Formation of Rydberg macrodimers and their properties

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We investigate the interaction between two rubidium atoms in highly excited Rydberg states, and find that very long-range potential wells exist. These wells are shown to support many bound states. We calculate the properties of the wells and bound levels, and show that their lifetimes are limited by that of the constituent Rydberg atoms. We also show how these \(\mu\)m-size bound states can be populated \textit{via} photoassociation (PA), and how the signature of the ad-mixing of various \(\ell\)-character producing the potential wells could be probed. We discuss how sharp variations of the PA rate could act as a switching mechanism with potential application to quantum information processing.

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In recent years, the strong interactions between Rydberg atoms, due to their exaggerated properties \cite{1}, have been detected experimentally \cite{2,3}, and led to proposals for quantum computing \cite{4}, \textit{e.g.} to achieve fast quantum gates \cite{5,6} or to study quantum random walks \cite{7}.

One effect of these interactions is the excitation blockade \cite{8}, where a Rydberg atom prevents the excitation of nearby atoms \cite{9–13}. Recently, dipole blockade between two atoms has been observed in microtraps \cite{14,15}, and a C-NOT gate implemented \cite{16}. Another signature of these strong interactions is the molecular resonance features in excitation spectra, first observed in Rb \cite{17} and in Cs atoms \cite{18}. Other molecular features involving Rydberg excitations, such as the so called \textit{trilobite} and \textit{butterfly} states, where one atom is in its ground state and another in a Rydberg state \cite{19}, have recently been detected \cite{20}, while polyatomic molecules involving Rydberg electrons have also been studied \cite{21,22}.

In this article, we investigate the existence of long-range potential wells produced by two atoms excited into Rydberg states. As opposed to previous predictions of \textit{macrodimers} \cite{23} – double-excited Rydberg molecules – with very shallow wells due to induced van der Waals (vdW) interactions, the macrodimers we study here are due to the strong mixing of \(\ell\)-characters of various Rydberg states. As we will see, these molecular wells are deeper (a few GHz), but still very extended with an equilibrium separation of 1 \(\mu\)m or so. We concentrate on wells existing in the vicinity of two rubidium (Rb) atoms excited to the \(np\) Rydberg state, for which the relevant molecular symmetries are \(0_1^+, 0_u^+\), and \(1_u\) \cite{24}.

We build our long-range molecular potential curves following the procedure described in \cite{24}: we consider two free Rydberg atoms in states \(|a\rangle \equiv |n, \ell, j, m_\ell\rangle\) and \(|a'\rangle \equiv |n', \ell', j', m_{\ell'}\rangle\), where \(n\) is the principal quantum number, \(\ell\) the orbital angular momentum, and \(m_\ell\) is the projection of the total angular momentum \(\vec{j} = \vec{\ell} + \vec{s}\) onto a quantization axis (chosen in the \(z\)-direction for convenience). The long-range molecular potential curves in Hund’s case (c) are calculated directly by diagonalizing the interaction Hamiltonian consisting of both the Rydberg-Rydberg interaction and the atomic fine structure \cite{24}. The basis states are built as follows:

\[
|a, a'; \Omega_{g/u}\rangle \sim |a\rangle_1 |a'\rangle_2 - p(-1)^{\ell+\ell'} |a'\rangle_1 |a\rangle_2
\]

where \(\Omega = m_\ell + m_{\ell'}\) is the projection of the total angular momentum on the molecular axis and is conserved. The quantum number \(p\), describing the symmetry property under inversion, is \(1(-1)\) for \(g(u)\) states. For \(\Omega = 0\), the states’ symmetry under reflection must also be considered \cite{24}. The basis \cite{1} assumes no overlap of charge distributions between the two atoms, so the long-range

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interaction can be expanded as an inverse power series of the nuclear separation distance $R$ [24].

