

Calculating Ground-State-Energy of LiCoO_2 by using Fragment molecular orbital-based Variational Quantum Eigensolver

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Abstract. The Variational Quantum Eigensolver (VQE) is a quantum computing algorithm designed to calculate ground-state energies and has been highly anticipated for its potential effectiveness in new material development fields, such as drug discovery and battery technology. However, the current quantum computers are limited by the number of available qubits, posing challenges for applying VQE to large and complex molecules used in industrial applications. To address this issue, this study proposes the use of the FMO/VQE approach. The Fragment Molecular Orbital (FMO) method divides a system into smaller fragments for processing, and FMO/VQE integrates this FMO method into the VQE framework [1]. In this study, the ground-state energy of LiCoO_2 , a cathode material for lithium-ion batteries, was calculated using the FMO/VQE method to demonstrate its utility. By employing this approach, the number of required qubits was reduced from 24 to a maximum of 14, while achieving accuracy comparable to results obtained through classical methods. This study demonstrates that the proposed FMO/VQE algorithm can mitigate size-related constraints, enabling the application of the VQE algorithm to larger molecular systems. This advancement paves the way for successful implementation of the VQE algorithm in fields such as battery technology and drug discovery.

Keywords: VQE, Quantum Computing, Lithium-ion batteries

1 Introduction

The ground state of a molecule and its energy are the most stable state and lowest energy that a molecule can have. Knowing this information can be used to understand the bonds between molecules and the structure of molecules, which can be used in various fields of chemical research, such as the development of new drugs or new anode materials for secondary batteries. Currently, the ground state energy of a molecule is mostly calculated on classical computers. However, when the number of atoms in a molecule increases or the atomic number of the atoms in a molecule is large, the number of particles to be considered increases, and the interactions of each particle to be considered increase exponentially. The interactions of each particle are considered in the Hamilto-

nian of the molecule, and representing this molecular Hamiltonian on a computer requires a large amount of memory and computation. Therefore, calculating the ground state energy of a molecule using a classical computer is difficult to apply to large molecules.

Quantum computers are a recently emerging field, and there is a lot of research going on. Quantum computers use different qubits than classical computers, i.e., quantum bits, which are qubits with superposition of 0 and 1 states, unlike bits that represent only 0 and 1 in classical computers. This is why, in calculating the ground state energy of a molecule, when the number of particles increases and the memory required doubles, a quantum computer can do the calculation with one more qubit. Where a classical computer would require an exponential increase in memory, a quantum computer can solve with a polynomial increase in qubits. Therefore, even large molecules that are currently incomputable can be computed by quantum computers. The quantum computer algorithm Quantum Phase Estimation (QPE) is an algorithm that calculates the eigenvalues and eigenstates of a unitary matrix on a quantum computer, which can be used to find the ground state energy of a molecule. However, QPE is known to be very accurate and requires many qubits to implement. Current quantum computers are said to be at the level of noisy intermediate-scale quantum devices (NISQs). NISQ means that the quantum computer is noisy and has intermediate-scale qubits. Current quantum computers are still noisy and lack accuracy, and the number of qubits is still low, in the tens or hundreds, making it difficult to implement QPEs.

Variational Quantum Eigensolver (VQE) is an algorithm for solving eigenvalue problems using these NISQ-level quantum computers, which uses both quantum and classical computers. VQE uses variational principles to construct a parameterized trial wave function to obtain the basis of the molecular Hamiltonian. The algorithm iterates by adjusting the parameters until this basis value converges to a constant value to obtain an upper bound on the ground state energy of the molecule. In this process, the molecular Hamiltonian and the test wavefunction are replaced by quantum circuits, and the process of obtaining the expectation value is performed by a quantum computer, and the optimization process of adjusting the parameters of the test wavefunction is performed by a classical computer. Unlike QPE, VQE does not require many qubits, and the optimization that is difficult for current quantum computers is calculated by a classical computer, so there are many expectations that quantum computers can be commercialized in the NISQ era. However, current quantum computers have too many errors, their accuracy decreases significantly as the circuit depth increases, and the number of qubits available is limited, which is insufficient to calculate very large molecules. Therefore, one of the various methods to solve this problem is the Fragment Molecular Orbital method based Variational Quantum Eigensolver (FMO/VQE). FMO/VQE was first devised by Hocheol Lim, et al. as an algorithm that applies the FMO method, a quantum chemistry method, to VQE to improve the limitations of VQE due to the limitations of current quantum computers. The FMO method approximates the ground-state energy of the whole molecule by dividing the whole system into small pieces, and then finds the ground-state energy of the small pieces (monomer) and the pair of small pieces (dimer) and uses them to approximate the ground-state energy of the whole molecule. By breaking the molecule into smaller pieces, the maximum size of the system

that needs to be considered in the overall calculation can be reduced, and this can be applied to VQE to reduce the number of cubits required and the depth of the quantum circuit. Therefore, FMO/VQE is expected to be a way to utilize quantum computers in the near future.

