# **QOSF Mentorship Assessment Task**

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## Question:

Find the lowest eigenvalue of the following matrix:

$$\begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 0 & -1 & 0 \\ 0 & -1 & 0 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix}.$$

using VQE-like circuits, created by yourself from scratch.

#### Solution:

My solution involves usage of Qiskit platform and is based on a very generalizable approach, i.e., not restrictive to the current Hamiltonian itself and can be outlined as follows:

- 1. Hamiltonian in Pauli Basis
- 2. Calculations of Expectation Value
- 3. Designing Ansatze
- 4. Running Simulations
- 5. Running Noisy Simulations
- 6. Extension to Excited Energy States
- 7. Interpreting the Results

import numpy as np

In [1]:

# **Necessary Imports**

```
import scipy as sp
import itertools
import functools as ft
%matplotlib notebook
import matplotlib.pyplot as plt
from mpl_toolkits import mplot3d
import re

In [65]:
#!pip install qiskit
from qiskit import *
from qiskit.providers.aer.noise import NoiseModel
from qiskit.providers.aer.noise import QuantumError, ReadoutError
from qiskit.providers.aer.noise import phase_amplitude_damping_error
from qiskit.providers.aer.noise import deplarizing_error
from qiskit.providers.aer.noise import thermal_relaxation_error
from qiskit.providers.aer.noise import complete_meas_cal, CompleteMeasFitter
```

## Hamiltonian in Pauli Basis

$$\sigma_0 = egin{pmatrix} 1 & 0 \ 0 & 1 \end{pmatrix} \quad \sigma_x = egin{pmatrix} 0 & 1 \ 1 & 0 \end{pmatrix} \quad \sigma_y = egin{pmatrix} 0 & -i \ i & 0 \end{pmatrix} \quad \sigma_z = egin{pmatrix} 1 & 0 \ 0 & -1 \end{pmatrix}$$

The normalized Pauli matrices  $\{\sigma_0, \sigma_x, \sigma_y, \sigma_z\}/\sqrt{2}$  form an orthogonal basis of  $\mathcal{M}_2$ , this vector space can be endowed with a scalar product called the Hilbert-Schmidt inner product:  $\langle A, B \rangle = Tr(A^\dagger B)$ . Since the Pauli matrices anticommute, their product is traceless, and since they are Hermitian this implies that they are orthogonal with respect to that scalar product. Hence this property can be used for decomposing a Hamiltonian as:

$$H = \sum_{i_1, \dots, i_n = \{0, x, y, z\}} h_{i_1, \dots, i_n} \cdot \frac{1}{2^n} \sigma_{i_1} \otimes \dots \otimes \sigma_{i_n}$$
  $h_{i_1, \dots, i_n} = \frac{1}{2^n} \mathrm{Tr} ig( (\sigma_{i_1} \otimes \dots \otimes \sigma_{i_n})^\dagger \cdot H ig) = \frac{1}{2^n} \mathrm{Tr} ig( (\sigma_{i_1} \otimes \dots \otimes \sigma_{i_n}) \cdot H ig)$ 

## Decomposing Hamiltonian into Pauli Terms

```
def decompose_ham_to_pauli(H):
    """Decomposes a Hermitian matrix into a linear combination of Pauli operators.

Args:
    H (array[complex]): a Hermitian matrix of dimension 2**n x 2**n.

Returns:
    tuple[list[float], list[string], list [ndarray]]: a list of coefficients,
    a list of corresponding string representation of tensor products of Pauli observables that decompose the Hamiltonian, and
    a list of their matrix representation as numpy arrays
    """
    n = int(np.log2(len(H)))
```

```
# Sanity Checks
if H.shape != (N. N):
    raise ValueError(
         "The Hamiltonian should have shape (2**n, 2**n), for any qubit number n>=1"
if not np.allclose(H, H.conj().T):
    raise ValueError("The Hamiltonian is not Hermitian")
sI = np.eye(2, 2, dtype=complex)
sX = np.array([[0, 1], [1, 0]], dtype=complex)
sZ = np.array([[1, 0], [0,-1]], dtype=complex)
sY = complex(0,-1)*np.matmul(sZ,sX)
paulis = [sI, sX, sY, sZ]
paulis_label = ['I', 'X', 'Y', 'Z']
obs = []
coeffs =
matrix = []
for term in itertools.product(paulis, repeat=n):
    matrices = [pauli for pauli in term]
    coeff = np.trace(ft.reduce(np.kron, matrices) @ H) / N
    coeff = np.real_if_close(coeff).item()
    # Hilbert-Schmidt-Product
    if not np.allclose(coeff, 0):
         coeffs.append(coeff)
         obs.append(''.join([paulis_label[[i for i, x in enumerate(paulis)
         if np.all(x == t)][0]]+str(idx) for idx, t in enumerate(reversed(term))]))
        matrix.append(ft.reduce(np.kron, matrices))
return obs, coeffs , matrix
```

## Composing Hamiltonian into Pauli Terms

```
In [4]:
          def compose_ham_from_pauli(terms, coeffs):
               """Composes a Hermitian matrix from a linear combination of Pauli operators.
                    tuple[list[float], list[string]]: a list of coefficients,
                    a list of corresponding string representation of tensor products of
                    Pauli observables that decompose the Hamiltonian.
               H (array[complex]): a Hermitian matrix of dimension 2**n x 2**n.
               pauli_qbs = [re.findall(r'[A-Za-z]|-?\d+\.\d+|\d+', x) for x in terms]
               qubits = max([max(list(map(int, x[1:][::2]))) for x in pauli_qbs])
               N = int(2**np.ceil(np.log2(qubits+1)))
               sI = np.eye(2, 2, dtype=complex)
               sX = np.array([[0, 1], [1, 0]], dtype=complex)
sZ = np.array([[1, 0], [0,-1]], dtype=complex)
               sY = complex(0, -1)*np.matmul(sZ, sX)
               paulis = [sI, sX, sY, sZ]
paulis_label = ['I', 'X', 'Y', 'Z']
paulis_term = {'I':sI, 'X':sX, 'Y':sY, 'Z':sZ}
hamil = np.zeros((2**N, 2**N), dtype=complex)
               for coeff, pauli_qb in zip(coeffs, pauli_qbs):
    term_str = ['I'] * N
                 for term, index in zip(pauli_qb[0:][::2], pauli_qb[1:][::2]):
                    term_str[int(index)] = term
                 matrices = [paulis_term[pauli] for pauli in term_str]
                 term_matrix = np.asarray(ft.reduce(np.kron, matrices[::-1]))
                 hamil += coeff*term_matrix
               # Sanity Check
               if not np.allclose(hamil, hamil.conj().T):
                    raise ValueError("The Hamiltonian formed is not Hermitian")
               return hamil
```

## Test Hamiltonian

0.

