Quantum computations of relativistic and many-body effects in atomic and molecular systems based on variational algorithms

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Inventing Harmonious Future

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Outline

- Variational Quantum Eigensolver (VQE):
	- Brief introduction to VQE.
	-
	- Dipole moments of molecules.
- Quantum Annealer Eigensolver (QAE):
	- Brief introduction to QAE.
	- Applications to fine structure splitting in atoms.

• Applications to ground state energies and hyperfine interactions in atoms.

The focus of both variational algorithms in this talk will be on quantum computations of relativistic and many-body effects.

Digital quantum computing

Problem

Quantum Algorithim

Figure: Flowchart for Quantum Computation

- Qubits: Quantum states $a |0\rangle + b |1\rangle$.
- CNOT: an example of a 2-qubit gate.
- Quantum computation using quantum circuits (qubits and quantum gates):

• Quantum gates: Unitary operators.
$$
R_X(\theta) = e^{-i\frac{\theta}{2}X}
$$
, $R_Y(\theta) = e^{-i\frac{\theta}{2}Y}$, and $R_Z(\theta) = e^{-i\frac{\theta}{2}Z}$: examples of 1-qubit gates.

$$
|00\rangle \xrightarrow{R_X(\theta)} cos\left(\frac{\theta}{2}\right)|00\rangle - i sin\left(\frac{\theta}{2}\right)|10\rangle \xrightarrow{CNOT} cos\left(\frac{\theta}{2}\right)|10\rangle
$$

$$
|0\rangle
$$

• In our quantum computation, we use layers of rotations and CNOT gates.

Relativistic Effects in Atomic Systems

Non-Relativistic Hamiltonian

$$
H_{nr} = \sum_{i} \left(\frac{p_i^2}{2m} + V_{nuc}\right) + \sum_{i < j} \frac{e^2}{r_{ij}} \qquad \qquad \bigoplus \qquad \qquad \textbf{Z}, \textbf{N} \qquad \bigoplus
$$

For large Z, velocities of electrons increase and they must be treated relativistically

Relativistic Hamiltonian

$$
H_{DC} = \sum_i \left(c \alpha_i \cdot p_i + \beta_i mc^2 + V_{nuc} \right) + \sum_{i
$$

Leading order correction to the Coulomb interaction is the Breit interaction

$$
H_B=-\frac{e^2}{2}\sum_{i
$$

 α and β are (4x4) matrices. Other relativistic corrections are generally less important.

: Dirac-Coulomb Hamiltonian

 \bigcup

Hyperfine structure constant

- The hyperfine Hamiltonian is given by
-

- $\mathscr A$ is often a difficult quantity to evaluate, since it is determined by a complex interplay of several electron correlation effects, unlike the energy.
- Its computation requires accurate single particle wave functions in the nuclear region, and hence computing $\mathscr A$ is a sensitive test of relativistic and correlation effects in atoms and molecules.

$$
\mathcal{A} = \frac{1}{J} \mu_N g_I \langle JJ | \sum_i \frac{(\vec{r}_i \times \vec{\alpha}_i)_Z}{r_i^3} | JJ \rangle.
$$

•

$$
y H_{hf} = \overrightarrow{j_e} \cdot \overrightarrow{A_N}.
$$

 $\bullet\,$ The quantity can be represented as an effective Hamiltonian: $H^{ey}_{hf}=\mathscr{A} I\cdot\, J$. *Heff hf* $=$ $\mathscr{A}I \cdot J$

VQE algorithm

VQE Introduction

- Hybrid (classical-quantum) algorithm to calculate ground state energies and other properties of quantum many-body systems. It is suitable for NISQ computers (50-100) qubits).
- \bullet $|\Phi_0\rangle$ can be written in terms of orbitals, where each orbital is $|\phi_i\rangle = |n_i l_i m_{l,i} s_i m_{s,i}\rangle$ or $\langle n_i l_i s_i j_i m_i \rangle$ is a qubit. In qubit representation, $|\Phi_0\rangle = |1\rangle \otimes \cdots \otimes |1\rangle \otimes |0\rangle \otimes \cdots \otimes |0\rangle$ Occupied Unoccupied
- The variational principle is used to find the ground state energy of a system: $E(\theta) = \langle \Psi(\theta) | H | \Psi(\theta) \rangle$; $|\Psi(\theta) \rangle = U(\theta) | \Phi_0 \rangle$. Minimise $E(\theta) = \langle \Phi_0 | U(\theta)^\dagger H U(\theta) | \Phi_0 \rangle$ by varying till convergence is reached. Expectation value: $\langle \Phi_0 | U(\theta)^\dagger A U(\theta) | \Phi_0 \rangle$; $A = \sum_{pq} A_{pq} a_p^{\dagger} a_q$; A_{pq} (one particle matrix element) from classical computer.
- The choice of $U(\theta)$ is crucial for obtaining an accurate value for energy and other properties.
- \bullet Another choice is the hardware efficient ansatz (HEA)- e.g $U(\theta) = \prod [R(\theta)] \times CNOT$ (A Kandala et al, Nature 2017)

