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A randomized measurement toolbox for an interacting Rydberg-atom quantum simulator

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Abstract

We present a toolbox to probe quantum many-body states implemented on Rydberg-atoms quantum hardware via randomized measurements. We illustrate the efficacy of this measurement toolbox in the context of probing entanglement, via the estimation of the purity, and of verifying a ground-state preparation using measurements of the Hamiltonian variance. To achieve this goal, we develop and discuss in detail a protocol to realize independent, local unitary rotations. We benchmark the protocol by investigating the ground state of the one-dimensional Su–Schrieffer–Heeger model, recently realized on a chain of Rydberg atom, and the state resulting after a sudden quench in a staggered XY chain. We probe the robustness of our toolbox by taking into account experimental imperfections, such as pulse fluctuations and measurement errors.

1. Introduction

Synthetic quantum systems, composed of, e.g. neutral atoms [1, 2], ions [3], superconducting qubits [4], allow us to engineer spin-lattice models or implement quantum algorithms on qubit registers, with precise control over geometry and interactions. Among these platforms, Rydberg atoms have emerged as a promising system. They can be described in good approximation in terms of qubits, with the spin-up state $|\uparrow\rangle \equiv |1\rangle$ encoded by a Rydberg state, and the spin-down $|\downarrow\rangle \equiv |0\rangle$ encoded by another Rydberg state, or an atomic ground state [2]. One of the most relevant assets for Rydberg quantum technologies is the long qubit lifetime, which scales as $n^4$, where $n \sim 50 – 100$ is the atom principal quantum number. In addition, interactions between Rydberg qubits are naturally obtained via the dipole–dipole interactions, whose characteristic energy scales as $n^6$ in the resonant regime, and $n^{11}$ in the off-resonant van der Waals regime. Finally, Rydberg atoms can be placed on almost arbitrary geometries using optical forces generated, e.g. by optical tweezers [5]. In particular, recent experimental progresses in this direction allowed to experimentally study strongly correlated quantum states with hundreds of qubits in two dimensional lattice models [6, 7]. On the quantum computing side, Rydberg atom platforms have demonstrated remarkable performances in terms of scalability, connectivity, and gate fidelities [8–10].

In order to take advantage of all the promising aspects of Rydberg atoms for quantum technologies, it is desirable to equip such platforms with a measurement toolbox to extract key physical quantities, such as fidelities and entanglement, and in a state-agnostic way. A promising approach in this context consists in
and propose an experimental demonstration of the SSH model, in order to verify the adiabatic preparation of the ground state. We also show the measurement of the Hamiltonian variance, which can be used to verify experimentally ground state preparation (see figure 1) estimating the Hamiltonian variance of \( |\psi_0\rangle \) and the purity of system bipartitions with size \( \ell \).

Figure 1. Pictorial representation of the randomized measurements toolbox. Starting from a separable state (all sites in \(|\downarrow\rangle\)), an adiabatic state preparation \( P \) is applied to implement the target state of interest \( |\psi_0\rangle \) [11]. Then, we randomly choose and apply local unitaries for each lattice site, repeating the protocol for \( N_S \) samplings. For each sampling \( \{U_i\} \), we estimate the probabilities \( P_{\ell}(s) \) (see equation (2)) estimating the Hamiltonian variance of \( |\psi_0\rangle \) and the purity of system bipartitions with size \( \ell \).

Using randomized measurements (RMs) based on performing random single-qubit rotations followed by measurements in the computational basis [12–14] (see figure 1).

RMs [15] have been used to estimate the purity and the second Rényi entropy \( S_2 = -\log \mathrm{Tr}[\rho^2] \) of (sub-)systems consisting of up to ten qubits in a trapped ion [16–19] and superconducting qubit quantum simulator [20, 21]. An alternative purity estimator can be obtained via the classical shadows formalism [14, 18] (see also below). In addition, it has been proposed to use RMs to reveal other properties of many-body quantum states. This concerns for instance the fidelity of quantum states realized in different experiments [18, 22] and versus an ideal theoretical target state, as well as many-body topological invariants associated with (symmetry-protected) topological phases [23, 24]. Moreover, beyond the immediate opportunity to extract via RMs entanglement entropies and related quantities for quantum simulation, our toolbox enables the measurement of arbitrary observables based on the classical shadow formalism [14]. This is in particular relevant in the context of variational quantum optimization algorithms [25, 26], where an observable cost function is repetitively measured.

