Introduction to Event Generators

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1C techniques Quadratures Monte Carlo Simulation Summary

Topics of the lectures

1 Lecture 1: The Monte Carlo Principle

2 Lecture 2: Parton level event generation

1 Lecture 3: *Dressing the Partons*

Lecture 4: Modelling beyond Perturbation Theory

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Menu of lecture 1

- Prelude: Selecting from a distribution
- Standard textbook numerical integration (quadratures)
- Monte Carlo integration
- A basic simulation example



Prelude: Selecting from a distribution

The problem

- A typical Monte Carlo/simulation problem:
 Distribution of "usual" random numbers #:
 "flat" in [0, 1].
- But: Want random numbers $x \in [x_{\min}, x_{\max}]$, distributed according to (probability) density f(x).



The exact solution

- The first method applies if both the integral of the density f(x) and its inverse are known (i.e. practically never).
- To see how it works realise that the diff. probability $\mathcal{P}(x \in [x', x' + \mathrm{d}x']) = f(x')\mathrm{d}x'$.
- Therefore: x given by

$$\int_{x_{\min}}^{x} dx' f(x') = \# \int_{x_{\min}}^{x_{\max}} dx' f(x').$$

Since everything known:

$$x = F^{-1}[F(x_{\min}) + \#(F(x_{\max}) - F(x_{\min}))].$$

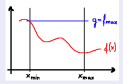


The work-around solution: "Hit-or-miss"

(Solution, if exact case does not work.)

- Builds on "over-estimator" g(x) (G and G^{-1} known): g(x) > f(x) $\forall x \in [x_{\min}, x_{\max}].$
- Select an x according to g (with exact algorithm);
- Accept with probability f(x)/g(x) (with another random number);
- Obvious fall-back choice for g(x):

$$g(x) = \operatorname{Max}_{[x_{\min}, x_{\max}]} \{ f(x) \}.$$



MC techniques

Quadratures: standard numerical integration

Reminder: Basic techniques

- Typical problem: Need to evaluate an integral, cannot do it in closed form.
- Example: nonlinear pendulum. Can calculate period T from E.o.M. $\ddot{\theta} = -g/I \sin \theta$:

$$T = \sqrt{\frac{8I}{g}} \int_{0}^{\theta_{\text{max}}} \frac{d\theta}{\sqrt{\cos \theta - \cos \theta_{\text{max}}}}$$

Elliptic integral, no closed solution known => entering (again) the realm of numerical solutions.



Numerical integration: Newton-Cotes method

- Nomenclature now: Want to evaluate $I_f^{(a,b)} = \int_f^b dx f(x)$.
- Basic idea: Divide interval [a, b] in N subintervals of size $\Delta x = (b-a)/N$ and approximate

$$I_f^{(a,b)} = \int_a^b \mathrm{d}x f(x) \approx \sum_{i=0}^{N-1} f(x_i) \Delta x = \sum_{i=0}^{N-1} f(a+i\Delta x) \Delta x,$$

i.e. replace integration by sum over rectangular panels.

Obvious issue: What is the error? How does it scale parametrically with "step-size" (or, better, number of function calls)? Answer: It is linear in Δx .



Improving on the error: Trapezoid, Simpson and all that

- A careful error estimate suggests that by replacing rectangles with trapezoids the error can be reduced to quadratic in Δx .
- This boils down to including a term [f(b) f(a)]/2:

$$I_f^{(a,b)} pprox \sum_{i=1}^{N-1} f(x_i) \Delta x + \frac{\Delta x}{2} [f(a) + f(b)]$$

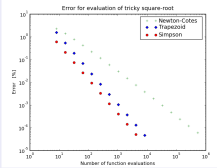
• Repeating the error-reducing exercise replaces the trapezoids by parabola: The Simpson rule. In so doing, the error decreases to $(\Delta x)^4$.



Numerical integration: Results

• Consider test function $f(x) = \sqrt{4 - x^2}$ in [0, 2].

$$(I_f^{(0,2)} = \int_0^2 \mathrm{d}x \sqrt{4 - x^2} = \pi).$$





Convergence of numerical integration: Summary

 First observation: Numerical integrations only yield estimators of the integral, with an estimated accuracy given by the error.

(Proviso: the function is sufficiently well behaved.)

