A comprehensive study on how to construct local bases in deep variational quantum eigensolver for molecular systems

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Current quantum computers are limited in the number of qubits and coherence time, constraining the algorithms executable with sufficient fidelity. Variational quantum eigensolver (VQE) is an algorithm to find an approximate ground state of a quantum system and expected to work on even such a device. The deep VQE [K. Fujii, et al., arXiv:2007.10917] is an extension of the original VQE algorithm, which takes a divide-and-conquer approach to relax the hardware requirement. While the deep VQE is successfully applied for spin models and periodic material, its validity on a molecule, where the Hamiltonian is highly non-local in the qubit basis, is still unexplored. Here, we discuss the performance of the deep VQE algorithm applied to quantum chemistry problems. Specifically, we examine different subspace forming methods and compare their accuracy and complexity on a ten H-atom tree-like molecule as well as a 13 H-atom version. Additionally, we propose multiple methods to lower the number of qubits required to calculate the ground state of a molecule. We find that the deep VQE can simulate the electron-correlation energy of the ground-state to an error of below 1%, thus helping us to reach chemical accuracy in some cases. The accuracy differences and qubits reduction highlights the basis creation method’s impact on the deep VQE.

I. INTRODUCTION

Quantum algorithms have better asymptotic behavior than classical alternatives for some of the most challenging and intriguing calculation problems today. Outstanding examples are factorization of large numbers using the Shor algorithm [1], linear algebraic processes (matrix inversion) [2–4], as well as promising results in quantum machine learning [5]. Quantum chemistry will, however, arguably be the research field that gets impacted most by quantum computing [6]. Simulating large molecules using a quantum computer will push our understanding of nature to new levels and significantly affect today’s society. A specific intent in quantum chemistry is to calculate the electronic ground state of a molecule. Since the dimension of the Hilbert space grows exponentially to the system size, a molecular electronic structure can be exceedingly complex to be solved on a classical computer.

Despite recent developments [7–9], we anticipate that quantum computers in the near future still have a limited number of qubits and only partial error resistance. Therefore the number of operations on a quantum device is limited, as the errors are building up to a point where we cannot receive meaningful results. These restrictions confine us to the area of so-called noisy intermediate-scale quantum (NISQ) algorithms [10].

A promising approach for overcoming these obstacles is the variational quantum eigensolver (VQE) [11] algorithm. One uses the parameterized variational circuit, or ansatz, on a quantum device to prepare a trial state and uses the classical computer to set the parameters for the ansatz. By looping between the quantum device and a classical computer, one tries to minimize the energy. Using such an approach, it is possible to find a good approximation of the ground state. The states created in such a way can be classically hard to represent. Depending on the ansatz, this only requires a circuit with a modest number of gates. There is, therefore, a possibility that VQE could be running on a NISQ device.

The system size solvable by the VQE is essentially limited by the number of qubits on the quantum device, which makes it challenging to apply it to large-scale systems. In classical quantum chemistry methods, fragmentation techniques [12] proved to be highly successful in reducing the computational requirement to handle large-scale molecules. Recently divide-and-conquer techniques in quantum computing also got much attention to potentially solve the problem of the limited quantum resource [13–16]. They aim to decrease the number of qubits needed to solve complex problems. In these techniques, we separate the original system into separate subsystems and first solve them individually. The subsystem solutions set the starting point to formulate a meaningful result of the original problem.

One such method is the deep VQE [13]. It first separates the target quantum system into subsystems and obtains their approximate ground states by usual VQE. The
next step constructs a basis set for each subsystem by applying specific excitation operators to the subsystem ground states. The basis set is later used to form an effective Hamiltonian of the whole system. Another VQE can then solve the effective Hamiltonian to obtain a ground state of the target system. This process can be repeated multiple times to solve increasingly large systems. The performance of the deep VQE has been analyzed for spin systems [13] and periodic materials [14], which have suggested that the deep VQE can produce accurate results while reducing the number of needed qubits simultaneously. However, only systems with minimal interactions between subsystems have been examined. Molecules are highly complex systems for which the Hamiltonian consists of many terms with various strengths. Such complex systems provide a unique set of challenges for the deep VQE. It is therefore vital to examine the performance of the deep VQE for such systems.