In the aforementioned local implementation of the minimal random unitaries, interactions were absent during the local rotations protocol. Analogously, recent Rydberg atom quantum simulation and computation platforms exploit a qubit implementation in which the \(|0\rangle\) and \(|1\rangle\) states correspond to hyperfine ground states [10, 27–30]. In this case, it is possible to locally manipulate the quantum many-body state in the absence of atom–atom interactions.

In this work, we answer the question of whether the local unitaries apparatus can also be exploited in cases when the atom–atom interactions cannot be avoided. To do so, we equip an interacting Rydberg atom quantum simulator with a tailored RM toolbox based on local unitaries, i.e. spin qubit rotations. Despite the protocol we present constitutes a receipt tailored for a particular Rydberg atom setup, it nevertheless demonstrates that the local unitaries apparatus can also be extended to the case of qubits where interactions cannot be turned off. Our approach complements a recent work employing quasi-local ‘scrambled’ unitaries generated by Rydberg interactions and used for benchmarking and fidelity estimation [31]. Our approach fully relies on experimental tools that are currently available. It consists in implementing RMs via random single-qubit rotations combining local light shifts and time-varying global microwave drives to the atoms. Our approach proposes directly operating on the Rydberg manifold, allowing to realize random single qubit rotations in parallel, i.e. with a duration that does not scale with the number of qubits. One can also straightforwardly adapt our protocols for hyperfine state qubits, e.g. replacing microwave drives by Raman pulses.

Importantly, we show that the effects of the interactions between Rydberg atoms during the generation of such single-qubit rotations can be made negligible using an optimized pulse sequence for the different drives. We illustrate the use of the RM toolbox for the measurement of the purity \( p_2 \) [giving access to the entanglement Rényi entropy \( S_2 = -\log_2(p_2) \)] in the Su–Schrieffer–Heeger (SSH) model, and of the Hamiltonian variance, which can be used to verify experimentally ground state preparation [14, 32]. We also estimate the purity of a state resulting from the dynamics after a sudden quench with a staggered XY model. Our simulations take into account the most important realistic error sources and analyze the role of statistical errors. We conclude that RMs can be implemented via the presented approach in existing Rydberg platforms.

In the following, we describe our general RM toolbox in section 2 and propose an experimental implementation in section 3. In section 4, we illustrate our approach in the context of characterization of topological phases with entanglement entropies. We also show the measurement of the Hamiltonian variance of the SSH model, in order to verify the adiabatic preparation of the ground state.
2. Local random unitaries toolbox

2.1. Probability estimation

RMs provide a powerful toolbox to investigate the properties of quantum many-body systems beyond standard low-order correlation functions [12–14, 16, 18, 23, 24, 33–46]. In the following, we outline the RM protocol employing local (single-spin) random unitary operations. To this end, we consider a quantum state $\rho$ defined on a lattice of qubits with size $L$ and associated Hilbert space of dimension $2^L$. We denote its computational $z$-basis with $\{|s\}\}$ by bitstrings $s = (s_1, \ldots, s_L)$ and $s_m = 0, 1$ for $m = 1, \ldots, L$. A RM comprises the following steps: (i) a random unitary $U = \bigotimes_{m=1}^{L} u_m$ is applied to $\rho$, where each $u_m$ is sampled independently from an appropriate ensemble of local (single-spin) unitary transformations, typically a unitary 2-design [47, 48]. Examples of such unitary 2-designs include continuous single-spin rotations which cover the Bloch sphere uniformly (the Haar measure on the unitary group $U(2)$) as well as the (discrete) single-qubit Clifford group [47, 48]. (ii) This is followed by a measurement in the computational $z$-basis with outcome bitstring $s = (s_1, \ldots, s_L)$. This procedure is then repeated with the same unitary $U$ to obtain an estimate of the probabilities $P_U(s) = \text{Tr} [U\rho U^\dagger |s\rangle \langle s|]$, and subsequently with newly sampled unitaries to estimate the average over the ensemble of unitary transformations. We denote the number of repetitions with the same random unitary with $N_{\text{meas}}$ and the number of applied unitaries $U$ with $N_U$ such that the rotations protocol is repeated $N_{\text{rot}} = N_U \times N_{\text{meas}}$ times in total.

