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Observation of dipole-quadrupole interaction in an ultracold gas of Rydberg atoms

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We observe the direct excitation of pairs of Cs atoms from the ground state to molecular states correlating asymptotically to \( nsn’f \) asymptotes. The molecular resonances are interpreted as originating from the dipole-quadrupole interaction between the \( nsn’f \) pair states and close-by \( npnp \) asymptotes (22 \( \leq n \leq 32 \)). This interpretation is supported by Stark spectroscopy of the pair states and a detailed modeling of the interaction potentials. The dipole-quadrupole interaction mixes electronic states of opposite parity and thus requires a coupling between electronic and nuclear motion to conserve the total parity of the system. This non-Born-Oppenheimer coupling is facilitated by the near-degeneracy of even and odd \( L \) partial waves in the atom-atom scattering which have opposite parity.

The excitation of ultracold atoms to interacting Rydberg states has given rise to the observation of a wide range of fundamental phenomena such as the formation of ultracold neutral plasmas [1, 2], the conditional blockade of excitation (dipole-blockade) that can be exploited to realize basic quantum gates [3, 4], and the formation of macrodimers [5, 6]. In a macrodimer, two atoms are bound by long-range dispersion forces between two highly excited Rydberg atoms [7]. Most observations could be interpreted as arising from dipole-dipole interactions, but occasionally quadrupole-quadrupole interactions have been found to be relevant [8, 9]. These interactions conserve the electronic parity of the system of two interacting Rydberg atoms and obey the condition

\[
(-1)^{\ell_A,\ell_f+f_{0,\ell}} = (-1)^{\ell_A,\ell_f+f_{0,\ell}},
\]

where \( \ell \) represents the orbital angular momentum quantum number of the Rydberg electron, the indices \( i \) and \( f \) the initial and final pair states coupled by the interaction, and \( A \) and \( B \) the two atoms. Typical examples include \( ss-pp’ \) interactions (using the notation \( \ell_A,\ell_B;\ell_f,\ell_f \)), which are exploited in dipole-blockade experiments [10].

We report on the observation of sharp \( pp-sf’ \) pair resonances in the Rydberg spectrum of a dense ultracold Cs atom sample following excitation with an intense narrowband pulsed UV laser. \( pp-sf \) pair resonances violate condition (1) and are attributed to dipole-quadrupole interactions between pairs of Rydberg atoms. The surprising observation is the strength and ubiquity of these resonances in the spectrum of Cs, which, in retrospect, makes it surprising that they have not been observed or discussed before. Dipole-quadrupole interactions violate some of the symmetries usually employed to designate the quantum states of a homonuclear diatomic molecule. These symmetries include (i) the inversion symmetry, or \( g/u \) symmetry, of the electronic wavefunction upon inversion through the symmetry center, (ii) the electronic parity, or \( \pm \) electronic symmetry, of the electronic wavefunction upon inversion of all coordinates through an arbitrary point of the space-fixed coordinate system - this operator is formally equivalent to a reflection \( \sigma_v \) of the electronic wavefunction through a plane containing the internuclear axis \([11, 12] \) - (iii) the total (rovibronic) parity, and (iv) the total angular momentum \( J \) (or \( F \) if nuclear spins are considered) and its projections \( hM \) on a chosen space-fixed axis and \( h\Omega \) on the internuclear axis. In free space and neglecting the electroweak force [13], only (iii) and (iv) are strictly conserved. Because the total parity can be obtained as a product of the parities of the electronic and rotational wavefunctions, the violation of condition (1) observed in our experiments implies the breakdown of the \( \pm \) and \( g/u \) symmetries and the resulting entanglement of rotational and electronic motions.

In the standard treatment of long-range interactions between two particles \( A \) and \( B \) [14, 15] and its recent application to the treatment of Rydberg pair resonances [5, 16, 17], the interaction term \( V_{\text{inter}} \) of the total Hamiltonian (in atomic units)

\[
H = H_A^0 + H_B^0 + V_{\text{inter}}
\]

\[
V_{\text{inter}} = \sum_{L_{A/B}=1}^{\infty} \sum_{\Omega=-L_{A/B}}^{L_{A/B}} \sum_{L_{\text{inter}}=0}^{\infty} \sum_{\Omega_{\text{inter}}=-L_{\text{inter}}}^{L_{\text{inter}}} \frac{(-1)^{L_{A/B},L_{\text{inter}},L_{A/B},L_{\text{inter}}}}{R^{L_{A/B}+L_{\text{inter}}+1}} Q_{L_{A/B}L_{\text{inter}}}(\vec{r}_{A/B}) Q_{L_{\text{inter}}}(\vec{r}_{\text{inter}})
\]

