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Advantages of discrete variable representation in variational quantum eigensolvers for vibrational energy calculations

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While quantum computing algorithms have been widely applied for electronic structure calculations, applications to molecular dynamics remain scarce. Complex and varied landscapes of molecular potential energy surfaces give rise to vibrational states with a wide range of properties, making it difficult to construct a general representation of ro-vibrational states by a quantum computer with a limited number of qubits and gates. Another challenge is the exponential growth of the computational complexity – for example, the number of terms required to expand a general Hamiltonian in Pauli strings increases exponentially with the number of qubits. Here, we show that discrete variable representation (DVR) can be leveraged to represent molecular Hamiltonians by the polynomial (in the number of qubits) number of quantum circuits. We then demonstrate that DVR Hamiltonians lead to very efficient quantum ansätze for vibrational states. For this purpose, we develop a compositional quantum ansatz search that adapts gate sequences in variational quantum eigensolvers (VQE) to a specific molecular state. We apply VQE to compute the vibrational energy levels of Cr₂ in seven electronic states, as well as those of van der Waals complexes Ar–HCl and Mg–NH. Our numerical results show that an accuracy of 1 cm⁻¹ can be achieved by very shallow quantum circuits with 2 to 9 entangling gates.

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1. Introduction

Accurate calculation of molecular properties is considered a promising application of quantum computing. The eigenstates of molecular Hamiltonians can be obtained on quantum computers by variational quantum eigensolvers (VQEs)^{1,2} that employ sequences of gates operating on qubits (quantum circuits) to prepare quantum states tailored for specific problems. VQEs have been applied for solving the electronic structure problem for molecules^{3–11} and lattice models.^{1,7,12} However, applications of VQEs to computations of ro-vibrational energies and states have been limited.^{13–20} Ref. 16 and 17 demonstrated a general approach for computing ro-vibrational energy levels of polyatomic molecules inspired by previous work on the electronic structure. However, the methods used in ref. 16 and 17 require extended quantum circuits including a large number of entangling gates. This makes the applications of VQEs to vibrational energy calculations challenging for quantum computers limited by errors and noise. The errors grow with the circuit size and make large quantum circuits impractical for implementation in current quantum

devices. This challenge is compounded by transpilation of quantum circuits into hardware-specific gate sequences, which often further extends the size of the quantum circuits.

A central goal for quantum computing of molecular dynamics can thus be formulated to develop a general approach that (i) is applicable to a broad range of molecules with widely varying ro-vibrational states, from deeply bound to van der Waals states; (ii) yields high accuracy with shallow quantum circuits; and (iii) exhibits at most a polynomial scaling with the dimensionality of the molecular configuration space or, alternatively, in the number of qubits if binary basis encoding is used. Within the VQE framework, requirement (i) demands an ansatz for quantum circuits that can represent molecular states across widely varying landscapes of potential energy surfaces; requirement (ii) is necessary due to the noise and hardware limitations of current quantum computers; and requirement (iii) is essential for realizing the quantum advantage over classical methods.

Here, we explore an approach for computing the vibrational energy levels of molecules with VQEs based on discrete variable representation (DVR) of molecular Hamiltonians.^{21,22} We first derive the theoretical bounds on the quantum measurement complexity with VQE based on Fourier grid DVR. Our analysis shows that the structure of the DVR matrices can be exploited to reduce the expansion of the molecular Hamiltonians in

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measurement operators to the desired polynomial scaling in the number of qubits. In the second part of this work, we explore an automated construction of the quantum circuits for the vibrational energy computations. Our goal is to build quantum circuits, yielding accurate VQE results, without any constraints on the quantum ansatz. Our results demonstrate that DVR Hamiltonians lead to very efficient (small number of qubits and gates) quantum circuits for representing vibrational states of molecules by states of a quantum computer. To probe the minimal gate count for accurate vibrational VQE computations, we develop a compositional search algorithm that incrementally grows quantum circuits to identify optimal sequences in the space of gate permutations.

To illustrate the generality of this approach and the efficiency of the resulting quantum circuit representations of vibrational states, we consider Cr₂ in seven different electronic states²³ and van der Waals complexes Ar-HCl(¹Σ) and Mg-NH(³Σ). These molecular systems exhibit vibrational states with widely different energies (from -55 to -15 000 cm⁻¹ from the dissociation threshold) and spatial variations of wave functions and energy level patterns. Our compositional search yields quantum circuits that produce VQE results with accuracy <1 cm⁻¹ for ground and excited vibrational energy levels, illustrating the ability of VQEs to compute the rotational constants and vibrational anharmonicity, with between 2 and 9 entangling gates, for diatomic and triatomic molecules. For reference, previous VQE calculations of vibrational energy levels required extended quantum circuits with >200 (for CO, COH and O₃ molecules¹⁷) or between 44 and 140 292 (for CO₂, H₂CO and HCOOH molecules¹⁶) entangling gates.

The remainder of this article is organized as follows. After a brief introduction of VQEs, section 2.1 presents an algorithm to evaluate the DVR Hamiltonians with a polynomial number of measurements. Section 2.2 describes the algorithm for the compositional ansatz optimization, which is followed by numerical results illustrating the efficiency and accuracy of the optimized Ansatz for Cr₂ in seven different electronic states²³ and van der Waals complexes Ar-HCl(¹Σ) and Mg-NH(³Σ) in section 3. The work is summarized in section 4.

II. Theory

In VQEs, a quantum computer estimates the expectation value $\langle \psi(\boldsymbol{\varphi}) | \hat{H} | \psi(\boldsymbol{\varphi}) \rangle$, which is minimized by varying $\boldsymbol{\varphi}$ to yield the lowest eigenvalue $E_{i=0}$ and an approximate representation of the corresponding eigenvector of \hat{H} . This method can be extended to compute excited states by optimizing²

$$\tilde{\boldsymbol{\varphi}}_v = \operatorname{argmin}_{\boldsymbol{\varphi}} \left[\langle \psi(\boldsymbol{\varphi}) | \hat{H} | \psi(\boldsymbol{\varphi}) \rangle + \sum_{i=0}^{v-1} \beta_i \langle \psi(\tilde{\boldsymbol{\varphi}}_i) | \psi(\boldsymbol{\varphi}) \rangle \right] \quad (1)$$

where $\beta_i \geq E_{i+1} - E_i$ and $\tilde{\boldsymbol{\varphi}}_i$ denotes an optimal solution for the corresponding quantum state.

The quantum states $|\psi(\boldsymbol{\varphi})\rangle$ are obtained by quantum circuits acting on qubits, and the Hamiltonian is expanded in quantum

operators. Most generally,

$$\hat{H} = \sum_{i=0}^{4^n} A_i K_1^i \otimes K_2^i \dots \otimes K_n^i \quad (2)$$

where n is the number of qubits, $K_j^i \in \{\sigma_x, \sigma_y, \sigma_z, I\}$ acting on qubit j , $\{\sigma_i\}$ are the Pauli matrices, and I is the identity matrix,

$$A_i = \frac{1}{2^n} \operatorname{Tr}[(K_1^i \otimes K_2^i \dots \otimes K_n^i) \cdot \mathbf{H}] \quad (3)$$

and \mathbf{H} is the Hamiltonian matrix in some basis. The computational complexity is determined by the number of non-zero terms in eqn (2), which for a general matrix is 4^n ; and the complexity of the quantum circuits yielding $|\psi(\boldsymbol{\varphi})\rangle$. In Section II.A, we show that the structure of DVR matrices allows an efficient quantum circuit representation of \hat{H} , scaling with n as $\operatorname{poly}(n)$.