, -1.1246, 0.182 ,

0.

$$H_0 = egin{bmatrix} 0.7056 & 0 & 0. & 0. & 0. \\ 0. & -1.1246 & 0.182 & 0. \\ 0. & 0.182 & 0.4318 & 0. \\ 0. & 0 & 0. & 0.888 \end{bmatrix}$$

```
In [5]:
        H0 = np.array([
                                    , 0.
                                                 , -0.
                          0.7056
                                                                0.
                                    , -1.1246
                                                 , 0.182
                                                             , 0.
                        [ 0.
                                    , 0.182
                                                 , 0.4318
                                                             , -0.
                        Γ-0.
                                    , 0.
                                                  , -0.
                                                               0.888
                        Γ 0.
                       ])
        НΘ
Out[5]: array([[ 0.7056, 0.
                               , -0.
```

```
, 0.182 , 0.4318, -0.
                     , 0. , -0.
                                    , 0.888 ]])
        a, b , c = decompose_ham_to_pauli(H0)
In [6]:
H_0 = 0.2252 \times I_0 I_1 + 0.3435 \times Z_0 - 0.4347 \times Z_1 + 0.91 \times X_0 X_1 + 0.91 \times Y_0 Y_1 + 0.5716
In [7]: compose_ham_from_pauli(a, b)
                                +0.j, 0.
Out[7]: array([[ 0.7056+0.j, 0.
                     +0.j, -1.1246+0.j, 0.182 +0.j,
                                                  0.
0.
               0.
                                                        +0.j],
                     +0.j, 0.182 +0.j, 0.4318+0.j,
               0.
                                                        +0.j],
                                            +0.j, 0.888 +0.j]])
                                +0.j, 0.
In [8]: | assert(np.isclose(compose_ham_from_pauli(a, b).real, H0).all())
```

#### Given Hamiltonian

Γ-0.

$$H_1 = egin{bmatrix} 1. & 0 & 0. & 0. \ 0. & 0 & -1 & 0. \ 0. & -1 & 0 & 0. \ 0. & 0 & 0. & 1. \end{bmatrix}$$

```
H1 = np.array([[1, 0, 0, 0],
                            [0, 0, -1, 0],
                             [0, -1, 0, 0],
                            [0, 0, 0, 1]])
 In [10]: | a, b, c = decompose_ham_to_pauli(H1)
Out[10]: (['I0I1', 'X0X1', 'Y0Y1', 'Z0Z1'], [0.5, -0.5, -0.5, 0.5])
                                                                         H_1 = 0.5 	imes (I_0I_1 - X_0X_1 - Y_0Y_1 + Z_0Z_1)
In [11]: compose_ham_from_pauli(a, b)
Out[11]: array([[ 1.+0.j, 0.+0.j, 0.+0.j, 0.+0.j], [ 0.+0.j, 0.+0.j, -1.+0.j, 0.+0.j], [ 0.+0.j, -1.+0.j, 0.+0.j, 0.+0.j], [ 0.+0.j, 0.+0.j, 0.+0.j, 0.+0.j, 1.+0.j]])
In [12]: assert(np.isclose(compose_ham_from_pauli(a, b).real, H1).all())
```

# Calculating the Expectation values

For an n-qubit quantum state, measuring one qubit corresponds to projecting the quantum state onto one of two half-spaces determined by the unique eigenvalues of our measurement operator.

By convention, performing a computational basis measurement is equivalent to measuring in measuring Pauli Z, which gives us two eigenvectors  $|0\rangle$  and  $|1\rangle$ , with corresponding eigenvalues  $\pm 1$ . Therefore, doing a computation measurement of a qubit and obtaining 0 means the state of our qubit is in the +1 eigenstate of the Z operator. In general, we can then use the definition of the tensor product to perform multi-qubit measurements as will be explained below.

```
For some qubit |\psi\rangle we have |\psi\rangle=\alpha|0\rangle+\beta|1\rangle, we can calculate expectation value of a single-qubit operator M=\sum_m m|m\rangle\langle m|, as
\langle M \rangle = \langle \psi | \sum_m m | m \rangle \langle m | \psi \rangle = \sum_m m \langle \psi | m^\dagger m | \psi \rangle = \sum_m m \ p(m) = (+1)p(0) + (-1)p(1)
. Similarly, for a two-qubit operator M we will get \langle M 
angle = p(00) + p(11) - p(01) - p(10)
```

## Change of Basis

Therefore, to measure a qubit, we can use any 2 imes 2 operator that is a unitary transformation of Z. That is, we could also use a measurement operator  $M = U^\dagger Z U$ , where U is an unitary operator and M gives two unique outcomes of a measurement in its  $\pm 1$  eigenvectors. Thereofre, to measure in a basis other than the computational (or Z), we need to find the corresponding unitary equivalence. Similar to the one-qubit case, all two-qubit Pauli-measurements can be written as  $(U_1^{\dagger} \otimes U_2^{\dagger})[Z \otimes Z](U_1 \otimes U_2)$  for  $2 \times 2$ unitary matrices  $U_1$  and  $U_2$ .

For X, and Y basis measurements, these unitary equivalence are given below:

$$X=(H)^\dagger Z H$$
  $Y=(HS^\dagger)^\dagger Z (HS^\dagger)$ 

This feature can be exploited to create circuits for both X basis and Y basis measurements from the standard Z basis. It is done by applying the corresponding unitary operation to the prepared quantum state and then performing a measurement in Z basis.

For example: In order to do a Y basis measurement, first apply  $HS^\dagger$  to the quantum state and then do a normal Z basis measurement.

Similar to the one-qubit case, all multi-qubit Pauli-measurements can be written as a generalization of above.

For example: For two-qubit X basis measurement,  $X \otimes X = (H \otimes H)[Z \otimes Z](H \otimes H)$ .

