Rotational and vibrational spectra of quantum rings

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One can confine the two-dimensional electron gas in semiconductor heterostructures electrostatically or by etching techniques such that a small electron island is formed. These man-made “artificial atoms” provide the experimental realization of a text-book example of many-particle physics: a finite number of quantum particles in a trap. Much effort was spent on making such “quantum dots” smaller and going from the mesoscopic to the quantum regime \[1,2\]. Far-reaching analogies to the physics of atoms, nuclei or metal clusters were obvious from the very beginning: the concepts of shell structure and Hund’s rules were found \[1\] to apply – just as in real atoms! In this Letter, we report the discovery that electrons confined in ring-shaped quantum dots form rather rigid molecules with antiferromagnetic order in the ground state. This can be seen best from an analysis of the rotational and vibrational excitations.

While the independent-particle picture was successful in describing the electronic structure for rather large particle densities, for more dilute systems or in stronger magnetic fields correlation effects are of crucial importance. Configuration-interaction (CI) calculations, which have a long tradition in quantum chemistry and cluster physics, were then much used \[3\]. Although these so-called “exact” calculations are numerically demanding and limited to the smallest sizes, they still are able to provide significant insight into the many-body phenomena that occur in these finite fermion systems with reduced dimensionality. In this Letter we apply CI techniques to investigate the electronic structure of quantum rings that contain up to seven electrons. Usually the confinement of small, two-dimensional quantum dots is to a very good approximation harmonic. Correspondingly, we model quantum rings as they are realized in the laboratory by a potential of the form \( V(r) = \frac{1}{2} m^* \omega_0^2 (r - r_0)^2 \). For moderate confinement this potential corresponds to a harmonic dot with its center removed. Ground and excited states of \( N \) electrons trapped in the potential \( V(r) \) are determined from numerical diagonalization as a function of the total angular momentum. Surprisingly, at electron densities and strengths of the ring confinement where one should expect electron liquid behavior, a model which assumes localization of the electrons in the ring is successful in analyzing the many-body spectra. Group-theoretical methods familiar from molecular physics provide the necessary tools to uncover rotational and vibrational structures in the spectra. The spin sequence and energies of the low-lying states for given angular momentum can be understood from the symmetry associated with the electronic ground state configuration. It is intriguing that the success of the simple rigid-rotor model for the low-lying states is not limited to a regime where the system becomes very one-dimensional. The fact that the electrons behave as if they were localized in the ring is also reflected in a remarkable agreement of the CI results with the Hubbard or Heisenberg model. Localization at large electron densities has earlier been discussed in parabolic quantum dots, where the interpretation is not yet conclusive \[4\]. We write for the Hamiltonian

\[
H = \sum_{i=1}^{N} \left[ -\frac{\hbar^2}{2m^*} \nabla_i^2 + V(r_i) \right] + \sum_{i<j}^{N} \frac{\epsilon^2}{4\pi \varepsilon |\mathbf{r}_i - \mathbf{r}_j|},
\]

where \( m^* \) and \( \varepsilon \) are the effective mass and the dielectric constant of the corresponding semiconductor material. The parameters that determine the properties of the quantum ring are the number of electrons \( N \), the radius of the ring \( r_0 \) and the strength \( \omega_0 \) of the harmonic confinement in the radial direction. The quantities \( r_0 \) and \( \omega_0 \) are related to the more fundamental quantities \( r_s \), the one-dimensional density parameter which describes the particle density \( n = 1/(2\pi r_s) \) along the ring (thus \( r_0 = N r_s/\pi \)) and \( C_F \), a dimensionless parameter that measures the degree of one-dimensionality. \( C_F \) essentially describes the excitation energy of the next radial mode \( \hbar \omega_0 \), which is defined to be \( C_F \) times the (1D) Fermi energy. We thus obtain \( \hbar \omega_0 = C_F \hbar^2 \pi^2/(32 m^* r_s^2) \). The higher the value of \( C_F \), the more the radial modes are frozen in their ground states. Thus, the ring is narrower for larger \( C_F \). For the CI calculation, the spatial single-particle states of the Fock space are chosen to be eigenstates of the single-particle part of the Hamiltonian \( H \). We expand them in the harmonic oscillator basis. According to their eigenenergies, from 30 to about 50 lowest single-particle states are selected to span the Fock space. Typically this means that for lower angular momentum states several radial quantum numbers \( n = 0, 1, 2, 3 \) are included, whereas the higher angular momentum states \( l = \pm \) have only \( n = 0 \). To set up the Fock states for diagonalization, we sample over the full space with a fixed number of spin down and spin up electrons, \( N_\downarrow + N_\uparrow = N \). From this sampling, only those states with a given total orbital angular momentum and a configuration energy (corresponding to the sum of occupied single-particle energies) less than the specified cutoff energy \( e_C \) are selected. The purpose was to choose only the most important Fock states from the full basis and thereby reducing the matrix dimension to a size \( d \approx 2 \times 10^6 \). To obtain all the eigenstates with different total spin, we have to set \( N_\downarrow = N_\uparrow = N/2 \) for even particle numbers \( (S_z = 0) \), all states with different total spin have this component, and analogously \( N_\downarrow = N_\uparrow = 1 \) for odd numbers. Once the active Fock states have been specified, the Hamiltonian matrix is calculated. For diagonalization we use the Arpack library \[6\] suitable for large, sparse matrices. Finally, the total spin of each eigenvector is determined
by calculating the expectation value of the $\hat{S}^2$ operator. The many-particle states are characterized by the total orbital angular momentum $M$ and the total spin $S$. The lowest energy at given angular momentum $M$ defines the so-called “yrast” line, a terminology that was introduced in nuclear physics many years ago and comes from the Swedish word for “the most dizzy”.

