NUMERICAL APPROACHES TO QUANTUM MANY-BODY PROBLEMS, IPAM, JANUARY 22, 2009

Introduction to quantum Monte Carlo: The Stochastic Series Expansion method Anders Sandvik, Boston University

- Illustration of concept; classical Monte Carlo example
- Detailed account of SSE for the S=1/2 Heisenberg model

This presentation is based on material available at http://physics.bu.edu/~sandvik/programs/

A simple SSE program (Fortran90) for the 2D Heisenberg model can be downloaded from this site

References:

- A. W. Sandvik, Phys. Rev. B 59, 14157 (1998),
- O. F. Syljuasen and A. W. Sandvik, Phys. Rev. E 66, 046701 (2002)

Quantum Monte Carlo for S=1/2 Heisenberg models

Classical Monte Carlo - we know the energies for all configurations

$$\langle A
angle = rac{1}{Z} \sum_n A_n \mathrm{e}^{-E_n/T}, \ \ Z = \sum_n \mathrm{e}^{-E_n/T}$$

Quantum systems - we normally don't know the eigenstates/energies

- often the goal is to find the ground state |0>
- for small N (<25), we can diagonalize the hamiltonian
- number of basis states = 2^{N} (size of the matrix)
- block diagonalize using symmetries; up to N=40 possible
- not enough to study N→∞ limit

QMC based on quantum mechanical thermal expectation value

$$egin{array}{rcl} \langle A
angle &=& rac{1}{Z} {
m Tr} \{A {
m e}^{-H/T} \} = rac{1}{Z} \sum_{s} \langle s | A {
m e}^{-H/T} | s
angle \ Z &=& {
m Tr} \{ {
m e}^{-H/T} \} = \sum_{s} \langle s | {
m e}^{-H/T} | s
angle \end{array}$$

The trace can be written in any basis

 But the exponential operator is in general difficult to evaluate Quantum Monte Carlo; rewrite the exponential operator

Can be done in different ways; many different algorithms

Warm-up: SSE for a classical problem

Classical thermal expectation value

$$\langle f \rangle = \frac{1}{Z} \sum_{\{\sigma\}} f(\sigma) e^{-\beta E(\sigma)}, \qquad Z = \sum_{\{\sigma\}} e^{-\beta E(\sigma)}$$

Classical (e.g., Ising) spins: $\sigma = \{\sigma_1, \sigma_2, \dots, \sigma_N\}$

Classical Monte Carlo: Importance sampling of spin configurations Probability of generating a configuration

$$P(\sigma) = \frac{1}{Z}W(\sigma), \quad W(\sigma) = e^{-\beta E(\sigma)}$$

Estimate of expectation value based on sampled configurations

$$\langle f \rangle = \langle f \rangle_W = \frac{1}{N_{\text{samples}}} \sum_i f(\sigma[i])$$

Imagine that we are not able to evaluate the exponential function How could we proceed then?

Use Taylor expansion of the exponential function

$$\langle f \rangle = \frac{1}{Z} \sum_{\{\sigma\}} \sum_{n=0}^{\infty} f(\sigma) \frac{(-\beta E)^n}{n!}, \qquad Z = \sum_{\{\sigma\}} \sum_{n=0}^{\infty} \frac{(-\beta E)^n}{n!}$$

Expansion power n is a new "dimension" of the configuration space To ensure positive-definitness we may have to shift E (must be < 0) $E(\sigma) \rightarrow E(\sigma) - \epsilon$

The sampling weight for the **configurations** (σ,n) is

$$W(\sigma, n) = \frac{\beta^n [\epsilon - E(\sigma)]^n}{n!}$$

The function to be averaged (estimator) $f(\sigma)$ is the same as before; it does not depend on n

$$\langle f \rangle = \langle f \rangle_W = \frac{1}{N_{\text{samples}}} \sum_i f(\sigma[i])$$

However, if $f(\sigma)$ is a function of the energy it can be rewritten as a function of n only!