## **Constructing Measurement Circuits**

```
def measure_circuit(basis, num_qubits):
In [13]:
              Generate measurement circuit according to the Pauli observable
              string provided.
              basis (str): String representation of tensor products of Pauli
              observables
              num_qubits (int): Number of qubits in the circuit
              measure_qc (QuantumCircuit): Measurement Circuit for the
              corresponding basis.
              basis_qb = re.findall(r'[A-Za-z]|-?\d+\.\d+|\d+', basis)
              basis = basis_qb[0:][::2]
              qubit = list(map(int, basis_qb[1:][::2]))
              measure gc = QuantumCircuit(num gubits, num gubits)
              for base, qb in zip(basis, qubit):
                if base == 'I' or base == 'Z':
                  pass
                elif base == 'X':
                  measure qc.h(qb)
                elif base == 'Y
                  measure_qc.sdg(qb)
                  measure_qc.h(qb)
                else:
                  raise ValueError("Wrong Basis provided")
              for qb in range(num_qubits):
                  measure_qc.measure(qb, qb)
              return measure_qc
In [14]:
          def calculate_expecation_val(circuit, basis, shots=2048, backend='qasm_simulator'):
              Calculate expectation value for the measurement of a circuit in a
              given basis.
              Args:
              circuit (QuantumCircuit): Circuit using which expectation value
              will be calculated for a given basis.
              basis (str): String representation of tensor products of Pauli
              observables.
              shots (int): Number of times measurements needed to be done for calculating
              probability.
              backend (str): Backend for running the circuit.
              Returns:
              exp (float): Expectation value for the measurement of a circuit
              in a given basis.
              exp_circuit = circuit + measure_circuit(basis, circuit.num_qubits)
              result = execute(exp_circuit, backend=Aer.get_backend(backend),
                              shots=shots).result()
              exp = 0.0
              for key, counts in result.get_counts().items():
```

## Examples

return exp/shots

exp += (-1)\*\*(int(key[0])+int(key[1])) \* counts

```
Measurements in XX, YY, ZZ basis

In [15]: measure_circuit('X0X1', 2).draw()

Out[15]: q_0: H H M C: 2/ 0 1

In [16]: measure_circuit('Y0Y1', 2).draw()

Out[16]: q_0: SDG H M M M
```

# **Determining the Ansatz**

## **Designing Ansatze**

c: 2/=

In [17]: | measure\_circuit('Z0Z1', 2).draw()

Ansatze are simply a parameterized quantum circuits (PQC), which play an essential role in the performance of many variational hybrid quantum-classical (HQC) algorithms. Major challenge while designing an asatz is to choose an effective template circuit that well represents the solution space while maintaining a low circuit depth and number of parameters. Here, we make a choice of two ansatze, one randomly and another inspired from the given hint.

### Ansatz 1 (Random Choice)

```
In [21]:
          def ansatz1(params, num_qubits):
              Generate an templated ansatz with given parameters
              params (array[float]): Parameters to initialize the parameterized unitary.
              num_qubits (int): Number of qubits in the circuit.
              ansatz (QuantumCircuit): Generated ansatz circuit
              ansatz = QuantumCircuit(num_qubits, num_qubits)
              params = params.reshape(2,2)
              for idx in range(num_qubits):
                ansatz.rx(params[0][idx], idx)
              ansatz.cnot(0, 1)
              for idx in range(num_qubits):
                ansatz.rz(params[1][idx], idx)
              return ansatz
          ansatz1 (np.random.uniform(-np.pi, np.pi, (2,2)), 2).draw()\\
Out[21]: q_0: -
                 RX(3.0659)
                                      RZ(-0.23082)
```

### Ansatz 2 (From Hint)

q\_1: -

c: 2/=

RX(-2.9547)

```
Generate an templated ansatz with given parameters

Args:
    params (array[float]): Parameters to initialize the parameterized unitary.
    num_qubits (int): Number of qubits in the circuit.

Returns:
    ansatz (QuantumCircuit): Generated ansatz circuit

"""

ansatz = QuantumCircuit(num_qubits, num_qubits)
    ansatz.h(0)
    ansatz.cx(0, 1)
    ansatz.rx(params[0], 0)

return ansatz

ansatz2([np.pi], 2).draw()
```

Out[22]: q\_0: H RX(pi) q\_1: X

## Checking Expressibility of Ansatze

We quantify expressibility of ansatze using the Hilbert-Schmidt norm of  $\boldsymbol{A}$  defined as:

$$A = \int_{Haar} |\psi\rangle\langle\psi| d\psi - \int_{\theta} |\psi_{\theta}\rangle\langle\psi_{\theta}| d\theta$$

This quantity needs to be taken with a pinch of salt as it is an oversimplification of the A which actually has to be calculated with the definition of an  $\epsilon$ -approximate state t-design [1].

Here, the first term, i.e. a Haar integral, is the integral over a group of unitaries distributed randomly according to the Haar measure. Whereas, the second term, is taken over all states over the measure induced by uniformly sampling the parameters  $\theta$  of the PQC.

```
def random_unitary(N):
        Return a Haar distributed random unitary from U(N)
    Z = np.random.randn(N, N) + 1.0j * np.random.randn(N, N)
    [Q, R] = sp.linalg.qr(Z)
    D = np.diag(np.diagonal(R) / np.abs(np.diagonal(R)))
    return np.dot(Q, D)
def haar_integral(num_qubits, samples):
        Return calculation of Haar Integral for a specified number of samples.
    N = 2**num_qubits
    randunit_density = np.zeros((N, N), dtype=complex)
    zero_state = np.zeros(N, dtype=complex)
    zero_state[0] = 1
    for _ in range(samples):
    A = np.matmul(zero_state, random_unitary(N)).reshape(-1,1)
      randunit\_density += np.kron(A, A.conj().T)
    randunit density/=samples
    return randunit_density
def pqc_integral(num_qubits, ansatze, size, samples):
        Return calculation of Integral for a PQC over the uniformly sampled
        the parameters \boldsymbol{\theta} for the specified number of samples.
    N = num_qubits
    randunit_density = np.zeros((2**N, 2**N), dtype=complex)
    for _ in range(samples):
      params = np.random.uniform(-np.pi, np.pi, size)
      ansatz = ansatze(params, N)
                        backend=Aer.get_backend('statevector_simulator')).result()
      U = result.get_statevector(ansatz, decimals=5).reshape(-1,1)
      randunit_density += np.kron(U, U.conj().T)
    return randunit_density/samples
```

#### Sanity Check (Comparing Two Haar Integrals)

Out[24]: 0.029869558791692247

#### Ansatz 1 (Random Choice)

```
In [25]: np.linalg.norm(haar_integral(2, 2048) - pqc_integral(2, ansatz1, (2,2), 2048))
Out[25]: 0.025339519330263287
```

### Ansatz 2 (From Hint)

```
In [26]: np.linalg.norm(haar_integral(2, 2048) - pqc_integral(2, ansatz2, 1, 2048))
```

Out[26]: 0.5009773145594799

## Ansatz 3 (Empty Circuit)

```
def ansatz3(params, num_qubits):
    Generate an templated ansatz with no parameters
    params (array[float]): Parameters to initialize the parameterized unitary.
    num_qubits (int): Number of qubits in the circuit.
    ansatz (QuantumCircuit): Generated ansatz circuit
    ansatz = QuantumCircuit(num_qubits, num_qubits)
np.linalg.norm(haar_integral(2, 2048) - pqc_integral(2, ansatz3, 0, 2048))
```

Out[27]: 0.8647745124283421

Clearly, expressibility are in the order: Ansatz 3 < Ansatz 2 < Ansatz 1, i.e. the power to probe Hilbert space is much more for our randomly chosen ansatz, which is guessable.