Fig. 1 shows the results of the diagonalized interaction matrix for the $0^+_g$ symmetry near various $n\ell + n\ell$ asymptotes. The left panel shows curves for several states located around $np_{3/2} + np_{3/2}$ (in red) and $ns + ns$ (in red) near $n = 70$, 69, 68, and lower asymptotes as well ($n \sim 58$ and 40). These plots depict the strong mixing of different $\ell$-characters due mainly to the dipole-dipole interactions (although higher orders are also present). The right panel enlarges a major feature arising from the $\ell$-mixing near $70p_{3/2} + 70p_{3/2}$, namely a large potential well (between 0 and -2.5 GHz) with a depth $D_c \sim 1$ GHz measured from its minimum (equilibrium separation) $R_e$ at roughly 40,500 $a_0$ ($a_0$: Bohr radius), i.e. over 2 $\mu$m. The $R$-dependence for lower asymptotes is also shown on the same panel; as $n$ decreases the potential curves become flatter (for same $R$ interval and energy scale). This is particularly true as $n$ reaches 40 or so.

The long-range well correlated to the $(n-1)s + (n+2)s$ asymptote is a general feature for this system; we depict a few cases in Fig. 2(a). The well depth $D_c$ and equilibrium separation $R_e$ scale roughly as $n^{-3}$ and $n^{2.5}$ (Fig. 2(b)), respectively, in good agreement with the $D_c \propto n^{-3}$ and $R_e \propto n^{7/3}$ scalings expected for a dominantly dipolar interaction between states. This last result can be obtained by keeping only the major $n$-dependence, so that the interaction matrix between two asymptotes has the form $n^{-3}\tilde{h}_0 + n^3R^{-3}\tilde{h}_1$ where $\tilde{h}_0$ and $\tilde{h}_1$ contain the energy spacings between the asymptotes and the strengths of the dipole-dipole coupling, respectively. If we define $\rho \equiv n^3R^{-3}$ and rewrite the interaction matrix as $n^{-3}(\tilde{h}_0 + \rho \tilde{h}_1)$, it becomes clear that its eigenvalues $E_i$ take the form $E_i = n^{-3}f_i(\rho)$. Since $\partial E_i/\partial R = -3n^2R^{-4}f_i(\rho) = 0$ at an extremum, we must have $f'_i(\rho) = 0$, which is only fulfilled for some particular values $\rho_c$. In the parameter space of $R$ and $n$, this leads to $\rho_c = n^{-7/3}|R_e| = \text{const.}$, so that $R_e \propto n^{7/3}$, and $f_i(\rho_c) = n^{-3}E_i|_{\rho_c} = \text{const.}$, so that $D_c \propto n^{-3}$. The difference between the analytical and numerical scalings reflects the more complex nature of the coupling than the assumed dipolar interaction.

We return to the curve correlated to the $69s + 72s$ asymptote in the $0^+_g$ symmetry. Its well is deep enough to support several bound molecular levels, a few of which are listed in Table 1 and shown in Fig. 2(c). Since the levels are separated by a few MHz, they could be detected by spectroscopic means. Oscillation periods of a few $\mu$s, related to the MHz frequency of these deepest levels, are rapid enough to allow several oscillations during the lifetime of the Rydberg atoms (roughly a few 100 $\mu$s for $n \sim 70$). The classical inner and outer turning points of the levels indicate very extended molecules, hence we keep the term macrodimer to describe them.

The molecular electronic basis states $|\lambda(R)\rangle$ corresponding to the potential curves $U_\lambda(R)$ are composed of several molecular electronic basis states $|\Pi\rangle$; the exact mixing (after diagonalization) varying with $R$:

$$|\lambda(R)\rangle = \sum_j c_j(\lambda)(R)|j\rangle. \quad (2)$$

Here, $c_j(\lambda)(R)$ are the eigenvectors and $|j\rangle$ the electronic basis states. As an example, we illustrate the composition of the well correlated to the $69s + 72s$ asymptote in Fig. 3(a). If we write $|aa\rangle \equiv |a, a; 0^+_g\rangle$, this well is composed primarily of $|69s72s\rangle$, $|70s71s\rangle$, $|70p_{3/2}70p_{1/2}\rangle$, $|69p_{3/2}21p_{1/2}\rangle$, and $|69p_{1/2}21p_{3/2}\rangle$; these states are indicated by different colors in Fig. 3(a). Using the same color labeling, we show the corresponding probabilities $|c_j(\lambda)|^2$ of all five states against $R$ in Fig. 3(b).