In this work, we propose multiple methods to create the search-able subspace and compare their influence on the accuracy using a tree-like molecule as a testbed. The strategy for creating a subsystem basis ultimately limits the deep VQE performance as it determines which subspace of the entire Hilbert space can be explored to express the ground state. To examine the impact of different bases on the deep VQE, we tested three strategies to select excitation operators to form the subsystem basis. The first method followed the original paper and is based on the interaction operators. The second technique we examined uses single-qubit Pauli operators to create a basis set, whereas the third obtain a basis using single-electron excitation and deexcitation operators. Additionally to the different basis creation methods, we propose multiple techniques to control the needed qubits. We chose two molecular systems to test our methods, a 10-atom, and a 13-atom dendrimer-like molecule, consisting of only hydrogen atoms. This work shows the remarkable accuracy improvement and reduction of qubits the deep VQE offers for even complex quantum systems. We find that the deep VQE can simulate the electron-correlation energy of the ground-state to an error of below 1%, thus helping us reach chemical accuracy in some cases. Our understanding of the various basis creation methods and the comparison of their performance provides an essential recipe for determining the bases of subsystems for the use of deep VQE in larger molecules of even more practical importance.

II. THEORY

A. Deep VQE

The deep VQE [13] is an algorithm of the divide and conquer family. Its purpose is to calculate the ground state of a quantum system. We review the algorithm of the original deep VQE, which can treat spin Hamiltonians consisting of 2-local interactions. In deep VQE, we first divide the system into $M$ subsystems. The problem Hamiltonian with 2-local interactions can consequently be written in the form of

$$H = \sum_{i=1}^{M} H_i + \sum_{i,j=1}^{M} V_{ij},$$

(1)

where $H_i$ acts on subsystem $i$ and $V_{ij}$ on subsystems $i$ and $j$. The interaction term $V_{ij}$ decompose into operators $V_{\mu,i,j}$ acting only on single subsystems $i$:

$$V_{ij} = \sum_{\mu} A_{ij}^{\mu} V_{\mu,i} V_{\mu,j},$$

(2)

where $\mu$ indicates the different interaction terms. The deep VQE first finds ground states $|G_i\rangle$ of each subsystem Hamiltonian $H_i$ using a VQE algorithm. Then we build a subsystem basis $\{|b_{i,k}\rangle\}_{k=1}^{K_i}$ for each subsystem $i$ with dimensions $K_i$ by acting with certain excitation operators $B_i = \{B_{i,k}\}$ on $|G_i\rangle$. The selection of the excitation operators $B_{i,k}$ is a crucial step which determines the performance of the deep VQE, and we will discuss it in detail in Sec. III. To ensure the created basis set $\{|b_{i,k}\rangle\}_{k=1}^{K_i}$ is orthogonal a Gram-Schmidt orthogonalization is applied.

In the next step, we construct an effective Hamiltonian $H^{\text{eff}}$ by projecting the original Hamiltonian $H$ to the subspace spanned by $\bigotimes_i \{|b_{i,k}\rangle\}_{k=1}^{K_i}$. This can be achieved by measuring all matrix elements $\langle b_{i,k} | H_i | b_{i,i} \rangle$ and $\langle b_{i,k} | V_{ij} | b_{i,p} \rangle | b_{j,q} \rangle = \sum_{\mu} A_{ij}^{\mu} \langle b_{i,k} | V_{\mu,i} | b_{i,p} \rangle \langle b_{j,l} | V_{\mu,j} | b_{j,q} \rangle$. Notice that calculating the expectation values only involves one subsystem, which can be calculated separately on a quantum computer. Now, we represent each subsystem by $N_i = [\log_2 K_i]$ qubits, and we again utilize the VQE using $N_{\text{tot}} = \sum_{i} [\log_2 K_i]$ qubits to search for the ground state of $H^{\text{eff}}$. Even though $H^{\text{eff}}$ whose dimension is $d = \prod_i K_i$ could, in principle, be mapped to $[\log_2 d] \leq N_{\text{tot}}$ qubits, it is beneficial to map each subsystem individually since it preserves the locality of the Hamiltonian. The deep VQE algorithm can be repeated iteratively to treat larger and larger systems on a qubit-limited quantum computer.

B. Deep VQE for fermionic system

The central problem in quantum chemistry is finding the first-principles Hamiltonian’s ground state, which describes interacting electrons. In second quantization, the Hamiltonian can be written in the form

$$H = \sum_{\alpha,p=1}^{N} h_{\alpha,p} a_{\alpha,p}^\dagger a_{\alpha,p} + \sum_{q,r,s,t=1}^{N} h_{q,r,s,t} a_{q,r}^\dagger a_{s,t} a_{s,t} a_{q,r}^\dagger a_{s,t}.$$

(3)

where $N$ is the number of orbitals of the molecule. Here we consider how to apply the deep VQE to this Hamiltonian.
A frequent approach to treat the Hamiltonian of Eq. (3) on a quantum computer is to map the fermionic operators \( a_1^\dagger \) and \( a_s \) to qubit operators through, e.g., Jordan-Wigner transformation. This leads to a Hamiltonian in the form of,

\[
H = \sum_i h_i P_i,
\]

where \( P_i \) is a Pauli string, \( P_i \in \{ I, X, Y, Z \}^\otimes N \), and \( h_i \) is a real coefficient. The obstacle to apply the deep VQE is that \( P_i \) is not 2-local but can be nonlocal when using JW transformation as we will see below.

