Rydberg atom mediated polar molecule interactions: a tool for molecular-state conditional quantum gates and individual addressability

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We study the possibility to use interaction between a polar molecule in the ground electronic and vibrational state and a Rydberg atom to construct two-qubit gates between molecular qubits and to coherently control molecular states. A polar molecule within the electron orbit in a Rydberg atom can either shift the Rydberg state, or form Rydberg molecule. Both the atomic shift and the Rydberg molecule states depend on the initial internal state of the polar molecule, resulting in molecular state dependent van der Waals or dipole-dipole interaction between Rydberg atoms. Rydberg atoms mediated interaction between polar molecules can be enhanced up to $10^3$ times. We describe how the coupling between a polar molecule and a Rydberg atom can be applied to coherent control of molecular states, specifically, to individual addressing of molecules in an optical lattice and non-destructive readout of molecular qubits.

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I. INTRODUCTION

Ultracold polar molecules placed in periodic arrays of traps represent an attractive system for quantum information processing [1] and quantum simulation of condensed matter systems [2]. They offer a variety of long-lived states for qubit encoding, described by various quantum numbers; rotational, spin and hyperfine (if electronic and nuclear spins are non-zero), or projections of electronic angular momenta $\Lambda$ (orbital) and $Ω$ (orbital and spin) onto the molecular axis [3]. Cold polar molecules also offer scalability to a large number of qubits. Polar molecules can be easily controlled by DC electric and magnetic fields, as well as by microwave and optical fields allowing to design various traps [4, 5]. A strong appeal of cold polar molecules, however, is that they possess permanent electric dipole moments, allowing for long-range dipole-dipole interaction. The dipole-dipole interaction offers a tool to construct two-qubit gates, required for universal quantum computation [1, 6]. A naturally scalable setup is an optical lattice, i.e. a periodic trapping potential produced by an optical standing wave formed from two counterpropagating laser beams. With a typical focus diameter of the beams $d \sim 1$ mm and the lattice period $\lambda/2 \sim 300 – 500$ nm, this system can accommodate $\sim (d/\lambda/2)^2 \sim 10^5 – 10^7$ molecular qubits. A two-qubit gate, e.g. a phase gate $|\psi_{12}\rangle \rightarrow e^{i\phi}\psi_{12}\rangle$ $|\psi_{12}\rangle$, where $\psi_{1,2} = 0, 1$ are states of the first and second qubit), can be realized using dipole-dipole interaction when both molecules are in the qubit state $|1\rangle$ such that the two-qubit state $|11\rangle$ accumulates a phase $\phi = V_{int}T = \pi$ during the interaction. Polar molecules that can be currently produced at ultracold temperatures (required to place molecules in a lattice) by Feshbach and photo-association are limited to alkali diatoms, having dipole moments of the order of a Debye (1 Debye=10^{-18} esu cm in CGS units). The resulting dipole-dipole interaction strength scales as

$$V_{int} \sim \mu^2/R^3 \sim 10^{-9}/R(cm)^3 s^{-1}$$

(1)

Given the gate duration $T \leq 1$ μs, which is typically much shorter than the qubit decoherence time, one finds that the required interaction strength can be achieved at distances of the order of

$$R \sim (\mu^2 T/\pi)^{1/3} \sim 100 \text{ nm}$$

(2)

In an optical lattice, these molecules interact efficiently with only several nearest neighbours. To be able to take full advantage of the scalability offered by optical lattices, a mechanism has to be found to make molecules interact at greater distances, ideally at the opposite sides of the lattice.

The interaction strength can be significantly increased by coupling the state of a polar molecule to a Rydberg atom excitation (see Fig.1). One way that this can be achieved is by placing a polar molecule within a Rydberg electron’s orbit where the ground state molecule can act as a perturber, shifting the electron energy. Alternatively the coupling can be realized through the creation of a new type of ultralong-range Rydberg molecule proposed recently [7], where a molecule is made of a polar molecule in the ground electronic state and a Rydberg atom. These Rydberg molecules can have extremely large dipole moments, on the order of a thousand Debye, comparable to dipole moments of Rydberg atoms. Moreover, the Rydberg molecules have different sizes and binding energies depending on the state of the molecular perturber. This means that by coupling to Rydberg atoms, interaction between polar molecules can be, first, strongly enhanced and, second, state-dependent. These two conditions facilitate implementation of two-qubit gates between polar molecules at large distances.