## Checking Entangling Capability of Ansatze

We quantify entangling capability [1] of ansatze by calculating the average Meyer-Wallach entanglement, Q, of the states generated by it:

$$Q = rac{2}{|S|} \sum_{ heta_i \in S} \left( 1 - rac{1}{n} \sum_{k=1}^n Tr(
ho_k^2( heta_i)) 
ight)$$

Here,  $\rho_k$  is the density operator for the  $k^{th}$  qubit after tracing out the rest, and S is the set of sampled parameters. The quantity within the first summation can also be called as the average subsystem linear entropy for the system, and to calculate it we make use of qiskit's partial\_trace.

```
In [28]:
          def meyer_wallach(circuit, num_qubits, size, sample=1024):
                  Returns the meyer-wallach entanglement measure for the given circuit.
              res = np.zeros(sample, dtype=complex)
              N = num qubits
              for i in range(sample):
                  params = np.random.uniform(-np.pi, np.pi, size)
                  ansatz = circuit(params, N)
                  result = execute(ansatz,
                                 backend=Aer.get_backend('statevector_simulator')).result()
                  U = result.get_statevector(ansatz, decimals=5)
                  entropy = 0
                  qb = list(range(N))
                  for j in range(N):
                      dens = quantum_info.partial_trace(U, qb[:j]+qb[j+1:]).data
                      trace = np.trace(dens**2)
                      entropy += trace
                  entropy /= N
                  res[i] = 1 - entropy
              return 2*np.sum(res).real/sample
```

### Sanity Check (Empty circuit aka Ansatz 3)

```
In [29]: meyer_wallach(ansatz3, 2, 0)
```

Out[29]: 0.0

### Ansatz 1 (Random Choice)

```
In [30]: meyer_wallach(ansatz1, 2, (2,2))
Out[30]: 0.6274618290825067
```

Ansatz 2 (From Hint)

```
In [31]: meyer_wallach(ansatz2, 2, 1)

Out[31]: 1.0
```

Clearly, the entangling capability are in the order: Ansatz 3 < Ansatz 1 < Ansatz 2. Therefore, we can guess limited expressibility of Ansatz 2 is compensated by its higher entangling capability.

# **Running Simulations**

*Variational Quantum Eigensolver (VQE)* is based on *Rayleigh-Ritz variational principle*. To perform VQE, the first step is to enocde the problem into a Hermitian matrix M, whose expectation value w.r.t a trial wave function  $|\psi(\theta)\rangle$  which is yielded by an ansatz  $U(\theta)$ , i.e., nothing but a unitary paramterized by  $\{\vec{\theta}\}$ .

Here, we optimize these parameters  $\{\vec{\theta}\}$  using scipy optimize library based on a cost function  $C(\theta)$ , which is nothing but the calculated expectation value of Hamiltonian,  $H_1$  for  $|\psi(\theta)\rangle$ :

$$C(\theta) = \langle \psi(\theta) | H_1 | \psi(\theta) \rangle$$

```
def vqe(params, meas_basis, coeffs, circuit, num_qubits, shots=2048):
        Return the calculated energy scalar for a given asnatz and
        decomposed Hamiltoian.
        params (matrix(np.array)): Parameters for initializing the ansatz.
        {\tt meas\_basis} \ ({\tt list[str]}) \colon {\tt String} \ {\tt representation} \ {\tt of} \ {\tt measurement} \ {\tt basis}, \ {\tt i.e.}
        the decomposed pauli term with their corresponding qubits.
        coeffs (vector(np.array)): Coefficients for the decomposed Pauli Term
        circuit (QuantumCircuit): Template for Ansatz circuit
        num_qubits (int): Number of qubits in the given asatze
        shots (int): Number of shots to get the probability distribution.
        energy (float): Expectation value of the Hamiltonian whose
        decomposition was provided.
    N = num\_qubits
    circuit = circuit(params, num_qubits)
    energy = 0
    for basis, coeff in zip(meas_basis, coeffs):
      if basis.count('I') != N:
        energy += coeff*calculate_expecation_val(circuit, basis, shots)
    energy += 0.5
    return energy
```

## Ansatz 1 (Random Choice)

```
a, b, c = decompose_ham_to_pauli(H1)
           params = np.random.uniform(-np.pi, np.pi, (2,2))
           func = ft.partial(vqe, meas_basis=a, coeffs=b, circuit=ansatz1,
                              num_qubits=2, shots=2048)
In [34]:
          res = sp.optimize.minimize(func, params, method='Powell')
             direc: array([[1., 0., 0., 0.],
Out[34]:
                 [0., 1., 0., 0.],
[0., 0., 1., 0.],
               [0., 0., 0., 1.]])
fun: array(-1.)
           message: 'Optimization terminated successfully.'
              nfev: 203
               nit: 3
            status: 0
           success: True
                 x: array([-1.57402215, 3.11992864, -0.61255446, 0.98333759])
          assert(float(res.fun) == min(np.linalg.eig(H1)[0])) #Sanity Check
```

### Visualizing the Result

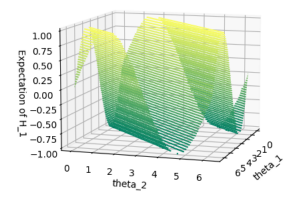
After looking at multiple results' res.x , we realize that first and second parameters always have the values  $\pm \pi/2$  and  $\pm \pi$  respectively. So, while keeping them fixed, we do a parameters search-scan in the range  $\theta_{3,4} \in [0,2\pi]$  and visualize how they affect our expectation value.

```
In [36]: def energy_expectation(x, y):
    """ Returns meshgrid values for plotting """
    energy = np.zeros(x.shape)
    for idx, thetas in enumerate(x):
        for ind, theta1 in enumerate(thetas):
            params = np.array([np.pi/2, np.pi, theta1, y[idx][ind]])
            energy[idx][ind] = func(params)
        return energy

    theta1 = np.linspace(0.0, 2*np.pi, 100)
    theta2 = np.linspace(0.0, 2*np.pi, 100)

    X, Y = np.meshgrid(theta1, theta2)
    Z = energy_expectation(X, Y)
```

```
fig = plt.figure()
ax = plt.axes(projection='3d')
ax.contour3D(X, Y, Z, 50, cmap='summer')
ax.set_xlabel('theta_1')
ax.set_ylabel('theta_2')
ax.set_zlabel('Expectation of H_1');
plt.show()
```



## Ansatz 2 (From Hint)

```
In [37]:
          a, b, c = decompose_ham_to_pauli(H1)
          params = np.random.uniform(-np.pi, np.pi, 1)
          func = ft.partial(vqe, meas_basis=a, coeffs=b, circuit=ansatz2,
                             num_qubits=2, shots=2048)
In [38]:
          res = sp.optimize.minimize(func, params, method='Powell')
          res
            direc: array([[0.00588462]])
Out[38]:
          fun: array(-1.)
message: 'Optimization terminated successfully.'
              nfev: 48
              nit: 2
           status: 0
           success: True
                x: array([3.14367297])
          assert(float(res.fun) == min(np.linalg.eig(H1)[0])) #Sanity Check
In [39]:
```