As expected, the right side of the well (near the $69s + 72s$ asymptote) is composed primarily of the $|69s72s\rangle$ state, while the left side contains mostly $|69p_{3/2}21p_{1/2}\rangle$, a state whose asymptote is above $69s + 72s$. Although one could have expected $|68p72p\rangle$ to provide most of the mixing, as suggested by Fig. 3(a), the proximity of the $69s + 72s$ and $69p_{1/2} + 71p_{3/2}$ asymptotes and their strong dipole coupling result in this larger mixing. The probability for $|68p_{3/2}72p_{1/2}\rangle$ is also included in Fig. 3(b).

The coupling to states correlated to lower asymptotes could lead to predissociation of the macrodimers and their decay into free atoms, heating up the sample with more energetic collisions and ionization. To estimate the predissociation lifetime of our bound states, and because of the large number of coupled electronic states, we assume that the total resonance width $\Gamma_1$ associated to the nonadiabatic transition from the molecular state 1 is just the sum over the widths $\Gamma_{1i}$ associated to transitions to individual adiabatic curves $i$, namely $\Gamma_1 = \Sigma_i \Gamma_{1i}$. To compute the widths, we use a simple
two-channel approach in which the nonadiabatic coupling $V_{ii}$, between the close channel corresponding to the molecular bound state (with electronic adiabatic basis state $|\chi_1(R)\rangle$), and the open channel corresponding to the dissociation state (with electronic adiabatic basis state $|\chi_i(R)\rangle$), is $V_{ii}(R) = -\frac{\hbar \alpha}{2 \mu} (\chi_1(R) | \partial_R | \chi_i(R) \rangle \langle \partial_R | \chi_1(R) \rangle$, where $\mu$ is the reduced mass of the system. We then obtain the width $\Gamma_{i1}$ using a Green’s function method \[25\]: $\Gamma_{i1} = 2\pi |\langle \phi_i | V_{ii} | \phi_{reg} \rangle|^2$, where $\phi_{reg}(R)$ is the regular, energy normalized solution of the open channel $i$ (in the absence of channel coupling), and the bound vibrational state $\phi_i(R)$ in the closed channel $1$. For the 69s72s $0^+_r$ potential curve, this approach give predissociation lifetimes that are extremely long, at least $10^8$ years, and we thus can conclude that these long-range Rydberg molecules are limited by the lifetimes of the Rydberg atoms themselves. This result could be expected, since the most important states contributing to the well are above the 69s + 72s asymptote (Fig. 3(b)). We note however that for other wells due to mixing with lower asymptotic electronic electronic states, predissociation would play a major role in the lifetime of macrodimers.

In order to detect and probe the properties of the macrodimers predicted above, one can photoassociate two ground state atoms using intermediate Rydberg states that provide good overlap with the target long-range molecular states. Exciting the two ground state atoms into an intermediate Rydberg state $ns$, $np$, or $nd$ will allow us to probe the different $\ell$-characters (Fig. 3) in the bound level. The photoassociation (PA) rate $K_v$ into a bound level $v$ can be calculated \[26\] using

$$K_v \propto I_1 I_2 \sum_j |\langle d_1 d_2 \rangle_j|^2 \int_0^\infty dR \phi_i^\ast(R) c_j^\ast(R) \phi_g(R),$$

where $d_1 = \langle n_j f_j | e_1 | n_g f_g \rangle$ and $d_2 = \langle n'_j f'_j | e_2 | n'_g f'_g \rangle$ are the electronic dipole moments between electronic states $|a_j, a'_j\rangle$ and $|a_g, a'_g\rangle$ for atom 1 and atom 2, respectively.

