Charge dynamics of a molecular ion immersed in a Rydberg-dressed atomic lattice gas

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Charge dynamics in an ultracold setup involving a laser-dressed atom and an ion is studied here. This transfer of charge is enabled through molecular Rydberg states that are accessed via a laser. The character of the charge exchange crucially depends on the coupling between the electronic dynamics and the motional dynamics of the atoms and ion. The molecular Rydberg states are characterized and a criterion for distinguishing coherent and incoherent regimes is formulated. Furthermore the concept is generalized to the many-body setup as the ion effectively propagates through a chain of atoms. Aspects of the transport, such as its direction, can be controlled by the excitation laser. This leads to new directions in the investigation of hybrid atom-ion systems that can be experimentally explored using optically trapped strontium atoms.

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I. INTRODUCTION

Ultracold atoms in optical lattices opened the door for experimental studies of a wide range of quantum many-body problems [\[1–4\]](#page-6-0). Similar breakthroughs have been achieved in trapped ion systems [\[5,6\]](#page-6-0). One of the main motivations of using such systems is to simulate spin models [\[7,8\]](#page-6-0) in a controlled environment including Hamiltonians which are intractable by conventional numerical methods [\[9\]](#page-6-0). Emerging from these efforts, there is a growing interest in exploring hybrid systems formed of trapped atoms and ions [\[10](#page-6-0)[–17\]](#page-7-0). This combination enables access to a plethora of novel phenomena such as strongly coupled polaron states [\[18–20\]](#page-7-0), longrange collisions $[21-23]$, the study of exciton transport $[24]$, electron-phonon coupling in Fermi gases [\[25\]](#page-7-0), many-body quantum dynamics $[26-28]$, the implementation of atom-ion quantum gates [\[29\]](#page-7-0), switches for information transfer [\[30\]](#page-7-0), quantum simulation of novel ultracold chemistry [\[31\]](#page-7-0), as well as the formation of mesoscopic molecular ions [\[32,33\]](#page-7-0).

Charge exchange are processes central in atom-ion systems and has relevance in the study of chemical reactions [\[22,](#page-7-0) [34–36\]](#page-7-0) as well as charge transport in the ultracold domain [\[37\]](#page-7-0). Resonant charge exchange in atom-ion setups plays a crucial role in the cooling of ions [\[38–41\]](#page-7-0). At sufficiently low temperatures, the mechanism for charge exchange involves electron hopping from neutral atoms to a neighboring ion. This process is highly suppressed for ground-state atoms due to the negligible overlap between the electron wave function with the orbital of a nearby ion. This unfavorable situation can change for highly excited (Rydberg)atoms [\[42\]](#page-7-0) where the large spatial extent of the electronic wave function enhances the probability for electron hopping onto the ion [\[43\]](#page-7-0).

The aim of this work is to investigate the charge dynamics in an atom-ion hybrid system that is formed by a deep optical lattice filled with a single atom per site out of which one is ionized. Before moving to the many-body problem, we first study the two-body problem involving the atom and an ion. The underlying key ingredient is the existence of electronic molecular Rydberg states that encompass the ion and an

adjacent atom. For a Rydberg atom, the wave function of the electron has a large spatial extension which enhances the hopping probability as depicted in Fig. $1(a)$. The tunneling rate is determined by the splitting between the corresponding potential curves of opposite symmetry as shown in Fig. [1\(b\).](#page-1-0) However, enhanced tunneling occurs at internuclear distances where the Rydberg atom is considerably polarized by the ion leading to strong l-mixing of the Rydberg states resulting in the formation of complex Rydberg molecular ion states as shown in Fig. $1(c)$. In the presence of a detuned laser, the coupling of the ground-state atom to Rydberg states is described by a dressed atom picture and the ion dynamics is effectively given by $J(R)$. The implicit dependence of $J(R)$ on interatomic distance entangles the electronic and motional dynamics which then introduces decoherence into the dynamics. Coherent charge dynamics can be achieved by trapping the atoms and the ion in an identical potential [see Fig. $1(d)$] similar to [\[44\]](#page-7-0). The natural extension of the two-particle picture to the many-body system leads to delocalized charge dynamics which is interesting in its own right. However, in certain regimes of the parameter space, the many-body charge dynamics is effectively dictated with nearest-neighbor hopping as shown schematically in Fig. [1\(e\).](#page-1-0)

II. RYDBERG MOLECULAR ION

The Rydberg molecular ion states are calculated for strontium (Sr) by adopting a linear combination of localized orbitals. The orbitals (ψ_{nl}) correspond to Rydberg states of Sr obtained using a single active electron approximation [\[45\]](#page-7-0). The electronic Hamiltonian describing the atom-ion system is discussed in Appendix [B.](#page-4-0) The interactions of the Rydberg molecular ion is invariant with respect to exchange in nuclear positions and it is always possible and often convenient to express the molecular states in the (un)gerade venient to express the molecular states in the (un)gerade
basis, $|e^{\alpha_1(\pm)}\rangle = 1/\sqrt{2}(|ie^{\alpha}\rangle \pm |e^{\alpha}i\rangle)$ and their corresponding energies $E^{\alpha, (\pm)}(R)$. $|ie^{\alpha}\rangle$ or $|e^{\alpha}i\rangle$ are defined depending on whether the Rydberg atom is to the right or left of the ion. Owing to the nonorthogonality between the Rydberg wave