In this work, we choose to use local random unitary operations $u_m$ which are sampled from the discrete, finite single-qubit Clifford group $U(2)$. Since for RMs, the application of a random unitary is directly followed by a computational $z$-basis measurement in the $z$-direction, the application of randomly sampled single-qubit Clifford gates is equivalent to sampling the $L$ measurement directions $v_m$ among a finite set of three mutually orthogonal directions, as for example $\{x, y, z\}$ (see [14]). In this case, the corresponding set of transformations is $R = \{e^{-i\pi/sx}, e^{-i\pi/4sy}, 1\}$, that rotates each direction onto the measurement axis $z$.

2.2. Purity estimation

A key application of RMs is the estimation of the purity of quantum states to characterize the coherence of the underlying quantum device and to reveal entanglement [12–14, 16, 33, 35, 49]. In the following, we consider a system with size $L$ and with basis $\{|s\}$, and a sub-system $A$ with size $N_A$. The purity $\text{Tr} [\rho_A^2]$ of the reduced density matrix $\rho_A$ of $A$ can be estimated, following the procedure presented in [16, 35]. Given the estimates of the probabilities $P_U(s)$, one obtains estimates of the probabilities $P_U(s_A) = \sum_{s_{A'} = s_A} P_U(s)$ of computational basis states $|s_A\rangle$ for any subsystem $A$ via post-processing. Then, the purity $\text{Tr} [\rho_A^2]$ is obtained from second-order correlations of the probabilities $P_U(s_A)$ via

$$\text{Tr} [\rho_A^2] = 2^{N_A} \sum_{s_A, s'_A} (-2)^{-D(s_A, s'_A)} P_U(s_A) P_U(s'_A) . \quad (1)$$

Here, $D(s_A, s'_A)$ denotes the Hamming distance of the bitstrings $s_A$ and $s'_A$ and ... the ensemble average over the local random unitaries. Equation (1) represents an exact relation in the limit of $N_{\text{meas}} \to \infty$ and when the local random unitaries are averaged over a complete unitary 2-design. In practice, statistical errors arise from a finite number of measurements $N_{\text{meas}}$ per unitary and a finite number $N_U$ of local random unitaries sampling the ensemble average. Numerical and analytical analysis of such statistical errors showed that the total number of experimental runs $N_U N_{\text{meas}}$ to estimate the purity with high confidence and probability scales approximately as $2^{N_{\text{meas}}} / b \approx 1$. The exact value of $b$ and the optimal ratio $N_U / N_{\text{meas}}$ depends on the state of interest and the required precision. We note that this represents a substantial improvement compared to full quantum state tomography, requiring at least $2^N / N$ experimental runs with $b' \gtrsim 2$ (see e.g. [50]). The scaling of statistical errors with system size can furthermore be substantially improved via importance sampling [49].

2.3. Estimating expectation values of arbitrary observables

The same RM data can be used to estimate expectation values $\text{Tr} [O \rho]$ of arbitrary observables $O$ [14]. Utilizing the tomographic completeness of RMs [14, 35, 51], the expectation value of arbitrary observables $O$ can be obtained via

$$\text{Tr} [O \rho] = 2^N \sum_{s, s'} (-2)^{-D(s, s')} P_U(s) \langle s'|OUO^\dagger |s'\rangle . \quad (2)$$

Differently from equation (1), this expression is linear in the experimentally estimated outcome probabilities $P_U(s)$. Hence, the procedure to estimate expectation values $\text{Tr} [O \rho]$ is as follows: In the experiment, we estimate outcome probabilities $P_U(s) = \langle s' | U \rho U^\dagger |s\rangle$, as in the case of the purity estimation. On a classical
computer, we calculate the corresponding matrix elements $\langle s | U O U^\dagger | s \rangle$ (for the same unitaries $U$ which have been applied in the experiment). Then, we cross-correlate according to equation (2).

Observable estimation with RMs has been formalized and rigorous error bounds have been obtained via the classical shadows formalism [14]. There, it has been shown that statistical errors depend on the set of observables $O$ of interest. Below, we consider the specific case of $O = H$ ($O = H^s$) with $H$ being a Hamiltonian with $k$-body interactions. Then, in the limiting case $N_{\text{pre}} = 1$, $N_U \sim 2^k \log(N)$ $(N_U \sim 2^k \log(N^2))$ random unitaries are required to estimate $\text{Tr}[H \rho] (\text{Tr}[H^s \rho])$ with high confidence and probability [14]. This number can be substantially further decreased using derandomization techniques [52]. We furthermore note that the equation (2) can be generalized to estimate expectation values of arbitrary multi-copy observables [14]. This enables, for instance, the detection of mixed state entanglement via higher-order moments of (the partial transpose) of the density matrix $\rho$ [18] (see also [38]), of symmetry-resolved entanglement entropies [53], and of the quantum Fisher information [41].