A physically motivated ansatz for $U(\theta)$ is based on **unitary coupled cluster** (UCC) method, where $U(\theta = t) = e^{T-T^{\dagger}}$. Where T and t is cluster operator and amplitudes. $T=T_1+T_2\cdots=\sum_{ia}t^a_ia^{\dagger}_aa_i+\sum_{ijab}t^{ab}_{ij}a^{\dagger}_aa^{\dagger}_ba_ja_i+\cdots$ (A Peruzzo et al, Nature Communications 2014)

VQE Flowchart

- Hyperfine structure constants computation for neutral Li and highly charged isoelectronic systems: Li-like Sc, Lilike Pr, and Li-like Bi.
- Four qubit computations on superconducting qubit hardware at RIKEN (Nakamura lab).
- Choice of hardware efficient ansatz: $R_X(\pi/2) R_Z(\theta) R_X(\pi/2) CNOT$ type. Linear entanglement strategy. Depth of one.
- Benchmarked with all-electron calculations in the complete Hilbert subspace that was considered.
- The major challenge: the same ansatz needs to capture dissimilar correlation effects involved in both properties.

• can occur from both these sources. $e^{exp} = \sum w_l^{hfs}$ *l ^l* ⟨*Pl* \rangle^{exp} ; w_j^{hfs} λ_l^{hfs} is evaluated on a classical computer, and $\langle P_l \rangle^{exp}$

tcg crest **VQE algorithm for relativistic calculations of ground state energies and Inventing Harmonious Future hyperfine structure costants**

• The accuracy in our calculations is about 99 percent for energies and vary between 60 and 80 percent for the hyperfine structure constants, all relative to the best classical computation (Full CI- exact diagonalization in a limited space).

State-of-the-art:

- Hardware efficient ansatz: Kandala et al (Nature 2017): best precision is 1.6 mHa for H2 in its equilibrium bond length.
- UCC Ansatz: Guo et al (Nature Physics 2024): Best precision is ~0.1 mHa for H2 in its equilibrium bond length.

; w_i^{nys} is evaluated on a classical computer, and $\langle P_i \rangle^{exp}$ on a quantum computer. Error

VQE algorithm for relativistic calculations of molecular electric dipole moments

- Computation of molecular electric dipole moments (PDMs) of single valence molecules.
- Choice of ansatz: unitary coupled cluster in the singles and doubles approximation (UCCSD): $|\Psi\rangle = e^{T-T^{\dagger}}|\Phi_0\rangle; T = T_1 + T_2.$ $|\Phi_0\rangle$; $T = T_1 + T_2$
- The set of amplitudes $\{t_{ia}, t_{ijab}\} \equiv \{\theta\}$ are the variational parameters.
- Eighteen qubit simulations: PDMs of BeH through RaH (3 occupied + 15 unoccupied). Relativistic effects can be as large as 25 percent for PDM of RaH.
- Six qubit computations on IonQ Aria-I device: PDMs of moderately heavy SrH and SrF $(3+3)$.
- Twelve qubit computations on IonQ Forte-I device: PDMs of moderately heavy SrH $(5+7)$.

arXiv 2406.04992 (2024)

strategies: use of point group symmetry, energy sort VQE procedure, pipeline based circuit optimization, RL-based ZX-calculus, cliques to reduce number of terms measured in PDM operator, particle number conserving post selection

- For quantum hardware computations, we used a suite of resource reduction scheme.
- Six qubit result: Accuracy of ~95 percent relative to the best classical computations after error mitigation.
- Twelve qubit result: Accuracy of ~99 percent relative to the best classical computations after error mitigation.

VQE algorithm for relativistic calculations of molecular electric dipole moments

arXiv 2406.04992 (2024)

Conclusion

• The VQE algorithm has been successfully used for computing ground state properties of lithium-like systems, in particular, ground state energies and hyperfine interactions on

• Correct trends for relativistic effects have been reproduced. More qubits are needed for

• For molecular electric dipole moments on lonQ hardware, we obtain ~95 percent and ~99 percent accuracies relative to the best classical computations for six- and twelve- qubit computations respectively using two different versions of IonQ. The circuit optimisation was

constants and accuracy between 60 and 80 percent has been obtained respectively relative to the best classical computations on a four-qubit superconducting quantum computer at

- superconducting and trapped ion quantum computers have been computed.
- accurate computations of many-body effects.
- For ground state energies, accuracy of about 99 percent and for hyperfine structure RIKEN.
- superior for the latter case.