A. Efficient measurement of DVR Hamiltonians

DVR is a finite basis representation, in which the coordinate operators (and consequently the potential energy) are diagonal. The DVR matrix of kinetic energy is not sparse. However, it has specific structure that is exploited here. We use the DVR introduced by Colbert and Miller²¹. Other DVR bases can be reduced to those in ref. 21 by coordinate transformations. As follows from ref. 21, the Hamiltonian matrix in the DVR representation has the following structure:

$$H_{ij} = \begin{cases} d(i), & i = j, \\ f(|i-j|) + g(i+j), & i \neq j \end{cases} \quad (4)$$

We use a number encoding to map the DVR basis states onto qubit states, where the indices of the flattened DVR basis states are mapped into their binary representation on the qubits. This allows VQE to compute the eigenvalues of \mathbf{H} of size $2^n \times 2^n$ using n qubits. As the number of DVR bases required for accurate results grows exponentially with the number of degrees of freedom of the system, the number of qubits necessary to represent the system grows polynomially with the size of the system.

As shown in Appendix A, the functions f and g in eqn (4) satisfy

$$\sum_{k=s}^{2^n-1} |f(k)| \leq O(s^{-\alpha}), \quad \alpha > 0, \quad (5)$$

$$\sum_{k=r}^{2^{n+1}-1-r} |g(k)| \leq O(r^{-\beta}), \quad \beta > 0, \quad (6)$$

for $1 \leq r \ll N = 2^n$ and $1 \leq s \ll N = 2^n$, where N is the number of DVR states, assumed to be large.

Given a quantum state $|\psi\rangle$, our aim is to show that

$$\tau = \langle \psi | \hat{H} | \psi \rangle + O(\epsilon) \quad (7)$$

can be measured with the number of quantum circuits that scales polynomially with n and $1/\epsilon$. We first use eqn (5) and (6)



to show that the expectation value (7) can be approximated as

$$\tau = \langle \psi | \hat{H}^{(s,r)} | \psi \rangle + O(\epsilon) \quad (8)$$

where $\hat{H}^{(s,r)}$ employs a truncated DVR matrix with the following elements:

$$H_{ij}^{(s,r)} = \begin{cases} d(i) & i = j, \\ f(|i-j|) + g(i+j) & i \neq j, |i-j| < s, i+j < r, \\ 0 & \text{otherwise,} \end{cases} \quad (9)$$

for

$$s \sim \epsilon^{-1/\alpha} \quad (10)$$

and

$$r \sim \epsilon^{-1/\beta}. \quad (11)$$

To prove eqn (8), we consider operators defined as a k th (anti)-diagonal matrix, with elements

$$K_{ij}^{k\pm\pm} \equiv \delta_{i\pm j, \pm k}, \quad (12)$$

where only $++$, $+-$ and $--$ combinations of signs are used. Since $K^{k\pm\pm}$ either permutes or shifts vector components, while either preserving the norm of vectors or truncating vectors in a finite Hilbert space, $\|K^{k\pm\pm}\psi\|_2 \leq \|\psi\|_2$ (and the inequality is tight). This can be used in the Cauchy-Schwarz inequality $|\langle \psi, K^{k\pm\pm}\psi \rangle| \leq \|\psi\|_2 \|K^{k\pm\pm}\psi\|_2$ to bound the expectation value of $K^{k\pm\pm}$ as

$$|\langle \psi, K^{k\pm\pm}\psi \rangle| \leq \|\psi\|_2^2. \quad (13)$$

Using the triangle inequality and $\|\psi\|_2^2 = 1$, the approximation error can be bounded by

$$\begin{aligned} & \left| \langle \psi | (\hat{H} - \hat{H}^{(s,r)}) | \psi \rangle \right| \\ &= \left| \langle \psi | \left(\sum_{k=s}^{2^n-1} f(k) [K^{k--} + K^{k--}] + \sum_{k=r}^{2^{n+1}-1-r} g(k) K^{k++} \right) | \psi \rangle \right| \\ &\leq \left(2 \sum_{k=s}^{2^n-1} |f(k)| + \sum_{k=r}^{2^{n+1}-1-r} |g(k)| \right) \end{aligned} \quad (14)$$

Due to eqn (5), (6) and (8), the relationship holds for s and r given by eqn (10) and (11).

We next seek to transform $|\psi\rangle$ by short-depth \hat{V}_i , so that

$$\tau = \sum_{i=1}^{\text{poly}(n,1/\epsilon)} \sum_{j=1}^{2^n} w_{ij} \left| \langle \psi | V_i^\dagger | j \rangle \right|^2 \quad (15)$$

where j is the index that enumerates the states of n qubits in the computational Z basis. We leverage eqn (8) to decompose τ into contributions from a diagonal matrix (D), $s \approx \epsilon^{-1/\alpha}$ diagonal

bands (t^k) and $r \approx \epsilon^{-1/\beta}$ anti-diagonal components (a^k), as follows:

$$\begin{aligned} \tau &= \langle D \rangle + \sum_{k=1}^{\left\lceil \epsilon^{-\frac{1}{\alpha}} \right\rceil} f(k) \langle t^{k[n]} \rangle \\ &+ \sum_{k=1}^{\left\lceil \epsilon^{-\frac{1}{\beta}} \right\rceil} \left(g(k) \langle a^{k[n]} \rangle + g(2^n - k) \langle a^{(2^n-k)[n]} \rangle \right) + O(\epsilon). \end{aligned} \quad (16)$$

We note that the construction of $t^{k[n]}$ and $a^{k[n]}$ described below produces contributions to the main diagonal upon measurements. This is offset by a corresponding change in the diagonal matrix elements, as specified in Appendix B. The computation of the diagonal contribution $\langle D \rangle$ is classically efficient. In this case, one can set $\{V_i\} = \{1\}$ and combine $w_{1i} = D_{ii}$ from Appendix B with the weights given by the measurements in the computational basis.

To construct $t^{k[n]}$, we note that $l \equiv \lceil \log_2(k+1) \rceil$ is the smallest number of qubits that allows the k -th band. We consider 1-qubit entangled states produced from the Z basis by a sequence of CNOT gates shown in Fig. 1 (left panel). The resulting state is

$$|j, p\rangle_l = \frac{|j\rangle + |p\rangle}{\sqrt{2}}, \quad (17)$$

where $|j\rangle = \otimes_i^l |j_i\rangle$ and $|p\rangle = \otimes_i^l |p_i\rangle$ are the binary representations of the row (p) and column (j) indices of the DVR matrix, with $p = j + k$, and j_i and p_i representing single-qubit states. The projection of an n -qubit state onto the $|j, p\rangle_l$ state has 2 non-zero off-diagonal matrix elements of magnitude $1/2$. The binary representation of the row and column positions of these off-diagonal elements differ by l bits. The position of the non-zero off-diagonal elements is controlled by the position of the CNOT gates in Fig. 1. Thus, the expectation value of $t^{k[l]}$ can be obtained by measurements with, at most, k l -qubit entangled states.

We now observe that $t^{k[m+1]} = \mathbb{1}_2 \otimes t^{k[m]} + \gamma_{k[m]}$, where $\gamma_{k[m]}$ represents k pairs of elements missing from the middle of the tensor product matrix (shown as pluses in Fig. 1, right panel). These elements can be directly targeted by additional measurements in an entangled basis, analogous to states in eqn (17) but with a different number q of qubits,

$$|j, p\rangle_{q \in [l,m]} = \frac{|j\rangle + |p\rangle}{\sqrt{2}}, \quad (18)$$

where $|j\rangle = \otimes_i^q |j_i\rangle$ and $|p\rangle = \otimes_i^q |p_i\rangle$. These states can be constructed in the same way as states (17). The missing elements can thus be obtained with, at most, k measurements if performed element-wise.