Not only Rydberg atoms can enhance the interaction, they can also provide individual control over polar molecules. One realization of this control would be...
individual addressing of molecules in an optical lattice, which still poses a problem. It means that one qubit rotations and readout of molecular states can be implemented without the need for experimentally challenging laser tight-focusing and single-site addressability. Another example would be the molecular control in small registers, which form the basis of quantum communication with quantum repeaters and distributed quantum computation. In summary, coupling of polar molecules to Rydberg atoms offers a way to enhance the range of efficient interaction as well as to realize coherent control over molecular states, including individual addressing of molecular qubits in a lattice.

The paper is organized as follows. In Section II we analyze two schemes allowing to increase the efficient interaction range, based on van der Waals (vdW) and dipole-dipole interaction between Rydberg molecules. We describe how a two-qubit phase gate can be realized with KRb and CH molecules, coupled to Rb atoms. In Section III, we discuss the possibility to selectively perform one-qubit rotations and read out qubit states of polar molecules by coupling to Rydberg atoms. We also describe how a controlled-NOT gate can be implemented between a polar molecule and a neutral atom mediated by an atomic Rydberg state, which is the main ingredient of computation in quantum repeaters and small quantum registers.

II. ENHANCEMENT OF POLAR MOLECULES INTERACTION STRENGTH

We envision a setup shown in Fig.1a, where polar molecules are placed in an optical lattice while ground state neutral atoms are kept either in an additional lattice or in individually controlled dipole traps. Polar molecules made of alkali atoms can be formed in an optical lattice by Feshbach associating pairs of atoms in each site and transferring the resulting Feshbach molecules to the stable ground rovibrational state by STIRAP (stimulated Raman adiabatic passage) [3]. As we already mentioned in the introduction polar molecules can interact efficiently with several neighbors via dipole-dipole interaction. To enable interaction between two distant molecules, atoms are brought sufficiently close to molecules so that the molecule is within a Rydberg electron orbit as is illustrated in Fig.1b. A phase gate then can be realized using state-dependent dipole-dipole interaction (Rydberg blockade) [3] in the following way: (i) a control atom is conditionally excited (dependent on the qubit state of the control polar molecule) by a \( \pi \)-pulse to a Rydberg state \( |r\rangle \); (ii) a \( 2\pi \)-pulse of the same wavelength is applied to an atom near the target polar molecule to excite it to the same Rydberg state. The doubly excited state \( |rr\rangle \) is however shifted by the energy of the Rydberg-Rydberg vdW or dipole-dipole interaction and is far off-resonance (the excitation is blockaded). As a result, the \( 2\pi \)-pulse has no effect on the target molecule except for a small accumulated phase; (iii) the control atom is deexcited back to the ground state by a second \( \pi \)-pulse. The excitation sequence is shown schematically in Fig.1a.