is expressed in the frame of the nonrotating molecule \( AB \) [18], in order to obtain potential-energy functions, which adequately describe the interaction at long range. Eq. (2) therefore neglects, by design, electronic-rotational interactions. In Eq. (2), \( H_0^A \) is the unperturbed Hamilton operator of atom \( i \), \( Q_{L_{i},\Omega}(\vec{r}_i) \) corresponds to the \( L_{i} \)-th multipole moment of atom \( i \) (\( L_i=1 \) and \( L_i=2 \) are the dipole and the quadrupole moment, respectively), and \( L_{\text{inter}} \) is the smaller value of \( L_A \) and \( L_B \). The dif-
ferent terms of the sum in $V_{\text{inter}}$ transform under $\sigma_z$ as $(-1)^{L_A+L_B}$ [19], which implies that only multipole interactions with even values of $L_A + L_B$ conserve the electronic parity.

The experiments were performed on Cs atoms released from a crossed optical dipole trap ($\lambda = 1064$ nm, $P = 10$ Watt). Using saturated absorption imaging, we determine typical particle numbers of $N = 10^6$, densities of $n = 10^{12}$ cm$^{-3}$, and temperatures of 40 $\mu$K. We employ a one-photon transition to excite atoms from the 6s$_{1/2}$ ground state to np Rydberg states. The UV light ($\lambda \approx 320$ nm) is provided by a cw ring dye laser (Coherent 899-21) which is frequency doubled after pulsed amplification [20]. The laser pulses are 4.4 ns long with a transform-limited bandwidth of 140 MHz and pulse energies of up to 100 $\mu$J. The laser is focused down to a waist of $\sim 100$ $\mu$m, still significantly larger than the size of the atomic sample ($\sigma \approx 30$ $\mu$m). Varying the intensity of the UV laser did not lead to an observable shift of the transitions. The laser frequency is calibrated using a wavemeter (High Finesse WS6-200). For detection, we rely on spontaneous ionization of the excited Rydberg sample by photoinitiated collisions (PIC) [17] and subsequent detection of the resulting ions. Alternatively, we can detect np states with $n \geq 27$ using pulsed field ionization (PFI) [21]. Spectra obtained with PIC and PFI are very similar, however the former have a better signal-to-noise ratio and thus only those are shown.

The dipole-quadrupole interaction is observed as the appearance of additional resonances in Rydberg spectra around np$_{3/2}$ ($22 \leq n \leq 32$), see Fig. 1, when the Rydberg-atom density is increased by increasing the UV laser intensity. The positions of these resonances correspond closely to the asymptotic energies of pairs of Rydberg atoms in ns$_{1/2}(n-3)f_f$ ($J=5/2,7/2$) and $(n+1)s_{1/2}(n-4)f_f$ states indicated in each spectrum by vertical lines. For $n < 22$, larger detunings and reduced matrix elements cause the ns$nf$ pair resonances to be too weak to be observable in our experiment. For $n > 32$, more strongly coupled pair resonances mask the ns$nf$ resonances. At $n = 23$, the two fine-structure components of the 24s$_{1/2}$18f$_f$ pair state are almost degenerate with the 23p$_{1/2}$23p$_{3/2}$ pair state, and at $n = 24, 25$, the detunings of the ns$_{1/2}(n-3)f_f$ pair states from the np$_{3/2}$np$_{3/2}$ pair states are too small to be resolved in our experiment.

The excitation of ns$nf$ pair states could in principle also result from $\ell$ mixing induced by an external electric field [9]. Using Stark spectroscopy of high-lying Rydberg states, we estimate residual stray electric fields to be less than 50 mV/cm. For the investigated quantum numbers $22 \leq n \leq 32$ such small electric fields lead to negligible $\ell$ mixing. We experimentally verified this expectation by intentionally applying larger electric fields and observing the shift of the ns$nf$ pair resonances (see Fig. 2 for a measurement at $n = 22$). The observed line shifts are in good agreement with the calculated asymptotic Stark shifts of the pair states, which are the sum of the Stark shifts of the corresponding atomic states, calculated following Ref. [22]. Although the spectral features are broadened because of the splitting of the magnetic sublevels by the field, the integrated line intensity remains approximately constant. If the observation of ns$nf$ pair resonances were caused by an external electric field, one would instead expect an increase in the line intensities by several orders of magnitude.