After partitioning the system into \( M \) subsystems and \( M_{\text{int}} \) interaction terms, we can write the Hamiltonian in Eq. (4) in the form of,

\[
H = \sum_{i=1}^{M} H_i + \sum_{\mu=1}^{M_{\text{int}}} \lambda^\mu V_{\mu,1} \otimes V_{\mu,2} \otimes \cdots \otimes V_{\mu,M}, \tag{5}
\]

where \( H_i \) and \( V_{\mu,i} \) is an operator acting only on the \( i \)-th subsystem, and \( \mu \) indexes different interaction terms. To perform the deep VQE, we construct the effective Hamiltonian \( H_{\text{eff}} \) using the subsystem basis \( \{ b_{i,k} \}_{k=1}^{K_i} \).

To do this, we measure the matrix elements \( \langle H_{\text{eff}} \rangle_{k,l} = \langle b_{i,k} | H_i | b_{i,l} \rangle \) and \( \langle V_{\mu,i} \rangle_{k,l} = \langle b_{i,k} | V_{\mu,i} | b_{i,l} \rangle \) for all combinations of \( i \), \( k \), \( l \), and \( \mu \). Let \( H_{\text{eff}} \) and \( V_{\mu,i} \) be \( n_i = \lceil \log_2 K_i \rceil \)-qubit operators with the evaluated matrix elements. The total effective Hamiltonian can be written as,

\[
H_{\text{eff}} = \sum_{i=1}^{M} H_i^{\text{eff}} + \sum_{\mu=1}^{M_{\text{int}}} \lambda^\mu V_{\mu,1}^{\text{eff}} \otimes V_{\mu,2}^{\text{eff}} \otimes \cdots \otimes V_{\mu,M}^{\text{eff}}. \tag{6}
\]

If we wish to perform the VQE of \( H_{\text{eff}} \), we must be able to efficiently measure the expectation value \( \langle H_{\text{eff}} \rangle = \langle \psi | H_{\text{eff}} | \psi \rangle \) for a given state \( | \psi \rangle \) prepared on a quantum computer. A usual approach to measure the expectation value of a Hamiltonian is to expand it into Pauli operators. However, if we do so in Eq. (6), the number of Pauli operators grows exponentially to \( M \) due to its nonlocality, which blocks efficient evaluation of \( \langle H_{\text{eff}} \rangle \).

We must avoid this exponential growth to apply the deep VQE to fermionic systems. To this end, we first diagonalize \( V_{\mu,i}^{\text{eff}} \) classically and obtain unitary \( U_{\mu,i} \), such that

\[
V_{\mu,i}^{\text{eff}} = U_{\mu,i}^\dagger \text{diag}(v_{\mu,i}) U_{\mu,i}, \tag{7}
\]

where \( v_{\mu,i} \) are eigenvalues of \( V_{\mu,i}^{\text{eff}} \). Note that this process can be performed in time \( O^{(n_i)} \) and that it is natural to assume \( n_i \) is a small constant in the deep VQE. We can also construct a quantum circuit to realize a \( 2^{n_i}\)-dimensional unitary \( U_{\mu,i} \) in time \( O^{(n_i)} \). Therefore, we can measure \( \langle V_{\mu,1}^{\text{eff}} \otimes \cdots \otimes V_{\mu,M}^{\text{eff}} \rangle \) for a given \( \sum_{i=1}^{M} n_i \)-qubit state \( | \psi \rangle \) by first applying \( U_{\mu,1} \otimes \cdots \otimes U_{\mu,M} \) to \( | \psi \rangle \) and then measuring it in the computational basis.

Therefore, the required measurements to determine the expectation value of the interaction term scale linearly with the number of interactions in the system. Thereby, we can efficiently measure \( \langle V_{\mu,1}^{\text{eff}} \otimes \cdots \otimes V_{\mu,M}^{\text{eff}} \rangle \) and hence \( \langle H_{\text{eff}} \rangle \) as well.

Note that we prefer JW transformation for applying deep VQE to fermionic systems because it directly maps the \( i \)-th orbital to the \( i \)-th qubit; if the \( i \)-th qubit is \( | 1 \rangle \), it indicates that an electron occupies the \( i \)-th orbital. This property allows us to split systems naturally into subsystems using localized orbitals. Although other techniques such as Bravyi-Kitaev (BK) transformation relaxes the locality of Pauli strings down to \( O(\log N) \), it makes the correspondence between fermion occupation and qubit configurations not straightforward. Note that the above technique may also be used for BK-transformation-based deep VQE to obtain the expectation values efficiently.