In this paper, we use FMO/VQE to calculate the ground state energy of LiCoO_2 , a representative material used as an anode material for secondary batteries, and compare it with classical computer calculations. We also compare the accuracy of FMO/VQE with classical VQE to confirm the feasibility of FMO/VQE in the NISQ era.

2 Method

2.1 Lithium-ion battery

Lithium-ion batteries are one of the secondary batteries used in various industries. A lithium-ion battery consists of an anode material, a cathode material, and an electrolyte, where lithium oxide is used as the anode material. This compound has a layered structure of CoO_2 oxide, with Li^+ ions bonded between each layer. The energy charge depends on the degree to which the lithium ions are oxidized in the oxide. Lithium is known to have a limited number of reserves in the world. Many methods have been proposed as solutions to this problem, one of which is an approach that seeks to maximize the energy charged per lithium ion. This is a critical situation, both in terms of lithium reserves and in terms of battery miniaturization. To calculate the amount of energy stored per lithium, it is essential to calculate the ground state of the molecule. However, this calculation is classically performed, and the ability to calculate larger molecules or more diverse combinations is limited. In this study, we show the results of applying the VQE algorithm to LiCoO_2 molecules and demonstrate the applicability of quantum computers in the field of battery development.

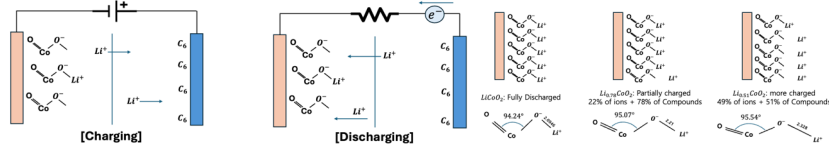


Fig. 1. Left) Schematic of battery charging and discharging, Right) Geometry of compounds by oxidation state

When a voltage is applied to the anode and cathode to supply electrons (i.e., charging), the Li^+ ions migrate to the cathode. When discharged from this charged state, the Li^+ ions from the negative electrode are recombined with the positive electrode, as shown in the figure below, and current flows through the resistor. At this time, depending on the oxidation state of lithium, the average geometry of the molecular structure changes as shown in **Fig.1**. so, the energy in the bulk can be estimated using the geometry. The energy of the molecule is calculated by assuming the gas-phase model, which considers the average oxide structure as a single molecule.

2.2 VQE(Variational Quantum Eigensolver)

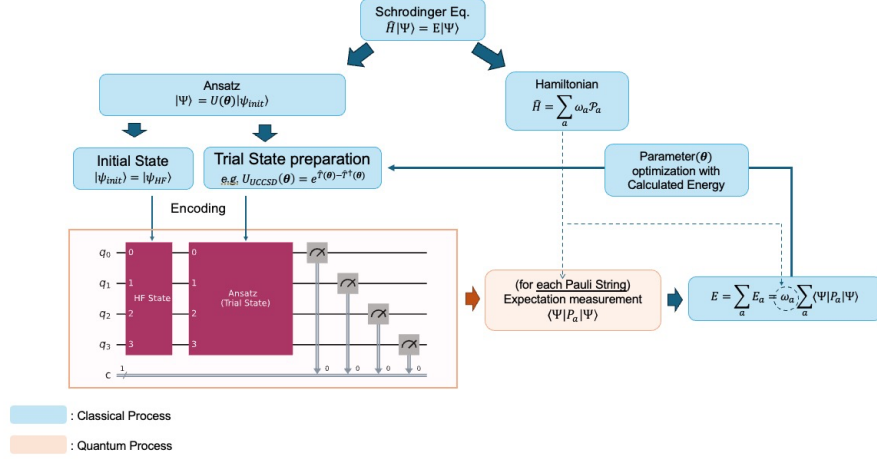


Fig. 2. VQE Pipeline

The VQE algorithm is an eigenvalue problem solving algorithm first proposed by Peruzzo et al. in [2] that works by computing the energy in Hilbert space that provides an upper bound on the ground state energy according to the variational principle. VQE computes the ground state energy of a system by measuring the energy on a quantum computer and performing optimization on a classical computer. To measure the energy of a system on a quantum computer, we need to map the Hamiltonian to a foliate, represent an arbitrary quantum state, and encode it in a quantum circuit.

