Assignment 8

https://github.com/jchryssanthacopoulos/quantum_information/tree/main/assignment_8

Quantum Information and Computing AA 2022–23

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Real-Space Renormalization Group



- Real-space renormalization group algorithm can be used to compute ground-state energy E_0 of quantum Ising model with traverse field in one dimension
- Following steps are repeated until $|E_0^{(i+1)} E_0^{(i)}| < \epsilon$, where i is iteration number:
 - **1** Starting with Hamiltonian of system with N sites, \hat{H}_N , construct Hamiltonian by replicating system:

$$\hat{H}_{2N} = \hat{H}_N \otimes \mathbb{1}_N + \mathbb{1}_N \otimes \hat{H}_N + \hat{H}_{int}$$

where $\hat{H}_{\text{int}} = \hat{A}_N \otimes \hat{B}_N$ is interaction between left and right bipartitions, initialized with $\hat{A}_N = \mathbb{1}_{N-1} \otimes \sigma^x$, $\hat{B}_N = \sigma^x \otimes \mathbb{1}_{N-1}$. \hat{H}_N is initialized to Hamiltonian of Ising model with N sites

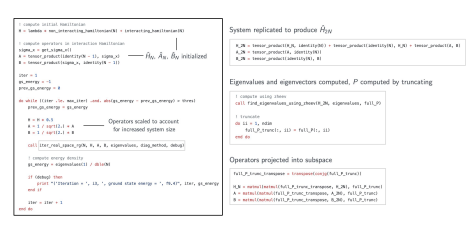
- 2 Diagonalize \hat{H}_{2N} , obtaining ground-state energy $E_0^{(i)}$. Construct projector using 2^N eigenvectors with lowest energy, $P = \sum_{i=1}^{2^N} |E_i\rangle \langle E_j|$
- 3 Project operators into subspace spanned by chosen eigenvectors:

$$\hat{H}_N = P^{\dagger} \hat{H}_{2N} P, \quad \hat{A}_N = P^{\dagger} (\hat{A}_N \otimes \mathbb{1}_N) P, \quad \hat{B}_N = P^{\dagger} (\mathbb{1}_N \otimes \hat{B}_N) P$$

Implementation



Program computes ground-state energy given number of sites N, interaction strength λ , termination threshold ϵ , and maximum number of iterations



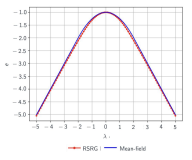
Results

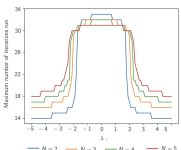


■ Energy density, $e \equiv E_0/N$, computed using RSRG compared to mean-field result:

$$e_{\mathsf{MF}} = \begin{cases} -1 - \lambda^2/4, & |\lambda| \leq 2\\ -|\lambda|, & |\lambda| > 2 \end{cases}$$

- Energy densities are very similar, but deviate the most when $0<|\lambda|<2$, when external field is present but not strong enough to coordinate all spins
- Experiments showed that energy density doesn't change with N, as expected
- lacktriangle Number of iterations increases with N, except when field is weak





Density Matrix Renormalization Group



Density matrix renormalization group algorithm improves truncation rule, resulting in more accurate final state while increasing computational complexity. Below are the steps:

1 Starting with Hamiltonian of system with N sites, \hat{H}_1 , add one site to right to get Hamiltonian of enlarged block:

$$\hat{H}_B = \hat{H}_1 \otimes \mathbb{1}_1 + \mathbb{1}_N \otimes \hat{H}_{single} + \hat{H}_{12} \otimes \sigma^x$$

where \hat{H}_{single} is Ising Hamiltonian with single site (i.e., $\lambda \sigma^z$) and \hat{H}_{12} is Hamiltonian representing rightmost spin of \hat{H}_1 (i.e., $\mathbb{1}_{N-1} \otimes \sigma^x$)

2 Replicate system to produce right enlarged block. Total Hamiltonian is:

$$\hat{H} = \hat{H}_B \otimes \mathbb{1}_{N+1} + \mathbb{1}_{N+1} \otimes \hat{H}_B + \hat{H}_{int}$$

where \hat{H}_{int} is interaction between left and right blocks (i.e., $\mathbb{1}_N \otimes \sigma^x \otimes \sigma^x \otimes \mathbb{1}_N$)

- 3 Diagonalize \hat{H} , obtaining $E_0^{(i)}$ with wavefunction $|\psi_0\rangle$. Construct density matrix $\rho = |\psi_0\rangle \langle \psi_0|$, and reduced density matrix of left bipartition, $\rho_L = \operatorname{Tr}_R \rho$. Construct projector using 2^N eigenvectors with highest eigenvalues, $P = \sum_{i=1}^{2^N} |w_j\rangle \langle w_j|$
- 4 Project operators into subspace spanned by chosen eigenvectors:

$$\hat{H}_1 = P^{\dagger} \hat{H}_B P, \quad \hat{H}_{12} = P^{\dagger} (\hat{H}_{12} \otimes \mathbb{1}_1) P$$

Implementation and Results



Ground-state energy with DMRG similar to that with RSRG, with percent difference between 1-4%

```
I enlarge the block
H_12 = tensor_product(A, sigma_x)
H_enlarged_1 = tensor_product(H_1, identity(1)) + lambda * tensor_product(identity(N), sigma_z) + H_12
! interaction (always the same?)
H_23 = tensor_product(tensor_product(identity(N), sigma_x), tensor_product(sigma_x, identity(N)))
! Hamiltonian of left and right enlarged blocks
H = tensor_product(H_enlarged_1, identity(N + 1)) + tensor_product(identity(N + 1), H_enlarged_1) + H_23
! diagonalize H
call find eigenvalues using zheev(H, eigenvalues, eigenvectors)
! compute density matrix of ground state
rho = compute_density_matrix(eigenvectors(:, 1))
! compute left reduced density matrix
rho_reduced_L = compute_left_reduced_density_matrix(rho, d, 2 * N + 2, N + 1)
! diagonalize reduced density matrix
call find eigenvalues_using_zheev(rho_reduced_L, eigenvalues_rho, eigenvectors_rho)
! truncate to get projection operator
do 11 = 1. m
     if (debug) then
        print *, "Using eigenvector with eigenvalue = ", eigenvalues_rho(ndim + 1 - ii)
     P(:, ii) = eigenvectors rhg(:, ndim + 1 - ii)
P transpose = transpose(conig(P))
```

! project into lower-dimensional subspace
H_1 = matmul(matmul(P_transpose, H_enlarged_1), P)
A = matmul(matmul(P_transpose, tensor product(A, identity(1))), P)