In contrast to the purity estimation formula, in equation (2) we exploit explicitly the knowledge of the random unitaries $U$ to calculate the required matrix elements $\langle s | U O U^\dagger | s \rangle$. Thus, any miscalibration between the local random unitaries actually applied in the experiment and those applied on the classical computer affects the estimation of $\text{Tr}[\rho O]$ [37, 54]. We will discuss the influence of such implementation errors in detail below. In addition, the robustness can be improved via calibration experiments with simple states which can be prepared with high fidelity [52, 54, 55]. In the next section, we describe our Rydberg quantum optics model, and the corresponding implementation of RMs.

3. Proposal to experimentally implement the RMs toolbox

3.1. The model

We consider an array of atoms (either one-dimensional as shown here, or two-dimensional), made to interact by exciting them to Rydberg states [2]. In particular, we focus on the setup used to observe symmetry-protected topological phases in a SSH chain (see, e.g., [11]). By encoding pseudo-spin-1/2 states in two dipole-coupled Rydberg levels (such as $nS$ for $| \downarrow \rangle \equiv | 0 \rangle$ and $nP$ for $| \uparrow \rangle \equiv | 1 \rangle$, with $n \sim 60$), the dipole–dipole interaction at work between the atoms implements the XY spin Hamiltonian $\sum_{i < j} J_{ij} \sigma_i^x \sigma_j^x + \text{h.c.}$, with $J_{ij}$ decaying as $1/r_{ij}^3$ with the distance $r_{ij}$ between the atoms $i$ and $j$, and $\sigma^\pm = (\sigma_x \pm i \sigma_y)/2$ are linear combinations of the usual Pauli matrices.

To manipulate the internal spin states and thus implement local rotations, we start from the experimental setup used in [11], sketched in figure 2. In particular, we can first manipulate them globally by using microwave pulses with a Rabi frequency $\Omega(t)$ and a detuning $\Delta(t)$. For local manipulation, we add a local light shift with a tightly focused laser beam (for instance coupling off-resonantly the $nS$ state to a low-lying $P$ state such as the $6P$ state for Rb) on a selected atom in order to tune the qubit frequency into (or out of) resonance with the microwave field [56]. A spatial light modulator (SLM) is used to program at will the spatial dependence of these addressing beams, while the (global) time dependence $f(t)$ of the intensity of the addressing beams is set with an acousto-optic modulator, placed before the SLM, and that allows for the generation of fast pulses.

By taking all the available terms into account, the experimental Rydberg Hamiltonian describing the local transformations is

$$H_{\text{prot}}(t) = \sum_{m=1}^{L} \left[ \frac{\Omega(t)}{2} \sigma_m^x - \frac{\Delta(t) - f(t)}{2} \delta_{m \alpha_m} n_m \right],$$

where $\sigma_m^\alpha$ are the Pauli matrices acting on site $m \in [1, L]$, $\alpha_m \in \{1, 2, 3\}$ and $n_m = (\sigma_m^3 + 1)/2$.

During the application of the rotation protocol, the state is evolving under the total Hamiltonian

$$H(t) = H_{\text{prot}}(t) + H_{\text{mod}},$$

where $H_{\text{mod}}$ is the static model Hamiltonian describing the interactions between Rydberg atoms, see equation (6) below. The interaction terms can create spurious correlations between the local rotations and affect the estimation of the probabilities $P_O(s)$. In the following, we determine the pulses of the rotation protocol after fixing the maximum amplitude of the Hamiltonian $H_{\text{mod}}$ parameters $J$ to be $< 1\text{MHz}$.