Hamiltonian

The Hamiltonian of a molecule is represented by the Electronic Structure Hamiltonian as shown below.

$$\hat{H}_{el}(r) = -\sum_i \frac{1}{2} \nabla_{r_i}^2 - \sum_I \sum_i \frac{Z_I}{|R_I - r_i|} + \sum_i \sum_{j>i} \frac{1}{|r_i - r_j|} \quad (1)$$

(where r_i denotes the position vector of the i^{th} electron, R_I denotes the position of the i^{th} nucleon, and Z_I is the atomic number of the I^{th} nucleon). To map this Hamiltonian to the Pauli gate, we write it in second quantized form, which is expressed through the fermionic creation/annihilation operator as follows

$$\hat{H}_{el} = \sum_{p,q} h_{pq} \hat{a}_p^\dagger \hat{a}_q + \frac{1}{2} \sum_{p,q,r,s} h_{pqrs} \hat{a}_p^\dagger \hat{a}_q^\dagger \hat{a}_r \hat{a}_s, \quad (2)$$

$$\text{where, } h_{pq} = \langle \phi_p | \hat{H}_{el} | \phi_q \rangle, \quad h_{pqrs} = \langle \phi_p \phi_q | \hat{H}_{el} | \phi_r \phi_s \rangle \quad (3)$$

Fermionic operators can be mapped to Pauli gates using Jordan-Wigner mapping, Parity mapping, and bravyi kitaev mapping. In this experiment, we will use the Parity mapping method. Parity mapping expresses the creation/annihilation operator through a

Pauli gate by corresponding the parity of the i^{th} qubit to the parity of the electron occupancy of the i^{th} orbital. In this case, the parity of the α -spin β -spin of the molecule can be utilized to reduce the number of qubits required by two. The resulting Hamiltonian of the mapping is represented by a Pauli string (\mathcal{P}_a) and its weights or linear coefficients (ω_a) as shown below.

$$\hat{H} = \sum_a \omega_a \mathcal{P}_a \quad (4)$$

Ansatz

To represent arbitrary quantum states, we construct a parameterized quantum circuit (PQC) with linear coefficients corresponding to each basis as parameters. The performance of the ansatz is difficult to evaluate, and it is in the a priori domain that it is not known which ansatz is better for different systems. Therefore, in this experiment, we use two different ansatzes to get better results.

Two-Local (Efficient $SU(2)$) Ansatz

The quantum state of a single qubit is represented by a rotation gate for two arbitrary angles on a Bloch sphere. If the angle of the rotation gate is a parameter, and the entanglement, which is the property of a qubit in a multi-qubit system, is expressed using a CNOT gate, the quantum state of an arbitrary multi-qubit system can be expressed using parameters.

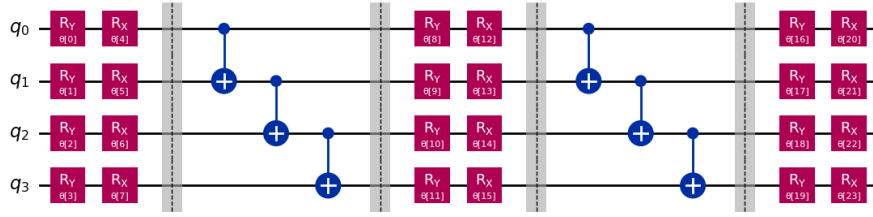


Fig. 3. Two_Local Ansatz Example on a 4-Qubit System

The construction of the Two-Local Ansatz does not involve a Hamiltonians interpretation. It is a representation of an arbitrary quantum state that is represented in hardware and is therefore hardware efficient.

