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Experimental error mitigation via symmetry verification in a variational quantum eigensolver

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Variational quantum eigensolvers offer a small-scale testbed to demonstrate the performance of error mitigation techniques with low experimental overhead. We present successful error mitigation by applying the recently proposed symmetry verification technique to the experimental estimation of the ground-state energy and ground state of the hydrogen molecule. A finely adjustable exchange interaction between two qubits in a circuit QED processor efficiently prepares variational ansatz states in the single-excitation subspace respecting the parity symmetry of the qubit-mapped Hamiltonian. Symmetry verification improves the energy and state estimates by mitigating the effects of qubit relaxation and residual qubit excitation, which violate the symmetry. A full-density-matrix simulation matching the experiment dissects the contribution of these mechanisms from other calibrated error sources. Enforcing positivity of the measured density matrix via scalable convex optimization correlates the energy and state estimate improvements when using symmetry verification, with interesting implications for determining system properties beyond the ground-state energy.

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Noisy intermediate-scale quantum (NISQ) devices [1], despite lacking layers of quantum error correction (QEC), may already be able to demonstrate quantum advantage over classical computers for select problems [2,3]. In particular, the hybrid quantum-classical variational quantum eigensolver (VQE) [4,5] may have sufficiently low experimental requirements to allow estimation of ground-state energies of quantum systems that are difficult to simulate purely classically [6–9]. To date, VQEs have been used to study small examples of systems that are difficult to simulate purely classically [10–15], strongly correlated magnetic models [15], and the Schwinger model [18]. Although these experimental efforts have achieved impressive coherent control of up to 20 qubits, the error in the resulting estimations has remained relatively high due to performance limitations in the NISQ hardware. Consequently, much focus has recently been placed on developing error mitigation techniques that offer order-of-magnitude accuracy improvement without the costly overhead of full QEC. This may be achieved by using known properties of the target state, e.g., by checking known symmetries in a manner inspired by QEC stabilizer measurements [19,20], or by expanding around the experimentally obtained state via a linear (or higher-order) response framework [21]. The former, termed symmetry verification (SV), is of particular interest because it is comparatively low-cost in terms of required hardware and additional measurements. Other mitigation techniques require understanding the underlying error models of the quantum device, allowing for an extrapolation of the calculation to the zero-error limit [22–24], or the summing of multiple calculations to probabilistically cancel errors [23,25,26].

In this Rapid Communication, we experimentally demonstrate the use of SV to reduce the error of a VQE estimating the ground-state energy and the ground state of the H2 molecule by one order of magnitude on average across the bond-dissociation curve. Using two qubits in a circuit QED processor, we prepare a variational ansatz state via an exchange gate that finely controls the transfer of population within the single-excitation subspace while respecting the underlying symmetry of the problem (odd two-qubit parity). We show that SV improves the energy and state estimates by mitigating the effect of processes changing total excitation number, specifically qubit relaxation and residual qubit excitation. We do this through a full density-matrix simulation that matches the experimental energy and state errors with and without SV, and then using this simulation to dissect the contribution of each error source. Finally, we explore the limitations of SV arising from statistical measurement noise, and find that enforcing the positivity of the fermionic 2-reduced density matrix ties the improvement in energy estimation from SV to the improvement in ground-state fidelity (which was previously not the case).

A VQE algorithm [4,5] approximates the ground state \( \rho^{(0)} \) of a Hamiltonian \( \hat{H} \) by a variational state \( \rho^{(\text{raw})}(\vec{\theta}) \), with \( \vec{\theta} \) a set of parameters that control the operation of a quantum device. These parameters are tuned by a classical optimization routine to minimize the variational energy \( E(\vec{\theta}) = \text{Tr}[\rho^{(\text{raw})}(\vec{\theta})\hat{H}] \). In practice, this is calculated by expanding \( \rho^{(\text{raw})}(\vec{\theta}) \) and \( \hat{H} \) over
the $N$-qubit Pauli basis $\mathbb{D}^N := \{I, X, Y, Z\}^\otimes N$, 
\begin{equation}
\rho^{(\text{raw})}(\vec{\theta}) = \frac{1}{2^N} \sum_{P \in \mathbb{D}^N} \rho^{(\text{raw})}_{P}(\vec{\theta}) \hat{P}, \quad \hat{H} = \sum_{P \in \mathbb{D}^N} h_P \hat{P},
\end{equation}
where the Pauli coefficients are given by $\rho^{(\text{raw})}_{P}(\vec{\theta}) = \text{Tr}[\hat{P} \rho^{(\text{raw})}(\vec{\theta})]$. The variational energy may then be calculated as 
\begin{equation}
E^{(\text{raw})}(\vec{\theta}) = \sum_{P \in \mathbb{D}^N} \rho^{(\text{raw})}_{P}(\vec{\theta}) h_P.
\end{equation}

