# 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

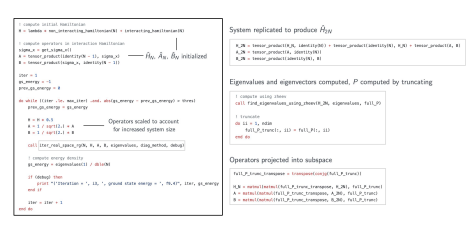
- 2 Diagonalize  $\hat{H}_{2N}$ , obtaining ground-state energy  $E_0^{(i)}$ . Construct projector using  $2^N$  eigenvectors with lowest energy,  $P = \sum_{i=1}^{2^N} |E_i\rangle \langle E_j|$
- 3 Project operators into subspace spanned by chosen eigenvectors:

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

### **Implementation**



Program computes ground-state energy given number of sites N, interaction strength  $\lambda$ , termination threshold  $\epsilon$ , and maximum number of iterations



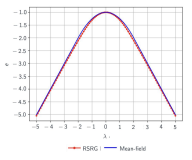
#### Results

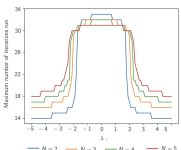


■ Energy density,  $e \equiv E_0/N$ , computed using RSRG compared to mean-field result:

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

- 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  $\hat{H}_{\text{single}}$  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)

