlo Methods in Boundary Value Problems*, Springer-Verlag, Berlin, Heidelberg, New York.

Sabelfeld, K. and N. Mozartova (2009) "Sparsified Randomization Algorithms for large systems of linear equations and a new version of the Random Walk on Boundary method "Monte Carlo Methods and Applications, 15(3): 257–284.

- McKean, H. P. (1988) Private communication.
- M. E. Muller, "Some Continuous Monte Carlo Methods for the Dirichlet Problem," The Annals of Mathematical Statistics, 27(3): 569-589, 1956.
- Mikhailov, G. A. (1995) New Monte Carlo Methods With Estimating Derivatives, V. S. P. Publishers.
- Niederreiter, H. (1978) "Quasi-Monte Carlo methods and pseudo-random numbers," *Bull. Amer. Math. Soc.*, **84**: 957–1041.
- Rubenstein, M. (1981) *Simulation and the Monte Carlo Method*, Wiley-Interscience: New York.
- Sabelfeld, K. K. (1991), *Monte Carlo Methods in Boundary Value Problems*, Springer-Verlag, Berlin, Heidelberg, New York.
- Sabelfeld, K. and N. Mozartova (2009) "Sparsified Randomization Algorithms for large systems of linear equations and a new version of the Random Walk on Boundary method."

  Monte Carlo Methods and Applications, 15(3): 257–284.

- McKean, H. P. (1988) Private communication.
- M. E. Muller, "Some Continuous Monte Carlo Methods for the Dirichlet Problem," The Annals of Mathematical Statistics, 27(3): 569-589, 1956.
- Mikhailov, G. A. (1995) New Monte Carlo Methods With Estimating Derivatives, V. S. P. Publishers.
- Niederreiter, H. (1978) "Quasi-Monte Carlo methods and pseudo-random numbers," *Bull. Amer. Math. Soc.*, **84**: 957–1041.
- Rubenstein, M. (1981) Simulation and the Monte Carlo Method, Wiley-Interscience: New York.
- Sabelfeld, K. K. (1991), *Monte Carlo Methods in Boundary Value Problems*, Springer-Verlag, Berlin, Heidelberg, New York.
- Sabelfeld, K. and N. Mozartova (2009) "Sparsified Randomization Algorithms for large systems of linear equations and a new version of the Random Walk on Boundary method."

  Monte Carlo Methods and Applications, 15(3): 257–284.

- McKean, H. P. (1988) Private communication.
- M. E. Muller, "Some Continuous Monte Carlo Methods for the Dirichlet Problem," The Annals of Mathematical Statistics, 27(3): 569-589, 1956.
- Mikhailov, G. A. (1995) New Monte Carlo Methods With Estimating Derivatives, V. S. P. Publishers.
- Niederreiter, H. (1978) "Quasi-Monte Carlo methods and pseudo-random numbers," *Bull. Amer. Math. Soc.*, **84**: 957–1041.
- Rubenstein, M. (1981) Simulation and the Monte Carlo Method, Wiley-Interscience: New York.
- Sabelfeld, K. K. (1991), *Monte Carlo Methods in Boundary Value Problems*, Springer-Verlag, Berlin, Heidelberg, New York.
- Sabelfeld, K. and N. Mozartova (2009) "Sparsified Randomization Algorithms for large systems of linear equations and a new version of the Random Walk on Boundary method," *Monte Carlo Methods and Applications*, **15(3)**: 257–284.

- Sherman, A. S., and C. S. Peskin (1986) "A Monte Carlo method for scalar reaction diffusion equations", *SIAM J. Sci. Stat. Comput.*, **7**: 1360–1372.
- Sherman, A. S., and C. S. Peskin (1988) "Solving the Hodgkin-Huxley equations by a random walk method", *SIAM J. Sci. Stat. Comput.*, **9**: 170–190.
- Shreider, Y. A. (1966) *The Monte Carlo Method. The Method of Statistical Trial*, Pergamon Press: New York.
- Spanier, J. and E. M. Gelbard (1969) *Monte Carlo Principles and Neutron Transport Problems*, Addison-Wesley: Reading, MA.
- Wasow, W. R. (1952), "A Note on the Inversion of Matrices by Random Walks," *Mathematical Tables and Other Aids to Computation*, **6(38)**: 78–81.