Define:
$$H(\sigma) = \epsilon - E(\sigma)$$

 $\langle H \rangle = \frac{1}{Z} \sum_{\sigma,n} H(\sigma) W(\sigma, n), \quad Z = \sum_{\sigma,n} W(\sigma, n), \quad W(\sigma, n) = \frac{\beta^n H(\sigma)^n}{n!}$

Shift summation index: m=n+1

$$\sum_{\sigma,n} H(\sigma) W(\sigma,n) = \sum_{\sigma,m} \frac{m}{\beta} W(\sigma,m)$$

Therefore the energy expectation value is

$$\langle H \rangle = \frac{1}{\beta} \langle n \rangle_W \Rightarrow E = \epsilon - \frac{1}{\beta} \langle n \rangle_W$$

We can also easily obtain

$$\langle H^2 \rangle = \frac{1}{\beta^2} \langle n(n-1) \rangle_W$$

And thus the specific heat $C = \beta^{-1} (\langle E^2 \rangle - \langle E \rangle^2)$ is

$$C = \frac{1}{\beta} (\langle n^2 \rangle - \langle n \rangle^2 - \langle n \rangle)$$

What range of expansion orders n is sampled?

From the preceding results we obtain

 $\langle n \rangle = \beta(\epsilon - E)$

 $\langle n^2 \rangle - \langle n \rangle^2 = \beta (C + \epsilon - E)$

Consider low T; $\mathbf{C} \rightarrow \mathbf{0}$

 $\langle n^2 \rangle - \langle n \rangle^2 = \langle n \rangle$

Thus, for a system with N spins:

Average expansion order $\propto \beta N$ Width of distribution $\propto \sqrt{\beta N}$

These results hold true for quantum systems as well H^n requires more complicated treatment

Quantum-mechanical SSE

Thermal expectation value

$$\langle A \rangle = \frac{1}{Z} \operatorname{Tr} \{ A e^{-\beta H} \}, \quad Z = \operatorname{Tr} \{ e^{-\beta H} \}$$

Choose a basis and Taylor expand the exponential operator

$$Z = \sum_{\alpha} \sum_{n=0}^{\infty} \frac{\beta^n}{n!} \left\langle \alpha | (-H)^n | \alpha \right\rangle$$

Write the hamiltonian as a sum of local operators

 $H = -\sum_{a,b} H_{a,b}$ **a** = operator type (e.g., 1=diagonal, 2=off-diagonal) **b** = lattice unit (e.g., bond connecting sites i,j)

such that for every a, b: $H_{a,b}|\alpha\rangle = h_{a,b}(\alpha)|\alpha'\rangle$ (no branching)

Write the powers of H in terms of "strings" of these operators

$$(-H)^n = \sum_{\{H_{ab}\}} \prod_{p=1}^n H_{a(p),b(p)}$$

Operator strings of varying length n

• as in the classical case $\langle n \rangle = -\beta \langle H \rangle$

Fixed-length operator strings: introduce unit operator: $H_{0,0} = 1$

Expansion cut-off M: add M-n unit operators to each string

• there are M!/n!(M-n)! ways of doing this \Rightarrow

$$(-H)^{n} = \sum_{\{H_{ab}\}} \frac{(M-n)!n!}{M!} \prod_{p=1}^{M} H_{a(p),b(p)} \qquad \begin{array}{l} \mathbf{n} = \text{number} \\ \text{of non-[0,0]} \\ \text{operators} \end{array}$$

The truncation should not be considered an approximation

• M can be chosen such that the **truncation error is negligible**

$$Z = \sum_{\alpha} \sum_{\{H_{ab}\}} \frac{\beta^n (M-n)!}{M!} \left\langle \alpha \left| \prod_{i=1}^M H_{a(i),b(i)} \right| \alpha \right\rangle$$

The terms $(\alpha, \{H_{ab}\})$ are sampled according to weight in this sum

- requires positive-definiteness
- to this end, a constant may have to be added to diagonal H_{ab}
- there can still be a "sign problem" arising from off-diagonal H_{ab}

SSE algorithm for the S=1/2 Heisenberg model

• The algorithm for this model is particularly simple and efficient $H = J \sum_{\langle i,j \rangle} \mathbf{S}_i \cdot \mathbf{S}_j$

Consider **bipartite lattice** (sign problem for frustrated systems)



Standard z-component basis:

$$|\alpha\rangle = |S_1^z, S_2^z, \dots, S_N^z\rangle, \qquad S_i^z = \pm \frac{1}{2}$$