FIG. 1. (a) Illustration of the key principle: While a low-lying state (shown in red) remains localized, a Rydberg state (shown in orange) can tunnel through the ionic potential barrier (black lines) at rate *T* . (b) A laser addresses the excited Rydberg molecular states with effective coupling $\Omega(R)$ and detuning $\Delta(R)$. Tunneling $[T(R)]$ is given by the splitting between the gerade $[E^+(R)]$ and ungerade [*E*[−](*R*)] states which are, in fact, the electronic molecular ion states. In the Rydberg-dressed picture, $J(R)$ is the effective hopping and $U(R)$ is the overall light shift of the relevant Rydberg-dressed states (denoted by $|i\tilde{g}\rangle$ and $|\tilde{g}i\rangle$). (c) Depicts the probability density for a particular Rydberg-dressed molecular ion wave function. (d) Coherent dynamics is facilitated by confining the ion and the Rydbergdressed atom in an identical double well optical trap. Initially prepared in their motional ground states $|0\rangle$ (shown in green), the coupling to higher motional states such as $|1\rangle$ (shown in aquamarine) are suppressed by choosing optimum optical and trapping conditions. (e) The two-particle picture is generalized to obtain the effective many-body charge transport model with nearest-neighbor hopping $\bar{J}_{k,k\pm 1}$ and on-site energy \bar{U}_k .

functions (ψ_{nl}) defined at either nuclei, there is a small but nonzero overlap function. However, for this work, the focus is on internuclear distances where these overlap integrals are small thereby obtaining a simplified eigenvalue problem for the electronic Hamiltonian

$$
\hat{H}_{\rm el}|e^{\alpha,\pm}\rangle = E_{\alpha}^{\pm}(R)|e^{\alpha,\pm}\rangle. \tag{1}
$$

The index $\alpha = 1, 2, \ldots$, represents the different excited states of the Rydberg molecular ion. Upon diagonalization, we have the Rydberg molecular states and energies. Compared to calculations of low-lying states, those for highly excited molecules prove very demanding due to the need for a large basis set and the highly oscillatory character of the involved atomic Rydberg states. Figure 2(a) depicts a characteristic pair of molecular potential curves $E^{\alpha, (\pm)}(R)$ around the Sr₂⁺(50*S*) asymptote, obtained for a basis set of $\sim 10^3$ atomic states. The numerical calculations used basis states with principal quantum number ranging from $n = 40-60$ including $l =$ 0 ... (*n* − 1) states for each *n*. At such high excitations, the ion-atom interaction leads to strong state mixing already at micrometer distances which is reflected in the molecular ion wave functions shown in Figs. $2(b)$ to $2(d)$. The charge exchange between the ion and Rydberg atom is determined

FIG. 2. (a) Potential curves for high-lying Rydberg states of a $Sr₂⁺$ molecular ion. The molecular energies are given relative to the Sr⁺(50*S*) asymptote. The relative energies, $\delta E^{\alpha,(\pm)} = E^{\alpha,(\pm)}$ – $(E^{\alpha,+} + E^{\alpha,-})/2$, of a selected pair ($\alpha = 243$) with (un)gerade symmetry is shown in the inset for which the tunnel splitting is as large as several 100 MHz at an internuclear distance of $R = 730$ nm. (b–d) Plotting the complex Rydberg molecular ion wave functions for selected pair of potentials shown in the inset for three different internuclear distances, 700 nm in (b), 670 nm in (c), and 640 nm in (d).

by the energy splitting given as

$$
T^{\alpha}(R) = \frac{E^{\alpha,+}(R) - E^{\alpha,-}(R)}{2}.
$$
 (2)

There is substantial tunnel splitting between the opposite symmetry states [see inset of Fig. 2] of up to several hundred MHz, even at distances for which the Rydberg electron remains well localized at either ionic core. The polarization of the Rydberg atom due to the ion is calculated from the slope of the molecular potential curves as a function of the internuclear distance.

III. OPTICAL COUPLING TO RYDBERG MOLECULAR ION STATES

Using the two-particle notation introduced in the previous section, the ion and the ground-state atom of Sr is denoted by $|i g\rangle$ or $|g i\rangle$ depending on the position of the respective particles, where $|g\rangle = |5s^2, {}^1S_0\rangle$. The optical coupling of $|ig\rangle$, $|gi\rangle$ to $|ie^{\alpha}\rangle$, $|e^{\alpha}i\rangle$, respectively, is a two-photon process via the intermediate triplet state, $5s5p$, ${}^{3}P_1$ with an effective Rabi frequency $\Omega^{\alpha}(R)$. The coupling is determined by the dipole matrix element, $\mu^{\alpha}(R) = \langle 5s5p|\mu|e^{\alpha}(R) \rangle$. To relate this coupling strength to that of neutral gas experiments, all Rabi frequencies are expressed in terms of a reference Rabi frequency, Ω_{5s}^{50s} , for an isolated atom which, for our purposes, is chosen to be 40 MHz. Detuning of the laser with respect to a particular molecular Rydberg state is given as $\Delta^{\alpha}(R)$ = $\omega_L - [E^{\alpha,-}(R) + E^{\alpha,+}(R)]/2$ where ω_L is the frequency of the second photon. Using the dipole approximation for the laser field and the rotating wave approximation, the resulting Hamiltonian is

$$
\hat{H}_{\text{opt}}^{\text{tp}}(R) = \sum_{\alpha} \left[-\Delta^{\alpha} (R)(|ie^{\alpha}\rangle \langle ie^{\alpha}| + |e^{\alpha}i\rangle \langle e^{\alpha}i|) + \frac{\Omega^{\alpha}(R)}{2} (|ig\rangle \langle ie^{\alpha}| + |gi\rangle \langle e^{\alpha}i| + \text{H.c.}) + T^{\alpha}(R)(|ie^{\alpha}\rangle \langle e^{\alpha}i| + \text{H.c.}) \right].
$$
\n(3)