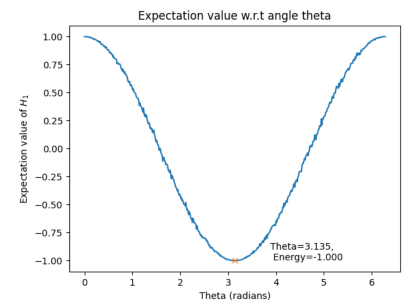
### Visualizing the Result

We scan over the paramter  $\theta \in [0, 2\pi]$ , to visualize the change in expectation value for the ansatz2, which is inspired from the hint given.

```
In [40]: thetas = np.linspace(0.0, 2*np.pi, 500)
energy = np.zeros(len(thetas))

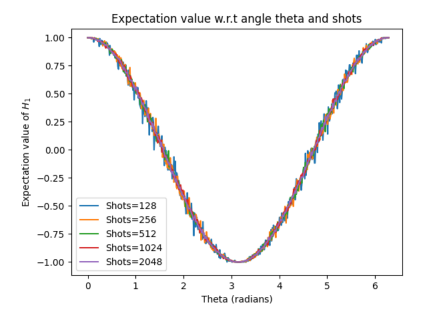
for idx, theta in enumerate(thetas):
    energy[idx] = func([theta])

fig = plt.figure()
plt.plot(thetas, energy)
plt.title('Expectation value w.r.t angle theta')
plt.ylabel('Expectation value of $H_1$')
plt.xlabel('Theta (radians)')
indices = [idx for idx, x in enumerate(energy) if x <= -1.0]
xmin = thetas[indices[len(indices)//2]]
ymin = energy.min()
text= "Theta={:.3f}, \n Energy={:.3f}".format(xmin, ymin)
plt.plot(xmin,ymin,'x')
plt.annotate(text, xy=(xmin+0.75, ymin))
fig.show()</pre>
```



Change of expectation value of  $H_1$  with the number of shots for ansatz 2.

```
In [63]:
          thetas = np.linspace(0.0, 2*np.pi, 500)
          shots = [128, 256, 512, 1024, 2048]
          shot_energy = np.zeros((len(shots), len(thetas)))
          for ind, shot in enumerate(shots):
              for idx, theta in enumerate(thetas):
                  func = ft.partial(vqe, meas_basis=a, coeffs=b, circuit=ansatz2,
                                 num_qubits=2, shots=shot)
                  shot\_energy[ind][idx] = func([theta])
In [64]:
          plt.figure()
          for ind, shot in enumerate(shots):
              plt.plot(thetas, shot_energy[ind], label='Shots=' + str(shot))
          plt.title('Expectation value w.r.t angle theta and shots')
          plt.ylabel('Expectation value of $H_1$')
          plt.xlabel('Theta (radians)')
          plt.legend()
          plt.show()
```



# **Noisy Simulation**

Computational power of NISQ processors suffers a lot from the limited capabilities of their physical qubits. This is essentially due to present of decoherence, limited connectivity, and absence of error-correction. Hence, it is essential to see how our ansatz would perform on a real device. Here, we replicate the noise model from a real device, and then test the performance of our VQE module.

```
In [42]: # Noise Model Preparation
# provider = IBMQ.load_account()
noisy_backend = provider.get_backend('ibmq_vigo')
coupling_map = noisy_backend.configuration().coupling_map
noise_model = providers.aer.noise.NoiseModel.from_backend(noisy_backend)
```

```
basis_gates = noise_model.basis_gates
def calculate_noisy_expecation_val(circuit, basis, mitigated=False, shots=2048, backend='qasm_simulator'):
    Calculate expectation value for the measurement of a circuit in a
   given basis in presence of noise.
   Aras:
    circuit (QuantumCircuit): Circuit using which expectation value
    will be calculated for a given basis.
    basis (str): String representation of tensor products of Pauli
    observables
    mitigated (bool): Whether mitigation has to be performed or not.
    shots (int): Number of times measurements needed to be done for calculating
    probability
    backend (str): Backend for running the circuit.
    Returns:
    exp (float): Expectation value for the measurement of a circuit
    in a given basis.
    exp_circuit = circuit + measure_circuit(basis, circuit.num_qubits)
    result = execute(exp_circuit, backend=Aer.get_backend(backend),
                      shots=shots, coupling_map=coupling_map,
                      basis_gates=basis_gates,
                      noise_model=noise_model).result()
    if mitigated:
        meas_calibs, state_labels = complete_meas_cal(qr=exp_circuit.qregs[0], cr=exp_circuit.cregs[0])
cal_results = qiskit.execute(meas_calibs, backend=Aer.get_backend(backend),
                                       coupling_map=coupling_map, basis_gates=basis_gates,
shots=shots, noise_model=noise_model).result()
        meas_fitter = CompleteMeasFitter(cal_results, state_labels)
        meas filter = meas fitter.filter
        result = meas_filter.apply(result)
    exp = 0.0
    for key, counts in result.get_counts().items():
        exp += (-1)**(int(key[0])+int(key[1])) * counts
    return exp/shots
def noisy_vqe(params, meas_basis, coeffs, circuit, num_qubits, mitigated=False, shots=2048):
        Return the calculated energy scalar for a given asnatz and
        decomposed Hamiltoian in presence of noise.
        params (matrix(np.array)): Parameters for initializing the ansatz.
        meas_basis (list[str]): String representation of measurement basis, i.e.
        the decomposed pauli term with their corresponding qubits
        coeffs (vector(np.array)): Coefficients for the decomposed Pauli Term
        circuit (QuantumCircuit): Template for Ansatz circuit
        num_qubits (int): Number of qubits in the given asatze.
        mitigated (bool): Whether mitigation has to be performed or not.
        shots (int): Number of shots to get the probability distribution.
        energy (float): Expectation value of the Hamiltonian whose
        decomposition was provided.
   N = num qubits
   circuit = circuit(params, num_qubits)
    energy = 0
   for basis, coeff in zip(meas_basis, coeffs):
   if basis.count('I') != N:
        energy += coeff*calculate_noisy_expecation_val(circuit, basis, mitigated, shots)
    energy += 0.5
    return energy
```

/home/whatsis/.local/lib/python3.8/site-packages/qiskit/providers/ibmq/ibmqfactory.py:192: UserWarning: Timestamps in IBMQ backend propertie s, jobs, and job results are all now in local time instead of UTC.
warnings.warn('Timestamps in IBMQ backend properties, jobs, and job results '