3.2. Experimental proposal

In the following, we exploit the Hamiltonian $H_{\text{prot}}(t)$ to simultaneously implement three transformations $R = \{ R_1, R_2, R_3 \}$ that rotate the measurement axis $z$ onto three mutually perpendicular directions. First, we consider the idealized limit of very short protocol time $T$ (i.e. $JT \ll 1$, to neglect the influence of
interactions) and an arbitrarily large detuning $\Delta(t)$ much larger than the Rabi frequency $\Omega(t)$. The local potentials $f(t)\delta_{\alpha n} \sim \Delta(t)$ are fixed to implement the different rotations. In this limit, we can present analytical pulse sequences to realize $R = \{e^{-i\pi/4\alpha\sigma_z}, e^{-i\pi/4\alpha\sigma_x}, 1\}$ exactly. We consider constant, square pulses. The Rabi frequency pulse $\Omega(t)$ has a duration of $T$. It implements two successive $\pi/2$ rotations and satisfies the condition $\Omega(t)T/2 = \pi/2$. Instead, the detuning term $\Delta(t)$ is null in the first half of the protocol, while we set $\Delta(t)T/2 = \pi/2$ in the second half (this choice will be clarified in the following).

The transformation $R_1$ is implemented by setting $Tf(t)\delta_1 = 0$. During the first half of the total time interval $T$, we have $T\Delta(t) = 0$ and the Rabi frequency term implements a $\pi/2$ pulse. In the second half-time, the detuning pulse by $\Delta(t) \gg \Omega(t)$ shifts the transition between the two levels off-resonant. Note that the $\Delta(t)$ pulse adds a phase that does not affect the measurement along the $z$ axis. In order to implement the $R_2$ transformation, we decompose the rotation around the $y$ axis into the sequence of rotations $e^{-i\pi/4\alpha\sigma_x}e^{-i\pi/4\sigma_z}e^{i\pi/4\alpha\sigma_x}$. Then, we ignore the last $\sigma_z$ rotation which does not influence the final measurement outcome, and resulting in the rotations $e^{-i\pi/4\alpha\sigma_x}e^{i\pi/4\alpha\sigma_x}$ to be implemented. The rotation around $z$ is realized by setting $Tf(t)\delta_2/2 = \pi/2$, with $f(t)\delta_2 \gg \Omega(t)$, in the first half of the protocol. In the second half, the choice of $\Delta(t)$ realizes the condition $\Delta(t) - f(t)\delta_2 = 0$, and the pulse $\Omega(t)$ implements the rotation around $x$.

Finally, the rotation $R_3$ is realized by the pulse with amplitude $f(t)\delta_3 \gg \Omega(t)$ in the first half of the protocol and $\Delta(t) - f(t)\delta_3 \gg \Omega(t)$ in the second half (recall that $T\Delta(t) = 0$ during the first half of the protocol). The effects of $\Omega(t)$ can be neglected as the pulse $f(t)\delta_3$ always guarantees the off-resonance condition.

Starting from this ideal setting described above, we now assess the role of finite time preparation and amplitude parameters. We keep the same general pulse sequence and investigate whether it implements the required transformations $R$ with high fidelity. In particular, the pulses implemented by the functions $\Omega(t)$ and $\Delta(t)$ are the same in all three cases, while the amplitudes $\delta_{\alpha}$ can be different and are set to three values $\delta_1, \delta_2, \delta_3$. The figure of merit we use to search the pulses is

$$A_{\alpha}(R) = \left| \varepsilon_{\alpha\beta\gamma} \langle \uparrow \mid R_{\beta}R_{\gamma}^\dagger \mid \uparrow \rangle \right|^2 = 1, \quad \alpha \in \{1, 2, 3\},$$

where $\varepsilon$ is the antisymmetric Levi-Civita tensor, and indexes $\beta, \gamma$ are implicitly summed. The $\sigma_z$ eigenstate $\mid \uparrow \rangle$ is one of the two possible measurement outcomes. Note that for short pulses, larger than the interactions strength, we compute the figure of merit on one-site rotations, independently on the size of the system. We numerically investigate the role of interactions below. To test each parameters choice, we consider $N_{\text{tot}} = 10^5$ copies of the local protocols and add random fluctuations in each pulses realization. We model them as independent Gaussian fluctuations. We assume their variance to be proportional to the pulse amplitude through the percentage coefficient $\varepsilon_{\text{rel}} = 3$ (the effect of considering different coefficients will be analyzed in the following). As a result, we obtained the protocol $R^*$ shown in figure 3, for which $A_{\alpha}(R^*)/2 = \{0.55 \pm 0.06, 0.56 \pm 0.05, 0.58 \pm 0.04\}$.