To model the experimental spectra, we determine the matrix elements of the electronic Hamiltonian (2) for $L_A, L_B = 1, 2$ and fixed $\Omega = \omega_A + \omega_B$ in the atomic basis $|\ell_A\ell_B\omega_A, \nu_B|\equiv |\gamma_{\ell_A\ell_B\omega_A, \gamma_B\nu_B}|$, following the general approach of Refs. [8, 23, 24]. The Hamiltonian omits interatomic electron-correlation. This omission is permitted if the internuclear distance exceeds the
to this work (e.g., $R \approx 0.23 \mu$m for $n = 22$, see Fig. 3 a)) and ultracold temperatures, $\Delta E_{\text{rot,max}}$ is always smaller than the linewidth of the pair states (approximated by the sum of atomic linewidths from Ref. [29]) and therefore the rotational states of opposite parity can be considered as degenerate. We therefore assume the coupling between electronic and rotational motion to be the same for interaction terms which require a change in $L$ (i.e., dipole-quadrupole) as for terms which do not require a change in $L$ (i.e., dipole-dipole and quadrupole-quadrupole). This assumption is justified a posteriori by the agreement between simulated and experimental spectra, as is now explained.

Diagonalization of $H$ for different internuclear separations $R$ yields the eigenstates $|\Psi(R)\rangle = \sum \alpha_{\alpha A J A \omega A, \gamma B J B \omega B}(R) |\alpha A J A \omega A, \gamma B J B \omega B\rangle$ with eigenenergies $E_{\Psi}(R)$ [30]. The resulting potential-energy curves are shown in Fig. 3a) for $n = 22$ (gray lines). The states correlating to the $22_{1/2}^{19}f_{1/2}$ asymptotes are all attractive, while the states correlating to the $23_{1/2}^{18}f_{1/2}$ asymptotes are all repulsive. The origin of this apparent repulsion between these asymptotes, which are not directly coupled by $H$, lies in the large number of coupled states with both negative and positive detuning, especially the dipole-dipole coupled asymptotes $20k23p$ and $21l22p$ at detunings of approximately $-60$ GHz and $+100$ GHz, respectively. The coupling of pair states with $pp$ and $sf$ character leads also to the appearance of minima in some potential energy curves. In Fig. 3a) such a minimum for a state with $\Omega = 2$ (blue curves) is visible below the $22p_{3/2}22p_{3/2}$ asymptote. This state has a strong $pp$ character and represents a bound macrodimer under field-free conditions. We estimate a vibrational frequency of about $20$ MHz, which is larger than the expected linewidth and we predict that the vibrational structure of this state should be observable.

The direct diagonalization of $H$ in the atomic basis results in a large number of potential-energy curves. However, very few of them can be excited from the $6s_{1/2}6s_{1/2}$ ground state. The excitation probability $p_{\Psi}(R)$ of an eigenstate $|\Psi\rangle$ is proportional to the square of the overlap of the molecular wavefunction with the laser-excited $np_{3/2}np_{3/2}$ pair state $|\phi\rangle = \sum a_{J A m A , J B m B} |j A m A , j B m B\rangle$ in the laboratory-fixed system

$$p_{\Psi}(\theta, R) = \left| \langle \Psi(R) | \phi \rangle \right|^2 = \left| \sum_{j A m A, j B m B} a_{j A m A, j B m B} d_{j A m A}^{\phi}(\theta) d_{j B m B}^{\phi}(\theta) |j A m A, j B m B\rangle \right|^2,$$

where $\theta$ is the angle between the internuclear and the laboratory-fixed $z$ axis and $d_{j m}^{\phi}(\theta)$ is Wigner’s $d$ matrix describing the rotation from the laboratory-fixed system to the molecular-fixed system [31]. Neglecting nuclear spin and assuming an initial random population of the $m_A$ and $m_B$ components of the $6s_{1/2}$ ground state, one
obtains four possible pair states $|\phi_i\rangle$ after excitation with circularly polarized laser radiation. We obtain $p_\Phi$ by summing over these four initial states with equal weights. After averaging $p_\Phi(\theta, R)$ over $\theta$ we obtain $\overline{p_\Phi}(R)$, which is shown as color shading in Fig. 3a) for $n = 22$. The spectral line profile is then given by [8]

$$s(E) = \frac{\omega^4_{\text{atom}}}{(E_{pp} - E)^2} \int_0^\infty \sum_\Psi G(E - E_\Psi(R))|\overline{p_\Psi}(R)|R^2dR ,$$

(3)

where $E = 2h\nu$, $E_{pp}$ is the energy of the np$_{3/2}$np$_{3/2}$ pair state, $\omega_{\text{atom}}$ is the atomic Rabi frequency, and $G$ is the spectral-density function of the laser, assumed to be a Gaussian with a full width at half maximum of 140 MHz. The resulting spectrum for $n = 22$ is shown in Fig. 3b) and in Fig. 1 for $22 \leq n \leq 32$. An analysis of the interaction matrix shows that the coupling relevant for the excitation of sf pair states is indeed the direct off-resonant dipole-quadrupole coupling to the p$_{3/2}$p$_{3/2}$ states.