#### Ansatz 1 (Random Choice)

```
nfev: 266
              nit: 3
            status: 0
                 x: array([1.64212384, 3.14620703, 2.15383843, 0.53103981])
          a, b, c = decompose_ham_to_pauli(H1)
In [45]:
           params = np.random.uniform(-np.pi, np.pi, (2,2))
           func = ft.partial(noisy_vqe, meas_basis=a, coeffs=b, circuit=ansatz1,
                              num_qubits=2, mitigated=True, shots=8192)
In [46]:
          res = sp.optimize.minimize(func, params, method='Powell')
          res
             direc: array([[1., 0., 0., 0.],
Out[46]:
          [0., 1., 0., 0.],

[0., 0., 1., 0.],

[0., 0., 0., 1.]])

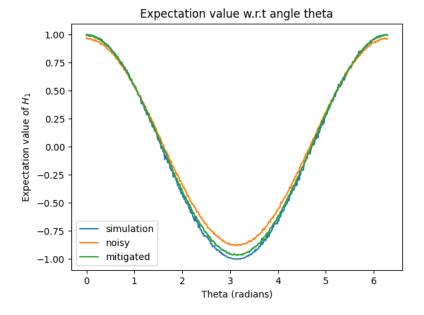
fun: -0.9619728377818637

message: 'Optimization terminated successfully.'
              nfev: 257
               nit: 3
            status: 0
           success: True
                 x: array([ 1.58409174, -3.11397559, -0.50929574, -1.96419365])
         Ansatz 2 (From Hint)
In [47]:
          a, b, c = decompose_ham_to_pauli(H1)
           params = np.random.uniform(-np.pi, np.pi, 1)
           func = ft.partial(noisy_vqe, meas_basis=a, coeffs=b, circuit=ansatz2,
                              num_qubits=2, mitigated=False, shots=8192)
In [48]: res = sp.optimize.minimize(func, params, method='Powell')
          res
            direc: array([[-0.1127984]])
  fun: array(-0.88122559)
Out[48]:
           message: 'Optimization terminated successfully.'
             nfev: 88
               nit: 5
            status: 0
           success: True
                x: array([3.17376448])
          a, b, c = decompose ham to pauli(H1)
          params = np.random.uniform(-np.pi, np.pi, 1)
func = ft.partial(noisy_vqe, meas_basis=a, coeffs=b, circuit=ansatz2,
num_qubits=2, mitigated=True, shots=8192)
In [50]: res = sp.optimize.minimize(func, params, method='Powell')
Out[50]:
             direc: array([[3.01441921e-09]])
           fun: -0.9702586386528804
message: 'Optimization terminated successfully.'
              nfev: 120
               nit: 5
            status: 0
           success: True
                 x: array([-3.12601157])
         Visualizing the Result (Noisy v/s Mitigated)
          thetas = np.linspace(0.0, 2*np.pi, 500)
           energy1 = np.zeros(len(thetas))
           energy2 = np.zeros(len(thetas))
           for idx, theta in enumerate(thetas):
              energy1[idx] = func1([theta])
energy2[idx] = func2([theta])
In [56]:
          #plt.figure()
           plt.plot(thetas, energy1, label='noisy')
           plt.plot(thetas, energy2, label='mitigated')
           plt.title('Expectation value w.r.t angle theta')
           plt.ylabel('Expectation value of $H_1$')
           plt.xlabel('Theta (radians)')
           plt.legend()
           plt.show()
```

## Expectation value w.r.t angle theta 1.00 0.75 0.50 Expectation value of H<sub>1</sub> 0.25 0.00 -0.25-0.50 -0.75noisy mitigated -1.002 0 5 6 Theta (radians)

## Visualizing the Result (Simulated v/s Noisy v/s Mitigated)

```
In [54]: plt.plot(thetas, energy, label='simulation')
  plt.plot(thetas, energy1, label='noisy')
  plt.plot(thetas, energy2, label='mitigated')
  plt.title('Expectation value w.r.t angle theta')
  plt.ylabel('Expectation value of $H_1$')
  plt.xlabel('Theta (radians)')
  plt.legend()
  plt.show()
```



Comparing the performance of VQE in presence of errors due to Readout, Thermal Relaxation, Amplitude-Phase damping and Depolarization

#### Readout

Describes classical readout errors

```
In [71]: # Measurement miss-assignement probabilities
p0given1 = 0.1
p1given0 = 0.05
noise_readout = ReadoutError([[1 - p1given0, p1given0], [p0given1, 1 - p0given1]])
noise_readout
```

#### Thermal Relaxation

Single qubit thermal relaxation error is characterized by relaxation time constants  $T_1$ ,  $T_2$ , and the gate time t.