The electronic ground-state energies are set to zero. The Hamiltonian $\hat{H}_{\text{opt}}^{\text{tp}}$ is diagonalized to obtain exact solutions for the laser-dressed molecular states $|d_{\beta}\rangle$ along with the energies $\omega^{\beta}(R)$ (*h* is set to 1). $|d_{\beta}\rangle$ are expressed in terms of a superposition of the electronic ground states $|ig\rangle$, $|gi\rangle$ as well as the molecular excited states $|ie^{\alpha}\rangle$, $|e^{\alpha}i\rangle$. The index $\beta = 1, 2, \ldots$, represents the different dressed states. Of major interest is the pair of molecular states that has the largest contribution of electronic ground states. These states correspond to states with large lifetimes and are denoted by $|\tilde{g}_{1,2}\rangle$ with energies $\omega_{1,2}^{\tilde{g}}(R)$. Expressing the electronic dynamics as effective hopping between the relevant dressed states, we have

$$
\hat{H}_{\text{effec}}^{\text{tp}}(R) = U(R)(|i\tilde{g}\rangle\langle i\tilde{g}| + |\tilde{g}i\rangle\langle \tilde{g}i|)
$$

+ $J(R)(|i\tilde{g}\rangle\langle \tilde{g}i| + \text{H.c.})$, (4)

where $U(R)$ is the light shift associated with an ion-dressed atom pair and $J(R)$ is the effective hopping (see Fig. [1\)](#page-1-0). The definitions of $U(R)$ and $J(R)$ contain the details of the admixture of Rydberg state to the ground-state atom as determined by the laser parameters (refer to Appendix \mathbf{C}).

IV. CHARGE DYNAMICS WITH CLASSICAL AND QUANTUM MOTION OF RYDBERG-DRESSED MOLECULAR ION

Here classical and quantum dynamics refer to the motional states of untrapped and trapped ion-atom pairs, respectively. For an unconfined pair of particles, one obtains different hopping rates corresponding to different internuclear distances which leads to dephasing in the overall charge dynamics. The instantaneous state for classical dynamics is given by $|\psi(R, t)\rangle = c_{i\tilde{g}}(R, t)|i\tilde{g}\rangle + c_{\tilde{g}i}(R, t)|\tilde{g}i\rangle$ and the corresponding equations of motion using Eq. (4) are

$$
i\partial_t c_{i\tilde{g}}(R,t) = U(R)c_{i\tilde{g}}(R,t) + J(R)c_{\tilde{g}i}(R,t) ,\qquad (5)
$$

$$
i\partial_t c_{\tilde{g}i}(R, t) = U(R)c_{\tilde{g}i}(R, t) + J(R)c_{i\tilde{g}}(R, t). \tag{6}
$$

For a fixed internuclear distance R , one obtains the probability to be in state $|i\tilde{g}\rangle$ or $|\tilde{g}i\rangle$ to be $\cos^2[J(R) t]$. On averaging over the internuclear distance, the probability to obtain a particular two-particle state, for example, $|i\tilde{g}\rangle$ is given by $|\bar{c}_{i\tilde{g}}(t)|^2$ and is shown to slowly decay as seen in Fig. $3(b)$. To control the uncertainty in the position of either particle (ion or atom), we propose to have an identical confinement for the ion and atom [see Fig. $1(d)$]. This is achievable for alkaline-earth atoms as suggested in [\[44\]](#page-7-0) since the dynamic polarizability of a singly ionized alkaline-earth atom is comparable to that of a singly excited Rydberg alkaline-earth atom. In a double well, the electronic and motional degrees of freedom are entangled in the overall state given as $|\psi\rangle = \sum_{n,n'} (c_{i\tilde{g}}^{n,n'}(t)|i\tilde{g}\rangle|nn'\rangle +$ $c_{\tilde{g}i}^{n,n'}(t)|\tilde{g}i\rangle|nn'\rangle$). Here $|nn'\rangle$ is the eigenstate of the Hamiltonian corresponding to the center-of-mass dynamics for two particles in a double-well harmonic trap [\[46\]](#page-7-0) (refer to Appendix E). For a typical trapping frequency in the range of hundred kHz, the nuclear dynamics within the trap is much slower than the electronic dynamics and is solved under the Born-Oppenheimer approximation. It is assumed that the system is prepared in the lowest motional state denoted by $|00\rangle$. The probability to excite the first motional state can be

FIG. 3. (a) Color density plot is shown for M [see Eq. (7)] distinguishing coherent dynamics ($\mathcal{M} \ll 1$) from incoherent dynamics $(M \ge 1)$ for a range of detunings and lattice spacings. Relevant (un)gerade pair of potential curves are shown in the background as (red) blue lines for reference. (b) Plot depicts classical motional dynamics for an unconfined ion-atom pair by plotting $|\bar{c}_{i\bar{g}}(t)|^2$ which is calculated by averaging $c_{i\tilde{g}}(R, t)$ over the internuclear distance. (c) Plot shows the probability for an ion to be at a given site for an ion-atom pair trapped in a double well with trap frequency ω_{tr} = $(2\pi)80$ kHz and lattice spacing as 796 nm. The chosen detuning is -0.7 GHz which corresponds to $\mathcal{M} = 0.044$.