III. METHOD

A. Basis creation strategies

Choosing a basis set impacts the performance of the deep VQE. It ultimately determines the number of qubits needed to run the algorithm and the accuracy of the result. It is therefore vital to make an educated decision for the used method.

Below we examine different procedures to generate the basis sets, which are summarized in Tab. I. In the single Pauli method, we use the set of the Pauli operators \( X,Y,Z \) acting on each qubit as \( B_i \). In the interactions approach, we apply all \( V_{\mu,i} \) in Eq. (2) to the subsystem \( i \) to create a basis, this leads to \( B_i = \{ V_{\mu,i} \}_{\mu=1}^{M_{\text{int}}} \). For the particle conserving approach, we apply the excitation \( a_1^\dagger \) or the de-excitation operators \( a_s \) to every qubit \( s \) of the \( n_i \) qubits. Additionally, to capture effects inside the subsystems, we applied swap gates between different qubits belonging to the same subsystem to create a basis. Therefore, in this approach, \( B_i = \{ a_s \}_{s=1}^{n_i} \cup \{ a_s^\dagger \}_{s=1}^{n_i} \cup \{ \text{SWAP}_{s,s'} \}_{s,s'=1}^{n_i} \).

We also use additional low lying energy eigenstates \( | G_i^c \rangle \) to generate the basis sets instead of using only \( | G_i \rangle \) as has been done in the original paper [13]. The superscript \( c \) marks the \( c \)-th excited state and we define \( | G_i^c \rangle := | G_i \rangle \). Note that if the subsystem Hamiltonian has degenerate ground states, \( | G_i^c \rangle \) does not necessarily have different energy than \( | G_i \rangle \). Such excited states can be constructed with an algorithm like the subspace-search VQE [17]. After applying \( B_i \) to \( \{ | G_i^c \rangle \}_{c=1}^{\ell} \), we use the Gram-Schmidt algorithm to ensure that the created basis is orthogonal and minimal with \( K_i \) elements.

The number of basis vectors determines how many qubits we need to run the deep VQE algorithm. We, therefore, also examine different approaches to reduce the dimensionality of the basis. For the interaction-based methods, we only consider interactions between subsystems with \( \lambda^\mu \) exceeding a certain threshold \( \varepsilon > 0 \). In
TABLE I. Description of basis sets with starting vector and excitations. \( A_i \in \{1, ..., n_i \} \) is a set of orbitals evolved in interactions with largest \( \lambda^\mu \). In Interactions fix \( \epsilon \) is selected such that the generated basis set is of fixed dimension.

| Method | Start vectors | Applied excitation \( \mathbf{r}_{i,j} \) |
|--------|---------------|--------------------------------------|
| Interactions | \( |G_i^0\rangle \) \( \forall s, s' \in \{1, ..., n_i\} \) \( \forall \mathbf{r}_{i,j} \in V_{i,j} \) | \( \sum_{\mu = 1}^{O} \lambda^\mu V_{\mu,i} \) |
| Interactions & excited | \( |G_i^1\rangle \) \( \forall s, s' \in \{1, ..., n_i\} \) \( \forall \mathbf{r}_{i,j} \in V_{i,j} \) | \( \sum_{\mu = 1}^{O} \lambda^\mu V_{\mu,i} \) |
| Interactions fix | \( |G_i^0\rangle \) \( \forall s, s' \in \{1, ..., n_i\} \) \( \lambda^\mu > \epsilon \) | \( \sum_{\mu = 1}^{O} \lambda^\mu V_{\mu,i} \) |
| Single Pauli | \( |G_i^0\rangle \) \( P_\mu s \in \{1, ..., n_i\} \) | \( \sum_{\mu = 1}^{O} \lambda^\mu V_{\mu,i} \) |
| Single Pauli & excited | \( |G_i^1\rangle \) \( \forall s, s' \in \{1, ..., n_i\} \) | \( \sum_{\mu = 1}^{O} \lambda^\mu V_{\mu,i} \) |
| Particle conserving | \( |G_i^0\rangle \) \( \forall s, s' \in \{1, ..., n_i\} \) \( s, s' \in \{1, ..., n_i\} \) \( \forall \mathbf{r}_{i,j} \in V_{i,j} \) | \( \sum_{\mu = 1}^{O} \lambda^\mu V_{\mu,i} \) |
| Single/Particle conserving edge | \( |G_i^1\rangle \) \( \forall s, s' \in \{1, ..., n_i\} \) \( s, s' \in \{1, ..., n_i\} \) \( \forall \mathbf{r}_{i,j} \in V_{i,j} \) | \( \sum_{\mu = 1}^{O} \lambda^\mu V_{\mu,i} \) |

The particle conserving and single Pauli methods as well can produce basis sets with a dimensionality that is challenging to represent on a quantum machine. We, therefore, also examine methods to truncate their basis sets. Our approach aimed to reduce the number of interaction terms between the subsystems for the molecular Hamiltonian.