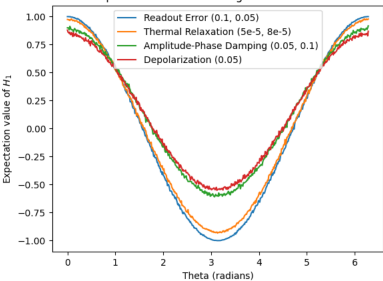
```
In [72]: num_qubits = 2
T1s = np.random.normal(50e3, 10e3, num_qubits) # Sampled from normal distribution
```

```
 T2s = np.array([min(T2s[j], 2*T1s[j]) \ \textit{for} \ j \ in \ range(num\_qubits)]) \ \textit{\# Ensuring T2s} < 2*T1s[j] 
                  # Instruction times (in nanoseconds)
time_u1 = 0  # virtual gate
                  time_u2 = 50 # (single X90 pulse)
                  time_u3 = 100 # (two X90 pulses)
                  time cx = 300
                  time_measure = 1000 # 1 microsecond
                  # Add depolarizing error to all single qubit u1, u2, u3, cx, measure gates
                  errors_thermal_u1 = [thermal_relaxation_error(t1, t2, time_u1)
                                          for t1, t2 in zip(T1s, T2s)]
                  errors_thermal_u2 = [thermal_relaxation_error(t1, t2, time_u2)
                                          for t1, t2 in zip(T1s, T2s)]
                  errors_thermal_u3 = [thermal_relaxation_error(t1, t2, time_u3)
                                          for t1, t2 in zip(T1s, T2s)]
                  errors_thermal_cx = [[thermal_relaxation_error(t1a, t2a, time_cx).expand(
                                         thermal_relaxation_error(t1b, t2b, time_cx))
                                          for t1a, t2a in zip(T1s, T2s)]
                                            for t1b, t2b in zip(T1s, T2s)]
                  errors_thermal_measure = [thermal_relaxation_error(t1, t2, time_measure)
                                                  for t1, t2 in zip(T1s, T2s)]
                  # Add errors to noise model
                  noise_thermal = NoiseModel()
                  for j in range(num_qubits):
                         noise_thermal.add_quantum_error(errors_thermal_u1[j], "u1", [j])
                         noise_thermal.add_quantum_error(errors_thermal_u2[j], "u2", [j]) noise_thermal.add_quantum_error(errors_thermal_u3[j], "u3", [j])
                         for k in range(num gubits):
                         noise_thermal.add_quantum_error(errors_thermal_cx[j][k], "cx", [j, k])
noise_thermal.add_quantum_error(errors_thermal_measure[j], "measure", [j])
                  noise thermal
Out[72]: NoiseModel:
                    Basis gates: ['cx', 'id', 'u2', 'u3']
Instructions with noise: ['measure', 'cx', 'u2', 'u3']
                Qubits with noise: [0, 1]
Specific qubit errors: [('u2', [0]), ('u2', [1]), ('u3', [0]), ('u3', [1]), ('cx', [0, 0]), ('cx', [0, 1]), ('cx', [1, 0]), ('cx', [1, 1]), ('measure', [0]), ('measure', [1])]
               Amplitude-Phase Damping
               Single-qubit generalized combined phase and amplitude damping error is given by an amplitude damping parameter \lambda, and a phase damping parameter \gamma.
In [70]:
                 lamb = 0.05
                  gamma = 0.1
                  errors_phase_amplitude_1 = phase_amplitude_damping_error(lamb, gamma)
                  errors\_phase\_amplitude\_2 = phase\_amplitude\_damping\_error(lamb, gamma). expand(phase\_amplitude\_damping\_error(lamb, gamma)) = phase\_amplitude\_damping\_error(lamb, gamma) = phase\_amplitude\_damping\_error(
                  # Add errors to noise model
                  noise_phase_amplitude = NoiseModel()
                  noise\_phase\_amplitude.add\_all\_qubit\_quantum\_error(errors\_phase\_amplitude\_1, \ ['u1', \ 'u2', \ 'u3', \ 'measure'])
                  noise_phase_amplitude.add_all_qubit_quantum_error(errors_phase_amplitude_2, ['cx'])
                  noise_phase_amplitude
Out[70]: NoiseModel:
                    Basis gates: ['cx', 'id', 'u1', 'u2', 'u3']
Instructions with noise: ['u1', 'u3', 'measure', 'cx', 'u2']
All-qubits errors: ['u1', 'u2', 'u3', 'measure', 'cx']
               Depolarization
               N-qubit depolarizing error is given by a depolarization probability p.
                 # Add depolarizing error to all single qubit u1, u2, u3, cx, measure gates
                  error_depol_1 = depolarizing_error(0.05, 1)
                  error_depol_2 = depolarizing_error(0.1, 2)
                  error_depol_3 = depolarizing_error(0.1, 1)
                  # Add errors to noise model
                  noise_depol = NoiseModel()
                  noise_depol.add_all_qubit_quantum_error(error_depol_1, ['u1', 'u2', 'u3'])
                  noise_depol.add_all_qubit_quantum_error(error_depol_2, ['cx'])
                  noise_depol.add_all_qubit_quantum_error(error_depol_3, ['measure'])
                  noise_depol
Out[73]: NoiseModel:
                   Basis gates: ['cx', 'id', 'u1', 'u2', 'u3']
Instructions with noise: ['u1', 'u3', 'measure', 'cx', 'u2']
All-qubits errors: ['u1', 'u2', 'u3', 'cx', 'measure']
                 def calculate_restricted_noisy_expecation_val(circuit, basis, noise_model, shots=2048, backend='qasm_simulator'):
In [74]:
                         Calculate expectation value for the measurement of a circuit in a
                         given basis in presence of a particular kind of noise.
                         circuit (QuantumCircuit): Circuit using which expectation value
                         will be calculated for a given basis.
                         basis (str): String representation of tensor products of Pauli
                         observables.
                         noise_model (NoiseModel): Noise Model for execution
                         shots (int): Number of times measurements needed to be done for calculating
                         probability.
                         backend (str): Backend for running the circuit.
```

T2s = np.random.normal(80e3, 20e3, num\_qubits) # Sampled from normal distribution

```
Returns:
                          exp (float): Expectation value for the measurement of a circuit
                          in a given basis.
                          exp_circuit = circuit + measure_circuit(basis, circuit.num_qubits)
                         result = execute(exp_circuit, backend=Aer.get_backend(backend),
                                                         shots=shots, noise_model=noise_model).result()
                          exp = 0.0
                          for key, counts in result.get_counts().items():
                                 exp += (-1)**(int(key[0])+int(key[1])) * counts
                          return exp/shots
                   def restricted_noisy_vqe(params, meas_basis, coeffs, circuit, num_qubits, noise_model, shots=2048):
                                 Return the calculated energy scalar for a given asnatz and
                                 decomposed Hamiltoian in presence of a particular kind of noise.
                                 params (matrix(np.array)): Parameters for initializing the ansatz.
                                 meas_basis (list[str]): String representation of measurement basis, i.e.
                                 the decomposed pauli term with their corresponding qubits.
                                 coeffs (vector(np.array)): Coefficients for the decomposed Pauli Term
                                 circuit (QuantumCircuit): Template for Ansatz circuit
                                 num_qubits (int): Number of qubits in the given asatze.
                                 noise_model (NoiseModel): Noise Model for execution
                                 shots (int): Number of shots to get the probability distribution.
                                 Return:
                                 energy (float): Expectation value of the Hamiltonian whose
                                 decomposition was provided.
                         N = num_qubits
                         circuit = circuit(params, num_qubits)
                          energy = 0
                          for basis, coeff in zip(meas_basis, coeffs):
                             if basis.count('I') != N:
                                 energy += coeff*calculate_restricted_noisy_expecation_val(circuit, basis, noise_model, shots)
                          energy += 0.5
                          return energy
                  noise_models = [noise_readout, noise_thermal, noise_phase_amplitude, noise_depol]
In [76]:
                   thetas = np.linspace(0.0, 2*np.pi, 500)
                   noisy_energy = np.zeros((len(noise_models), len(thetas)))
                   for ind, model in enumerate(noise_models):
                          func3 = ft.partial(restricted_noisy_vqe, meas_basis=a, coeffs=b, circuit=ansatz2,
                                                  num_qubits=2, noise_model=model, shots=4096)
                          for idx, theta in enumerate(thetas):
                                 noisy_energy[ind][idx] = func3([theta])
In [81]: noise_models_name = ['Readout Error (0.1, 0.05)', 'Thermal Relaxation (5e-5, 8e-5)', 'Amplitude-Phase Damping (0.05, 0.1)', 'Depolarization (0.1, 0.05)', 'Thermal Relaxation (5e-5, 8e-5)', 'Amplitude-Phase Damping (0.05, 0.1)', 'Depolarization (0.1, 0.05)', 'Thermal Relaxation (5e-5, 8e-5)', 'Amplitude-Phase Damping (0.05, 0.1)', 'Depolarization (0.1, 0.05)', 'Thermal Relaxation (5e-5, 8e-5)', 'Amplitude-Phase Damping (0.05, 0.1)', 'Depolarization (0.1, 0.05)', 'Thermal Relaxation (5e-5, 8e-5)', 'Amplitude-Phase Damping (0.05, 0.1)', 'Depolarization (0.1, 0.05)', 'Thermal Relaxation (5e-5, 8e-5)', 'Amplitude-Phase Damping (0.05, 0.1)', 'Depolarization (0.1, 0.05)', 'Thermal Relaxation (5e-5, 8e-5)', 'Amplitude-Phase Damping (0.05, 0.1)', 'Depolarization (0.1, 0.1)', 'Depolarizati
                   plt.figure()
                   for ind, model in enumerate(noise_models):
                          plt.plot(thetas, noisy_energy[ind], label=noise_models_name[ind])
                   plt.title('Expectation value w.r.t angle theta and shots')
                   plt.ylabel('Expectation value of $H_1$')
                   plt.xlabel('Theta (radians)')
                   plt.legend()
                   plt.show()
```