For example, consider the $\text{H}_2$ molecule studied in this work. Mapping the Hamiltonian of this system (in the STO-3G basis) onto four qubits via the Bravyi-Kitaev transformation [27] and further reducing dimensions by projecting out two noninteracting qubits [10] gives 
\begin{equation}
\hat{H}_{\text{H}_2} = h_{11} \hat{I} + h_{2Z} \hat{Z} + h_{IZ} \hat{Z}
+ h_{XX} \hat{X} + h_{YY} \hat{Y} + h_{ZZ} \hat{Z},
\end{equation}
where coefficients $h_P$ depend on the interatomic distance $R$. These coefficients may be determined classically using the OPENFermion [28] and PSI4 [29] packages. The Pauli coefficients $\rho^{(\text{raw})}_{P}$ of the density matrix $\rho^{(\text{raw})}$ are extracted by repeated preparation and (partial) tomographic measurements of the ansatz state. As one only needs those Pauli coefficients $\rho^{(\text{raw})}_{P}$ with nonzero corresponding Hamiltonian coefficients $h_P$, one need not perform full tomography of $\rho^{(\text{raw})}$. However, in a small-scale experiment, full state tomography of $\rho^{(\text{raw})}$ may still be feasible, and may provide useful information for the purposes of benchmarking. In particular, the fidelity of $\rho^{(\text{raw})}$ to $\rho^{(0)}$, 
\begin{equation}
E^{(\text{raw})} = \text{Tr}[\rho^{(\text{raw})} \rho^{(0)}],
\end{equation}
is a more rigorous measure of the ability to prepare the ground state than the energy error, 
\begin{equation}
\Delta E^{(\text{raw})} = \text{Tr}[\rho^{(\text{raw})} - \rho^{(0)} \hat{H}].
\end{equation}
Error mechanisms such as decoherence pull $\rho^{(\text{raw})}$ away from $\rho^{(0)}$, decreasing $F$ and increasing $\Delta E$.

These errors may be mitigated by using internal symmetries $\hat{S} \in \mathbb{D}^N$ [30] of the target problem, such as parity checks [19,20]. These checks project $\rho^{(\text{raw})}$ to a symmetry-verified matrix $\rho^{(\text{SV})}$ that lies in the $\{\hat{S}\} = s$ subspace of the symmetry. This projection could be performed via direct measurement of $\hat{S}$ on the quantum device, but one may instead extract the relevant terms of the density matrix $\rho^{(\text{SV})}$ in postprocessing:
\begin{equation}
\rho^{(\text{SV})}_{P} = \frac{\rho^{(\text{raw})}_{P} + S \rho^{(\text{raw})}_{S}}{1 + S \rho^{(\text{raw})}_{S}}.
\end{equation}
The right-hand side may be obtained by partial tomographic measurement of the ansatz state, with at most twice the number of Pauli coefficients that need to be measured. This upper bound is not always achieved. For example, the $\hat{H}_{\text{H}_2}$ Hamiltonian has a $\hat{S} = ZZ$ symmetry, which maps the nonzero Pauli terms in $\hat{H}_{\text{H}_2}$ to other nonzero Pauli terms in $\hat{H}_{\text{H}_2}$. Symmetry verification in this problem then does not require any additional measurements to estimate $E^{(\text{SV})}$ beyond those already required to estimate $E^{(\text{raw})}$. Even when it does require additional measurements, SV remains attractive because it does not require additional quantum hardware or knowledge of the underlying error model. One can show that the SV state $\rho^{(\text{SV})}$ may be equivalently obtained via a variant of the quantum subspace expansion (QSE) [21], suggesting an alternative name of S-QSE [19].