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The numerical experiments that follow, we examine the results for \( \epsilon = 10^{-2} \) if not stated otherwise. This selection was necessary since otherwise, the interaction-based methods would lead to basis sets that resemble no reduction from the entire Hilbert space. This was caused due to the numerous interaction terms between the subsystems for the molecular Hamiltonian.

We apply the deep VQE to two different tree-like molecules in Fig. 1. Nakatani and Chan have used this type of molecule for benchmarking tree tensor network [19]. Since a tree-like molecule naturally separates into different branch-like subsystems, it is also an ideal benchmark molecule for the deep VQE. These molecules are the first step to calculating dendrimers or Cayley tree-like molecules. Today’s industry frequently uses dendrimers, e.g., for pharmaceuticals [20]. Understanding dendrimers better would allow us to tailor them specifically to the applications. We describe the geometry of the tree-like molecules in the Appendix A. We repeated the deep VQE simulation multiple times, enlarging the distance between atoms with a stretching factor ranging from 0.9 to 2.0.

Using STO-3G minimal basis set, the molecules in Fig. 1a and Fig. 1b respectively have 20 and 26 spin-orbitals. The Hamiltonians of the molecules are obtained with PySCF [21]. We map it to a qubit system by JW transformation [22] implemented in OpenFermion [23], resulting in 20- and 26-qubit Hamiltonians. We use the full

not find a unique ground state. The found ground state would be semi-random and depends on multiple factors, such as the starting condition of the VQE and the noise in the quantum device. In principle, all linear combinations of the degenerate ground states are possible solutions to the VQE without any treatment. This results in an instability of the deep VQE, since the random starting vectors, in general, leads to different basis sets and consequently to different \( H^{\text{eff.}} \).

This dependency on the starting vector set is tricky as, without supplementary knowledge about the system, there is no justification for favoring one sub-vector set over the other without running it through the deep VQE first. Therefore, the deep VQE must consider all degenerated vectors as starting vectors to ensure that the result is independent of the randomness of the first VQE. Alternatively, we should design the first VQE to return a unique solution for the reproducibility of the experiment by, for example, unfolding the degeneracy with constrained VQE [18].

**IV. NUMERICAL SIMULATION**

**A. Setup**

We apply the deep VQE to two different tree-like molecules in Fig. 1. Nakatani and Chan have used this type of molecule for benchmarking tree tensor network [19]. Since a tree-like molecule naturally separates into different branch-like subsystems, it is also an ideal benchmark molecule for the deep VQE. These molecules are the first step to calculating dendrimers or Cayley tree-like molecules. Today’s industry frequently uses dendrimers, e.g., for pharmaceuticals [20]. Understanding dendrimers better would allow us to tailor them specifically to the applications. We describe the geometry of the tree-like molecules in the Appendix A. We repeated the deep VQE simulation multiple times, enlarging the distance between atoms with a stretching factor ranging from 0.9 to 2.0.

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configuration interaction (FCI) solution of the molecules as the reference. Additionally, we show the energy of the “combined subsystem” solution, which is the product state of all individual subsystem solutions, as well as the restricted Hartree-Fock solution. Hartree-Fock is a standard mean-field method and is not able to account for the electron-correlation energy in the molecule. In the case of the 10-atom molecule, the ground state is a quintet, whereas, in the 13-case, the ground state is a doublet. For simulating the quantum states of these molecules on a classical machine, we use the python library Qulacs [24].

Fig. 1 shows the separation of the tree-like molecule into subsystems. We use localized orbitals to define the subsystems based on the distance between their associated atoms. The Löwdin orthogonalization method is employed to create well-localized orbitals.

The performance of deep VQE depends partially on the subroutine VQE algorithm employed in it. However, this study aims to compare the different basis creation methods. Therefore we replaced the VQE subroutines with direct diagonalizations of the operators. This replacement assures us that we find accurate eigenstates of the systems. Additionally, the direct diagonalization replacement allows us to compare the different basis sets without the additional effects a realistic VQE algorithm would add, such as the noise of the quantum system. We find that the ground state of each subsystem of the 10-atom molecule (Fig. 1a) has a two-fold degeneracy for all stretching factors. For the 13-atom molecule (Fig. 1b), each subsystem, except for the central hydrogen atom, has a unique ground state, but their first excited states are three-fold degenerated. We also consider how this degeneracy affects the overall result in Sec. IV B. The ground state solution of the subsystems as well as for the \( H^{\text{eff}} \) was determined using the exact diagonalization with SciPy [25]. The following subsections show the energy difference between the different basis sets and the FCI solution for the 10 and 13-atom molecules. To better compare the different methods, we also show the weighted mean error of correlation energies over the different stretching factors and the required number of qubits to run the deep VQE on a quantum machine. We define the weighted mean error of correlation energies as,