calculated from the off-diagonal couplings $\langle 00|U(R)|01\rangle$ and $\langle 00|J(R)|01\rangle$. If the off-diagonal couplings are smaller than the trapping frequency ω_{tr} and the corresponding light shifts, then we have coherent dynamics. To quantify the degree to which we couple the lowest motional states to their next higher motional state, the following parameter is introduced:

$$
\mathcal{M} = \frac{\left| U_{00}^{01} + J_{00}^{01} \right|}{\left| U_{01}^{01} - U_{00}^{00} + \omega_{\text{tr}} \right|} ,\tag{7}
$$

where $A^{nm}_{n'm'} = \langle nm | A(R) | n'm' \rangle$ and $A \in \{U, J\}$. If the coupling of $|00\rangle$ to $|01\rangle$ is small enough then the coupling to higher motional states such as $|02\rangle$ (or $|20\rangle$) are suppressed as well since they are higher-order processes. Thus, the lower the value of M, the more coherent is the charge dynamics. As expected, for sufficiently large trapping frequencies, it is possible to suppress the population of higher motional states. However, there are experimental limitations to how large the optical trap frequency can be and the typical values of $(U_{01}^{01} - U_{00}^{00})$ are comparable to ω_{tr} . Thus to have lower values of \mathcal{M} , we need $|U_{00}^{01}|, |J_{00}^{01}| \ll |U_{01}^{01} - U_{00}^{00}|$, which is easily satisfied for large-enough lattice spacings due to lower values of tunneling. In Fig. $3(a)$, M is represented in a two-dimensional color plot for different values of detuning and lattice spacing for a fixed trapping frequency of $\omega_{tr} = (2\pi)80$ kHz. It is always possible to find suitable lattice spacing and detuning to obtain coherent dynamics for our chosen trap frequency as done in Fig. 3(c). Whenever we work in the regime where the higher motional states are not populated, we can replace $U(R)$ and $J(R)$ by $\bar{U} = U_{00}^{00}$ and $\bar{J} = J_{00}^{00}$, respectively.

V. GENERALIZATION TO MANY-BODY CHARGE TRANSPORT

Here the two-particle picture will be generalized to many particles involving a deep optical lattice filled with a single atom per site out of which one is ionized. In the many-particle system, we assume that the excitation laser parameters and lattice parameter are selectively chosen such that the atoms that are nearest neighbors to the ion have the highest probability of Rydberg excitation. We also assume that the nearest-neighboring atom on either side of the ion remain in the blockade regime despite the polarization shifts [\[47\]](#page-7-0). The shift in the energy levels is strongest for atoms that are nearest neighbors to the ion than compared to those further away from it. To this effect, we have the following reduced basis for the many-particle picture: all atoms in the ground state with an ion at site $k \tbinom{|I_k|}{=}$ |*g*¹ ... *gk*[−]¹ *ik gk*⁺¹ ... *gN*-), a Rydberg atom to the right of the ion ($|R_k^{\alpha}\rangle = |g_1 \dots g_{k-1} i_k e_{k+1}^{\alpha} \dots g_N \rangle$) and a Rydberg atom to the left of the ion $(|L_k^{\alpha}\rangle = |g_1 \dots e_{k-1}^{\alpha} i_k g_{k+1} \dots g_N\rangle$). Any accidental Rydberg excitation for atoms further away from the ion do not contribute to the overall ion dynamics as the tunneling rate is negligible and it couples back to $|I_k\rangle$. The Hamiltonian for the optical coupling of the many-body system is given as

$$
\hat{H}_{\text{opt}}^{\text{mp}} = \sum_{\alpha} \sum_{k=1}^{N} \left[-\Delta_{k}^{\alpha} |R_{k}^{\alpha}\rangle \langle R_{k}^{\alpha}| + \frac{\Omega_{k}^{\alpha}}{2} (|R_{k}^{\alpha}\rangle \langle I_{k}| + \text{H.c}) - \Delta_{k}^{\alpha} |L_{k}^{\alpha}\rangle \langle L_{k}^{\alpha}| + \frac{\Omega_{k}^{\alpha}}{2} (|L_{k}^{\alpha}\rangle \langle I_{k}| + \text{H.c}) \right] + \sum_{\alpha} \sum_{k=1}^{N-1} \frac{T_{k}^{\alpha}}{2} [|R_{k}^{\alpha}\rangle \langle L_{k+1}^{\alpha}| + \text{H.c}]. \tag{8}
$$

On comparing with Eq. (3) , we find that the dependence of the optical parameters and the tunneling on distance *R* has been replaced by subscript k , which denotes the site number of the ion placed within the atomic lattice. Similar to the two-particle picture, we can diagonalize \hat{H}_{opt}^{mp} to obtain dressed states and focus on the many-body Rydberg-dressed ground states denoted as $|\tilde{I}^k\rangle = |\tilde{g}_1 \dots \tilde{g}_{k-1} i_k \tilde{g}_{k+1} \dots \tilde{g}_N\rangle$. Unlike in the two-particle picture, in the many-particle setup, the electron can, in principle, tunnel multiple times across the lattice before it couples back to the ground-state atom. Although we assume nearest-neighbor tunneling and work in the reduced basis, we find that the effective ion dynamics is delocalized in the Rydberg-dressed picture. This implies that the effective equations of motion couple $|\tilde{I}_k\rangle$ to $|\tilde{I}_{k\pm 2}\rangle$ as well and so on. Thus the effective ion dynamics in the many-body dressed atoms cannot be described simply by its nearest-neighbor exchange term unless we include additional constraints. However, using time-independent perturbation theory where $T_k \ll \Omega_k$ for all *k*, it is possible to derive the effective nearest-neighbor hopping term. The effective Hamiltonian obtained in this limit describes charge dynamics for an ion in the Rydberg-dressed atomic lattice,