One may further minimize the error in a quantum algorithm by tailoring the quantum circuit or the gates within. In a VQE, one wishes to choose a variational ansatz motivated by the problem itself [10,31] while minimizing the required quantum hardware [14]. To balance these considerations, we suggest constructing an ansatz from an initial gate set that is relevant to the problem at hand. For example, in the electronic structure problem, the quantum state is generally an eigenstate of the fermion number. When mapped onto qubits, this often corresponds to a conservation of the total qubit excitation number. Gates such as single-qubit $Z$ rotations, two-qubit C-phase [32], and two-qubit iSWAP [33] gates preserve this number, making these gates a good universal gate set (within the target subspace [34]) for quantum simulation of electronic structure. In the example of $\text{H}_2$, the total two-qubit parity $(ZZ)$ is indeed conserved and the ground state at any $R$ may be generated by applying to $|01\rangle$ or $|10\rangle$ an exchange gate 
\begin{equation}
U_\theta = \begin{pmatrix} 1 & 0 & 0 & 0 \\
0 & \cos \theta & i \sin \theta & 0 \\
0 & i \sin \theta & \cos \theta & 0 \\
0 & 0 & 0 & 1 \end{pmatrix}
\end{equation}
with $R$-dependent optimal exchange angle $\theta$ and a follow-up phase correction on one qubit.

We now experimentally investigate the benefits of SV in the VQE of $\text{H}_2$ using two of three transmon qubits in a circuit QED quantum processor (see details in [35]). The two qubits ($Q_0$ and $Q_1$) are coupled by a common bus resonator, and have dedicated microwave lines for single-qubit gating, flux bias lines for local and nanosecond-scale control of their frequency, and dedicated readout resonators coupling to a common feedline for independent readout by frequency multiplexing. We prepare the ansatz state with an efficient circuit [Fig. 1(a)] that first excites $Q_1$ with a $\pi$ pulse to produce the state $|10\rangle$, and then applies a square flux pulse of fixed duration and amplitude to $Q_0$, bringing it into or near resonance with $Q_1$ to coherently exchange the excitation population. A plot of population exchange as a function of flux-pulse amplitude and duration [Fig. 1(b)] reveals the expected chevron pattern that is the hallmark of coherent population exchange between the two qubits, albeit with some asymmetry arising from the bandwidth limitation of the flux-control line. We make use of the square-pulse duration (1 ns resolution) and amplitude (0.5 mV resolution) as coarse and fine knobs, respectively, to control population exchange. We choose 1500 combinations of pulse duration and amplitude settings to parametrize an experimental knob $\hat{\theta}$ [Fig. 1(d)] capable of finely controlling population exchange like $\theta$ in Eq. (7) over the range $[0, \pi/4]$ [Fig. 1(c)]. The circuit concludes with simultaneous prerotation gates on both qubits followed by simultaneous measurement of both qubits, in order to perform tomography of the prepared ansatz state. To fully reconstruct the state, we use an overcomplete set of 36 prerotation pairs and extract estimates of the average measurement for each qubit as well as their shot-to-shot.
correlation using $N_{\text{meas}}$ measurements per prerotation. Note that while the flux pulse implements the exchange gate of Eq. (7) with additional single-qubit phase rotations, the correction for these phase rotations can be performed virtually from the fully reconstructed state.

We now optimize the VQE to approximate the ground-state energy and ground state of H$_2$. At each chosen $R$, we employ the covariance matrix adaptation evolution strategy (CMA-ES) optimization algorithm [36], using $E^{(\theta)}$ as cost function and $\bar{\theta}$ as single variational parameter. The evolutionary strategy optimizes $\bar{\theta}$ over repeated generations of $N_{\text{pop}} = 10$ samples of $E^{(\theta)}$, each calculated from a raw density matrix $\rho^{(\theta)}$, using linear inversion of $N_{\text{meas}} = 10^5$ [37]. Optimizations have a hard-stop criterion of $[\text{Fig. 2(a)}\text{ inset}]$ 20 generations ($\sim 2$ h). The converged state is finally reconstructed with greater precision, using $N_{\text{meas}} = 10^5$. Figure 2 shows the resulting energy estimate for 12 values of $R$ and the reconstructed optimized state at three such distances. These tomographs show that the optimal solutions are concentrated in the single-excitation subspace of the two qubits, with two-qubit entanglement increasing as a function of $R$.