The full algorithm to compute $t^{k[n]}$ thus includes: (i) at most $2^l - k \leq k$ expectation values *via* circuits of depth $\leq l + 1$ to obtain $t^{k[l]}$ (yielding elements represented by open circles in Fig. 1); (ii) filling the gap (pluses in Fig. 1) in $\mathbb{1}_2 \otimes t^{k[m]}$ to



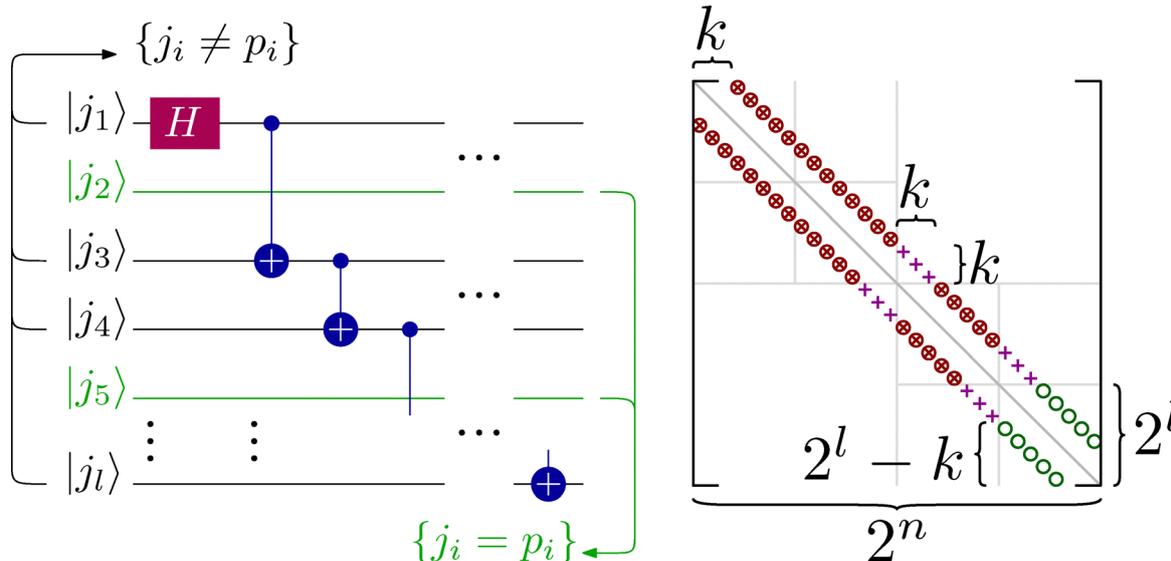


Fig. 1 Left panel: Preparation of state (17) from the computational basis. H denotes the Hadamard gate and the circles – the CNOT gates. Right: Measuring $t^{k[m]}$ that includes matrix elements on the main and k th diagonals. The open circles \otimes are obtained from projections of (17), \otimes – via $\mathbb{1}_2 \otimes t^{k[m]}$, and $+$ – by additional measurements in the entangled basis.

produce $t^{k[m+1]}$, which requires measurements of at most k projections on entangled states *via* circuits of depth at most $\lceil \log_2 2k \rceil$; (iii) a total of $n - l$ iterations to fill the entire band. The total number of measurement circuits for $t^{k[m]}$ is thus

$$\text{Comp}(t^{k[m]}) < (n + 1 - \log_2 k)k. \quad (19)$$

To construct the anti-diagonals $a^{k[l]}$, we first note that the expectation values of $a^{1[1]} = \begin{bmatrix} 1 & 0 \\ 0 & 0 \end{bmatrix}$, $a^{2[1]} = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix}$ and $a^{3[1]} = \begin{bmatrix} 0 & 0 \\ 0 & 1 \end{bmatrix}$ can be obtained by sampling one-qubit measurements of ψ in Z to get $\langle a^{1[1]} \rangle_\psi$ and $\langle a^{3[1]} \rangle_\psi$ and in X to get $\langle a^{2[1]} \rangle_\psi$. The elements $a^{k[m+1]}$ can be obtained by combining $\mathbb{1}_2 \otimes a^{k[m]}$ and $a^{k[m]} \otimes \mathbb{1}_2$, which at most doubles the number of measurements.²⁴

We limit the construction to $r = \log_2 \epsilon / \beta$ anti-diagonals. eqn (5) and (6) ensure that the remaining anti-diagonals will contribute less than ϵ to the expectation value (16). The circuits can be constructed from $a^{k[1]}$ by incrementally increasing the number of qubits to $a^{k[r]}$, which requires $\leq 2^r$ bases in total. The contributions from the anti-diagonals are obtained by measuring $n - r$ qubits in the computational basis and $a^{k[r]}$, thus constructed. For the $n - s$ qubits, one needs to take into account only the all- \uparrow and all- \downarrow outputs, yielding the up-most and down-most 2^r anti-diagonals. The total number of bases for this protocol is bound by

$$2^r < 2^{1 - \frac{\log_2 \epsilon}{\beta}} = 2\epsilon^{-\frac{1}{\beta}}. \quad (20)$$

B. Algorithm for compositional ansatz optimization

In the second part of this work, we demonstrate that DVR Hamiltonians also lead to very efficient quantum circuits for representing vibrational states of molecules. In order to apply

VQE, it is necessary to find an appropriate ansatz for $|\psi\rangle$. It is not always clear how to select the ansatz for $|\psi\rangle$. Previous work on the electronic structure proposed various types of Ansatzes for VQE with both fixed^{3,7,25–29} and adaptive structure.^{30–39} Unlike electronic structure problems, where interactions are pairwise additive, vibrational energy calculations are determined by a wide range of potential energy landscapes, which are highly molecule-specific. In order to obtain the most efficient quantum circuit representations of $|\psi\rangle$ for VQE with DVR matrices, we develop and illustrate an iterative algorithm for ansatz construction that minimizes the number of entangling gates for each specific molecule. This algorithm is inspired by work in ref. 40–42.

Our starting point is:⁷

$$\begin{aligned} \psi(\varphi) = & \prod_{d=0}^{k-1} \left[\prod_{q=0}^{n-1} U^{q,d}(\varphi_d^q) \times U_{\text{ent}}^d \right] \\ & \times \prod_{q=0}^{N-1} U^{q,k}(\varphi_k^q) |0^n\rangle, \end{aligned} \quad (21)$$

where $U^{q,d}(\varphi)$ represent $R_Y = \exp(-i\varphi\sigma_Y/2)$ for qubit q , and k is the number of repetitions of the ansatz blocks. The operator U_{ent}^d introduces entanglement between qubits. The form of U_{ent}^d is determined by the ansatz construction algorithm that incrementally increases the complexity of the quantum circuits. More specifically, the ansatz construction starts with a non-entangled quantum state given by eqn (21) with a predetermined number of blocks k and U_{ent}^d set to identity. The method considers $\text{CNOT}(q, p) \forall q < p$ as candidate gates for U_{ent}^d , with each d segment treated independently. In each optimization step, the candidate gate that lowers the VQE energy is added without replacement until convergence. Here, we aim to converge the VQE calculation of the ground state either to 1 cm^{-1}



or 0.01 cm^{-1} , which yields quantum circuits of different complexity, denoted as C_1 and $C_{0.01}$. This convergence error is with respect to the lowest eigenstate of the DVR matrix with the same number of DVR bases.