On the one hand, we chose the shape of the pulses to optimize the figure of merit $A_{\alpha}(R)$ defined in equation (5). On the other, we determined their duration and amplitude to minimize the effects of interactions. By fixing the total phase of the pulses, we balanced between the minimum duration of the pulse and the maximum pulse increasing speed. We got a rotation protocol time $T_R \simeq 0.15 \mu s$, with the largest

![Figure 2](image-url)
pulse amplitude of \(\sim 7\) MHz and an increasing speed of \(\sim 35\) MHz/10 ns. The condition \(T_R \ll 1/J_e \approx 2\) \(\mu s\), with \(J_e\) being the largest coupling in the Hamiltonian defined in section 4.1, allows to control the interactions spurious effects.

To benchmark the rotations \(\mathbf{R}^*\), we consider the SSH chain with size \(L\). Each random unitary \(U\) is sampled by randomly attributing to each atom a label \(\alpha_m = 1, 2, 3\) with equal probability \(1/3\), corresponding to a parametrization of the system with light shift \(f(t)\delta_{\alpha_m}\). To make our analysis realistic, we include measurement errors. In particular, we assume the probability of errors occurring during the readout process. We model them as a 1% error to detect a false \(|\uparrow\rangle\) state and 3% error to detect a false \(|\downarrow\rangle\) state [57]. We set \(N_{\text{tot}} = N_U \times N_{\text{meas}} < 10^5\) to make our estimations compatible with experimental typical capabilities.

4. Numerical illustration with the SSH model

4.1. Presentation of the model and measured quantities

As a testbed, we consider the SSH 1D chain described by the Hamiltonian

\[
H_{\text{mod}} = -J_e \sum_{\text{even},x} \sigma_x^+ \sigma_{x+1}^- - J_o \sum_{\text{odd},x} \sigma_x^+ \sigma_{x+1}^- + \text{H.c.} + H_{\text{nnn}}
\]  

with \((J_e, J_o) = (0.484, -0.18)\) MHz. By arranging the atoms as in [11], next-nearest neighbor interactions are suppressed, while \(H_{\text{nnn}} = -J_{\text{nnn}} \sum_x \sigma_x^+ \sigma_{x+3}^- + \text{H.c.}\) describes spurious next–next nearest neighbor exchange terms, with \(J_{\text{nnn}} \approx 0.04\) MHz. We neglect residual van der Waals interactions between excited atoms, while we artificially break the degeneracy of the ground state by adding a local chemical potential term to one of the extreme sites of the chain. Starting from the experimental realization of this model Hamiltonian, we apply the toolbox, and show that we can access particular entanglement entropies and Hamiltonian variances.

The model ground state exhibits two phases, a topological one for \(|J_e| \gg |J_o|\), with localized edge excitations, and a trivial one for \(|J_e| \ll |J_o|\). In both phases, the ground state bulk is composed by separable nearest-neighbors dimers sharing one excitation. The dimers form on those sites connected by the stronger interaction term leading, in the topological phase, to the localized boundary excitations. Figure 4(a) shows the two phases for the SSH model ground state, with thicker lines indicating the stronger coupling. Given a system bipartition with size \(\ell\), the purity of the reduced density matrix can assume two values: they are \(p_1 = 1\), if the boundary does not cross any dimer and \(p_\ell\) describes a pure state, and \(p_1 = 1/2\), if the boundary crosses a dimer: measuring the purity of a given subsystem allows to distinguish the topological from the trivial phase. Moreover, RMs can also be used to extract the quantized topological invariants [23, 58].

Finally, we benchmark the ground state preparation by measuring the Hamiltonian variance

\[
\langle \Delta H_{\text{mod}}^2 \rangle = \text{Tr} \left[ \rho H_{\text{mod}}^2 \right] - \text{Tr} \left[ \rho H_{\text{mod}} \right]^2.
\]  

(7)

In the results, we show the renormalized quantity \(\langle \Delta H^2 \rangle = \langle \Delta H_{\text{mod}}^2 \rangle / \langle H_{\text{mod}}^2 \rangle\). We first consider the exact ground state \(|GS\rangle\), for which ideally we would measure \(\Delta H_{\text{mod}}^2 = 0\), neglecting the effects due to the state preparation process. Then, we benchmark the rotation protocol on the state \(|GS\rangle = P[|\downarrow \downarrow \ldots \rangle]\), where \(P\) is the adiabatic state preparation protocol presented in [11] and the initial state is a fully ferromagnetic one. We also compare the variances obtained for the ground states \(|GS\rangle\) and \(|\bar{GS}\rangle\) with that of a separable anti-ferromagnetic state \(|AF\rangle = |\uparrow \downarrow \ldots \rangle\).