$$
\hat{H}_{\text{effec}}^{\text{mp}} = \sum_{k} U_{k} (|\tilde{I}_{k}\rangle \langle \tilde{I}_{k}|) + J_{k,k+1} (|\tilde{I}_{k+1}\rangle \langle |\tilde{I}_{k}| + \text{H.c.})
$$

$$
+ J_{k,k-1} (|\tilde{I}_{k-1}\rangle \langle |\tilde{I}_{k}| + \text{H.c.}), \qquad (9)
$$

where $U_k = \langle \tilde{I}_k | \hat{H}_{\text{effec}}^{\text{mp}} | \tilde{I}_k \rangle$ and $J_{k,k+1} = \langle \tilde{I}_k | \hat{H}_{\text{effec}}^{\text{mp}} | \tilde{I}_{k+1} \rangle$. To identify regimes in the parameter space where the perturbation theory is valid, we resort back to the two-particle picture. We derive dynamical parameters (U^{pert} , J^{pert}) by solving $\hat{H}^{\text{tp}}_{\text{opt}}$ perturbatively in the limit $\Omega \gg T$. Averaging over the motional

FIG. 4. (a) Density plot shows the relative difference in the dynamical parameters defined in C [see Eq. (10)] calculated using exact as well as perturbative methods (refer to text) for different laser parameters. The light regions in the density plot correspond to regimes where the perturbation theory is valid. Relevant (un)gerade pair of potential curves are shown as (red) blue lines for reference. (b)–(c) Averaged values of *U* and *J* from different methods are shown for a particular detuning $\Delta = -0.7$ GHz.

states, U^{pert} , J^{pert} are compared to \bar{U} , \bar{J} which were obtained by solving Eq. [\(4\)](#page-2-0) without any approximation [referred to as the exact method in Figs. $4(b)$ and $4(c)$]. This is numerically quantified by the following parameter:

$$
\mathcal{C} = \left| \frac{\bar{U} - U^{\text{pert}}}{\bar{U}} \right| + \left| \frac{\bar{J} - J^{\text{pert}}}{\bar{J}} \right|.
$$
 (10)

Figure $4(a)$ shows the different values of C for different laser parameters and lattice spacing. As expected, for larger lattice spacings, the overall *T* is smaller which easily satisfies our condition for perturbation theory and corresponds to lower values of C . This is further confirmed in Figs. $4(b)$ to $4(c)$ where we compare the dynamical parameters from two different methods. We include the motional states in next section, where we work in the coherent regime of the many-body setup.

VI. COHERENT MANY-BODY CHARGE TRANSPORT WITH NEAREST-NEIGHBOR HOPPING

The theory for charge transport over many-sites can be understood using the simple model of pair-wise charge exchange at two sites involving the ion and its neighboring atom. Having identified the optimum optical parameters (0.7–1 GHz with respect to 50*s* Rydberg state) and lattice spacing (750– 850 nm) in prior sections to have nearest-neighbor coherent hopping, we use them in our numerical simulation for charge dynamics involving a single ion and $N - 1$ atoms in a one-dimensional optical lattice. The typical lifetime of the Rydberg states is estimated to be in the order of hundred μ s [\[48\]](#page-7-0) which is enhanced to 250–350 ms by averaging over the Rydberg-dressed states whose major contribution is from the electronic ground state. Thus the main time constraint on the overall dynamics is the decay of the intermediate state, $5s5p³P₁$, which is 21 μ s. By increasing the detuning with respect to the intermediate state, this is increased to 8.4 ms. This requires the effective Rabi frequency for the two-photon excitation scheme to be in the order of tens of MHz, which remains experimentally achievable [\[49–51\]](#page-7-0). We ignore the accidental resonances to Rydberg states or doubly excited

FIG. 5. Density plot showing ion probability during its dynamics in Rydberg-dressed lattice with a spacing of 796 nm and laser detuning −0.7 GHz. Top panels show the corresponding Rabi profile of the excitation laser. (a) Constant Rabi frequency, $\Omega_c = 40 \text{ MHz}$, where $J_0 = \bar{J}_{k,k+1} = 27$ kHz and $U_0 = \bar{U}_k = 15.9$ kHz for all values of *k*. (b) Varying Rabi profile along the *x* direction, $\Omega_k(x) =$ $\Omega_c \sqrt{(N - kx)(k - 1)x}$ such that $\bar{J}_{k,k+1} = (2J_0/N\sqrt{(N - k)k})$ and $\bar{U}_{k,k+1} = (2U_0/N\sqrt{(N-k)k}).$

states since the probability for it to occur is small (\sim 10⁻²) particularly when averaged over the motional states. Solving for $\Psi = \sum_{k=1}^{N} \tilde{C}_{k}^{I} | \tilde{I}_{k}$ using the many-body Hamiltonian Eq. [\(9\)](#page-3-0) gives

$$
i\partial_t \tilde{C}_k^I = \bar{U}_k \tilde{C}_k^I + \bar{J}_{k,k-1} \tilde{C}_{k-1}^I + \bar{J}_{k,k+1} \tilde{C}_{k+1}^I ,\qquad (11)
$$