In order to benchmark the quantum circuits determined by this algorithm, we also use the following ansatz

$$U_{\text{ent}} = \prod_{q=0}^{n-1} \text{CNOT}(q, q+1). \quad (22)$$

instead of U_{ent}^d . The ansatz (22) is denoted hereafter as linear, as it linearly entangles adjacent qubits.

III. Numerical results

We calculate the vibrational energy levels of diatomic (Cr_2) and triatomic (Ar-HCl and Mg-NH) systems. We consider these molecular systems because they exhibit vibrational states with widely different energies (from -55 to $-15\,000 \text{ cm}^{-1}$ from the dissociation threshold) and spatial variations of wave functions and energy level patterns. Our goal is to demonstrate that the same approach can be applied to these widely different molecular systems. For diatomic molecules, we use the DVR Hamiltonian from ref. 21. For triatomic complexes, we use the DVR approach by Choi and Light.²² We use two classical constrained optimization methods to optimize quantum circuit parameters:

Table 1 Vibrational energy (in cm^{-1}) of Cr_2 ($v = 0-5$) in different electronic states. The benchmark (BM) results are obtained with a converged DVR basis. Exact energies obtained from the truncated DVR basis are displayed as E_v (DVR). VQEs use quantum circuits as shown in Fig. 2 (for C_1) and Fig. 3 (for $C_{0.01}$)

Electronic state	v	BM	E_v (DVR)	E_v		
				C_1	$C_{0.01}$	Linear
$1\Sigma_g^+$	0	-15358.94	-15358.99	-15358.87	-15358.99	-15358.99
	1	-14846.67	-14846.96	-14838.70	-14846.75	-14846.96
	2	-14333.21	-14332.81	-14310.29	-14332.80	-14332.82
	3	-13826.93	-13827.29	-13797.65	-13826.87	-13827.29
	4	-13334.02	-13335.45	-13275.12	-13318.77	-13335.45
	5	-12861.37	-12868.75	-12897.70	-12871.32	-12868.75
$3\Sigma_u^+$	0	-9862.07	-9862.14	-9861.37	-9862.14	-9862.14
	1	-9559.46	-9559.41	-9538.97	-9556.67	-9559.41
	2	-9300.92	-9300.88	-9240.74	-9266.82	-9300.88
	3	-9080.60	-9080.42	9068.09	-9079.26	-9085.40
	4	-8897.58	-8896.82	-8866.09	-8870.57	-8896.82
	5	-8747.53	-8742.68	-8729.41	-8750.18	—
$5\Sigma_g^+$	0	-7566.53	-7566.50	-7565.88	-7566.50	-7566.50
	1	-7416.28	-7416.29	-7397.28	-7416.28	-7416.29
	2	-7264.92	-7264.69	-7181.98	-7264.60	-7264.69
	3	-7114.40	-7118.43	-7075.88	-7117.83	-7118.43
	4	-6965.91	-6958.08	-7001.98	-6953.74	-6958.08
	5	-6820.04	-6840.02	-6873.37	-6837.02	-6840.02
$7\Sigma_u^+$	0	-6519.01	-6519.04		-6519.04	-6519.04
	1	-6350.36	-6350.11		-6350.11	-6350.11
	2	-6183.38	-6185.17		-6185.17	-6185.17
	3	-6018.09	-6018.28		-6018.12	-6018.28
	4	-5854.50	-5849.10		-5848.69	-5849.10
	5	-5692.63	-5731.33		-5728.00	-5731.33
$9\Sigma_g^+$	0	-5348.79	-5348.82		-5348.82	-5348.82
	1	-5175.81	-5175.51		-5175.51	-5175.51
	2	-5005.17	-5008.56		-5008.55	-5008.56
	3	-4836.85	-4829.92		-4829.80	-4829.92
	4	-4670.91	-4683.68		-4683.42	-4683.68
	5	-4507.31	-4530.16		-4526.72	-4612.83
$11\Sigma_u^+$	0	-3677.68	-3677.68	-3677.00	-3677.68	-3677.68
	1	-3507.89	-3507.82	-3489.51	-3507.82	-3507.82
	2	-3341.77	-3341.40	-3253.24	-3341.26	-3341.40
	3	-3180.16	-3186.93	-3133.59	-3185.51	-3186.93
	4	-3023.07	-3012.82	-3061.61	-3001.04	-3012.82
	5	-2870.22	-2874.49	-2904.46	-2866.05	-2874.54
$13\Sigma_g^+$	0	-548.68	-548.68	-548.65	-548.67	-548.68
	1	-497.16	-497.15	-496.47	-496.84	-497.15
	2	-449.18	-449.26	-443.48	-448.36	-449.26
	3	-404.71	-404.67	-382.88	-390.96	-404.67
	4	-363.58	-362.99	-369.09	-360.62	-362.99
	5	-325.67	-325.72	-315.46	-310.29	-325.72



the bounded limited memory method of Broyden, Fletcher, Goldfarb, and Shanno,^{48,49} and sequential least squares programming.⁵⁰ We benchmark the VQE results by the vibrational levels calculated using direct diagonalization with the converged DVR basis and previous literature results, where available.

Table 1 demonstrates the performance of VQE for vibrational states $\nu = 0-5$ of seven electronic states of Cr_2 with zero rotational angular momentum. We use the interaction potentials from ref. 23, illustrated in Fig. 2. The VQE calculations use 16 DVR points placed to span the range, including the minimum of the potential energy. Apart from the DVR parameters, the circuit optimization algorithm is applied identically to all seven electronic states. The Hamiltonian is represented by expansion (2) including ≈ 130 Pauli terms. Table 1 displays VQE results obtained with three types of quantum circuits: the linear ansatz (22) with 3 repetitions, and optimized circuits \mathcal{C}_1 (shown in Fig. 2) and $\mathcal{C}_{0.01}$ (shown in Fig. 3).

The results in Table 1 and Fig. 2, 3 show that VQE can be used to compute the vibrational levels of diatomic molecules with high precision using shallow quantum circuits. It is particularly instructive to analyze the difference between the quantum circuits displayed in Fig. 2 and 3. The quantum circuits in Fig. 3 yield a much better convergence (within 0.01 cm^{-1}) and a significantly higher accuracy than the quantum circuits in Fig. 2 at the expense of a small number of additional quantum gates. For example, for the ground vibrational state of Cr_2 in the $^3\Sigma_g^+$ electronic state, adding two entanglement gates reduces the error of the computation from 0.69 cm^{-1} to 0.07 cm^{-1} , with respect to the benchmark calculation result (labeled BM).

These results also illustrate the utility of the ansatz optimization algorithm developed here. By construction, the algorithm aims to produce quantum circuits of the lowest complexity for a particular accuracy target or particular convergence threshold.