\[
\text{Weighted mean error of correlation energies} = \sum_{x \in X} \frac{1}{|X|} \frac{E(x) - E_{\text{FCI}}(x)}{E_{\text{subsystems}}(x) - E_{\text{FCI}}(x)},
\]

where \( E(x) \), \( E_{\text{FCI}}(x) \), and \( E_{\text{subsystems}}(x) \) are the energy obtained by the deep VQE using particular strategies, the FCI energy, and the energy of the "combined subsystem" solution at a stretching factor \( x \). \( X \) denotes the set of stretching factors for which we perform the calculations. Here, \( X \) is \( \{0.9, 1.0, 1.1, 1.2, 1.3, 1.4, 2.0\} \). We applied the weight \( E_{\text{subsystems}}(x) - E_{\text{FCI}}(x) \) to the correlation energy error to take into account the difference in magnitude over the stretching factors.

B. Influence of starting vector for degenerated subsystems

First, we discuss the dependency of the deep VQE on the starting vector. For the sake of readability, we only show the results using the particle conserving method to produce the basis in the deep VQE. We show the results for the 10-atom molecule in Fig. 2. Due to spin symmetry, the ground states of all the subsystems of the 10-atom molecule are doubly degenerated. The ground state found by the VQE or, in our case, the direct diagonalization is therefore not unique. We use the spin state to distinguish the degenerate states. We mark the spin-up solution with \( \uparrow \) and the spin-down solution with \( \downarrow \). We only show a selection of all possible spin configurations due to redundancy resulting from the symmetry in the molecule. We observe two distinct resulting energy levels, indicating that the stating vector can considerably influence which Hilbert space can be searched for the overall ground state and, therefore, the performance of the deep VQE.

The choice of the starting vector is not trivial, and without further knowledge of the system, all possible starting vectors have to be considered. Additional starting vectors, however, come at the cost of requiring additional qubits to represent \( H^{\text{eff}} \).

C. Different Basis methods

Next, we compare the performance of different basis methods with different starting vectors introduced in Sec. III. We compare two choices of starting vectors. The first choice is using single ground states of each subsystem, whereas the second is using additional eigenstates of each subsystem. In the former case, we chose the spin configuration \( \downarrow \downarrow \downarrow \downarrow \) for the 10-atom molecule. In the case of the 13-atom molecule, only the central subsystem had a two-fold degeneracy, and we chose the spin \( \downarrow \) solution as well. For the interaction method, an \( \epsilon = 10^{-2} \) was
chosen for the 10-atom molecule and an $\epsilon = 10^{-3}$ for the 13-atom molecule.

Figures 4 and 5 shows the corresponding results for the 10-atom system. We also show the number of qubits needed for each method in Tab. III. We note that we are unable to perform the calculation for the particle conserving method with additional starting vectors due to its sizeable computational requirement. The particle conserving methods with only one starting vector per subsystem performed exceptionally well in terms of accuracy. For all tested basis creation methods, including an additional eigenstate significantly increased the accuracy of the deep VQE. This is a consequence of the expanded Hilbert space that can be explored.

Unlike the other methods, we observe that the interaction methods with a fixed truncation have a decreasing accuracy with increasing stretching factors. More concretely, the energy obtained by interaction methods jumps to the combined subsystem’s energy at the stretching factor of 1.4. The interaction methods also display different behavior regarding the saved qubits, showing a significant dependence on the stretching factor as shown in Tab. III. This is due to our selection process of the interactions to generate a basis set. We set a fixed cut-off of the interaction strength $\lambda^p$ for the interactions involved in the basis creation step. Consequently, as the interactions between subsystems become weaker by stretching the molecule, more interactions are disregarded. Lowering the dimensionality of the basis sets allows us to save more qubits but comes at the cost of lowering the method’s accuracy.

Fig. 6 shows the results for the 13-atom molecule. Here, only the central subsystem was doubly degenerated, and the first excited state of the branch subsystems for the 13-atom molecule is a triplet. To see the effect of adding additional excited states as starting vectors on the performance of the deep VQE, we considered all three states as additional starting vectors. Due to the high computational cost for the other methods, only the single Pauli edges and the particle conserving edges methods could be prepared with additional starting vectors. For both methods, the additional starting vectors were able to increase the accuracy of the deep VQE, but this came at the expense of requiring extra qubits (see Tab. IV for the exact number of qubits required for each method). The particle conserving edges methods performed slightly better than others. This is consistent with the case of the 10-atom molecule, indicating an advantage of using the particle conserving approach in deep VQE for chemistry problems.