QAE algorithm

Quantum annealing

Kadowaki and Nishimori, Phys Rev E 1998

- $\sigma_i^Z | s_i \rangle = s_i | s_i \rangle$. Thus, with $| \Psi \rangle = \otimes_{i=1}^N | s_i \rangle$ as the ground state wave function of the final Hamiltonian, an energy functional $\epsilon(s_i) = \langle \Psi | H | \Psi \rangle = \sum h_i s_i + \sum j_{ij} s_i s_j.$ *i i*,*j*
- With $q_i = \frac{1}{2}$, we get the QUBO form: . $q_i =$ $s_i + 1$ 2 $\varepsilon(q_i)_{Q} = \sum_{i} Q_i q_i + \sum_{j} Q_{ij} q_i q_j$ *i i*,*j*
- The energy functional: obtained by using $|\Psi\rangle = \sum_{i} c_i |\Phi_i\rangle$ can be expressed in QUBO form via a floating point encoding scheme given by $c_i^{\{r\}} = c_i^{\{r-1\}} + 2^{\frac{1-r}{2}} \sum f_k 2^{-k} q_k^i$ $\epsilon(c_i) = \langle \Psi | H | \Psi \rangle - \lambda \langle \Psi | \Psi \rangle = \sum (H_{ii} - \lambda) + \sum H_{ij} c_i c_j$ *i ij* $|\Psi\rangle = \sum c_i |\Phi_i\rangle$ *i i* $= c_i^{\{r-1\}}$ i^{r-1} + 2 $1 - r$ 2 *K*−1 ∑ $k=0$ $f_k 2^{-k} q_k^i$ *k*

Energy functional

Energy

 $\overline{|\langle \Psi|\overline{H}_{DCB}|\Psi\rangle}|$

Construct $\ket{\Psi}$ **from qubit configuration**

D-Wave Advantage 5000Q (Embedding, Annealing)

 $\epsilon(\lambda, H_{DCB})$

Optimization

Post-Processing

Construct QUBO functional

 ϵ_{Q}

Initialization

 $H_{DCB} = H_{DC} + H_{B}$

Final ground state energy

$\langle \Psi | H_{DCB} | \Psi \rangle$

Quantum annealer

Computation of Relativistic effects using QAE: Fine structure splitting for Boron-like ions

Excitation Energy (Fine Structure Splitting (FSS))

 $\Psi(J = 3/2)$) = $E_{3/2}$ | $\Psi(J = 3/2)$

E1/2

 $1s^22s^22p(L=1,S=1/2, J=1/2, 3/2)$

$$
H|\Psi(J = 1/2)\rangle = E_{1/2}|\Psi(J = 1/2)\rangle
$$

QAE is applied to J=1/2 and J=3/2 states separately on D-Wave quantum annealer

Results

FSS values for boron-like ions. 'relCI' refers to numerical relativistic CI calculations, 'Hardware' gives our mean (over five repetitions) of relativistic QAE performed on the D-Wave Advantage machine. The quantity in bracket is the percentage fraction difference, $\frac{E_{\text{relCI}} - E_{\text{hardware}}}{\text{relCI}} \times 100$. 'Expt' stands for the experimental value (in Ha).

(Kumar et al, Phys Rev A (2024))

Conclusions and Outlook

- ➔ We have performed computations of the Fine Structure Splitting in Boron-like ions using the Quantum Annealer Eigensolver using D-Wave 5000Q.
- ➔ We have obtained an accuracy of 99% compared to high precision spectroscopic measurements of the fine structure splitting of these ions.
- ➔ The accuracy was achieved by improving the workflow of the QAE algorithm and inclusion of important physical effects.
- \rightarrow This is the first step in carrying out high accuracy quantum annealing computations of atomic quantities that have a wide range of applications including the probing of new physical phenomena beyond the Standard Model of particle physics.

Hyperfine structure constant

-
- . μ_I is the nuclear magnetic moment. Thus, .
. • *μI* ∫ *f*(*r*) $\frac{1}{r^3}dV$, μ_I $\langle \Psi | H_{hf}^{eff}$ $\mathscr{A}_{hf}^{eff}|\Psi\rangle = \mathscr{A}\langle \Psi | I \cdot J | \Psi \rangle = \mathscr{A}IJ$ $\langle \Psi | H_{hf} | \Psi \rangle = \langle \Psi | H_{hf}^{eff}$ $\mathcal{H}_{hf}^{eff}|\Psi\rangle$. Thus, $\mathscr{A}=0$
- electron correlation effects, unlike the energy.
-

• The hyperfine Hamiltonian is given by
$$
H_{hf} = \vec{j_e} \cdot \vec{A_N}
$$
. Thus, $\langle \Psi | H_{hf} | \Psi \rangle = \langle \Psi | \vec{j_e} \cdot \vec{A_N} | \Psi \rangle$.