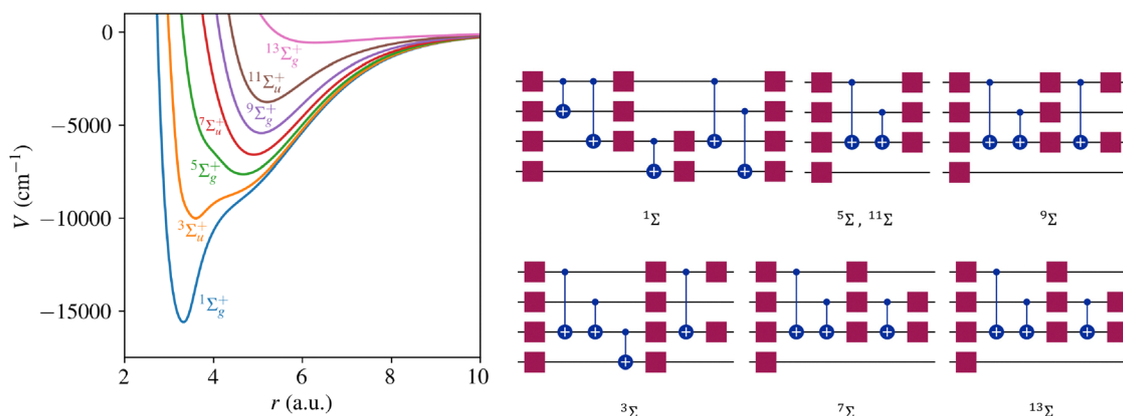


Fig. 2 Left panel: Potential energy for Cr_2 from ref. 23. Right: Quantum circuits for VQE yielding the ground state energy with error $\leq 1 \text{ cm}^{-1}$. The squares represent the R_Y gates and the circles show the entangling CNOT gates.

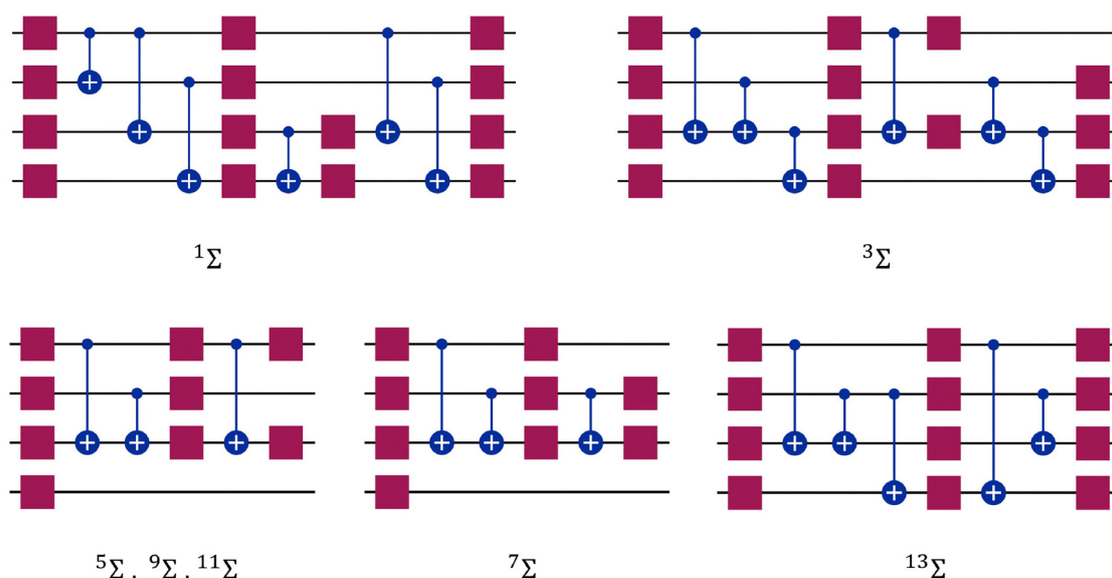


Fig. 3 Quantum circuits for VQE yielding the ground state energy with error $\leq 0.01 \text{ cm}^{-1}$. The squares represent the R_Y gates and the circles show the entangling CNOT gates.



This can be exploited to examine the role of specific gates or gate combinations in determining the expressivity of quantum circuits for representing the vibrational states of molecules. Given that the optimized quantum circuits illustrated in Fig. 2 and 3 are very shallow, the results of Table 1 suggest that current quantum computing devices can already be used for high-precision computation of vibrational energy levels.

For tri-atomic complexes, we use 32 DVR basis states and accurate atom-molecule potential energy surfaces by Hutson for Ar-HCl⁴³ and by Soldán *et al.* for Mg-NH.⁵¹ We keep both HCl and NH in the ground vibrational state and compute the vibrational states supported by the atom-molecule interaction potential. We obtain the DVR points for the triatomic systems by diagonalizing the coordinate representations. The parameters to generate the DVR points are selected to cover the low-energy regions of the potential energy surface. The DVR Hamiltonians are represented by 165 (Ar-HCl) and 170 (Mg-NH) Pauli terms in eqn (2) acting on 5 qubits. As above, we construct three types of quantum circuits: \mathcal{C}_1 , $\mathcal{C}_{0,01}$ and the linear ansatz (22) with k repetitions.

We note that the triatomic molecular systems considered here represent van der Waals complexes. Accurate calculations

of ro-vibrational energies for van der Waals complexes are generally challenging due to the weak binding of the underlying potential energy surface and a large spatial extent of the corresponding vibrational states. Typically, ro-vibrational energy levels of van der Waals complexes are computed by solving a system of coupled differential equations, using, for example, a log-derivative propagation method. Fig. 4 and Table 2 show that accurate results for these weakly bound molecular complexes can be obtained by VQE with very shallow circuits. Fig. 5 shows the fidelity of the ground and excited states obtained from \mathcal{C}_1 and $\mathcal{C}_{0,01}$, with respect to the corresponding DVR eigenstates and demonstrates that the quantum circuits not only provide accurate energies, but also a good representation of the underlying eigenstates. We observe that a small number of entangling gates is sufficient to ensure accurate calculations for both the ground and excited state energy. We also observe that some of the optimized circuits in Fig. 2 and 4 are only partially entangled, which indicates that accurate VQE results can be obtained with ensembles of unentangled quantum circuits. This further decreases the complexity of the quantum computations.

In order to assess whether the present method is practical for NISQ-era devices, we also ran the circuit optimization

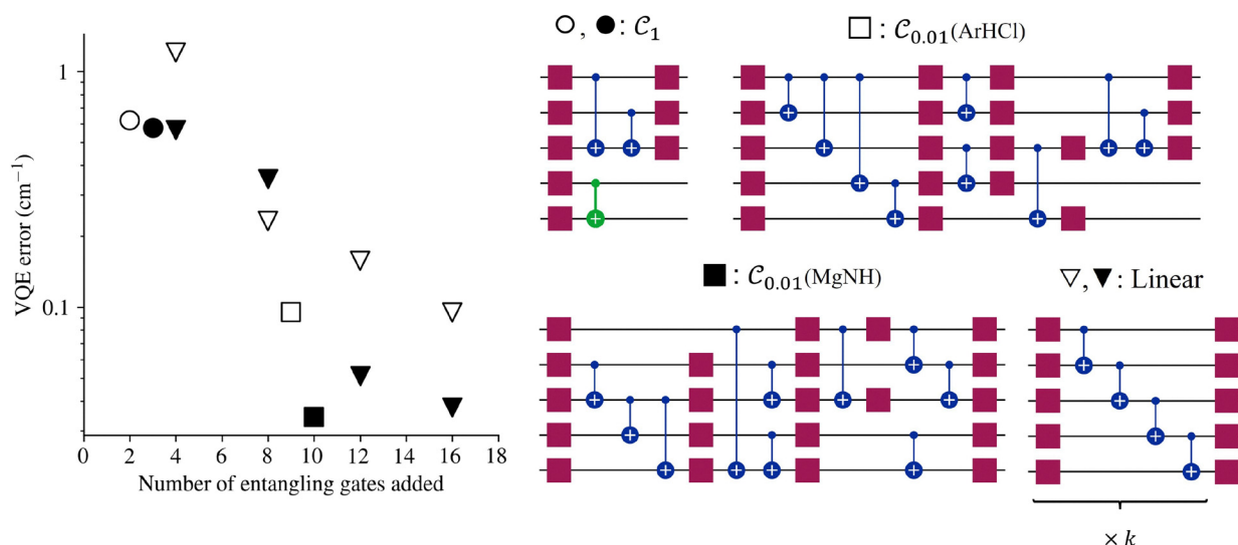


Fig. 4 Left panel: VQE error for the lowest energy of Ar-HCl (open symbols) and Mg-NH (full symbols) computed with optimized quantum circuits in the right panel. Circles – with \mathcal{C}_1 circuits; squares – with $\mathcal{C}_{0,01}$ circuits; triangles – with ansatz (21) using (left to right) $k = 1, 2, 3$ and 4 and eqn (22). The \mathcal{C}_1 Ansatz for Mg-NH excludes the gate shown in green.