UCCSD Ansatz

Through the second quantization, we express the Hamiltonian operator as the basis of the molecular orbitals of the system. In the end, the state we are looking for is expressed in the same Hilbert space as the Hamiltonian, so using the same spinorbit as the basis, we can express any quantum state in the Hilbert space where the Hamiltonian exists. In this way, the quantum state is expressed through Coupled-Cluster Theory using the spin-orbit wavefunction of the molecule as the basis, and considering only the 2nd-order excitation, it is expressed as a unitary one, UCCSD (Unitary Coupled Cluster Singles and Doubles), and the expression is as follows.

$$\hat{T} = \hat{T}_1 + \hat{T}_2, \text{ (Cluster operator)} \quad (5)$$

$$\text{where, } \hat{T}_1 = \sum_{i,a} c_i^a \hat{a}_a^\dagger \hat{a}_i, \quad \hat{T}_2 = \sum_{ijab} t_{ij}^{ab} \hat{a}_a^\dagger \hat{a}_b^\dagger \hat{a}_j \hat{a}_i$$

$$|\Psi\rangle_{UCCSD} = e^{\hat{T}-\hat{T}^\dagger} |\psi_{ref}\rangle \quad (6)$$

The states are represented by creation and destruction operators, which can be encoded in quantum circuits via Jordan-Wigner mapping or parity mapping.

The energy of the system is calculated by measuring the expectation value for each Pauli string in the Hamiltonian for these quantum circuits. By iterating over these measurements and optimizing the parameters of the ansatz so that the calculated energy is minimized, an upper bound on the ground state energy is found. The resulting minimum corresponds to the ground state energy calculated by the VQE algorithm for the system.

2.3 FMO(Fragmental Molecular orbital)

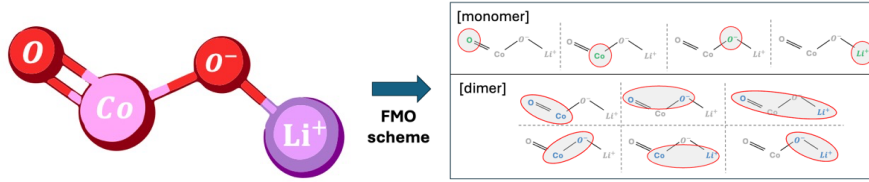


Fig. 4. An example of the FMO scheme in LiCoO_2

The FMO method is a method that divides a whole molecular system into smaller system fragments and approximates the energy of the whole system by using the energy of one fragment (monomer) and the energy of pairs of fragments (dimers). When calculating the ground state energy of a molecule by the usual method, the computational complexity is exponential in the size of the system. However, the FMO method can effectively reduce the computational complexity by dividing a large system into parts. The FMO method has two main steps. The first is the FMO-based restricted Hartree-Fock (FMO-RHF) process, which calculates the RHF through the Hamiltonian of monomers and dimers. The Hamiltonian of monomer(\hat{H}_A) and the Hamiltonian of dimer(\hat{H}_{AB}) are as follows.

$$\hat{H}_A = \sum_{i \in A} \left[-\frac{\nabla_i^2}{m_e} - \sum_I \frac{Z_I}{|r_i - R_I|} + \sum_{C \neq A}^{N_{tot}} \int dr' \frac{\rho_C(r')}{|r_i - r'|} \right] + \sum_{i \in A} \sum_{j \in A} \frac{1}{|r_i - r_j|} \quad (7)$$

$$\hat{H}_{AB} = \sum_{i \in A, B} \left[-\frac{\nabla_i^2}{m_e} - \sum_I \frac{Z_I}{|r_i - R_I|} + \sum_{C \neq A, B}^{N_{tot}} \int dr' \frac{\rho_C(r')}{|r_i - r'|} \right] + \sum_{i \in A, B} \sum_{j \in A, B} \frac{1}{|r_i - r_j|} \quad (8)$$

(Where A,B are the respective fragments. C refers to the fragments that are not included in monomer A or dimer AB, and the term $\rho_j(r')$ is the electric charge density of the fragments outside the monomer or dimer.) By calculating the RHF of each monomer and dimer through the Hamiltonian above, we can get $E^{FMO-RHF}$, which can be expressed as follows.

$$E^{FMO2-RHF} = E^{FMO1-RHF} + \Delta E^{FMO2-RHF} \quad (9)$$

$$E^{FMO1-RHF} = \sum_A^N E_A \quad (10)$$

$$\Delta E^{FMO2-RHF} = \sum_{A>B}^N [E_{IJ} - E_I - E_J] \quad (11)$$

In the above equations, N refers to all fragments. From the above equations, we can get the HF energy including the electrostatic potential of the whole molecule. The next step is to proceed with FMO based coupled-cluster (FMO-CC). This process is performed to obtain a more accurate value of the FMO-RHF energy, and since this process does not include the electrostatic potential, the Hamiltonian can be calculated using the Electronic Structure Hamiltonian defined earlier. The process of calculating the total energy (E^{FMO-CC}) and correlation energy ($E^{FMO-corr}$) obtained through FMO-CC is as follows.

