

SIGN OUT

Quantum Coding Challenges











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5. Hi, Hydrogen!

0 points

Welcome to the QHack 2023 daily challenges! Every day for the next four days, you will receive two new challenges to complete. These challenges are worth no points — they are specifically designed to get your brain active and into the right mindset for the competition. You will also learn about various aspects of PennyLane that are essential to quantum computing, quantum machine learning, and quantum chemistry. Have fun!

Tutorial #5 — Hi, Hydrogen!

The Coding Challenge Competition is now closed, but you are welcome to continue working on the challenges until QHack ends on Feb 28 @5pm ET.

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you will calculate the energy of the hydrogen molecule for various molecular charges and bond length combinations.

Challenge code

In the code below, you are given a few functions:

- hydrogen_hamiltonian: This function will return the qubit Hamiltonian of the hydrogen molecule, H2, given the coordinates of both hydrogen atoms and the net molecular charge. You'll usually find H2 with a charge of 0, but here we'll spice it up with a non-zero charge!
- num_electrons: In subsequent functions, we'll need the total number of electrons in the hydrogen molecule we're looking at. With a charge of 0, H₂ usually has just 2 electrons, one per hydrogen atom. Given the charge, how many electrons should H₂ have? **You must complete this function.**
- Inf: The "HF" stands for Hartree–Fock. This function's purpose is calculate the HF approximation treat every electron as independent, electrons move under a Coulomb potential from the positively charged nuclei, and there's a mean field from the other electrons for the ground state of the hydrogen molecule we're interested in. We'll use this later, so *you must complete this function*.
- run_VQE: This function takes the coordinates, charge, generates the HF state, defines a cost function and minimizes it. *You must complete this function* by:
 - defining the gates within the cost function, using the qml.AllSinglesDoubles template with singles and doubles arguments defined below; and
 - returning what we want to minimize, namely the expectation value of the hydrogen Hamiltonian!

Here are some helpful resources:

• Building molecular Hamiltonians

• Quantum Chemistry documentation

Input

As input to this problem, you are given:

- coordinates (list(float)): the x, y, and z coordinates of each hydrogen atom
- charge (int): the charge of the hydrogen molecule. It could be positive, negative, or zero!

Output

This code must output the ground state [energy] (float) of the hydrogen molecule in question.

If your solution matches the correct one within the given tolerance specified in Check (in this case it's a 1e-3 relative error tolerance), the output will be "Correct!"

Otherwise, you will receive a "Wrong answer" prompt.

Good luck!

```
Code

1 import json
2 import pennylane as qml
3 import pennylane.numpy as np
```

```
4 , def hydrogen hamiltonian(coordinates, charge):
        """Calculates the qubit Hamiltonian of the hydrogen molecule.
 5
 6
 7
        Args:
 8
             coordinates (list(float)): Cartesian coordinates of each hydrogen
 9
             charge (int): The electric charge given to the hydrogen molecule
10
11
        Returns:
12
             (qml.Hamiltonian): A PennyLane Hamiltonian.
13
        return qml.qchem.molecular_hamiltonian(
14
15
             ["H", "H"], coordinates, charge, basis="STO-3G"
16
        [0]
17
18 , def num electrons(charge):
        """The total number of electrons in the hydrogen molecule.
19
20
21
        Args:
22
             charge (int): The electric charge given to the hydrogen molecule
23
24
        Returns:
25
             (int): The number of electrons.
26
27
28
                                                                                 ٠
        # Put your solution here #
29
         return
30
                                                                                 ا
31 , def hf(electrons, num qubits):
32
        """Calculates the Hartree-Fock state of the hydrogen molecule.
33
34
        Args:
35
             electrons (int): The number of electrons in the hydrogen molecule
             num qubits (int): The number of qubits needed to represent the hy
36
37
38
        Returns:
39
             (numpy.tensor): The HF state.
        .....
40
41
42
        # Put your solution here #
```

```
45 def run VQE(coordinates, charge):
        """Performs a VQE routine for the given hydrogen molecule.
46
47
48
        Args:
49
            coordinates (list(float)): Cartesian coordinates of each hydrogen
50
            charge (int): The electric charge given to the hydrogen molecule
51
52
        Returns:
53
             (float): The expectation value of the hydrogen Hamiltonian.
54
55
        hamiltonian = hydrogen_hamiltonian(np.array(coordinates), charge)
56
57
        electrons = num_electrons(charge)
58
59
        num qubits = len(hamiltonian.wires)
60
61
        hf_state = hf(electrons, num_qubits)
62
        # singles and doubles are used to make the AllSinglesDoubles template
63
        singles, doubles = qml.qchem.excitations(electrons, num qubits)
64
65
        dev = qml.device("default.qubit", wires=num_qubits)
66
67
        @qml.qnode(dev)
68 ,
        def cost(weights):
             """A circuit with tunable parameters/weights that measures the ex
69
70
71
            Args:
72
                weights (numpy.array): An array of tunable parameters.
73
74
            Returns:
75
                 (float): The expectation value of the hydrogen Hamiltonian.
76
77
78
             # Put your solution here #
79
             return
80
```

```
81
         np.random.seed = 1234
 82
         weights = np.random.normal(
 83
             0, np.pi, len(singles) + len(doubles), requires grad=True
 84
 85
         opt = qml.AdamOptimizer(0.5)
 86
 87 ,
         for in range(200):
 88
             weights = opt.step(cost, weights)
 89
 90
         return cost(weights)
 91
 92
     # These functions are responsible for testing the solution.
 93 def run(test case input: str) -> str:
 94
         coordinates, charge = json.loads(test case input)
 95
         energy = run_VQE(coordinates, charge)
 96
 97
         return str(energy)
 98
 99 def check(solution output: str, expected output: str) -> None:
         solution output = json.loads(solution output)
100
101
         expected output = json.loads(expected output)
102
         assert np.allclose(solution output, expected output, rtol=1e-3)
103
     test cases = [['[0.0, 0.0, -0.8, 0.0, 0.0, 0.8], -1]', '-0.53168359']
104
                                                                                 105 for i, (input, expected output) in enumerate(test cases):
         print(f"Running test case {i} with input '{input }'...")
106
107
108 ,
         try:
109
             output = run(input )
110
         except Exception as exc:
111_{\vee}
              print(f"Runtime Error. {exc}")
112
113
         else:
114 ,
             if message := check(output, expected output):
115 ,
116
                  print(f"Wrong Answer. Have: '{output}'. Want: '{expected_output}'.
117
118 ,
             else:
                  print("Correct!")
119
```