## Expectation value w.r.t angle theta and shots



## **Excited States of the Hamiltonian**

One of the way to find the  $k^{th}$  exicted states of a Hamiltonian would be to update the Hamiltonian H to  $H_k$  as follows:

$$H_k = H + \sum_{i=0}^{k-1} eta_i |i
angle \langle i|$$

For example: To find the first excited state, we find  $H_1=H+eta_0|g\rangle\langle g|$ , where g is the ground state of the Hamiltonian. This works because of spectral decomposition.

This can be used to update our cost function,  $C_k$ , as follows:

$$C( heta_k) = \langle \psi( heta_k) | H | \psi( heta_k) 
angle + \sum_{i=0}^{k-1} eta_i | \langle \psi( heta_k) | \psi( heta_i) 
angle |^2$$

We'd already calculated the first term in VQE, but to calculate the next summation term, we refer to [2]. It suggests us to rewrite the overlap term  $|\langle \psi(\theta_k)|\psi(\theta_i)\rangle|^2$  as  $|\langle 00|U(\theta_k)^\dagger U(\theta_i)|00\rangle|^2$ . Therefore, we can prepare the state  $U(\theta_k)^\dagger U(\theta_i)|00\rangle$  using the trial state preparation circuit for current state and  $i^{th}$  previously-computed state.

This technique is known as Variational Quantum Deflation and requires the same number of qubits as Variational Quantum Eigensolver and around twice the circuit depth.

```
In [185.
          def vqd_cost(thetak, thetai, func, circuit, num_qubits, beta, shots=2048):
                   Returns cost for VQD
              energy = func(thetak)
              for idx, theta in enumerate(thetai):
                  ansatz = circuit(theta, num_qubits) + circuit(thetak, num_qubits).inverse()
                  ansatz.measure_all()
                  backend = Aer.get_backend('qasm_simulator')
                  result = execute(ansatz, backend, shots=shots).result()
                  energy += beta[idx]*result.get_counts().get('00 00', 0)/shots
              return energy
          def vqd(hamiltonian, circuit, num_qubits, params_shape, beta):
                  Runs Variational Quantum Deflation algorithm on a given Hamiltonian
                  to give its excited eigenvalues.
                  Aras:
                  hamiltonian (matrix(ndarray)): Hamiltonian for
                  circuit (Quantum Circuit): ansatz template
                  num_qubits (int): number of qubits to make an ansatz
                  params_shape (tuple): shape tuple for parameters in ansatz
                  beta (int): beta coefficent for VOD
                  energy (vector(ndarray)): Calculated eigen energies for the Hamiltonian.
              a, b, c = decompose_ham_to_pauli(hamiltonian)
              initial_params = np.random.uniform(-np.pi, np.pi, params_shape)
              func = ft.partial(vqe, meas_basis=a, coeffs=b, circuit=circuit,
                            num_qubits=num_qubits, shots=2048)
              res1 = sp.optimize.minimize(func, initial_params, method='Powell')
              energies = [float(res1.fun)]
              thetas = [np.reshape(res1.x, params_shape)]
              if len(beta) != np.shape(hamiltonian)[0]-1:
                  raise ValueError('Length mismatch! Provide sufficient amount of beta_{i}')
              for eigst in range(np.shape(hamiltonian)[0]-1):
                  cost = ft.partial(vqd_cost, thetai=thetas, func=func, circuit=circuit,
                            num_qubits=num_qubits, beta=beta, shots=2048)
```

```
res2 = sp.optimize.minimize(cost, initial_params, method='Powell')
if res2.success:
    thetas.append(np.reshape(res2.x, params_shape))
    energies.append(float(res2.fun))
else:
    raise ValueError('Minimization was not successful.')

return energies
```

## Ansatz 1

```
In [194... excited_energies = vqd(H1, ansatz1, 2, (2,2), [3.5, 1e-1, 1e-3])
excited_energies

Out[194... [-1.0, 0.9921875, 0.994189453125, 1.0003808593749999]

In [195... assert(np.allclose(np.asarray(excited_energies), np.sort(np.linalg.eig(H1)[0]), atol=1e-2))

Ansatz 2

In [196... excited_energies = vqd(H1, ansatz2, 2, 1, [2.7, 1e-2, 1e-3])
excited_energies

Out[196... [-1.0, 0.99365234375, 1.0012158203125, 1.0074306640625]

In [197... assert(np.allclose(np.asarray(excited_energies), np.sort(np.linalg.eig(H1)[0]), atol=1e-2))
```

# **Interpreting the Results**

- 1. Both of our ansatz produced accurate results despite of ansatz 2 being more expressible than ansatz 1. One reason for this can be that the the ansatz 2 expressibility was covered the interested solution subspace. Another reason could be due to its more entangling capability.
- 2. Convergence of VQE depends upon both the type of optimizer (gradient-based or point-search) and the choice of ansatz. In general more the number of parameters, and the depth of ansatz, more difficult will be the convergence.
- 3. Number of shots did matter for non-optimal values of theta in ansatz 2. However, at optimal values of theta, less shots also gave good results.
- 4. Ansatz 2 performed better than ansatz 1 under Noisy simulation. However, mitigation improved the result for both of them.
- 5. Out of all the all the noise models tested, we saw VQE still gave the (deviated) minima at the optimal value of theta, and performed worst for errors due to damping and depolarization.
- 6. VQD was able to perform better as we adjusted the values of  $\beta_i$  based on the difference between the eigen-energies. In general, values of these  $\beta_i$  should be larger than the largest difference between the consecutive eigen-energies.

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