$$E^{FMO n-CC} = E^{FMO n-RHF} + E^{FMO n-corr} \quad (12)$$

$$E^{FMO2-corr} = E^{FMO1-corr} + \Delta E^{FMO2-corr} \quad (13)$$

$$E^{FMO1-corr} = \sum_A^N E_I^{corr} \quad (14)$$

$$\Delta E^{FMO2-corr} = \sum_{A>B}^N (E_{IJ}^{corr} - E_I^{corr} - E_J^{corr}) \quad (15)$$

In the above, the FMO-RHF and FMO-CC processes should be separated because the Hamiltonian used in the two processes may be different, and the FMO-CC process should proceed after the SCF calculation of FMO-RHF converges. However, in this experiment, the electrostatic potential term in the FMO-RHF Hamiltonian was approximated by ignoring it, so each process was carried out consecutively and the total energy was calculated from the obtained monomer and dimer energies as follows.

2.4 FMO/VQE (Fragment molecular orbital-based Variational Quantum Eigensolver)

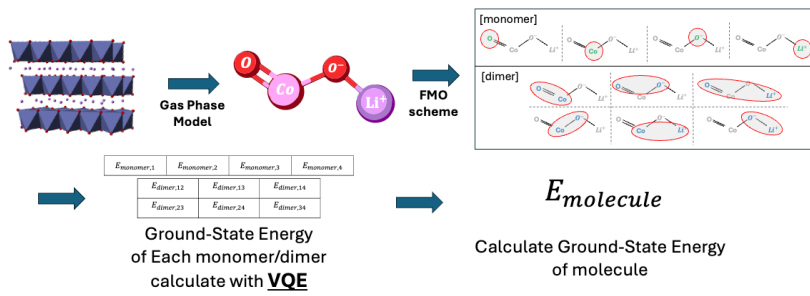


Fig. 5. An illustration of applying the FMO/VQE method assuming a gas phase model

FMO/VQE is an algorithm first introduced by Hochel Lim et al. that combines the VQE algorithm with the FMO method. In the VQE algorithm, the Hamiltonian is represented based on the molecular spinorbitals, so it requires as many qubits as the number of

molecular spinorbitals. However, the number of qubits is currently limited, so the number of molecules that can be simulated in the existing VQE algorithm is limited. However, if the FMO Method is applied to the existing VQE algorithm, the number of qubits required for a single calculation can be reduced by parallelizing the calculation and simulating the total system in pieces.

Let's compare the number of qubits required with and without the FMO method. Both calculations use Active Space to reduce the number of qubits required. Given the dynamics of the system, excitations in the core orbital of each atom will have a smaller probability than excitations in the valence orbital. Therefore, we will calculate the core orbital, which is less likely to be involved in bonding, as a separate constant, and assume that the space spanned by the wavefunction of the valence orbital of each atom is a Hilbert space. Under these assumptions, the number of qubits required for a conventional VQE calculation is as follows.

Table 1. Orbital structure of each atom

atom	Number of Valence orbital	Number of Valence electron
Li	2	1
O	6	8
Co	10	7

Table 2. Orbital structure of each Fragment

Configuration	Number of Valence orbital	Number of Valence electron
monomer		
Li	2	1
O	6	4
Co	10	7
Dimer		
Li – O	6	5
Li – Co	12	8
O – O	12	8
Co – O	16	11

Therefore, the number of spinorbitals required for the VQE calculation is 24, considering that oxygen has 2. If we use the parity mapper here, we can reduce the number of qubits needed for the calculation by 2, so 22 qubits are needed to solve this system via conventional VQE.

Let's apply the FMO method, where each atom is a fragment, which means that 4 monomers are created and 6 dimer systems need to be calculated. The number of spin-orbitals required for each calculation is shown below (only one is shown for configurations with the same composition).

Therefore, the maximum number of qubits required in the calculation is 14 (using Parity mapper). By applying the FMO method, we can reduce the number of qubits required

for the calculation by 8. Furthermore, the potential of this method is in larger systems. When simulating larger molecules such as Ni, Mn, etc. added to the popular LiCoO_2 molecule, conventional VQE requires more qubits than the number of valence orbitals of those atoms. However, in FMO/VQE, the increase in the size of the system leads to an increase in the number of fragments, so it is possible to calculate the energy of larger molecules using the same number of qubits.