We calculate $K_v$ by assuming that the Rb 5s atoms
are first excited to an intermediate state that we con-
side-er to be the electronic “ground” states $|x_0⟩$, before being excited to their final electronic states. For sim-
plicity, we consider lower asymptotes in the vicinity of $n = 40$ for which $|x_0⟩$ does not vary with $R$ (Fig. 1). As shown in Fig. 3, the final electronic state contain mostly $|np, n'p⟩$ or $|ns, n's⟩$; thus, we assume transitions from $40p_{3/2} + 40p_{3/2}$ to the $ns + n's$ components and from $41s + 41s$ to the $np + n'p$ components. In Fig. 4 we show $K_v$ against the detuning $Δ$ from the atomic $70p_{3/2}$ levels. For simplicity, we set $φ(K)$ constant, although other choices could be made to enhance $K_v$, such as a gaussian corresponding to the motion of atoms trapped in optical tweezers. The PA rate starting from both atoms in $40p_{3/2}$ is shown in red, and from $41s$ in turquoise, respectively, on a linear scale in (a) and a logarithmic scale in (b). In both cases, the rapid oscillation between a large rate for an even bound level ($v = 0, 2, 4, \ldots$) and a small rate for odd levels ($v = 1, 3, 5, \ldots$) gives the apparent envelop of the signal. This is illustrated in Figs. (c) and (d), where a zoom of the deepest levels $v$ is shown for both “ground” states (on a log-scale). This can be understood from the integral in Eq. (4), which is near zero for an odd wave function $φ_v(R)$.

The signature of a macrodimer would manifest itself by the appearance of a signal starting at $Δ \sim -0.93$ GHz red-detuned from the $70p_{3/2}$ atomic level, and ending abruptly at $\sim -1.36$ GHz. In addition, the PA rate from either “ground” state $40p_{3/2}$ or $41s$ can reveal the details of the $ℓ$-mixing in the potential well. First, the overall larger signal for $41s$ reflects the scaling of $d_1d_2$ appearing in Eq. (4), with $41s$ being closer than $40p_{3/2}$ to most atomic states appearing in the electronic molecular curve (see Fig. 3). As $Δ$ increases from $-1.36$ GHz, $K_v$ mimic the probabilities $|c_ℓ(R)|^2$. For $41s$, the progressive decrease followed by sharp increases between $-1.22$ and $-1.16$ GHz correspond to the slow decreases of most $|np, n'p⟩$ components between $R \sim 40 - 50$ a.u. and their sharp increases around $R \sim 33 - 35$ a.u. (especially from $|69p_{1/2}, 71p_{3/2}⟩$, $|69p_{3/2}, 71p_{1/2}⟩$, and $|70p_{3/2}, 70p_{1/2}⟩$; see Fig. 3). This is followed by another large feature between $-1.13$ and $-1.09$ GHz (corresponding to the sharp increase in $|69p_{1/2}, 71p_{3/2}⟩$ and $|69p_{1/2}, 71p_{1/2}⟩$ around $R \sim 53,000$ a.u.), and a “steady” rate before the signal drops to basically zero. For $40p_{1/2}$, we obtain two obvious features: first, a significant drop in $K_v$ between $-1.13$ and $-1.09$ GHz which mirrors the increase mainly in $|69s, 72s⟩$, and second, a steady growth corresponding to the rising probability of $|69s, 72s⟩$ beginning at about 55,000 a.u. (before the signal drops to basically zero).

In conclusion, we have shown the existence of potential wells for the $0^+_1$ symmetry of doubly-excited atoms due to $ℓ$-mixing. These wells support several bound levels separated by a few MHz with lifetimes limited by the Rydberg atoms themselves. These vibrational levels could be populated and detected by photoassociation.

We showed that the signature of macrodimers would be the appearance of a signal beginning and ending abruptly, and that features in that signal could be used to probe the $ℓ$ character of the potential well. The detection of such extended molecules with a lot of internal energy is in itself a goal, and would allow to study how macrodimers are perturbed by surrounding ground state atoms. In addition to interesting exotic chemistry (e.g., by an approaching Rydberg atom leading to a strong ultra-long van der Waals complex), the dependence of the PA rate on the exact $ℓ$-mixing (Fig. 4) could potentially be used as a switch with application in quantum information (e.g., by turning on and off the blockade effect). The work of N.S. was supported by the National Science Foundation, and the work of R.C. in part by the Department of Energy, Office of Basic Energy Sciences.

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