The blockaded phase gate can be realized provided that the interaction strength \( V_{\text{int}} \gg \Omega_{2\pi} \), where \( \Omega_{2\pi} \) is the Rabi frequency of the \( 2\pi \)-pulse. The Rabi frequency defines the speed of the gate, since the total gate time \( T_{\text{gate}} = 2\pi/\Omega_x + 2\pi/\Omega_{2\pi} \) is a sum of durations of two \( \pi \)- and one \( 2\pi \)-pulse. The gate time has to be much shorter than a qubit decoherence time and the Rydberg lifetime \( (T_{\text{Ryd}}) \). If a qubit is encoded in rotational states of a polar molecule, qubit decoherence times \( \sim 1 \) ms can be achieved [3], limited by electric field fluctuations. Even longer qubit coherence times (greater than a second) can be realized if electronic spin or hyperfine states are used [10], since these states are insensitive to electric field fluctuations in the first order. For sufficiently highly-excited Rydberg states, \( T_{\text{Ryd}} \sim 100-500 \) ms are expected, setting the main limit on gate duration and, therefore, on the Rabi frequencies. If we assume a gate duration \( T_{\text{gate}} \sim 1 \) \( \mu \)s, short enough compared to \( T_{\text{Ryd}} \), it will correspond to the Rabi frequency \( \Omega_x, \Omega_{2\pi} \sim 100 \) kHz. The blockade radius, which determines the range of efficient interaction, is estimated as \( R_b = (C_b/\Omega)^{1/6} \sim (n^{11}/\Omega)^{1/6} \) (in a.u.) for vdW interaction. For \( n \sim 50, R_b \approx 5 \) \( \mu \)m, about fifty times larger than compared to direct interaction between polar molecules. Even larger blockade radius of tens \( \mu \)m is possible if Rydberg molecules interact via dipole-dipole interaction. In this case \( R_b \sim (\mu^2/\Omega)^{1/3} \), where \( \mu \sim n^2 \) is the dipole moment of a Rydberg molecular state. Assuming again \( n \sim 50 \) and \( \Omega \sim 100 \) kHz, the resulting blockade radius \( R_b \approx 40 \) \( \mu \)m. At distances larger than the blockade radius the phase gate can be realized by simultaneously exciting both control and target molecules to the Rydberg molecular state, letting them interact via dipole-dipole interaction and accumulate a \( \pi \) phase shift, and deexciting them back [3]. Therefore, extending the blockade radius from \( R_b \sim 100 \) nm to \( R_b \sim 100 \) \( \mu \)m allows molecules separated by \( 10^3 \) lattice sites, to interact instead of only few nearest neighbors.

A qubit can be encoded into long-lived rotational states of the ground electronic and vibrational molecular state. Rotational transitions of polar molecules have dipole moments equal to the permanent one and frequencies in the GHz range, allowing for fast one-qubit gates with microwave pulses. Rotational states are shifted by an electric field and as a result a qubit will decohere in the presence of electric field fluctuations. The coherence time can be significantly increased if a qubit is encoded into spin or hyperfine sublevels and is swapped onto rotational states for one and two-qubit operations. If we encode a qubit into the first rotational states \( \left| 0 \right>, \left| 1 \right> = \left| J = 0,1,2 \right> \), where \( J \) is the rotational angular momentum, we can selectively excite a molecule plus an atom from a specific \( J \) state to the Rydberg molecular state, in which molecules interact via dipole-dipole interaction. Another possibility is to couple the \( J \) states to isotropic ns Ry-
FIG. 1: (a) Polar molecules in an optical lattice can be coupled with Rydberg atoms held in a separate lattice or individual optical dipole traps; (b) Coupling of a polar molecule to a Rydberg electron and a positive core. The molecule is separated from the core by \( \vec{R} \), the electron-core distance is \( \vec{r} \). The polar molecule is within the electron’s orbit, schematically shown by a dotted ellips; (c) Two-qubit phase gate between polar molecules based on the Rydberg blockade mediated by atomic Rydberg-Rydberg interaction (see text for details).

Rydberg atomic states, in which Rydberg atoms interact via vdW interaction. Here we analyze KRb+Rb as an example system with vdW interaction between Rydberg atoms dependent on the state of polar molecules.

We may find the shift in the atomic Rydberg excitation frequency by considering the shift of the polar molecule energy in a rotational state \( J = 0, 1, 2, \ldots \) due to the presence of a Rb(ns) Rydberg atom. In effect, the polar molecule’s energy is shifted by the internal field of the positive core shielded by a highly excited s-wave Rydberg electron. The internal electric fields of a Rydberg atom are small (on the order of \( 10^{-7} \) atomic units or several 100 V/cm) so that the field induced shift is small compared to the rotational splitting of the molecule (on the order of \( \sim 10 \) GHz). This means that there is very little mixing between different molecular rotational states, and their energy shifts quadratically in the field strength. The shift of the rotational states can be found by diagonalizing a Hamiltonian whose matrix elements are [2]

\[
H_{J,J'} = B \frac{J(J+1)}{2} \delta_{J,J'} - \left| J \right| \vec{F}_{\text{Ryd}} \cdot \vec{d}_0 \left| J' \right>, \tag{3}
\]

\[
\vec{F}_{\text{Ryd}} = \epsilon \hat{\mathbf{z}} \left\langle \psi_{ns} \left| \cos \theta \vec{R}_{\vec{R}_d} \right| \psi_{ns} \right\rangle,
\]

where \( B \) and \( \vec{d}_0 \) is the polar molecule rotational constant and body fixed permanent dipole moment. Here \( \vec{F}_{\text{Ryd}} \) is the electric field at the position of the polar molecule, \( \vec{r} \), due to the positive core of the Rydberg atom shielded by an s-wave electron, \( \vec{r} \) is the position of the Rydberg electron (see Fig. 1), \( \psi_{ns} \) is the Rydberg electron wavefunction.