3 Result

3.1 Monomer/Dimer Energy

Table 3. Energy of Monomer for each Ansatz/Optimizer

System	UCCSD		Two-Local	
	COBYLA	SPSA	COBYLA	SPSA
"Li"	-7.315526	-7.315526	-7.315526	-7.315526
"O"	-73.804150	-73.804150	-73.804150	-73.804150
"Co"	-1365.943920	-1366.002044	-1366.006785	-1365.740246

Table 4. Energy of Dimer for each Ansatz/Optimizer for each Oxidation State of Li_xCoO_2

[x=1] Configuration	UCCSD		Two-Local	
	COBYLA	SPSA	COBYLA	SPSA
"Li-O ¹ "	-81.036530	-81.062982	-81.069739	-81.023626
"Li-O ² "	-81.117531	-81.117951	-81.117956	-81.094238
"Li-Co"	-1373.429758	-1373.567946	-1373.618739	-1373.576316
"O-O"	-147.432250	-147.485749	-147.599156	-147.528891
"Co-O ¹ "	-1439.355819	-1439.796790	-1439.957262	-1439.315189
"Co-O ² "	-1439.459214	-1439.788926	-1439.888880	-1439.396154

[x=0.94] Configuration	UCCSD		Two-Local	
	COBYLA	SPSA	COBYLA	SPSA
"Li-O ¹ "	-81.079066	-81.071382	-81.084085	-81.014315
"Li-O ² "	-81.119992	-81.119699	-81.120071	-81.117758
"Li-Co"	-1373.434678	-1373.568648	-1373.569509	-1373.347301
"O-O"	-147.391974	-147.374393	-147.608042	-147.551759
"Co-O ¹ "	-1439.423661	-1439.843317	-1439.952027	-1439.169880
"Co-O ² "	-1439.393635	-1439.485720	-1439.898848	-1439.210821

[x=0.78] Configuration	UCCSD		Two-Local	
	COBYLA	SPSA	COBYLA	SPSA
"Li-O ¹ "	-81.074588	-81.076294	-81.086310	-81.070688
"Li-O ² "	-81.118325	-81.119637	-81.120088	-81.097025
"Li-Co"	-1373.206240	-1373.319480	-1373.354329	-1373.591995
"O-O"	-147.412124	-147.551726	-147.607067	-147.517407
"Co-O ¹ "	-1439.226044	-1439.778362	-1439.840047	-1439.319849
"Co-O ² "	-1439.547179	-1439.828032	-1439.944688	-1439.300309

[x=0.75]		UCCSD				Two-Local	
Configuration	COBYLA	SPSA	COBYLA	SPSA	COBYLA	SPSA	
"Li-O ¹ "	-81.065682	-81.081421	<u>-81.086310</u>	-81.003117	-81.023070	-81.012210	
"Li-O ² "	-81.119826	-81.120074	<u>-81.120088</u>	-81.095257	-81.071903	<u>-81.120088</u>	
"Li-Co"	-1373.223660	-1373.354256	-1373.328463	-1373.589681	-1373.585003	<u>-1373.790507</u>	
"O-O"	-147.413053	-147.281268	<u>-147.607526</u>	-147.445853	-147.442307	-147.462992	
"Co-O ¹ "	-1439.226044	-1439.778362	<u>-1439.840047</u>	-1439.319849	-1439.523202	-1439.714842	
"Co-O ² "	-1439.388348	-1439.758178	<u>-1439.857075</u>	-1439.354511	-1439.479002	-1439.722998	

[x=0.66]		UCCSD				Two-Local	
Configuration	COBYLA	SPSA	COBYLA	SPSA	COBYLA	SPSA	
"Li-O ¹ "	-81.053296	-80.984815	<u>-81.084085</u>	-81.014315	-81.042272	-81.084060	
"Li-O ² "	-81.119992	-81.119699	<u>-81.120071</u>	-81.117758	-81.066772	<u>-81.120071</u>	
"Li-Co"	-1373.434678	-1373.568648	-1373.569509	-1373.347301	-1373.529010	<u>-1373.584236</u>	
"O-O"	-147.391974	-147.374393	<u>-147.608042</u>	-147.551759	-147.523597	-147.489992	
"Co-O ¹ "	-1439.423661	-1439.843317	<u>-1439.952027</u>	-1439.169880	-1439.485884	-1439.803393	
"Co-O ² "	-1439.393635	-1439.485720	-1439.898848	-1439.210821	-1439.564247	<u>-1439.986058</u>	

*All values in the table are in Ha units. The underlined values represent the lowest energy convergence values for each configuration. The configurations with superscripts refer to different dimers with the same composition but different geometric structures.