Table 2 Vibrational energy (in cm⁻¹) of Ar-HCl and Mg-NH by VQEs with 32 DVR points and 5-qubit circuits

Molecule	ν	E_ν (experiment)	E_ν (computed in ref. 43)	E_ν (classical, present)	E_ν (DVR)	E_ν (VQE)		
						\mathcal{C}_1	$\mathcal{C}_{0,01}$	Linear
ArHCl	0	-114.7 ⁴⁴	-115.151	-115.265	-115.178	-114.645	-115.169	-115.171
	1	-91.04 ^{45,46}	-91.485	-91.642	-90.959	-80.824	-90.485	-90.929
	2	-82.26 ⁴⁷	-82.717	-82.825	-82.727	-75.900	-82.986	-82.650
MgNH	0	—	—	-88.227	-88.196	-87.650	-88.191	-88.190
	1	—	—	-63.603	-62.928	-56.050	-62.730	-62.664
	2	—	—	-55.461	-54.961	-55.145	-54.850	-54.866



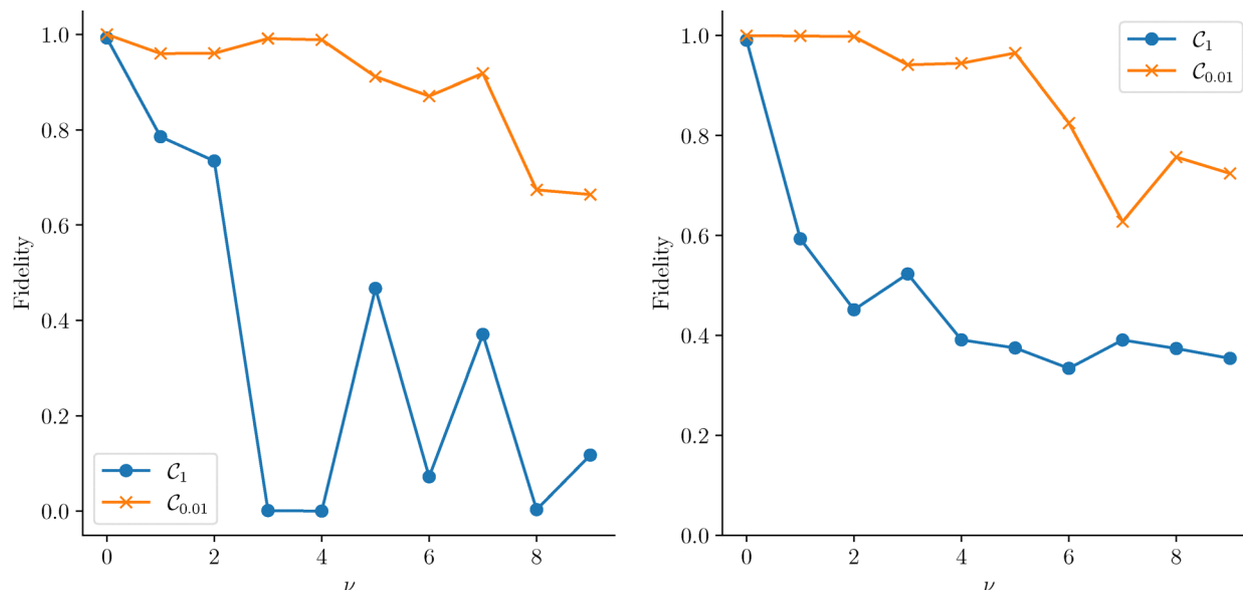


Fig. 5 Fidelity of the vibrational ground and excited states calculated with respect to the corresponding DVR wave functions for Ar-HCl (left panel) and Mg-NH (right panel).

algorithm for the Ar-HCl complex with a realistic noise model. Specifically, we repeat the full compositional ansatz search using the Sherbrooke noise model in Qiskit, which captures gate and readout errors of a superconducting quantum processor. Fig. 6 shows the error of the ground state VQE calculation during the circuit optimization algorithm for circuits with $k = 1$ in eqn (21), and two circuits obtained after 4 steps of the greedy search algorithm. We find that the greedy optimization continues to identify shallow circuits yielding spectroscopic accuracy for multiple excited vibrational states. While the circuits obtained under noise may differ from those found in noiseless simulations, their accuracy is comparable, indicating the existence of multiple noise-resilient shallow representations of the same vibrational eigenstates. The persistence of successful optimization under realistic noise and the existence of multiple shallow optima suggest that the DVR-based ansatz space avoids the effects typically associated with barren plateaus in deep, unstructured circuits. Further analysis of the robustness of the method to noise is left for future work.

IV. Conclusions

We have shown that the structure of DVR matrices can be leveraged to represent molecular Hamiltonians by the number of quantum circuits that grows polynomially with the number n of qubits. Given the binary basis encoding yielding $N = 2^n$, this offers an exponential reduction of the measurement complexity for matrices with N DVR basis states. This has significant implications for the potential quantum advantage of VQEs for ro-vibrational computations. Exponential scaling of VQE is a significant and common challenge, particularly relevant for unstructured Hamiltonians such as the vibrational eigenvalue

problem in an unstructured basis. For quantum chemistry calculations as well as for ro-vibrational problems, this is usually dealt with by second quantization. However, these approaches often lead to extended quantum circuits that require a large number of qubits and gates.

We have also demonstrated that DVR leads to efficient quantum circuits for VQE computations of vibrational energy levels. To show this, we have introduced a general approach for constructing the quantum ansatz by combining DVR with VQE and a greedy search in the space of gate permutations. The results yield compact representations of vibrational states by quantum circuits of a gate-based quantum computer. We have shown that both the ground and excited vibrational energies can be computed with a relative accuracy of $< 1\%$ using very simple, in some cases, partially entangled circuits. The accuracy of 1 cm^{-1} can be achieved with < 20 (< 5 entangling) gates and 4 qubits for diatomic molecules and < 30 (< 9 entangling) gates with 5 qubits for triatomic van der Waals complexes. This should be compared with previous VQE calculations of vibrational energy levels that required extended quantum circuits with > 200 (for CO, COH and O_3 molecules¹⁷) or between 44 and 140 292 (for CO_2 , H_2CO and HCOOH molecules¹⁶) entangling gates.