Performing the described symmetry verification procedure on the converged states shows improvement across the entire bond-dissociation curve. To quantify the improvement, we focus on the energy error $\Delta E$ and the infidelity $1 - F$ to the true ground state, with and without SV (Fig. 3). SV reduces the energy error by an average factor $\sim 10$ and reduces the infidelity by an average factor $\sim 9$. In order to quantitatively understand the limits of the VQE optimization, and to clearly pinpoint the origin of the SV improvement, we simulate the experiment via the density-matrix simulator $\text{quantumsim}$ [38], using an error model built from independently measured experimental parameters [35]. We build the error model incrementally, progressively adding optimization inaccuracy (the difference between the state ideally produced by the converged $\theta$ and the true ground state); dephasing on both qubits (quantified

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**FIG. 1.** Quantum circuit and energy landscape of the variational eigensolver. (a) Quantum circuit for generating and measuring the variational ansatz state. (b) Coherent excitation exchange, produced as $Q_0$ is fluxed into resonance with $Q_1$ by a square flux pulse of fixed amplitude ($x$ axis) and duration ($y$ axis). The amplitude controls the frequency to which $Q_0$ is pulsed ($\sim 1.428$ V bringing it on resonance with $Q_1$). (c) Zoom-in of (b) into the region used in the experiment to control the exchange of population between $Q_0$ and $Q_1$. Colored lines illustrate the combinations of square-pulse amplitudes and duration used to achieve fine adjustment of $\bar{\theta}$. (d) Excitation of $Q_0$ for the combinations of pulse amplitudes and duration marked by colored lines in (c), showing the matching of the experimentally defined $\bar{\theta}$ to the target $\bar{\theta}$ defined in Eq. (7) (black dashed curve). Colors [matching (c)] correspond to pulse duration. (e) Landscape of energies $E^{(\theta)}(\bar{\theta}, R)$ as a function of the experimentally defined $\bar{\theta}$ angle and the interatomic distance $R$.

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**FIG. 2.** Convergence of the VQE algorithm. (a) Experimental VQE estimate of H$_2$ ground-state energy as a function of interatomic distance $R$. At each chosen $R$, we minimize the raw energy $E^{(\theta)}$ (blue data points) over the variational parameter $\bar{\theta}$ using the CMA-ES evolutionary algorithm [36]. Applying SV to the converged solution (orange data points) lowers the energy estimate toward the exact solution (dashed curve). Inset: A typical optimization trace for the convergence of the energy estimate. (b)-(d) The reconstructed density matrices of the converged states at (b) $R = 0.25$ Å, (c) $R = 0.80$ Å, and (d) $R = 2.00$ Å, showing that the converged states lie mostly in the single-excitation subspace, and that entanglement increases with the interatomic distance $R$. 

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FIG. 3. Impact of SV in ground-state energy and state fidelity, and dissected error budget. (a) Experimental (solid circles) energy error \( \Delta E \) without and with SV compared to the result (empty circles and dashed line) of a full density-matrix simulation using the full error model. The contributions from optimizer inaccuracy, qubit dephasing, residual qubit excitations, and increased Qo dephasing during the exchange gate are shown as shaded regions for the case of no SV applied. Without SV, \( \Delta E \) is clearly dominated by residual qubit excitation. (b) Zoom-in on experimental and simulated \( \Delta E \) with SV and corresponding error budget. With SV, the effects of residual excitation and qubit relaxation are successfully mitigated, as predicted in Ref. [19]. The remaining energy error is dominated by optimizer inaccuracy. Simulation error bars are obtained by modeling measured fluctuations of \( T_1, T_2^*, \) and residual excitation. (c) Experimental (solid circles with error bars) infidelity to the true ground state without and with SV compared to simulation using the full error model (empty circles and dashed line). Error bars are propagated through the linear inversion procedure for experiment and calculated from sampling noise for simulations. For simulations, error bars are smaller than the markers.

by the measured Ramsey dephasing times \( T_2^* \); relaxation on both qubits (quantified by the measured relaxation times \( T_1 \)); residual qubit excitations (measured from single-shot histograms with each qubit prepared in \( |0\rangle \)); and increased dephasing of \( Q_0 \) during the exchange gate (quantified by its reduced \( T_2^* \) when tuned into the exchange interaction zone). By plotting the errors from each increment of the model, we are able to dissect the observed experimental error into its separate components without [Fig. 3(c)] and with [Fig. 3(b)] SV. Measured temporal fluctuations of dephasing, relaxation, and residual excitation are used to obtain simulation error bars.