The predicted shift in the Rydberg excitation frequency due to a KRb molecule perturber in several of the lowest rotational states is shown in Fig. 2 as a function of the core molecule separation, \( R \). We have subtracted out the zero field energies of the KRb rotational states as we are only interested in the net shift in the Rydberg excitation energy. Due to the small internal fields the mixing between different rotational states is very small (less than 2%), so even though the polar molecule is not in a pure \( J \) state, we have labeled each state by its dominant angular momentum component. Fig. 2 shows that for separation distances of the order of 100 nm, the shift in excitation frequency is in the range of several MHz to ten MHz. This shift can be resolved by a laser with a linewidth of several kHz and a Rabi frequency of the order of 100 kHz. The Rabi frequency of 100 kHz results in a gate duration \( T_{\text{gate}} \sim 1 \mu s \ll T_{\text{Ryd}} \). In a recent work, [11] a lifetime of a Rydberg molecular \( 5s_{1/2} - 35s \) state of two Rb atoms has been measured \( T_{\text{Ryd}} = 6.4 \mu s \). The lifetime is probably limited by collisions with ground state Rb atoms and is expected to be in the range of 10–100 \( \mu s \) for an isolated Rydberg atom, which means that a gate time of 1 \( \mu s \) is short enough compared to \( T_{\text{Ryd}} \).

The analysis presented above is appropriate for molecules with fairly small dipole moments, less than 0.4 a.u. (1 Debye). It was shown in Ref. [12] that for larger dipole moments the electron-dipole interaction is strong enough to mix several higher electron angular
momentum states which can significantly change the energy landscape. Ref. [12] describes the potential energy surface that arises due to the presence of a Λ-doublet molecular perturber (which can be modelled as a two-level system) with a larger dipole moment. Describing the intricate 2D energy surface that arises in the case of rigid rotor type molecules (having a multilevel structure of rotational states), similar to KRb or RbCs, is in progress.

As was shown in Fig. 2, ns states Rydberg atoms can interact via isotropic vdW interaction, allowing for the effective interaction range between polar molecules to be extended to several μm. A larger interaction range can be realized using dipole-dipole interaction between Rydberg molecules. As it has been shown, [7] states of a Rydberg atom with angular momenta \( l \geq 3 \) (those with negligible quantum defects) are hybridized in the presence of a Λ-doublet polar molecule perturber forming ultralong-range Rydberg molecule states which can acquire dipole moments of several kiloDebyes (kD). Furthermore, the vibrationally bound states formed in this system correspond to a polar molecule perturber that is in an internal state which is fully polarized with a dipole moment pointing either toward or away from the positive Rydberg core. The Rydberg molecule states for different internal configurations of the polar molecule perturber are also separated spatially with potential wells separated by 100s of nm.

This theory was further refined in [12], where an extended set of Rydberg atom states was considered by including all electron angular momentum states for several principal quantum numbers \( n \). For molecules with permanent dipole moments larger than \( d_0 \sim 0.4 \) a.u., the resulting Rydberg molecules acquire significant s-wave electron contribution (30 - 40%) making the molecular state accessible by a standard two-photon Rydberg excitation schemes. The resulting Rydberg molecule size and binding energy is dependent on the internal orientation of the perturbing polar molecule, and hence conditioned on the initial state of the polar molecule. Excitation from a desired internal state can be tailored by tuning a Rydberg excitation to the appropriate binding energy and by modifying the spatial separation of the ground state atom from the polar molecule.