Reducing the complexity of quantum circuits is particularly important for VQE approaches that are known to be affected by barren plateaus, which limit the quantum advantage of variational quantum computations. Although barren plateaus are formally an asymptotic effect, they can become relevant already for systems of moderate size. While a comprehensive treatment of barren plateaus is beyond the scope of the present work, our results provide several encouraging indications that this issue may be mitigated for vibrational energy calculations using the present framework. In particular, we observe no signs of barren



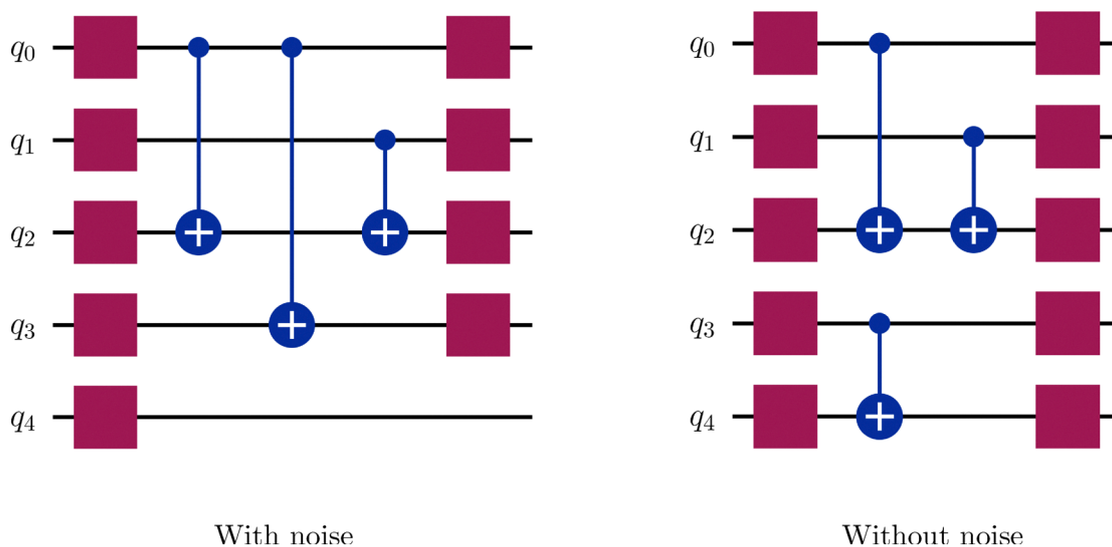
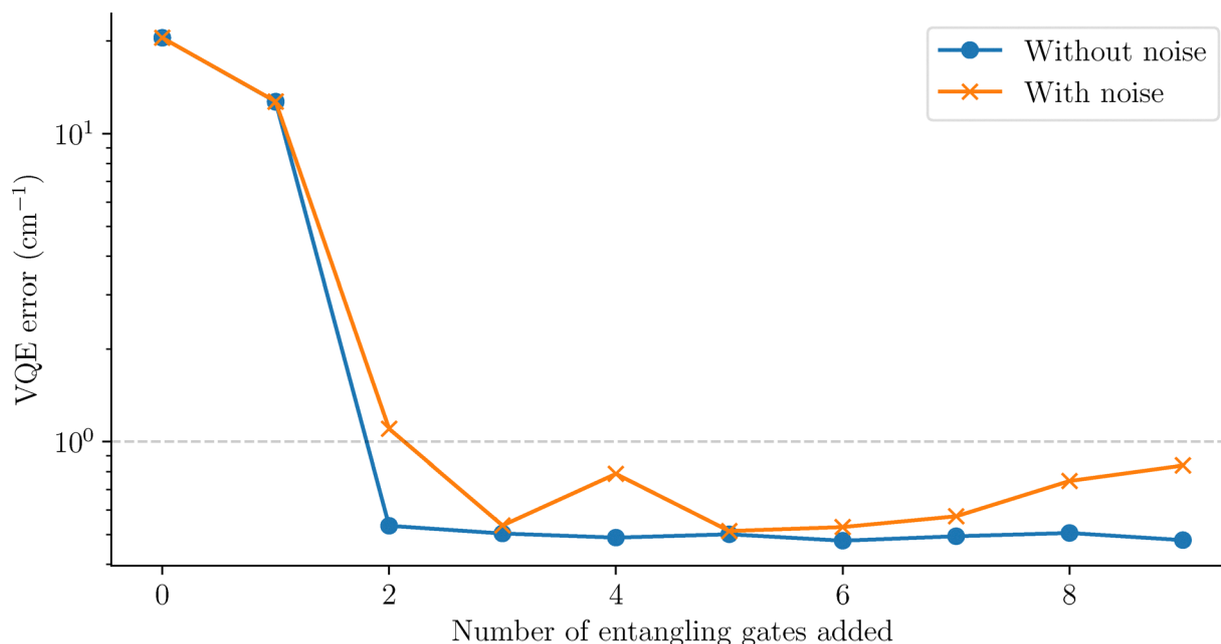


Fig. 6 (Top panel) VQE error during the greedy circuit optimization for Ar–HCl using $k = 1$ in eqn (21). (Bottom panel) The circuits obtained after adding 3 entangling gates with and without the noise model for Ar–HCl.

plateaus for the systems considered here, including in simulations that explicitly incorporate realistic noise. Moreover, the ansatz construction algorithm introduced in this work consistently yields accurate vibrational energies and wave functions using extremely shallow circuits, some of which are only partially entangled. These observations suggest that VQE computations of vibrational energies may be scalable to larger systems than can be anticipated based on the previous calculations.^{16,17} This must be confirmed by a more systematic numerical investigation, which we leave for future work.

Finally, we note that DVR does not require global fits of potential energy surfaces or integrals over the potential energy

for the construction of the Hamiltonian matrix. The implementation of VQEs with DVR in the first quantization does not require normal-mode analysis for the construction of the quantum ansatz. For these reasons, the present approach can be readily extended beyond molecular dynamics. For example, this method can be directly applied to designing efficient quantum circuits for variational computations of the eigenspectra of semiconductor quantum dots.⁵² Finally, our calculations suggest that quantum circuit representations of vibrational states can be used as efficient (small number of free parameters) Ansatz for classical variational calculations.



Author contributions

All authors have designed the concept of this work and co-wrote the manuscript. The algorithm presented in Section IIA was developed by DB. The algorithm for Ansatz optimization was developed by KA and RVK and the numerical calculations were performed by KA.

Conflicts of interest

There are no conflicts to declare.

Data availability

The data that support the findings of this study are available within the article.

Appendices

A Scaling of DVR matrix bands

The goal of this Appendix is to prove eqn (5) and (6). Specifically, we aim to show that

$$\sum_{k=s}^{2^n-1} |f(k)| \leq O(s^{-\alpha}), \quad \alpha > 0, \quad (\text{A1})$$

$$\sum_{k=r}^{2^{n+1}-1-r} |g(k)| \leq O(r^{-\beta}), \quad \beta > 0, \quad (\text{A2})$$

where $r \ll N$ and $s \ll N$, and $N = 2^n$ is assumed to be large.