The simulation using the full error model shows fairly good matching with experiment for both the ground-state energy error [Figs. 3(a) and 3(b)] and the state infidelity [Fig. 3(c)], without and with SV. The error model dissection shows that the energy error when not using SV is dominated by residual qubit excitations. This is remarkable as the calibrated residual excitations are only 0.25% for \( Q_0 \) and 1.34% for \( Q_1 \) [35]. The improvement from SV results from the mitigation of errors arising from these residual excitations and from qubit relaxation. This is precisely as expected: These error mechanisms change total qubit excitation number and violate the underlying ZZ symmetry. Using SV changes the dominant error mechanism from residual qubit excitation to optimization inaccuracy, which is bounded by the sampling noise during the optimization itself (where \( N_{\text{meas}} = 10^5 \), rather than the sampling noise from the final step (where \( N_{\text{meas}} = 10^7 \)). This error could be reduced experimentally by increasing \( N_{\text{meas}} \) during the optimization, at the cost of increased convergence time. The improvement in state infidelity by SV can be explained along similar lines. We observe some increased deviations between the observed and simulated state infidelity at large \( R \). We attribute these to limitations in our modeling of error during the exchange gate (whose duration increases with \( R \)).

VQEs rely on variational bounding to ensure that the obtained approximation to the ground-state energy is accurate, but this is only guaranteed when the experimental results correspond to a physical state. Our method for calculating the ground-state energy [Eq. (1)] independently estimates each Pauli coefficient of the density matrix with error \( \propto N_{\text{meas}}^{-1/2} \). Such estimation cannot guarantee a set of Pauli coefficients that could have come from a positive density matrix. This in turn breaks the variational lower bound on the energy estimate, and increases the error in estimates of other properties of the true ground state [39,40]. As experimental error is reduced, \( \rho^{(\text{raw})} \) tends toward a rank-1 density matrix, increasing its chance of being unphysical [40]. Moreover, \( \rho^{(\text{SV})} \) is a lower-rank density matrix than \( \rho^{(\text{raw})} \) (being projected onto a subspace of the Hilbert space), which implies that unphysicality may be enhanced by SV. The variance in a given term \( \rho_{F} \) post-SV can be calculated as

\[
\text{Var}(\rho_{F}^{(\text{SV})}) \approx \frac{3N_{\text{meas}}}{N_{\text{meas}}(1 + \text{Tr}(\rho^{(\text{raw})}S))}.
\]

SV has maximal impact on the quantum state precisely when this denominator is small, so this represents a natural bound for the power of SV as an error mitigation strategy.

The effect of sampling noise may be mitigated somewhat by restricting the fermionic 2-reduced density matrix to be positive (which may be completed in polynomial time) [39]. To investigate the effect of such mitigation, we bin the data used for final tomography of converged states to construct 100 density matrices with \( N_{\text{meas}} = 10^3 \) at each \( R \), thus increasing the sampling noise by a factor of 10. We wish to study the relative improvement of SV in the two figures of merit, which we quantify as

\[
\eta_E = \frac{\Delta E^{(\text{raw})}}{\Delta E^{(\text{SV})}} \quad \text{and} \quad \eta_F = \frac{1 - F^{(\text{raw})}}{1 - F^{(\text{SV})}},
\]

when physicality of the raw density matrices is enforced and not. To enforce physicality, we employ a convex optimization routine to find the closest positive semidefinite matrix to the experimentally measured \( \rho^{(\text{raw})} \) (closest in the \( L^1 \) norm sense on the space induced by the the Pauli basis). We then apply
Fig. 3 is split into 100 sample simulations for each $R$, increasing the sampling noise by a factor of 10 and making it comparable to other sources of experimental error. For each sample, we plot (red) the relative energy error and infidelity [Eq. (9)]. Values below 1 (dashed lines) indicate that SV has not provided an improvement, as may be the case when the density matrix has negative eigenvalues. We restore the improvement from SV by constraining the positivity of the 2-reduced density matrix [39] (green). Histograms on the top and right axes show the marginal distribution of the two scatter plots. When the density matrices are constrained to be positive, we observe the points fall along the line $y = x$ (blue dashed line), indicating that SV improves both metrics by the same amount.

Fig. 4. Constraining positivity with symmetry verification to mitigate the effect of sampling noise. The experimental data from Fig. 3 is split into 100 sample simulations for each $R$, increasing the sampling noise by a factor of 10 and making it comparable to other sources of experimental error. For each sample, we plot (red) the relative energy error and infidelity [Eq. (9)]. Values below 1 (dashed lines) indicate that SV has not provided an improvement, as may be the case when the density matrix has negative eigenvalues. We restore the improvement from SV by constraining the positivity of the 2-reduced density matrix [39] (green). Histograms on the top and right axes show the marginal distribution of the two scatter plots. When the density matrices are constrained to be positive, we observe the points fall along the line $y = x$ (blue dashed line), indicating that SV improves both metrics by the same amount.

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