where $\bar{U}_k = \langle 00 | U_k | 00 \rangle$ and $\bar{J}_{k,k+1} = \langle 00 | J_{k,k+1} | 00 \rangle$. In general, $\bar{J}_{k,k+1}$ does not have to be equal to $\bar{J}_{k+1,k}$. Figure 5 depicts the results of the numerical simulation for 13 sites with different laser profiles. Focusing on the excitation laser with constant Rabi profile [see Fig. $5(a)$], we have an ion initially located at site 7 which then propagates in both directions symmetrically as it has equal probability to hop in either direction at every instant. A scenario involving spatially varying Rabi profile is depicted in Fig. 5(b). In this case $\bar{J}_{k,k+1} \neq \bar{J}_{k+1,k}$ and the Rabi profile has been chosen in such a manner that it mimics motion of a particle in harmonic well [\[52\]](#page-7-0) with its minima at the center (site 7 in this case). Hence an ion situated at site 1 is akin to starting at one end of the well which then propagates through site 7 with maximum kinetic energy till it reaches the other end. The role of the light shift \bar{U}_k is simply an additional energy shift experienced by the atoms or ion in the lattice. This can be compensated by choosing an appropriate profile for the trapping laser.

VII. CONCLUSION

In this work, we propose and model the effective charge dynamics of an ion within trapped Sr atoms in an optical lattice. We conclude that optically trapped alkaline-earth atomsion systems can naturally serve as a platform for the study of charge transfer in a controlled many-body environment. Enhanced coherent charge dynamics requires the dressing of ground-state atoms to their Rydberg states [\[53,54\]](#page-7-0) and the provision of identical confinement for both the ion and the Rydberg-dressed atom [\[44\]](#page-7-0), both of which are potentially attainable with ongoing experiments with alkaline-earth atoms [\[55–60\]](#page-7-0). Recent experiments on optical trapping of ions [\[61\]](#page-7-0),

ion-Rydberg atoms [\[47,62\]](#page-7-0), and ion-dressed Rydberg atoms [\[63\]](#page-7-0) are all promising endeavors in realizing different aspects of this work.

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APPENDIX A: FULL HAMILTONIAN

In the simple picture of two particles where a Sr atom is next to an ion, the full Hamiltonian consists of three parts

$$
\hat{H} = (\hat{H}_{\rm el} + \hat{H}_{\rm opt}^{\rm tp} + \hat{H}_{\rm CoM}), \tag{A1}
$$

where \hat{H}_{el} is the electronic part, $\hat{H}_{\text{opt}}^{\text{tp}}$ represents the excitation of the atom to its Rydberg state and \hat{H}_{CoM} corresponds to the motion of the trapped particles in the lattice. We discuss each Hamiltonian in some detail in the following sections.

APPENDIX B: ELECTRONIC HAMILTONIAN

Similar to alkali atoms, we assume an effective model potential for the singly ionized alkaline-earth atom, reducing the many-electron problem to an effective two-electron atom problem. The model potential in atomic units is given as

$$
V_{\text{eff}}^{Sr^{+}}(r) = -\frac{1}{r} [2 + (Z - 2)e^{-a_1(l)r} + a_2(l)re^{-a_3(l)r}] - \frac{\alpha_c}{2r^4} [1 - e^{-(r/r_l)^6}], \tag{B1}
$$

where parameters $[a_1(l), a_2(l), a_3(l), \alpha_c, r_l]$ of the model potential are determined from fits to experimental data for low and intermediate levels of Sr^+ energies $[64]$. *l* is the orbital angular momentum. Using the singly ionized Rydberg wave functions as a basis, the atomic Sr Rydberg wave functions $\psi_{n,l}(\mathbf{r})$ were calculated using mean-field theory similar to Hartree-Fock theory. The mixing between Rydberg series are ignored and is justified in certain cases from experimental observations which show no high-lying perturbers in the Rydberg series.

For singly excited Rydberg states of strontium with very large principal quantum numbers $(n > 20)$, there is a large asymmetry in the orbit size of the Rydberg electron and the ground-state electron. Due to this asymmetry, we can treat the exchange interaction between the two valence electrons perturbatively. Since the effect of the inner valence electron (in its ground state) is negligible on the Rydberg electron, we write the electronic Hamiltonian in terms of the Rydberg electron, given as

$$
\hat{H}_{\rm el} = -\frac{\nabla^2}{2} + V_{\rm eff}^{Sr^{+}}(\mathbf{r}_{1}) + V_{\rm eff}^{Sr^{+}}(\mathbf{r}_{2}) + \frac{1}{\mathbf{R}},\qquad(B2)
$$

where $r_{i=1,2}$ is the relative position of the Rydberg electron with respect to either nucleus (see Fig. [6\)](#page-5-0) and *R* is the internuclear distance between the nuclei. We note that the basis states are not orthonormal given the small nonzero overlap

FIG. 6. The figure depicts a Sr Rydberg molecular ion trapped in a double well. Here $r_{i=1,2}$ is the relative position of the Rydberg electron with respect to either nucleus and *R* is the internuclear distance between the nuclei. Z_i is the relative displacement of the corresponding nucleus with respect to the center of lattice site *i*.