• The quantity can be rep as an effective Hamiltonian: $H_{hf}^{eff}=\mathscr{A}\vec{I}\cdot\overrightarrow{J}$, where \mathscr{A} is given by *hf* $=$ $\mathscr{A}I \cdot J$

Thus,
$$
\mathcal{A} = \frac{\langle \Psi | H_{hf} | \Psi \rangle}{IJ} = \frac{1}{IJ} \mu_N g_I I \langle JJ | \sum_i \frac{(\vec{r}_i \times \vec{\alpha}_i)_Z}{r_i^3} | JJ \rangle.
$$

 \bullet a is often a hard quantity to evaluate, since it is determined by a complex interplay of several

• Its computation requires accurate single particle wave functions in the nuclear region, and hence computing a is a sensitive test of relativistic and correlation effects in atoms and molecules.

•

TABLE III: Table presenting the results for ground state energies of the considered systems in the current work (in Hartree). 'DHF' refers to the Dirac-Hartree-Fock results, while 'CAS-CI' column gives the complete active space configuration interaction results. 'Experiment' presents our main results with error bars, and the 'Spectroscopy' column presents reference values for comparison. The energies have been rounded off to the sixth decimal place.

TABLE IV: Table of results for the hyperfine structure constants (in MHz). The notation followed is the same as in the preceding table, Table III. The hyperfine structure constants have been rounded off to the third decimal place.

Figure 1. The top panel shows the percentage relativistic effects = $\frac{A_{Rel}-A_{NR}}{A_{Rel}}$ × 100 for ground state energy (E) and PDM from our 18-qubit VQE simulations. Our results are benchmarked against CASCI calculations. The bottom panel shows the percentage correlation effects = $\frac{A_X - A_{MF}}{A_X} \times 100$, where X can be correlation energy (circular markers) or the PDM (square markers) relative to respective quantity at VQE and CASCI levels.

gate count by one half.

reduction

ource

Figure 2. (a) Our workflow for quantum hardware execution of SrH 12-qubit PDM calculation on the IonQ Forte device, which leads to reducing quantum resources while retaining precision. (b) Percentage reduction in resources (two-qubit gates, denoted as 2qq in the sub-figure, and circuit depth) with each step of our workflow: U_{ES-VQE} is the UCCSD circuit post-ES-VQE and $U_{ES, Pipopt}$ is the state after pipeline-based optimization (denoted as pipopt). (c) The loss of precision in predicting PDM after each step in our workflow, with ' 1^{st} Clique' indicating the selection of the dominant clique for the PDM operator (See Table $\boxed{S2}$ of the Supplemental Material). Sub-figure (d) illustrates the step ratio, which is the ratio of the number of 2qq before and after the current step in our workflow. It is important to stress that the compound strategy of our RL-ZX based agent followed by the causal flow deterministic algorithm (both based on ZX -Calculus, denoted as RL - ZX + Cflow) reduces the already small

QAE: subQUBO

Number of Repeats: 75 (30 for J=1/2 and 45 for J=3/2), QUBO size: 90 and 160, subQUBO size: 30 (110 qubits) and 40 (190 qubits). Anneal time: 20 microseconds. Individual energies: Best agreement with relCI: ~0.05 mHa, and the worst~1 mHa.

$$
\epsilon = \frac{\langle \Psi | H | \Psi \rangle}{\langle \Psi | \Psi \rangle} - \lambda \langle \Psi | \Psi \rangle
$$
\n
$$
= \sum_{i} c_{i} |\Phi_{i}(M\Pi)|
$$
\n
$$
= \sum_{i} (H_{ii} - \lambda)c_{i}^{2} - \sum_{i < j} H_{ij}c_{i}c_{j}
$$
\n
$$
H_{ij} = \langle \Phi_{i} | H | \Phi_{j} \rangle
$$
\n
$$
= \frac{\sum_{i} (H_{ii} - \lambda)c_{i}^{2} - \sum_{i < j} H_{ij}c_{i}c_{j}}{\sum_{i} H_{ij}^{2}} \frac{A^{(r+1)}, B^{(r+1)} - \sum_{i=1}^{r} A^{(r+1)} - \
$$

Connection between atomic physics and quantum annealing

$$
C_1 \sim \frac{H_{10}}{(H_{00} - H_{11})} \quad , \quad C_2 \sim \frac{H_{20}}{(H_{00} - H_{22})} \quad , \quad \cdots \quad .
$$

Repeat: involves updating λ and subQUBO!