We consider a particle with mass m on a grid of $x = [a, b]$ with the uniform spacing Δx , yielding

$$x_j = b + j\Delta x, \quad j = 1, \dots, N-1. \quad (\text{A3})$$

Given an orthonormal basis $\{\phi_n\}$, the matrix elements of the kinetic energy can be written as

$$T_{ij} = -\frac{\hbar^2}{2m}\Delta x \sum_{n=0}^{N-1} \phi_n(x_i)\phi_n''(x_j). \quad (\text{A4})$$

Fourier grid DVR uses the basis²¹

$$\phi_n(x) = \left(\frac{2}{b-a}\right)^{1/2} \sin\left[\frac{n\pi(x-a)}{b-a}\right]. \quad (\text{A5})$$

For this basis, the sum can be evaluated analytically, yielding²¹

$$T_{ij} = \begin{cases} \frac{\hbar^2}{2m} \frac{1}{(b-a)^2} \frac{\pi^2}{2} \left[\frac{2N^2+1}{3} - \frac{1}{\sin^2(\pi j/N)} \right], & i=j, \\ \frac{\hbar^2}{2m} \frac{(-1)^{i-j}}{(b-a)^2} \frac{\pi^2}{2} \left\{ \frac{1}{\sin^2[\pi(i-j)/2N]} - \frac{1}{\sin^2[\pi(i+j)/2N]} \right\}, & i \neq j. \end{cases} \quad (\text{A6})$$

Following Colbert and Miller,²¹ we now consider the cases of infinite lattices, relevant for radial molecular coordinates, and a case of finite a and b .

1. Infinite lattice [$a = -\infty, b = \infty$]. The finite grid spacing $\Delta x = \frac{b-a}{N}$ requires $N \rightarrow \infty$. In this limit, eqn (A6) becomes

$$T_{ij} = \frac{\hbar^2}{2m\Delta x^2} (-1)^{i-j} \begin{cases} \frac{\pi^2}{3}, & i=j, \\ \frac{2}{(i-j)^2}, & i \neq j. \end{cases} \quad (\text{A7})$$

Note that in this case $g(i+j)$ vanishes. It is evident that

$$|f(k)| = \frac{2}{(i-j)^2} \quad (\text{A8})$$

satisfies eqn (A1). To prove this, we note that for functions that are (strictly) monotonically decaying in absolute value

$$\int_s^u |f(k)| dk < \sum_{k=s}^u |f(k)| < \int_{s-1}^{u-1} |f(k)| dk. \quad (\text{A9})$$

For our choice of $f(\cdot)$ this yields the bound

$$\sum_{k=s}^{2^n-1} |f(k)| < 2 \int_{s-1}^{2^n-2} \frac{dk}{k^2} = 2 \left(\frac{1}{s-1} - \frac{1}{2^n-2} \right) < 3s^{-1}. \quad (\text{A10})$$

2. Infinite lattice [$a = 0, b = \infty$]. For the lattice with $[a = 0, b = \infty]$, the matrix elements

$$T_{ij} = \frac{\hbar^2}{2m\Delta x^2} (-1)^{i-j} \begin{cases} \frac{\pi^2}{3} - \frac{1}{2i^2}, & i=j, \\ \frac{2}{(i-j)^2} - \frac{2}{(i+j)^2}, & i \neq j. \end{cases} \quad (\text{A11})$$

include $g(k)$. Since eqn (A10) applies to $g(k)$, the arguments in subsection (8.1) apply to eqn (12).

3. Finite lattice [a, b]. We note that the prefactor in eqn (A6) satisfies $\frac{\hbar^2}{2m} \frac{1}{(b-a)^2} \frac{\pi^2}{2} \cdot \frac{2m\Delta x^2}{\hbar^2} = \frac{\pi^2}{2N^2}$. For the following functions

$$|\tilde{f}(k)| = \frac{\pi^2}{2N^2} \left[\frac{1}{\sin^2(\pi k/2N)} - 1 \right], \quad (\text{A12})$$

$$|\tilde{g}(k)| = \frac{\pi^2}{2N^2} \left[\frac{1}{\sin^2(\pi k/2N)} - 1 \right], \quad (\text{A13})$$

Eqn (A9) yields

$$\sum_{k=s}^{N-1} \frac{1}{\sin^2(\pi k/2N)} = \frac{2N}{\pi} \sum_{k=s} \frac{\pi}{2N \sin^2(\pi k/2N)} \quad (\text{A14})$$

$$< \frac{2N}{\pi} \int_{\frac{\pi(s-1)}{2N}}^{\frac{\pi}{2}} \frac{dx}{\sin^2(x)} \quad (\text{A15})$$

$$= \frac{2N}{\pi} \cot \left[\frac{\pi(s-1)}{2N} \right]. \quad (\text{A16})$$



Using the Laurent series for $\cot(y)$ around $y = 0$, we can write

$$\frac{2N}{\pi} \cot\left[\frac{\pi(s-1)}{2N}\right] = \frac{4N^2}{\pi^2(s-1)} - \frac{(s-1)}{3} + O\left(\frac{s^3}{N^2}\right). \quad (\text{A17})$$

This yields

$$\sum_{k=s}^{2^n-1} |\tilde{f}(k)| < \frac{\pi^2}{2N^2} \left[\frac{4N^2}{\pi^2(s-1)} - \frac{(s-1)}{3} - (N-1-s) + O\left(\frac{s^3}{N^2}\right) \right] \quad (\text{A18})$$

$$< \frac{2}{s-1} + O\left(\frac{s^3}{N^4}\right) \quad (\text{A19})$$

$$< 3s^{-1}. \quad (\text{A20})$$

For $g(k)$, we exploit the symmetry of $\sin(\cdot)$ to write

$$\sum_{k=r}^{2^{n+1}-1-r} \frac{1}{\sin^2(\pi k/2N)} = 1 + 2 \sum_{k=r}^{2^n-1} \frac{1}{\sin^2(\pi k/2N)}, \quad (\text{A21})$$

and apply eqn (A19)–(A21).

B Representation of $H^{(s,r)}$ in terms of $t^{k[n]}$ and $a^{k[n]}$

We leverage the structure of the truncated DRV matrix by representing it as a sum of a diagonal matrix D and k th diagonal and anti-diagonal components,

$$H^{(s,r)} = D + \sum_{k=1}^{s-1} f(k) t^{k[n]} + \sum_{k=1}^{r-1} \left(g(k) a^{k[n]} + g(2^n - k) a^{(2^n - k)[n]} \right), \quad (\text{B1})$$

where

$$D_{ij} = \delta_{ij} \left(d(i) - g(2i) - \sum_{k=1}^{s-1} f(k) q^{k[n]}(i) \right) \quad (\text{B2})$$

$$t_{ij}^{k[n]} = \delta_{|i-j|,k} + \delta_{ij} q^{k[n]}(i), \quad (\text{B3})$$

$$a_{ij}^{k[n]} = \delta_{i+j,k} \quad (\text{B4})$$

and $q^{k[n]}(i)$ can be computed by an efficient Algorithm 1. Our strategy is to measure the expectation values of each of the $\{D, t^{k[n]}, a^{k[n]}\}$ separately using the induction over n and sum them with the corresponding weights.

In addition to k th diagonals, the construction of $t^{k[n]}$ adopted in this work introduces additional contributions $q^{k[n]}(i)$ to the matrix elements on the main diagonal. These contributions can be deduced from the measurement protocol for $t^{k[n]}$ as described in Algorithm 1.

Algorithm 1 Computing $q^{k[n]}(i)$ from binary representation of i

```

define  $l \leftarrow \lceil \log_2(k+1) \rceil$ ,
          $last\_l\_bits \leftarrow i \bmod 2^l$ ,
          $anti\_last\_l\_bits \leftarrow 2^l - last\_l\_bits$ ;
initialize  $output \leftarrow 1$ ,
            $j \leftarrow l + 1$ ;
while  $j \leq n$  do
  if  $j$ th bit of  $i$  is 1 then
    if  $last\_l\_bits < k$  or  $anti\_last\_l\_bits < l$  then
       $output \rightarrow out + 1$ ;
     $j \rightarrow j + 1$ ;
return  $output$ .

```

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