Li-O¹: A dimer consisting of O and Li atoms that form a bond.

Li-O²: A dimer consisting of O and Li atoms that do not form a bond.

Co-O¹: A dimer consisting of Co and O atoms forming a single bond.

Co-O²: A dimer consisting of Co and O atoms forming a double bond.

Tables 1 and 2 present the VQE calculation results for each monomer and dimer of the LiCoO₂ molecule, divided into fragments. A total of six cases were computed, combining two types of Ansatz and three types of optimizers. Under the conditions of this experiment, the energy of the monomer does not depend on the relative coordinates of each fragment and is therefore independent of the molecular geometry. In other words, even if the oxidation state changes, the calculation results remain the same, so the calculation is performed only once.

In terms of optimizer selection, the difference between L-BFGS-B and other optimizers is approximately on the order of 0.1 Ha. This represents a significant error compared to the desired precision. While COBYLA and SPSA are much faster in terms of computation time, L-BFGS-B demonstrates better results when precision is prioritized.

Several configurations are noteworthy when selecting an Ansatz. For the Co-O¹ and Li-O¹ configurations, the UCCSD Ansatz consistently produced the lowest energy values across all calculations. In contrast, for the Li-Co configuration, Two_Local showed better convergence values than UCCSD in oxidation states other than $x = 1$. However, for configurations such as Co-O², Li-O and O-O the optimal Ansatz varied despite the identical fragment structures in each oxidation state. For instance, in the results for Co-O² the lowest energies were achieved with UCCSD at $x = 0.78$ and $x=0.75$, while Two_Local outperformed at $x = 1.00$, $x = 0.94$, and $x = 0.66$. This indicates that even for systems with similar characteristics, the choice of Ansatz that best represents the system can change depending on the specific configuration.

The choice of Ansatz ultimately determines how the Hilbert space is represented in each basis. In cases where the space represented by spin orbitals is not large, the difference between Ansatz selections is minimal. However, as the dimension of the Hilbert space increases, the expressive power of different Ansatz types can diverge significantly.

For systems such as Li-O^1 and Li-O^2 which are represented using approximately 8 spin-orbital wavefunctions as the basis, the energy difference between Ansatz is typically on the order of $10^{-3} Ha$, which is relatively small. However, as the system size grows, the difference becomes more pronounced. For example, in systems like Li-Co and O-O^2 represented by 12 spin-orbital wavefunctions, or Co-O^1 and Co-O^2 represented by 16 spin-orbital wavefunctions, the difference increases to the order of $10^{-1} Ha$. Such differences can significantly impact the precision of the calculated values.

Therefore, the choice of Ansatz can have a substantial effect depending on the system, particularly for experiments requiring high precision. It is essential to evaluate multiple Ansätze and select the optimal one for each specific system to ensure the best possible results.

3.2 Molecular Energy

The $E_{FMO/VQE}$ values in Table 5, 6 represent the ground-state energies of Li_xCoO_2 for each oxidation state(x) calculated using the FMO/VQE methodology proposed in this paper. The E_{FCI} and E_{CCSD} values are obtained using the traditional computational chemistry methods with the GAMESS simulator, corresponding to the Full Configuration Interaction (FCI) and Coupled-Cluster with Single and Double excitations (CCSD) methods, respectively. In the FCI calculations, the Active Core Scheme was used, whereas it was not used in the CCSD calculations. The energy values represent the results obtained from these classical simulation methods, and the error refers to the difference between the FMO/VQE calculation results and those obtained from the traditional methods. Additionally, the accuracy is expressed as the percentage of this error relative to the traditional energy values.