Bond operators: bond b connects sites i(b),j(b)

$$H = \sum_{b=1}^{B} \left[S_{i(b)}^{z} S_{j(b)}^{z} + \frac{1}{2} \left(S_{i(b)}^{+} S_{j(b)}^{-} + S_{i(b)}^{-} S_{j(b)}^{+} \right) \right]$$

1

$$H = -\sum_{b=1}^{B} \sum_{a=1}^{2} H_{a,b}$$

Diagonal and off-diagonal bond operators

$$H_{1,b} = \frac{1}{4} - S_{i(b)}^{z} S_{j(b)}^{z}, \qquad H_{2,b} = \frac{1}{2} \left(S_{i(b)}^{+} S_{j(b)}^{-} + S_{i(b)}^{-} S_{j(b)}^{+} \right)$$

A minus sign in front of the off-diagonal H_{2b} is neglected

- this corresponds to a **sublattice rotation**; 180 degree rotation in the xy-plane of the spin operators on sublattice B
- The sign is irrelevant for a bipartite lattice (will be shown later) M

SSE operator string $\prod_{p=1} H_{a(p),b(p)}$

Represented in the computer program by

 $\operatorname{opstring}[\mathbf{p}] = 2b(p) + a(p) - 1$

Spin state $|\alpha>$ represented by

 ${\tt spin}[{\tt i}]=2S_i^z$



SSE partition function $Z = \sum_{\alpha} \sum_{\{H_{ab}\}} \frac{\beta^n (M-n)!}{M!} \left\langle \alpha \left| \prod_{i=1}^M H_{a(i),b(i)} \right| \alpha \right\rangle$

Both H_{1b} and H_{2b} give 0 when acting on parallel spins • non-zero matrix element = 1/2 in both cases

Define propagated states

$$|\alpha(p)\rangle = \prod_{j=1}^{P} H_{a(j),b(j)} |\alpha\rangle \qquad |\alpha\rangle = |\alpha(0)\rangle$$

For a contributing configuration: $|\alpha(M)\rangle = |\alpha(0)\rangle$ (periodic) The **configuration weight** is then

$$W(\alpha, \{H_{ab}\}) = \left(\frac{\beta}{2}\right)^n \frac{(M-n)!}{M!}$$

Periodicity requires an even number of spin flips

- This is why the sign of H_{2b} is irrelevant for a bipartite lattice
- For a frustrated lattice an odd number of flips is possible

Graphical representation

• 1D example; 8 spins, M=12							1D: bond b connects sites b and b+1				
i = 1	2	3	4	5	6	7	8				
spin[i] = +1	+1	-1	-1	+1	-1	+1	-1	q	a(p)	b(p)	opstring[p]
•	•	0	0	٠	0	•	0	1	1	2	1 3.11
•	•	0	0		0		0	Ŧ	Ŧ	2	4
		0	0		0		0	2	0	0	0
•		Ŭ						3	2	4	9
•	•	0	•	0	0	_	0	4	2	6	13
•	•	0		0	٠	0	0	E	1	2	G
•	•	0		0		0	0	Э	Ŧ	3	0
		0		0		0	0	6	0	0	0
•		•						7	0	0	0
•			•	0	•	0	0	8	1	2	4
•		0	•	0	•	0	0	٥	2	6	10
•		0		0	0		0	9	2	0	10
		0		0	0		0	10	0	0	0
•				_			Ŭ	11	2	4	9
•		0	0		0			12	1	7	14
•		0	0		0		0		–	•	T A

Linked-list representation

• vertex: operator and spins before and after the operator has acted

• replace spins between vertices by **links**



• linked vertex list used in some parts of the program

A vertex has 4 "legs", numbered l=0,1,2,3:



position **p** of operator in operator string opstring[p], vertex leg $l \Rightarrow position v$ in linked vertex list: v=1+l+4*(p-1)

vertexlist[v] contains the element # to which v is linked

		1 =	Θ		1		2		3	р
<pre>[v] vertexlist[v]</pre>	<pre>vertexlist[v]:</pre>	[1]	31	[2]	32	[3]	29	[4]	17	1
		[5]	Θ	[6]	0	[7]	0	[8]	Θ	2
		[9]	43	[10]	44	[11]	18	[12]	42	3
		[13]	35	[14]	47	[15]	33	[16]	34	4
		[17]	4	[18]	11	[19]	30	[20]	41	5
		[21]	Θ	[22]	Θ	[23]	Θ	[24]	Θ	6
		[25]	Θ	[26]	0	[27]	0	[28]	0	7
		[29]	3	[30]	18	[31]	1	[32]	2	8
		[33]	15	[34]	16	[35]	13	[36]	45	9
		[37]	0	[38]	0	[39]	0	[40]	0	10
		[41]	20	[42]	12	[43]	9	[44]	10	11
	[45]	36	[46]	48	[47]	14	[48]	46	12	