D. Interaction based deep VQE with fixed qubit numbers

Finally, we show the results for the fixed qubit numbers approach in Figs. 8 to 11. We fix the number of qubits to 11, 14, and 17. For the 13-atom molecule, we
TABLE III. Resources needed to represent the $H_{\text{eff}}$ of the 10-atom molecule for the different basis creation methods. First number indicates the needed qubits and the values in the bracket are the individual dimension of the subsystem basis sets.

| Stretching factor                  | 0.9 | 1.0 | 1.1 | 1.2 | 1.3 | 1.4 | 2.0 |
|------------------------------------|-----|-----|-----|-----|-----|-----|-----|
| Interaction(1111)($\lambda > 10^2$) | 17  | 14  | 11  | 7   |     |     |     |
| Interaction(2222)($\lambda > 10^2$) | 17  | 14  | 10  |     |     |     |     |
| Particle conserving(1111)          | 17  |     |     |     |     |     |     |
| Particle conserving edges(1111)    | 11  |     |     |     |     |     |     |
| Particle conserving edges(2222)    | 14  |     |     |     |     |     |     |
| Single Pauli(1111)                 | 17  | 14  |     |     |     |     |     |
| Single Pauli edges(1111)           | 11  |     |     |     |     |     |     |
| Single Pauli edges(2222)           | 14  |     |     |     |     |     |     |

TABLE IV. The numbers of qubits needed to represent the $H_{\text{eff}}$ of the 13-atom molecule for the different basis creation methods. First number indicates the needed qubits and the values in the bracket are the individual dimension of the subsystem basis sets.

| Stretching factor                  | 0.9 | 1.0 | 1.1 | 1.2 | 1.3 | 1.4 | 2.0 |
|------------------------------------|-----|-----|-----|-----|-----|-----|-----|
| Interaction(1111)($\lambda > 10^3$) | 17  | 11  | 7   |     |     |     |     |
| Particle conserving edges(1111)    | 11  |     |     |     |     |     |     |
| Particle conserving edges(2444)    | 17  |     |     |     |     |     |     |
| Single Pauli(1111)                 | 17  |     |     |     |     |     |     |
| Single Pauli edges(1111)           | 11  |     |     |     |     |     |     |
| Single Pauli edges(2444)           | 17  |     |     |     |     |     |     |
FIG. 4. The energy difference between FCI and the deep VQE for the 10-atom molecule. We compare the different basis creation methods when using a single starting vector marked as (1111) and a complete degenerate basis set indicated by (2222). The first digit indicates the number of starting vectors for the central subsystem, where the three following digits count the starting vectors for the branch-like subsystems.

FIG. 6. The energy difference between FCI and the deep VQE for the 13-atom molecule. We compare the different basis creation methods when using a single starting vector marked as (1111) and a full degenerate basis set indicated by (2444). The first digit indicates the number of starting vectors for the central subsystem, where the three following digits count the starting vectors for the branch-like subsystems.

FIG. 5. Weighted mean error of correlation energies for all methods for the 10-atom molecule.

FIG. 7. Weighted mean error of correlation energies for all methods for the 13-atom molecule.
FIG. 8. The energy difference between FCI and the deep VQE for the 10-atom molecule. We compare the interaction fixed methods with a different number of qubits. The fully degenerate basis set is indicated by (2222). The first digit indicates the number of starting vectors for the central subsystem, where the three following digits count the starting vectors for the branch-like subsystems.

FIG. 9. Weighted mean error of correlation energies for the interaction based method with fixed number of qubits for the 10-atom molecule.

FIG. 10. The energy difference between FCI and the deep VQE for the 13-atom molecule. We compare the interaction fixed methods with a different number of qubits. The fully degenerate basis set is indicated by (2444). The first digit indicates the number of starting vectors for the central subsystem, where the three following digits count the starting vectors for the branch-like subsystems.

FIG. 11. Weighted mean error of correlation energies for the interaction based method with fixed number of qubits for the 13-atom molecule.
could not produce an 11-qubit version as already including only the strongest interaction strength results in a basis set requiring more than 11 qubits. The qubit number is increased by three per step to take into account that there are three equivalent branch-like subsystems of the molecules.

In Figs. 8 and 10, we see that if we define a fixed number of qubits, we can avoid the decreasing accuracy for increasing stretching factors. However, if the number of qubits is too restrictive, we cannot improve the result from the "combined subsystems". We see this behavior for the interaction(2222) method with 11 qubits for the 10-atom molecule. Another interesting point is that if the number of qubits is limited, the deep VQE performs better using only single starting vectors. We expect this because the method with a single starting vector can consider more interactions in the basis creation step with the same number of qubits compared to when we include multiple starting vectors.