- Sherman, A. S., and C. S. Peskin (1986) "A Monte Carlo method for scalar reaction diffusion equations", *SIAM J. Sci. Stat. Comput.*, **7**: 1360–1372.
- Sherman, A. S., and C. S. Peskin (1988) "Solving the Hodgkin-Huxley equations by a random walk method", *SIAM J. Sci. Stat. Comput.*, **9**: 170–190.
- Shreider, Y. A. (1966) *The Monte Carlo Method. The Method of Statistical Trial*, Pergamon Press: New York.
- Spanier, J. and E. M. Gelbard (1969) *Monte Carlo Principles and Neutron Transport Problems*, Addison-Wesley: Reading, MA.
- Wasow, W. R. (1952), "A Note on the Inversion of Matrices by Random Walks," *Mathematical Tables and Other Aids to Computation*, **6(38)**: 78–81.



- Sherman, A. S., and C. S. Peskin (1986) "A Monte Carlo method for scalar reaction diffusion equations", *SIAM J. Sci. Stat. Comput.*, **7**: 1360–1372.
- Sherman, A. S., and C. S. Peskin (1988) "Solving the Hodgkin-Huxley equations by a random walk method", *SIAM J. Sci. Stat. Comput.*, **9**: 170–190.
- Shreider, Y. A. (1966) The Monte Carlo Method. The Method of Statistical Trial, Pergamon Press: New York.
- Spanier, J. and E. M. Gelbard (1969) *Monte Carlo Principles and Neutron Transport Problems*, Addison-Wesley: Reading, MA.
- Wasow, W. R. (1952), "A Note on the Inversion of Matrices by Random Walks," *Mathematical Tables and Other Aids to Computation*, **6(38)**: 78–81.



- Sherman, A. S., and C. S. Peskin (1986) "A Monte Carlo method for scalar reaction diffusion equations", *SIAM J. Sci. Stat. Comput.*, **7**: 1360–1372.
- Sherman, A. S., and C. S. Peskin (1988) "Solving the Hodgkin-Huxley equations by a random walk method", *SIAM J. Sci. Stat. Comput.*, **9**: 170–190.
- Shreider, Y. A. (1966) *The Monte Carlo Method. The Method of Statistical Trial*, Pergamon Press: New York.
- Spanier, J. and E. M. Gelbard (1969) *Monte Carlo Principles and Neutron Transport Problems*, Addison-Wesley: Reading, MA.
- Wasow, W. R. (1952), "A Note on the Inversion of Matrices by Random Walks," *Mathematical Tables and Other Aids to Computation*, **6(38)**: 78–81.





- Sherman, A. S., and C. S. Peskin (1986) "A Monte Carlo method for scalar reaction diffusion equations", *SIAM J. Sci. Stat. Comput.*, **7**: 1360–1372.
- Sherman, A. S., and C. S. Peskin (1988) "Solving the Hodgkin-Huxley equations by a random walk method", *SIAM J. Sci. Stat. Comput.*, **9**: 170–190.
- Shreider, Y. A. (1966) *The Monte Carlo Method. The Method of Statistical Trial*, Pergamon Press: New York.
- Spanier, J. and E. M. Gelbard (1969) *Monte Carlo Principles and Neutron Transport Problems*, Addison-Wesley: Reading, MA.
- Wasow, W. R. (1952), "A Note on the Inversion of Matrices by Random Walks," *Mathematical Tables and Other Aids to Computation*, **6(38)**: 78–81.



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