Λ-doublet molecules with two opposite parity states \((f, e)\) far separated from higher rotational states fit this analysis particularly well. One good example is CH with a \(^2\Pi_{3/2}\) metastable state having lifetime > 10³ s [13], split into Λ-doublet \((f, e)\) states with a 700 MHz splitting, as shown in Fig. 3. Each Λ-doublet state is further split into two hyperfine sublevels due to the hydrogen nuclear spin (see Fig. 3). Therefore, it offers an opportunity to encode a qubit into hyperfine sublevels \(|F, m_F, p\rangle\), where \( p = f, e \) is the parity of the state. As was observed in the ground state OH (which has a similar Λ-doublet and hyperfine structure) in [14], two combinations of hyperfine states, namely, \(|F = 2, m_F = 1, e/f\rangle\) and \(|F = 1, m_F = -1, e/f\rangle\) are insensitive to fluctuations of a magnetic field in the first order at \( B = 3.01 \text{ G} \) and \( B = 7.53 \text{ G} \), respectively, similar to hyperfine clock states \(|F, m_F = 0\rangle\) and \(|F = 2, m_F = 1, |F = 1, m_F = -1\rangle\) in Rb atoms. The insensitivity to magnetic field fluctuations makes these states good candidates for qubit encoding.

We carried out similar analysis for the \(^2\Pi_{3/2}\) state of CH. The \((f, e)\) hyperfine states shift in the magnetic field as

\[
E_{f,e} = -\frac{\Delta E_{f,e}}{2(2J + 1)} + \mu_B g_f^e m_F B \pm \frac{|\Delta E_{f,e}^B|}{2} \sqrt{1 - \frac{4\mu_B g_f^e m_F B}{\Delta E_{f,e}^B (2J + 1)}} + \left(\frac{\mu_B g_f^e B}{\Delta E_{f,e}^B}\right)^2 \tag{4}
\]

where the ± sign before the square root refers to \( F = J + 1/2 \) for the \( e \), and to \( F = J - 1/2 \) for the \( f \) state; \( \Delta E_{f,e} = -29.908 \text{ MHz} \) and \( \Delta E_{f,e}^B = 2.593 \text{ MHz} \) are the hyperfine splittings of the \((f, e)\) states [12] (the hyperfine structure in the \( e \) state is inverted), \( g_f^e = 0.819537 \) and \( g_f^e = 0.817829 \) are the slightly different \( g \)-factors of the \((f, e)\) states, \( \mu_B \) is the Bohr magneton and \( B \) is the magnetic field. We found two transitions, \(|F = 1, m_F = 1, e\rangle - |F = 2, m_F = 1, f\rangle\) and \(|F = 2, m_F = 1, e\rangle - |F = 1, m_F = 1, f\rangle\) that become insensitive to the magnetic field at \( B = 2.4 \text{ G} \) (see Fig. 3). The \( g \)-factor values are correct up to \( \Delta g_{1/2} \approx 0.05g_J \), but this variation does not change the results appreciably. The splitting of the CH Λ-doublet is 700 MHz and the dipole moment for the transition between the \((f, e)\) states is 1.47 D [17], allowing to apply fast one-qubit rotations with microwave pulses.

Two-photon excitation to the high dipole moment states of a Rydberg molecule, which are composed of \( l > 2 \) angular momentum states of a Rydberg atom can be realized due to fortunate overlap of \( n(l > 2) \) and \((n + \nu)s\) states, where \( \nu \) is some number of the order of a quantum defect for the \( s \) state (\( \nu = 3 \) for Rb). This overlap mixes the \( nl \) and \((n + \nu)s\) states which result in the mixed state wavefunction

\[
\psi = a(d_0)\psi_d(\vec{r})|\xi_d\rangle + b(d_0)\psi_s(\vec{r})|\xi_s\rangle, \tag{5}
\]