E. Performance comparison

Overall we achieved similar accuracy for the 10 and the 13-atom molecules (see Figs. 4 and 6). We expect this from the minor influence the newly added outer atoms have on the overall ground state. This behavior would also explain the remarkable success the edge methods provide, saving up to 15 qubits with comparable accuracy to the other methods Tab. IV. All methods were able to produce lower energies than the Hartree-Fock method. Multiple methods were able to approximate the ground state energies within an error of below 1% of the electron correlation energy of the molecule. The electron correlation energy is equivalent to the error of the Hartree-Fock method shown in Figs. 5 and 6 and represents the error due to the mean-field approximation of Hartree-Fock. This improvement helped us reach chemical accuracy for some stretching factors. The particle conserving edges method performed exceptionally well. Using an edge method allows us to focus on changes affecting the orbitals involved in the most substantial interaction. We expect electrons occupying such orbitals to experience the most dramatic changes from the individual subsystem solutions when forming a bond with the other subsystems. To use the computational resources offered most effectively, it is advisable to use basis sets that focus on exploring the changes to the occupation of the most involved orbitals.

In contrast to the other methods, the interaction methods with a fixed truncation for the participating interactions significantly depended on the stretching factor. The number of qubits decreased with increasing distance. This reduction allows us to save more and more qubits but comes at the cost of decreasing accuracy. However, we consider this is not preferable if we have access to a quantum device with a fixed number of qubits. It seems unreasonable to not use all of them. The fixed number of qubits method we propose in this paper seems a more reasonable approach as it allows the use of all qubits. It also performs more stable for less interacting systems not suffering from decreasing accuracy.

V. CONCLUSION AND DISCUSSION

The deep VQE approach successfully reduced the number of qubits to calculate a ground state of a complex chemical molecule. All different basis creation methods we tested could create a ground state energy within a few mHa of the FCI solution. However, they showed a significant difference in the accuracy and the number of qubits they could save. We discussed the challenges of degenerated subsystems and provided a solution in the form of additional starting vectors or marking the single starting vector by its spin state. We also showed that adding additional low-lying states as starting vectors for the basis-creation method can improve the accuracy. However, the approximation of such states can be costly. Therefore, further research is needed to determine if this accuracy can be achieved using other basis creation methods. A possible approach would be to consider double Pauli excitation compared to the here used single Pauli method. An additional exciting way to think about a new basis creation method is their similarity to VQE ansatzes. A new basis creation method could be created using a VQE ansatz with discrete fixed parameters. We also proposed methods to upper bound the number of qubits by using edge methods or selection of interaction in the interaction-based approach. These methods have proven to be effective strategies to reduce the number of needed qubits for the deep VQE. With this modifications to the deep VQE and the right basis set, we believe that the use of deep VQE in a quantum chemistry setting can be highly beneficial.

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Appendix A: Geometry of molecule

In Tabs. V and IV, we show the geometry of the molecules used in this work. The distances are given in angstrom.

TABLE V. XYZ-coordinate of the 10 hydrogen molecule with stretching factor 1. This molecule’s geometry is the same as described in the appendix of [19].

| X            | Y            | Z            |
|--------------|--------------|--------------|
| H 0.000000   | 0.000000     | 0.000000     |
| H -1.732051  | -0.000000    | -1.000000    |
| H -1.848076  | -0.866025    | -2.799038    |
| H -3.348076  | 0.866025     | -0.200962    |
| H 0.000000   | 0.000000     | 2.000000     |
| H -1.500000  | -0.866025    | 3.000000     |
| H 1.500000   | 0.866025     | 3.000000     |
| H 1.732051   | 0.000000     | -1.000000    |
| H 3.348076   | -0.866025    | -0.200962    |
| H 1.848076   | 0.866025     | -2.799038    |

Table VI. XYZ-coordinate of the 13 hydrogen molecule with stretching factor 1.

| X            | Y            | Z            |
|--------------|--------------|--------------|
| H 0.000000   | 0.000000     | 0.000000     |
| H -1.732050  | -0.000000    | -1.000000    |
| H -1.848080  | -0.866030    | -2.799040    |
| H -3.348080  | 0.866030     | -0.200960    |
| H -3.464100  | -0.000000    | -2.000000    |
| H 0.000000   | 0.000000     | 2.000000     |
| H -1.500000  | -0.866030    | 3.000000     |
| H 1.500000   | 0.866030     | 3.000000     |
| H 0.000000   | 0.000000     | 4.000000     |
| H 1.732050   | 0.000000     | -1.000000    |
| H 3.348080   | -0.866030    | -0.200960    |
| H 1.848080   | 0.866030     | -2.799040    |
| H 3.464100   | 0.000000     | -2.000000    |

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