function, i.e., $\langle \psi_{n,l}(\mathbf{r}_{(1,2)}) | \psi_{n,l}(\mathbf{r}_{(2,1)}) \rangle \neq 0$. However for the internuclear distances considered in this work, the overlap function can be considered negligible. We solve the (approximate) eigenvalue problem by diagonalizing the following matrix Hamiltonian for \hat{H}_{el} :

$$
\begin{pmatrix} [P(r_1)] & [T(r_1, r_2)] \\ [T(r_2, r_1)] & [P(r_2)] \end{pmatrix}, \tag{B3}
$$

where $[P(r_i)]$ and $[T(r_i, r_j)]$ are block matrices given as

$$
[P(r_i)] = \begin{pmatrix} P_{nl}^{nl}(r_i) & P_{nl}^{n'l'}(r_i) & \cdots \\ P_{n'l'}^{nl}(r_i) & \ddots & \cdots \\ \vdots & \cdots & \ddots \end{pmatrix}, \quad (B4)
$$

$$
[T(r_i, r_j)] = \begin{pmatrix} T_{nl}^{nl}(r_i, r_j) & T_{nl}^{n'l'}(r_i, r_j) & \cdots \\ T_{n'l'}^{nl}(r_i, r_j) & \ddots & \vdots \\ \vdots & \vdots & \vdots \end{pmatrix}, \quad (B5)
$$

which are defined using the two center integrals $P_{n'l'}^{nl}(r_{i=1,2}) = \langle \psi_{n,l}(\mathbf{r_{i=1,2}}) | \hat{H}_{el} | \psi_{n',l'}(\mathbf{r_{i=1,2}}) \rangle$ and $T_{n'l'}^{nl}(r_1, r_2) = \langle \psi_{n,l}(\mathbf{r}_{(1,2)}) | \hat{H}_{\text{el}} | \psi_{n',l'}(\mathbf{r}_{(2,1)}) \rangle$. Equation [\(4\)](#page-2-0) given above can be a very large matrix to diagonalize for a large basis set and needs to be computed for internuclear distances that are finely resolved to obtain smooth potential curves. This is numerically cumbersome. However, using the "symmetry" between \mathbf{r}_1 and \mathbf{r}_2 in the electronic Hamiltonian, we express the Hamiltonian in the *gerade-ungerade* basis representation. The advantage of this representation is that the set of eigenvalue equations decouple between the states belonging to the two symmetry groups. Upon diagonalization, one obtains the following eigenvalue equation for the Rydberg states:

$$
\hat{H}_{\rm el}|e^{\alpha,\pm}\rangle = E_{\alpha}^{\pm}(R)|e^{\alpha,\pm}\rangle,\tag{B6}
$$

where α is the index for the Rydberg excited molecular ion state which in terms of the basis functions is expressed as

$$
|e^{\alpha, \pm}\rangle = \frac{1}{\sqrt{2}} (|ie^{\alpha}\rangle \pm |e^{\alpha}i\rangle)
$$

=
$$
\sum_{n,l} c_{n,l}^{\alpha, \pm} (|\psi_{n,l}(\mathbf{r}_1)\rangle \pm |\psi_{n,l}(\mathbf{r}_2)\rangle), \qquad (B7)
$$

along with the normalization condition $\sum_{n,l} |c_{n,l}^{\alpha,\pm}|^2 = 1$.

APPENDIX C: HAMILTONIAN FOR OPTICAL COUPLING TO RYDBERG MOLECULAR ION STATES

The laser-dressed molecular ion states are obtained by diagonalizing the following matrix Hamiltonian of $\hat{H}_{\text{opt}}^{\text{tp}}$

$$
\begin{pmatrix}\n0 & \frac{\Omega^{\alpha}(R)}{2} & \dots & 0 & 0 & \dots \\
\frac{\Omega^{\alpha}(R)}{2} & -\Delta^{\alpha}(R) & \vdots & 0 & T^{\alpha}(R) & \vdots \\
\vdots & \dots & \ddots & \vdots & \dots & \ddots \\
0 & 0 & \dots & 0 & \frac{\Omega^{\alpha}(R)}{2} & \dots \\
0 & T^{\alpha}(R) & \vdots & \frac{\Omega^{\alpha}(R)}{2} & -\Delta^{\alpha}(R) & \vdots \\
\vdots & \dots & \ddots & \vdots & \dots & \ddots\n\end{pmatrix}.
$$
\n(C1)

The size of the matrix Hamiltonian is $(2 + \alpha) \times (2 + \alpha)$ where α is the number of Rydberg molecular ion states. The eigenvalues obatined $\omega^{\beta}(R)$ were ordered in ascending magnitude of energies along with their corresponding eigenstates

$$
|d_{\beta}(R)\rangle = c_{\beta}^{g_1}(t)|g_1\rangle + c_{\beta}^{g_2}(t)|g_2\rangle
$$

+
$$
\sum_{\alpha=1} (c_{\beta}^{e_1,\alpha}(t)|e_1^{\alpha}\rangle + c_{\beta}^{e_2,\alpha}(t)|e_2^{\alpha}\rangle), \qquad (C2)
$$

where β represents different dressed molecular ion states. The states with lowest energy in magnitude are selected states and denoted as $|\tilde{g}_{1,2}\rangle$ with energies $\omega_{1,2}^{\tilde{g}}(R)$. These states would correspond to states with longest lifetimes depending on the fraction of the Rydberg population in them which in turn depends on the Rabi frequency and the detuning of the excitation laser.