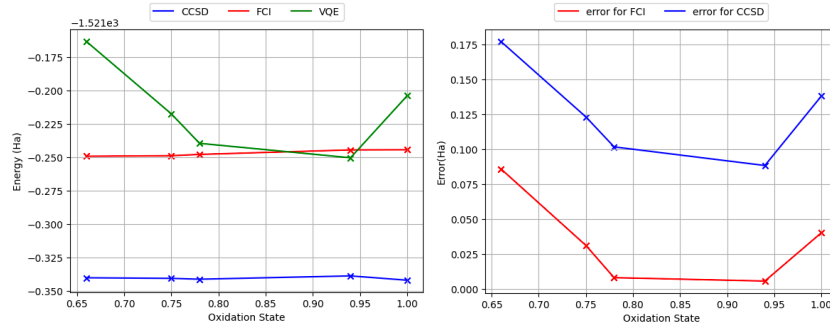
First, let's examine the comparison between VQE and FCI calculations. FCI is considered the most accurate theoretical calculation method and is often used as a reference value in many chemical calculations. In this experiment, both VQE and FCI used the same Active Core Scheme to approximate the system before calculating the energy, and it was confirmed that the VQE algorithm produced energy values very similar to those obtained from FCI. The smallest error was observed at $x = 0.94$, where the error was -0.005892 , resulting in a lower energy value compared to FCI, indicating that the VQE algorithm is highly accurate. On the other hand, at $x = 0.66$ the error was 0.086063 , and at $x = 1$ the error was 0.040462 , showing a relatively larger difference.

Table 5. Comparison of the energy, error, and accuracy between FMO/VQE and FCI

Oxidation Number (Li_xCoO_2)	$E_{\text{FMO/VQE}}$	E_{FCI}	Error	Accuracy(%)
1.00	-1521.203868	-1521.244330	0.040462	99.9973
0.94	-1521.250344	-1521.244451	-0.005892	99.9996
0.78	-1521.239517	-1521.247824	0.008307	99.9995
0.75	-1521.217377	-1521.248807	0.031431	99.9979
0.66	-1521.163133	-1521.249196	0.086063	99.9943

Table 6. Comparison of the energy, error, and accuracy between FMO/VQE and CCSD

Oxidation Number (Li_xCoO_2)	$E_{\text{FMO/VQE}}$	E_{CCSD}	Error	Accuracy(%)
1.00	-1521.203868	-1521.342101	0.138232	99.9884
0.94	-1521.250344	-1521.338869	0.088525	99.9919
0.78	-1521.239517	-1521.341358	0.101841	99.9933
0.75	-1521.217377	-1521.340712	0.123336	99.9942
0.66	-1521.163133	-1521.340283	0.177150	99.9909

**Fig. 6.** Energy of FMOVQE, FCI, and CCSD for each oxidation state and the error graph between FMOVQE and FCI, FMOVQE and CCSD

Next, compare VQE, FCI, and CCSD. FCI is generally regarded as the most precise calculation method and typically provides lower energy convergence values than CCSD. However, in this study, CCSD calculated energy values that were approximately 0.1 Ha lower than FCI for all oxidation states. This difference is related to the application of the Active Core Scheme. While FCI used the Active Core Scheme, CCSD performed calculations on the entire molecule, which led to lower energy values.

Through the comparison of the FMO/VQE algorithm with FCI, it was confirmed that the precision of FMO/VQE closely matches that of FCI. This suggests that the FMO/VQE method achieves an accuracy similar to that of traditional computational

methods, and quantum computing can deliver practical levels of accuracy. This result indicates that, with further advancements in quantum computing technology, it may provide more efficient and accurate calculations compared to traditional computational chemistry methods. Therefore, this paper suggests that the FMO/VQE method demonstrates accuracy nearly identical to that of classical methods and, with future improvements in quantum computers, could potentially achieve even higher accuracy. This highlights a significant advancement for chemical calculations using quantum computing, demonstrating that it may enable faster and more precise predictions than existing classical methods.

4 Conclusions

In this study, the quantum algorithm FMO/VQE was used to calculate the ground-state energy of the Li_xCoO_2 molecule with high accuracy, compared to classical simulators. By comparing the results with those from FCI calculations, the precision of the FMO/VQE algorithm was assessed, and errors due to the use of the Active Core Scheme were identified through comparisons with CCSD calculations. This result demonstrates the potential applicability of the FMO/VQE method for future drug synthesis and new material development.

Classical computing methods are not capable of handling the complex molecules required for drug synthesis and new material development. The FMO/VQE algorithm proposed in this study reduces the number of qubits needed for calculations through the FMO method. This allows for the resolution of complex molecular systems using quantum computing with VQE. These results suggest that the FMO/VQE algorithm could become a new paradigm in fields like drug synthesis and new material development, where the calculation of complex molecules is crucial.

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