- Scaling behaviour of the error translates into scaling behaviour for the number of function calls necessary to achieve a certain precision.
- In one dimension/per dimension, therefore, the convergence scales like
 - Trapezium rule: $\simeq 1/N^2$
 - Simpson's rule $\simeq 1/N^4$

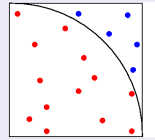
with the number N of function calls.



Monte Carlo integration

The underlying idea: Determination of π

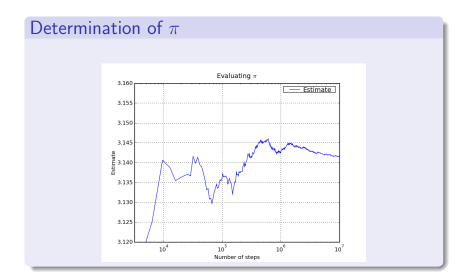
• Use random number generator!





Throw random points (x,y), with x, y in [0,1]

For hits: $(x^2 + y^2) < r^2 = 1$





Error estimate in Monte Carlo integration

• MC integration: Estimate integral by N probes

$$I_f^{(a,b)} = \int_a^b dx f(x)$$

$$\longrightarrow \langle I_f^{(a,b)} \rangle = \frac{b-a}{N} \sum_{i=1}^N f(x_i) = \langle f \rangle_{a,b},$$

where x_i homogeneously distributed in [a, b]

Basic idea for error estimate: statistical sample
 use standard deviation as error estimate

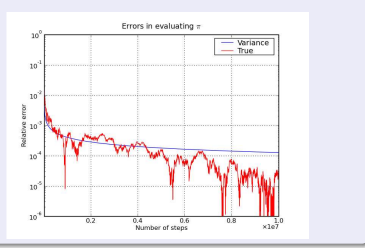
$$\langle E_f^{(a,b)}(N) \rangle = \sigma = \left[\frac{\langle f^2 \rangle_{a,b} - \langle f \rangle_{a,b}^2}{N} \right]^{1/2}.$$

Independent of the number of integration dimensions!
 Method of choice for high-dimensional integrals.



MC techniques Quadratures **Monte Carlo** Simulation Summary

Determination of π : Errors





Improve convergence: Importance sampling

Want to minimise number of function calls.

(They are potentially CPU-expensive.)

- \Longrightarrow Need to improve convergence of MC integration.
- First basic idea: Samples in regions, where f largest

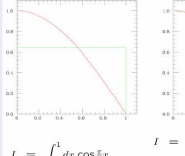
(\Longrightarrow corresponds to a Jacobian transformation of integral.)

- Algorithm:
 - Assume a function g(x) similar to f(x).
 - Obviously f(x)/g(x) is smooth $\Longrightarrow \langle E(f/g) \rangle$ is small.
 - Must sample according to dx g(x) rather than dx: g(x) plays role of probability distribution; we know already how to deal with this!
- Works, if f(x) is well-known. Hard to generalise.



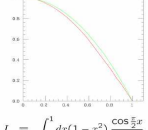
Importance sampling: Example results

• Consider $f(x) = \cos \frac{\pi x}{2}$ and $g(x) = 1 - x^2$:



$$I = \int_0^1 dx \cos \frac{\pi}{2} x$$

= 0.637 \pm 0.308/\sqrt{N}



$$I = \int_0^1 dx (1 - x^2) \frac{\cos \frac{\pi}{2} x}{1 - x^2}$$
$$= \int d\rho \frac{\cos \frac{\pi}{2} x}{1 - x^2} [x(\rho)]$$
$$= 0.637 \pm 0.032 / \sqrt{N}$$



Improve convergence: Stratified sampling

• Want to minimise number of function calls.

(They are potentially CPU-expensive.)

- \implies Need to improve convergence of MC integration.
- Basic idea here: Decompose integral in M sub-integrals

$$\langle I(f) \rangle = \sum_{j=1}^{M} \langle I_j(f) \rangle, \quad \langle E(f) \rangle^2 = \sum_{j=1}^{M} \langle E_j(f) \rangle^2$$

Then: Overall variance smallest, if "equally distributed".

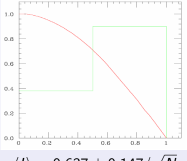
 $(\Longrightarrow \mathsf{Sample}, \mathsf{where} \mathsf{\ the} \mathsf{\ fluctuations} \mathsf{\ are.})$

- Algorithm:
 - Divide interval in bins (variable bin-size or weight);
 - adjust such that variance identical in all bins.