APPENDIX D: ELECTRONIC DYNAMICS IN THE RYDBERG DRESSED MOLECULAR ION

We solve the time-dependent Schrödinger equation for $|\Psi\rangle$

$$
i\frac{d|\Psi\rangle}{dt} = (\hat{H}_{\text{el}} + \hat{H}_{\text{opt}}^{\text{tp}})|\Psi\rangle, \tag{D1}
$$

where $|\Psi\rangle = c_1^{\tilde{g}}(R, t) |\tilde{g}_1\rangle + c_2^{\tilde{g}}(R, t) |\tilde{g}_2\rangle$ is expressed in terms of the Rydberg-dressed ground states $|\tilde{g}_{1,2}(R)\rangle$ obtained in the previous section. The dynamical equations for the electron between the Rydberg dressed ground states are

$$
i\partial_t c_1^{\tilde{g}}(R,t) = \omega_1^{\tilde{g}}(R)c_1^{\tilde{g}}(R,t) , \qquad (D2)
$$

$$
i\partial_t c_2^{\tilde{g}}(R,t) = \omega_2^{\tilde{g}}(R)c_2^{\tilde{g}}(R,t). \tag{D3}
$$

The dressed states expressed in the left or right basis are defined as

$$
|i\tilde{g}\rangle = \frac{1}{\sqrt{2}}(\tilde{g}_1 + \tilde{g}_2), \qquad (D4)
$$

$$
|\tilde{g}i\rangle = \frac{1}{\sqrt{2}}(\tilde{g}_1 - \tilde{g}_2). \tag{D5}
$$

Using $|\Psi\rangle = c_{i\tilde{g}}(R, t)|i\tilde{g}\rangle + c_{\tilde{g}i}(R, t)|\tilde{g}i\rangle$ we have the following dynamical equations in the left or right basis,

$$
i\partial_t c_{i\tilde{g}}(R,t) = U(R)c_{i\tilde{g}}(R,t) + J(R)c_{\tilde{g}i}(R,t) , \qquad (D6)
$$

$$
i\partial_t c_{\tilde{g}i}(R,t) = U(R)c_{\tilde{g}i}(R,t) + J(R)c_{i\tilde{g}}(R,t) , \qquad (D7)
$$

where the dynamical parameters, on-site energy *U* (*R*), and the hopping rate $J(R)$ are defined as

$$
U(R) = \frac{\omega_1^{\tilde{g}}(R) + \omega_2^{\tilde{g}}(R)}{2},
$$
 (D8)

$$
J(R) = \frac{\omega_1^{\tilde{g}}(R) - \omega_2^{\tilde{g}}(R)}{2}.
$$
 (D9)

 $U(R)$ and $J(R)$ are the dynamical parameters that correspond to the light shift and the effective hopping rate, respectively, between the Rydberg-dressed ground states for an ion and an atom.

APPENDIX E: HAMILTONIAN FOR CENTER-OF-MASS DYNAMICS

Each ion core is trapped in its own lattice site in a onedimensional lattice along the *z* axis (see Fig. [6\)](#page-5-0). The Hamiltonian for the nuclear motion for our two-site model is given as

$$
\hat{H}_{\text{CoM}}|n_1 n_2\rangle = \sum_{i=1,2} \left[-\frac{\hbar^2 \nabla_{Z_i}^2}{2M} + \frac{M}{2} \omega^2 Z_i^2 \right] |n_1 n_2\rangle
$$
\n
$$
= \hbar (n_1 + n_2) \omega |n_1 n_2\rangle, \tag{E1}
$$

where $M = 87.2$ is the mass of Sr in atomic units. and $|n_{i=1,2}\rangle$ is the motional state at the corresponding site and $|n_1n_2\rangle$ is the two-particle motional eigenstate. $Z_{(1,2)}$ is the relative motion of the corresponding trapped nuclei at each site. The full wave function $|\Psi\rangle$ is a product state of the electronic eigenstates and the motional states,

$$
|\Psi\rangle = \sum_{n_1, n_2} \left(c_{n_1 n_2}^{i\tilde{g}}(t) \, |i\tilde{g}\rangle + c_{n_1 n_2}^{\tilde{g}i}(t) \, | \tilde{g}i\rangle \right) |n_1 n_2\rangle \;, \tag{E2}
$$

and solving the Schrödinger equation with the full Hamiltoian [see Eq. $(A1)$] and multiplying with $\langle n_1 n_2 |$ throughout we get

$$
i\partial_t c_{n_1 n_2}^{i\bar{g}} = \left(\langle n_1 n_2 | U(R) | n_1 n_2 \rangle + \omega_{n_1 n_2} \right) c_{n_1 n_2}^{i\bar{g}} + \left(\langle n_1 n_2 | J(R) | n_1 n_2 \rangle \right) c_{n_1 n_2}^{\bar{g}i} + \sum_{n_1, n_2 \neq n'_1 n'_2} \left[\langle n'_1 n'_2 | U(R) | n_1 n_2 \rangle \right] c_{n'_2 n'_2}^{i\bar{g}} + \sum_{n_1, n_2 \neq n'_1 n'_2} \left[\langle n'_1 n'_2 | J(R) | n_1 n_2 \rangle \right] c_{n'_2 n'_2}^{\bar{g}i}, \tag{E3}
$$

$$
i\partial_t c_{n_1 n_2}^{\tilde{g}i} = ((n_1 n_2 | U(R) | n_1 n_2) + \omega_{n_1 n_2}) c_{n_1 n_2}^{\tilde{g}i} + ((n_1 n_2 | J(R) | n_1 n_2)) c_{n_1 n_2}^{\tilde{g}i} + \sum_{n_1, n_2 \neq n'_1 n'_2} [(n'_1 n'_2 | U(R) | n_1 n_2)] c_{n'_2 n'_2}^{\tilde{g}i} + \sum_{n_1, n_2 \neq n'_1 n'_2} [(n'_1 n'_2 | J(R) | n_1 n_2)] c_{n'_2 n'_2}^{\tilde{g}i}.
$$
 (E4)

On averaging over the spatial variation of the dynamical parameters, we get couplings between the motional ground states but also to other higher motional states. The criterion for coherent charge dynamics is discussed in the article and deals with finding optimal parameters that minimizes this coupling to higher motional states. Here we also numerically verified that the variation of the Rydberg-dressed ground-state wave function over the internuclear distance is negligible.

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