where \( \psi_d \) and \( \xi_d \) are the electronic wave function and internal polar molecule state, that include higher electron angular momentum with \( l > 2 \); \( \psi_s \) and \( \xi_s \) are the s-wave Rydberg electron wave function and the corresponding molecular state, respectively. The expansion coefficients \( a(d_0) \) and \( b(d_0) \) depend on the dipole moment of the polar perturber as well as its position \( R \). The \( |a(d_0)|^2 \) determines the dipole moment of the state scaling as \( d_{Ryd} \approx 1.3|a(d_0)|^2 n^2 \), which can be in kD range (for \( n = 25 \) and \( a(d_0) \approx 0.6 \) the dipole moment \( d_{Ryd} \approx 1.4 \text{ kD} \) [12]. On the other hand, the sizable \( b(d_0) \) coefficient would allow a convenient two-photon excitation from the ground atomic state. While for a two-state molecule, such as a Λ-doublet molecule, this theory works quite well, for a rigid rotor molecule, such as KRb, the formal-
Fig. 3: (a) Ground $^2\Pi$ state of CH split into $J = 1/2, 3/2$ states by spin-orbit interaction with $\sim 0.5$ THz splitting; (b) the hyperfine structure of the $^2\Pi_{3/2}$ state due to the hydrogen spin; (3) Zeeman shifts of the $\pi$ transition frequencies between satellite lines $|F = 1, m_F, e\rangle - |F = 2, m_F, f\rangle$, $|F = 2, m_F, e\rangle - |F = 1, m_F, f\rangle$, $m_F = 1$ transitions are magnetic field insensitive at $B = 2.4$ G and can be used for qubit encoding.

III. COHERENT CONTROL OF POLAR MOLECULES BY COUPLING TO RYDBERG ATOMS

Interaction between a polar molecule and a Rydberg atom opens the possibility to coherently control molecular states. In particular, we will show below that individual addressing and readout can be realized without resorting to tightly focused laser beams. This is similar to the idea of a "marker" atom for individual manipulation of neutral atoms in a lattice (atom microscope) [18, 19]. In our case the Rydberg atom mediates coherent control realizes a "molecule microscope".

Individual addressing of atoms and molecules in an optical lattice is a long-standing problem due to the difficulty of focusing laser beams to a single lattice site. Recently, individual addressing of single atoms in a lattice has been achieved by tightly focusing a laser beam to a single site in a spin-dependent lattice [20]. An alternative approach proposed in [18] is to use "marker" atoms that can selectively interact with atoms in a lattice. "Marker" atoms, held in a separate optical lattice, can be brought to the same lattice site with addressed atoms and interact via state-dependent collisions. Qubit states acquire interaction-induced shifts, which offer a way to address the atom by applying microwave or Raman pulses resonant with the shifted qubit states. In the same spirit, addressing can be realized for molecular qubits by coupling them to Rydberg atoms. The main difference is that the interaction is not relying on atom-molecule collisions (which can be inelastic or result in a chemical reaction), but on a long-range interaction. Rotational states of a polar molecule will be shifted due to the interaction with the Rydberg atom, which will allow to perform one-qubit gates individually.

It is illustrated in Fig. 4, and b: when an atom is in the ground electronic state there is no interaction between the atom and the molecule, and molecular rotational states are not shifted. Once the atom is excited to the Rydberg state, molecular states shift and the qubit transition frequency changes. A microwave pulse resonant to the shifted frequency will act on the addressed qubit without affecting the rest. As was shown in Fig. 2 at distances of $\sim 100$ nm between a polar molecule and a Rydberg atom the rotational states shift by several MHz. A microwave pulse resonant to the shifted qubit transition with a Rabi frequency of $\sim 100$ kHz, much smaller than the shift, will selectively interact with the addressed molecule. The resulting one-qubit gate time is again in the $\mu$s range, much shorter than the Rydberg state lifetime $T_{\text{Ryd}}$. Similarly, molecular qubits can be read out by molecular state dependent excitation of a Rydberg atom, followed by e.g. ionization of the Rydberg state (see Fig. 3). We note that qubit-carrying molecules are not destroyed in the process, although the Rydberg atom gets ionized and has to be replaced. Completely non-destructive readout maybe possible if after the selective excitation to the Rydberg molecular state the atom can
be transferred to a state that is a part of a cycling transition (Fig. 4). The readout is then realized by detecting resonant fluorescent photons. Once a photon is detected, both the molecule and the atom can be recycled.