Stratified sampling: Example results

• Consider $f(x) = \cos \frac{\pi x}{2}$ and $g(x) = 1 - x^2$:

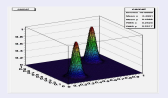


$$\langle I \rangle = 0.637 \pm 0.147 / \sqrt{N}$$

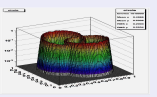


Example for stratified sampling: VEGAS

 Good for Vegas: Singularity "parallel" to integration axes



 Bad for Vegas: Singularity forms ridge along integration axes





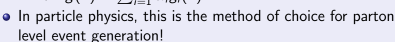
Improve convergence: Multichannel sampling

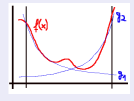
• Want to minimise number of function calls.

(They are potentially CPU-expensive.)

⇒ Need to improve convergence of MC integration.

- Basic idea: Best of both worlds: Hybrid between importance and stratified sampling.
- Have "bins" weight α_i of "eigenfunctions" $g_i(x)$: $\implies g(\vec{x}) = \sum_{i=1}^{N} \alpha_i g_i(\vec{x}).$





Basic simulation paradigm

An example from thermodynamics

• Consider two-dimensional Ising model:

$$\mathcal{H}=-J\sum_{\langle ij
angle} s_i s_j$$
 (Spins fixed on 2-D lattice with nearest neighbour interactions.)

- Traditional model to understand (spontaneous) magnetisation & phase transitions.
- To evaluate an observable \mathcal{O} , sum over all micro states $\phi_{\{i\}}$, given by the individual spins. (Similar to path integral in QFT.)

$$\langle \mathcal{O} \rangle = \int \mathcal{D}\phi_{\{i\}} \operatorname{Tr} \left\{ \mathcal{O}(\phi_{\{i\}}) \exp \left[-\frac{\mathcal{H}(\phi_{\{i\}})}{k_B T} \right] \right\}$$

Typical problem in such calculations (integrations!):
 Phase space too large

need to sample.



chniques Quadratures Monte Carlo **Simulation** Summary

Metropolis-Algorithm

- Metropolis algorithm simulates the canonical ensemble, summing/integrating over micro-states with MC method.
- Necessary ingredient: Interactions among spins in probabilistic language (will come back to us.)
- Algorithm will look like: Go over the spins, check whether they flip (compare $\mathcal{P}_{\mathrm{flip}}$ with random number), repeat to equilibrate.
- To calculate $\mathcal{P}_{\mathrm{flip}}$: Use energy of the two micro-states (before and after flip) and Boltzmann factors.
- While running, evaluate observables directly and take thermal average (average over many steps).



Why Metropolis is correct: Detailed balance

- Consider one spin flip, connecting micro-states 1 and 2.
- ullet Rate of transitions given by the transition probabilities ${\cal W}$
- If $E_1 > E_2$ then $\mathcal{W}_{1 o 2} = 1$ and $\mathcal{W}_{2 o 1} = \exp\left(-\frac{E_1 E_2}{k_B T}\right)$
- In thermal equilibrium, both transitions equally often:

$$\mathcal{P}_2 \mathcal{W}_{2 \to 1} = \mathcal{P}_1 \mathcal{W}_{1 \to 2}$$

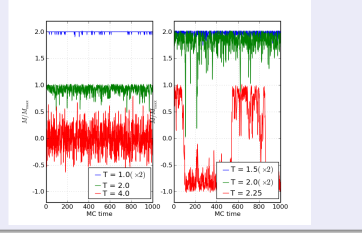
This takes into account that the respective states are occupied according to their Boltzmann factors.

$$(\mathcal{P}_i \sim \exp(-E_i/k_BT))$$

 In principle, all systems in thermal equilibrium can be studied with Metropolis - just need to write transition probabilities in accordance with detailed balance, as above
 general simulation strategy in thermodynamics.

Some example results

ullet Fix temperature, use a 10×10 lattice



Summary of lecture 1

- Discussed some basic numerical techniques.
- Introduced Monte Carlo integration as the method of choice for high-dimensional integration space (phase space).
- Introduced some standard improvement strategies to the convergence of Monte Carlo integration.
- Discussed connections between simulations and Monte Carlo integration with the example of the Ising model.