4.2. Numerical results

We benchmark now the set of rotations \(\mathbf{R}^*\). We start our analysis by considering the exact ground-state, with three different scenarios. In the first, we apply the rotation protocol by evolving the ground-state with the pulses shown in figure 3. We consider only the Hamiltonian \(H_{\text{prot}}\), thus for the moment ignoring interactions. Moreover, we do not add any fluctuations and the probabilities \(P_U(s)\) are computed exactly. We
compute the exact ground state $|GS\rangle$ for different sizes $L$ and estimate the purity $p_\ell$ of the reduced density matrix $\rho_\ell$ for a bipartition with sizes $\ell = L/2, L/2 + 1$ (see figure 4(b)). We fix the number of global unitary samplings $N_U = 100$ and repeat the whole process $N_{\text{mean}} = 20$ times. All the results shown hereafter are averaged over these repetitions and the error bars are estimated by taking the standard deviation computed over 20 repetitions of the protocol. In particular, the unbiased

\begin{align*}
\text{Figure 4.} & \ (a) \text{ SSH model ground state for } L = 8 \text{ and } \ell = L/2, \text{ in the trivial (left) and in the topological (right) phases. Thicker lines indicate the larger interaction terms and thus where dimers form. Thus, the purity is } \approx 1/2 \text{ (1) in the topological (trivial) phase.} \\
& \ (b) \text{ Purity for different values of } L \text{ obtained with the rotation protocol } R^\ast \text{ ignoring the interactions and every error source.} \\
& \ (c) \text{ Estimation of the energy variance renormalized with respect to the ground state energies for each value of } L. \ (d)\text{--}(e) \text{ Estimation of the purity and the Hamiltonian variance for } L = 8 \text{ by applying the rotation protocol } R^\ast \text{ with random fluctuation for different relative amplitude of the fluctuations } \epsilon_{\text{fluc}}. \text{ The fluctuations change at each unitary transformation sample. We set } N_U = 100.
\end{align*}
Figure 5. The experimental protocol is applied to the SSH ground state \(|\text{GS}\rangle\) to estimate the purity (a) and the Hamiltonian variance (b) as a function of the number of sampled unitaries \(N_U\). We consider the topological phase and \(L = 8\). We plot the average over 20 repetitions of the full estimation process (for each repetition we prepare the target state and evolve under \(H_{\text{tot}}(t)\) for \(N_{\text{meas}} \times N_U\) times). The colored areas correspond to the standard deviations. The inset shows square-root decreasing of the standard deviation of \(\text{Tr}[\rho_\ell^2]\) for \(\ell = L/2\). Analogous behaviors are observed for \(\ell = L/2 + 1\) and for the energy fluctuations. The Hamiltonian variances are computed both for the ground state and for a separable, antiferromagnetic state. All energy variances are renormalized with respect to the respective ground state energies. The dashed lines show the exact values for the purities (a) and the variances (b). To estimate the probabilities we set \(N_{\text{meas}} = 400\). We set the relative variance of pulse fluctuations \(\varepsilon_{\%} = 3\%\).

Figure 6. (a) Estimation of the purity (up to \(L = 14\)) and (b) the energy variance (\(L = 6, 8\)) via the experimental protocol. The systems is respectively prepared in the trivial phase (\(L = 6, 10, 14\) phase) and in the topological one (\(L = 8, 12\)). Squares and triangles are used to distinguish between the topological and trivial SSH ground states while pentagons refer to the antiferromagnetic state. (c)–(d) Test of the experimental protocol robustness as a function of \(N_{\text{meas}}\) for \(L = 8\). \(L = 6,8 : N_U = 100, N_{\text{meas}} = 400, L = 10, 12, 14 : N_U = 50, N_{\text{meas}} = 800, \varepsilon_{\%} = 3\%\).

Purity \(\text{Tr}[\rho^2] = x N_{\text{meas}} / (N_{\text{meas}} - 1) - 2^L / (N_{\text{meas}} - 1)\), where \(x\) is the biased result obtained from directly inserting the estimate \(\tilde{P}\) into equation (1).