The simulations reveal a small shift of the sf resonances from the asymptotic pair energies (see Fig. 3b)), which is also visible in the experimental spectra (Fig. 1 and Fig. 2), in contrast to the dipole-dipole-coupled pair states observed in previous works [5, 8]. The reason for the shift of the dipole-quadrupole resonances lies in the $R^{-6}$ dependence of the off-resonant dipole-dipole interaction (which dominates the $R$-dependent energy shifts of the sf pair states) and the $R^{-8}$ dependence of the off-resonant dipole-quadrupole interaction (which admixes pp character to the sf pair states). At distances where the latter interaction leads to a significant excitation probability ($R < 0.25\mu m$ for $n = 22$), the energy shifts caused by the former interactions are already significant.

The relative intensities of the dipole-dipole coupled ss' features (marked by asterisks for $30 \leq n \leq 32$ in Fig. 1) and the dipole-quadrupole coupled sf features are well captured by our model, justifying the equal weighting of L-changing and L-conserving interaction terms (see discussion above). The dependence of the visibility of the pair resonances on the principal quantum number $n$ is also well captured by the model, although we did not investigate this dependence systematically in our experiments. Combining the relative electronic transition strengths from our model with Monte-Carlo simulations of the atom distribution in the ground state sample we can estimate the relative strength of pair resonances and atomic transitions. For the 23s$_{1/2}$18f$_5$ resonance (see e.g. Fig. 2) and the 22p$_{3/2}$ atomic transition we find a ratio of about $10^{-4}:1$ which is consistent with our experimental observations, though the dynamic range of our intensity measurement prevents a quantitative assessment. The quadratic dependence of the pair excitation (see Eq. (3)) on the laser intensity, which we also observe experimentally, together with the large detunings of these pair resonances (see Fig. 1) necessitate large atomic Rabi frequencies and thus the use of an intense laser for the excitation. However, at $n = 23$, where the energy of the 23p$_{3/2}$23p$_{3/2}$ pair state lies in between the energies of the 24s$_{1/2}$19f$_5$ and 24s$_{1/2}$19f$_7$ asymptotes, we expect that the molecular resonance should also be observable in spectra recorded with a (less intense) cw laser. At this $n$, an electric field around 10 V/cm would be sufficient to bring the 24s$_{1/2}$19f$_5$ pair state into exact resonance with the 23p$_{3/2}$23p$_{3/2}$ pair state, allowing for the investigation of Förster resonant energy transfer with a $1/R^4$ dependence of the coupling (instead of the well studied $1/R^3$ dependence [32, 33]).

As discussed above, the dipole-quadrupole interaction couples electronic and rotational degrees of freedom, effectively entangling the electronic and rotational motion of the two atoms. Dipole-dipole interactions between Rydberg atoms have been suggested as means to manipulate their internuclear separation [34]. In analogy, the dipole-quadrupole interaction might allow for the control of the rotational motion of two ultracold atoms. The significance of our results is (i) that they provide empirical evidence for the inadequacy of Eq. (2) and of g/u and electronic ± symmetry to describe, in general, interacting Rydberg atoms near pair resonances violating condition (1), and (ii) that they suggest and justify a simple procedure with which the rotational-electronic interactions for odd values of $L_A+L_B$ may be incorporated as effective potential terms in $V_{\text{inter}}$ when the rotational levels can be regarded as “degenerate”. A similar assump-
tion, though not explicitly declared as such, appears to have been made in the recent discussion of the permanent dipole moment of Rydberg atoms interacting with ground-state atoms [35]. The effective dipole-quadrupole terms in $V_{\text{inter}}$ give rise to potential-energy contributions scaling as $R^{-8}$ and it appears that whenever such dispersion terms are reported for homonuclear diatomic molecules (see e.g. Ref. [15] for the $X^1\Sigma^+_g$ ground state of $H_2$) this assumption is also implicitly made.

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[19] The $Y_{l0}$ functions transform as $(-1)^{l-\Omega}$ under $\sigma_v$ [11], regardless of the position of their center along the inter-nuclear axis.
[30] In order to reproduce the experimental spectrum around the asymptote npnp we found it necessary to include the pair states $n_A s-n_B s$, $n_A p-n_B p$, $n_A d-n_B d$, $n_A f-n_B f$, $n_A s-n_B f$, and $n_A p-n_B d$ with $|n_{A/B}^* - n_p^*| \leq 1$ and $|E_{n_A s,n_B f} - E_{n_p p}| \leq 0.6|E_{n_p p,n_B p}|$ in the basis set.
[31] Because of a Cooper-minimum in the photoexcitation cross section from the 6s1/2 state to the np1/2 (np1/2) states (continuum) of Cs just above the ionization threshold [36], only the photoexcitation to np3/2np3/2 states is of relevance.