Coherent coupling between Rydberg atoms and polar molecules can be utilized to build small quantum registers, which form the basis of quantum repeaters [21, 22] and distributed quantum computing [23]. Let us briefly summarize the idea of a quantum repeater as was originally proposed in [21]. In quantum communication to teleport a quantum state a Bell state has to be shared at each node in the Bell state basis. This is followed by entanglement purification in the obtained entangled pairs to compensate for local one and two-qubit and measurement errors. In the third step nodes $A$ and $C_{L_2}$, $C_{L_2}$ and $C_{2L_2}$, etc. are connected in the same way. The process is repeated until $A$ is connected to $B$ directly without any nodes in between. A key ingredient in connecting two entangled pairs of qubits into a single one and in entanglement purification is the ability to perform a CNOT gate between two qubits at each node. To illustrate how the CNOT gate is used in entanglement purification, we describe as an example an entanglement swapping between two pairs of qubits. Suppose we have two qubits (qubits 1 and 2) in a $|\Phi^+\rangle = (|00\rangle + |11\rangle)/\sqrt{2}$ Bell state and two qubits (qubits 3 and 4) in a $|0\rangle$ state, and we want to swap the entanglement between the two pairs. To this end, first a CNOT is applied to the qubits in $|0\rangle$ state controlled by the qubits forming the entangled pairs

$$
(|00\rangle^{12} + |11\rangle^{12}) |00\rangle^{34} \rightarrow \frac{1}{\sqrt{2}} (|00\rangle^{12} + |11\rangle^{12}) |00\rangle^{34} + |11\rangle^{12} |11\rangle^{34} / \sqrt{2},
$$

where qubits 1, 2 control qubits 3, 4 respectively. As a next step, qubits 1, 2 are measured in the $|\pm\rangle = (|0\rangle \pm |1\rangle)/\sqrt{2}$ basis with the measurement outcomes $|++\rangle^{12}$, $|--\rangle^{12}$ producing $|\Phi^+\rangle^{34}$, while measurement outcomes $|+\rangle^{12}$ and $|--\rangle^{12}$ producing $|\Phi^-\rangle^{34}$ Bell states.

A ground state polar molecule and a neutral atom can form a quantum register with the molecule serving as a memory and the atom that can serve as a communication qubit to create initial entanglement between nearest nodes and for measurement. The quantum repeater protocol therefore can be implemented in this system provided a CNOT operation can be realized on a polar molecule controlled by the atomic qubit and vice versa. In Fig. 5 we illustrate how this can be done using intermediate Rydberg atomic states. A conditional flip of the molecular qubit can be realized using the slightly modified scheme of Fig. 4b: we selectively excite the atom to a Rydberg state only from the qubits state $|1_n\rangle$ and bring it at a distance $R$ of efficient interaction with the polar molecule, which will shift the molecular qubit frequency. The molecular qubit state then can be flipped by a microwave $\pi$ pulse resonant to the shifted qubit frequency. If initially the atom was in the $|0_n\rangle$ state it will not be excited to the Rydberg state, the molecular qubit states will not shift and the $\pi$-pulse will be off-resonant. The result is the molecular qubit flip if the atom is in the $|1_n\rangle$ state (see Fig. 5a). A conditional flip of the atomic qubit can be realized by a Raman $\pi$-pulse resonant to the atomic Rydberg state, corresponding to a specific state of the molecular qubit, e.g. the molecular $|1_m\rangle$ state as shown in Fig. 5b.

IV. CONCLUSIONS

We propose and study interactions between ground state polar molecules and Rydberg atoms as a tool in quantum information processing with polar molecules in
optical lattices. We show that this interaction results in a molecular state dependent shift of ns Rydberg atom states, which can be used to enhance the interaction range between molecules by many folds. The interaction range can be enhanced by excitation of ground state atoms to Rydberg states conditioned on the molecular state and allowing them to interact via long-range vdW interaction, increasing the interaction range up to several μm. Even larger interaction range of tens μm can be realized if the molecule and the Rydberg atom form a Rydberg molecule. Rydberg molecular states are expected to have dipole moments of several kD, allowing for strong long-range dipole-dipole interaction. The size of a Rydberg molecule as well as the energies of the bound vibrational states depend on the state of the polar molecule, resulting in molecular state-dependent interaction. We illustrate our proposal using rigid rotor KRb+Rb and A-doublet CH+Rb as examples of vdW and dipole-dipole interactions in Rydberg states. We also analyze the possibility to use the molecule-atom coupling to coherently control the state of the polar molecule with applications to individual addressing and molecular state readout as well as building small quantum registers composed of a polar molecule and a neutral atom.

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