In the estimation of the Hamiltonian variance, we compare the value relative to the model ground state with that of an antiferromagnetic separable state. The obtained values are separated within the error bars, confirming once again the robustness of our RM toolbox to distinguish between states with different properties. We execute this procedure for \(L = 6, \ldots, 14\), as shown in figures 6(a) and (b). Furthermore, we benchmark our toolbox by changing the number of measurements \(N_{\text{meas}}\) for each global unitary sample. We consider the cases with \(L = 8\). In figures 6(c) and (d), we observe that the results obtained for different values of \(N_{\text{meas}}\) do not differ substantially, allowing us to consistently reduce the number of required iterations of the protocol for small lattice sizes.

As an application of the measurement of the energy variance, we show that the protocol can be used to check the adiabatic preparation of the ground state. We consider the adiabatic sequence presented in [11], suitably scaled to the interaction strength adopted here. We obtain an imperfect ground state \(|\text{GS}\rangle\) for...
Figure 7. The purity (a) and the Hamiltonian variances (b) are estimated for the state $|f_{GS}\rangle = P(\otimes |\downarrow\rangle_i)$, where $P$ is the protocol presented in [11], as a function of the preparation time $T_P$ for $L = 8$, $N_U = 100$, $N_{\text{meas}} = 400$, $\epsilon_\% = 3$. In (a), continuous lines show the expected purities relative to $|f_{GS}\rangle$, while the dashed ones correspond to the values relative to the exact ground state. In the log-scale plot (b), the continuous lines show the expected variances relative to $|f_{GS}\rangle$.

Figure 8. The purity respect to the bipartition of size $\ell$ is estimated for a state evolved for $T = 1\mu s$ after a sudden quench. The dashed lines show the exact values. Numerical parameters: $L = 8$, $N_U = 50$, $N_{\text{meas}} = 800$, $N_{\text{ave}} = 10$.

different preparation times $T_P$ and compute the purities and the Hamiltonian variances for $L = 8$. In figure 7, we report the purities (figure 7(a)) and the Hamiltonian variances (figure 7(b)) relative to the experimental numerical simulation as a function of $T_P$. In figure 7(a), the dashed lines represent the values for the purities computed on the exact ground state $|GS\rangle$, while the continuous ones correspond to the imperfect ground state $|f_{GS}\rangle$. As expected, the preparation protocol is not adiabatic for shorter processes, and the ground state properties are affected. The low interaction strengths we considered here require $\sim 10\mu s$ to prepare the ground state. On the one hand, such a time scale also requires to consider incoherent effects. On the other hand, optimal control techniques allow to go beyond adiabatic protocols for state preparations and adopt faster ones [59, 60].

Finally, we test our protocol in a different scenario. First, we set $J_e = -J_o = 0.18\text{MHz}$ in the Hamiltonian in equation (4), implementing a staggered XY model. Then, we consider the separable $L = 8$ state with all spins down, except a single spin up the middle of the lattice. By evolving the system for a time $T = 1\mu s$, the domain wall spreads over the lattice until it reaches the boundary of the lattice. We measure the purity for different bipartitions of the lattice. The purity values estimated for different subsystem sizes $\ell = 1, \ldots, L/2$ are shown in figure 8. The values obtained from the protocol are compared with the exact one, represented by the dashed lines. We notice that the estimated values capture the entanglement growth as the subsystem boundary is shifted toward the center of the lattice. This result shows that our protocol can be used to infer different entanglement profiles in lattice models.

5. Conclusion

We have proposed a protocol to implement simultaneous local, independent unitary rotations in an interacting Rydberg quantum simulator. To benchmark it, we have investigated the ground state properties of the SSH chain and the XY model after a sudden quench. We have shown the effectiveness of the local random unitary rotations protocol, despite the presence of interactions. It allows us to estimate quantities such as the purity of a system bipartition and the Hamiltonian variance, taking into account realistic experimental parameters, the influence of residual interactions, and imperfections such as finite read-out fidelities. The presented results provide a complete RM toolbox to probe entanglement [13, 16, 18], many-body topological invariants [23] and quantum state fidelities [37], but also to measure any quantum observable from classical shadows [14], in Rydberg quantum simulators and quantum computers.
Data availability statements

The data that support the findings of this study are available upon reasonable request from the authors.

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