Quantum computing TP Phase estimation for chemistry

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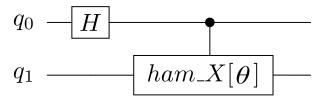
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Abstract

You need to fill up some notebook cells with your implementation according to the questions. Send the notebooks and helper files in a zip by next Thursday, January 14 2021. They will be corrected and graded.

1 Quantum Programming Basics

Question 1 Implement the following circuit, simulate it for $\theta = 0.3$, and print all states along with their amplitudes and probabilities.



Where
$$ham_{-}X(\theta) = e^{-i\theta X}$$
 with $X = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}$

Hint Define ham_X as an "AbstractGate", then apply it using ".ctrl()" to get a controlled version, as suggested in the "minimal notebook". Don't hesitate to also refer to the slides that were sent to you, for instance for the definition of what a controlled gate is.

2 Iterative quantum phase estimation, reproducing results from [1]

2.1 Hamiltonian simulation

The Hamiltonians whose ground state energy we are trying to compute are of the form:

$$H(R) = g_0 I + g_1 Z I + g_2 I Z + g_3 Z Z + g_4 Y Y + g_5 X X$$

Because we use **Trotterization**, we will only have to implement the unitary evolutions generated by each of the individual terms: e^{-iXXdt} , e^{-iZIdt} , e^{-iIZdt} , e^{-iZXdt} , e^{-iXXdt} and e^{-iXXdt} .

We could implement them using Abstract gates and a numerical computation of the matrix exponential. But the computation of the matrix exponential does not scale well with the number of qubits, and thus this method cannot be used in the "general case", when working with a large molecule.

What we will do therefore instead is find small parametrized circuits implementing each of the unitary evolutions. The small circuits may only use "usual gates", such as CNOTs and single-qubit Pauli rotations, that are already preimplemented in PyAQASM/myqlm. We will then use **QRoutines** to paste these circuits into the final algorithm. Such a method does yield circuits of polynomial size for arbitrary Pauli products over n qubits.

Example: You can check that: $Z \otimes Z = CNOT_{0 \to 1} \cdot I \otimes Z \cdot CNOT_{0 \to 1}$

with
$$CNOT_{0\to 1} = \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \end{pmatrix}$$
, \otimes the Kronecker product and \cdot the standard

matrix product.

It allows us to write:

$$\begin{split} e^{-iZZdt} &= e^{-i(CNOT_{0\to 1})\cdot (IZ)\cdot (CNOT_{0\to 1})dt} \\ &= CNOT_{0\to 1}\cdot e^{-iIZdt}CNOT_{0\to 1} \\ &= CNOT_{0\to 1}\cdot \left(I\otimes e^{-iZdt}\right)\cdot CNOT_{0\to 1} \end{split}$$

i.e:

$$q_0 = e^{-iZZdt} = q_0 = q_1 = e^{-iZdt}$$
 where $e^{-iZdt} = \begin{pmatrix} e^{-idt} & 0 \\ 0 & e^{idt} \end{pmatrix} = RZ(-dt)$.

RZ is a "standard gate", pre-implemented in PyAQASM. Likewise, $RX(\theta) = e^{i\theta X}$ and $RY(\theta) = e^{i\theta Y}$ are pre-implemented. You can look at their precise definitions here.

Question 2 Following the example above, which is already implemented in the notebook, and knowing in addition that:

$$X\otimes X=CNOT_{0\to 1}\cdot X\otimes I\cdot CNOT_{0\to 1}$$

$$SXS^\dagger=Y$$
 with $S=\begin{pmatrix}1&0\\0&i\end{pmatrix}$ and
$$Y\otimes X=CNOT_{0\to 1}\cdot Y\otimes I\cdot CNOT_{0\to 1}$$

write down the QRoutines carrying out each of the required Hamiltonian simulations, by filling the templates already written in the notebook.

2.2 Iterative Phase estimation

2.2.1 Trotterization

Question 3 Write a function taking as input hamiltonian coefficients, an interval of time dt, and a Trotter number p, and returning a Qroutine which implements the corresponding Trotterized implementation of the Hamiltonian.

We recall that Trotterization consists in approximating $e^{-idt\left(\sum_{l}c_{l}H_{l}\right)}$ as $\left(\prod_{l}e^{-i\frac{dtc_{l}H_{l}}{p}}\right)^{p}$. Therefore, your Qroutine should make use of the other Qroutines you implemented in Question 2. As for the coefficients weighting the influence of each individual terms, they are available in the list of dictionaries loaded at the beginning of the notebook, from hamiltonian_data.json. As specified in the notebook, your function should also take an "energy shift" as input parameter. It consists in adding a $+shift \cdot I$ term to your Hamiltonian. We will use it to ensure the energies we compute are > 0. Take a look at the slides for more explanations.

2.2.2 Iterative Phase estimation

2.3 final plots

References

[1] Peter JJ O'Malley, Ryan Babbush, Ian D Kivlichan, Jonathan Romero, Jarrod R McClean, Rami Barends, Julian Kelly, Pedram Roushan, Andrew Tranter, Nan Ding, et al. Scalable quantum simulation of molecular energies. *Physical Review X*, 6(3):031007, 2016.