Sampling the SSE configurations; updates

1) Diagonal update

- replace unit operator by diagonal operator, and vice versa $H_{0,0} \leftrightarrow H_{1,b}$

2) **Off-diagonal update** (local or loop)

- change the operator type, diagonal off-diagonal, for two (local) or several (loop) operators $\{H_{a_1,b_1}, H_{a_2,b_2}, \dots, H_{a_m,b_m}\} \leftrightarrow \{H_{3-a_1,b_1}, H_{3-a_2,b_2}, \dots, H_{3-a_m,b_m}\}$
- 3) Flip spins in the state $|\alpha>$
 - unconstrained "free" spins; weight unchanged after flip
 - only possible at high temperatures; strictly not necessary $S^z_i \to -S^z_i$

Updates satisfy detailed balance:

$$P_{\text{accept}}(A \to B) = \min\left(\frac{W(B)P_{\text{select}}(B \to A)}{W(A)P_{\text{select}}(A \to B)}, 1\right)$$

Diagonal update

- Carried out in opstring[p] for p=1,...,M
- State $|\alpha(p-1)\rangle$ stored in spin[]



Insertion of a diagonal operator if opstring[p]=0

Generate bond index b at random, attempt opstring[p]=2*b

- can only be done if spin[i(b)] ≠ spin[j(b)]
- n increases by 1; weight ratio

 $\frac{W(n+1)}{W(n)} = \frac{\beta/2}{M-n}$

<u>Removal of a diagonal operator if opstring[p]≠0</u>

• n decreases by 1; weight ratio $\frac{W(n-1)}{W(n)} = \frac{M-n+1}{\beta/2}$

B ways of selecting b but only one way of removing an operator; D = (b + 0)

$$\frac{P_{\text{select}}(0 \to 0)}{P_{\text{select}}(0 \to b)} = B$$

Accept probabilities: $P_{\text{accept}}(n \to n+1) = \min\left(\frac{B\beta/2}{M-n}, 1\right)$

 $P_{\text{accept}}(n \to n-1) = \min\left(\frac{M-n+1}{B\beta/2}, 1\right)$

Local off-diagonal update (obsolete)

Change type of 2 operators on the same bond

- cannot always be done; check for constraining operators
- no weight change; accept with fixed probability (e.g., P=1)



Note: periodic boundary conditions in the "propagation" direction • update spanning across the boundary affects the stored state $|\alpha>$



Local updates typically are **not very efficient**

- critical slowing-down
- no winding-number or particle-number fluctuations

Loop update

- carried out in the linked-vertex-list representation
- move "vertically" along links and "horizontally" on the same operator
- spins flipped at all vertex-legs visited; operator type changes; weight unchanged
- construct all loops, flip with probability 1/2 (as in Swendsen-Wang)



Monte Carlo step

- a cycle of diagonal updates (p=1,...,M in opstring[p])
- construction of the linked vertex list
- construct all loops, flip each with probability 1/2
- map updated vertex list back to opstring[], spin[]

Starting the simulation

- "empty" perator string, opstring[p]=0, p=1,...,M
- M is arbitrary, e.g., M=20
- random spin state; spin[p]=+1,-1

Determining the cut-off M

- after each, MC step, compare expansion order n with M
- if M-n<n/a, with, e.g., a=3, then M=n+n/a



Generalization of loop update; directed loops

In the case of the isotropic S=1/2 model

- There are only 4 non-0 vertices
- The operators uniquely define all loops
- Loops are non-self-intersecting

Directed loops

- In general, there are more than 4 allowed vertices
- A vertex is entered at some entrance leg
- The path can proceed (exit) through any of the 4 legs
- Exit probabilities are obtained from directed-loop equations
- Loops can back-track ("bounce") and self-intersect
- Bounces can be avoided for some models (more efficient)

Can be used for spins, bsons, 1D fermions,...