To begin with, let us look at the energy spectra of a ring with six electrons at a density corresponding to $r_s = 2a_B$. (Throughout this paper, we use effective atomic units. Taking GaAs as an example, the units of length and energy are $a_B = 9.8\text{nm}$ and a.u.$^* = 12\text{meV}$.) For $C_F = 4$, the strength of the ring confinement is moderate, as it can be seen from the density distribution displayed in Fig. 1.

The system parameters $N$, $r_s$ and $C_F$ in this case are chosen such that the rings resemble quite closely structures as they can nowadays be made in the laboratory. For the six electron ring, Fig. 2 shows the 50 lowest states for all angular momenta from $M = 0$ up to $M = 6$. The spin configurations are given for the low-lying states. The ground state for $M = 0$ has spin $S = 0$ and is followed by a state with $S = 2$, then $S = 1$ and, close in energy, another $S = 0$ state. A large energy gap separates these low-lying states from higher bands with a much increased density of states. We note that the sequence of spins of the states below the gap for $M = 0$ is repeated at $M = 6$. (The energy difference between the two $S = 0$ states at $M = 6$ is slightly reduced, as in a not very narrow ring rotation expands the ring.) Inspecting the grouping of the states below the gap for each of the different $M$-values more closely, we see that in a similar way, the states for $M = 1$ follow the same sequence as those for $M = 5$, and the $M = 2$ spectrum is repeated at $M = 4$.

These sequences of states clearly reveal the spectrum of a dynamical system that can be described, at least approximately, in terms of independent rotation, vibration and intrinsic spin degrees of freedom. This becomes particularly clear when repeating the above calculation for a much lower particle density in a more narrow ring. We show in Fig. 3 the 6-particle spectra for $r_s = 6a_B$ and $C_F = 25$. (The particle density for these parameters is displayed in Fig. 1(b).) The spectra now indeed consist of narrow bands. The lowest band is the vibrational ground state and consist of the rotational levels. The different spin levels become almost degenerate, as localization has
reached a degree where spin-spin interactions become less important. Nevertheless, the sequence of spins and their relative energetic order is similar to the higher density results shown in Fig. (1), as a plot of the low-lying levels at an enlarged energy scale for $M = 0$ demonstrates. In the following we analyze the above spectra in terms of the standard effective hamiltonian employed in the interpretation of the rotational and vibrational spectra of planar polygonal molecules composed of $N$ identical spin $1/2$ fermionic atoms: the standard description is very slightly modified in order enforce the planarity and the one-dimensionality of the molecule considered here:

$$H_{\text{eff}} = AM^2 + \sum_a \hbar \omega_a n_a + J \sum_{i,j} S_i \cdot S_j$$  \hspace{1cm} (2)

where $A$ is the rigid moment of inertia of the $N$ "atoms" located at the $N$ vertices of an equilateral polygon, $\omega_a$ are the frequencies and $n_a (= 0, 1, 2, \ldots)$ the number of excitation quanta of the different normal modes of vibrations; for odd $N$ there are $\frac{1}{2}(N-1)$ two fold degenerate normal modes, while for even $N$ there are $\frac{1}{2}N-1$ twofold degenerate and one non-degenerate modes. The last term in Eq. (2) describes the nearest neighbor spin-spin interaction that is a result of the exchange term in the interaction of the neighboring fermions. In the expression (3) we have ignored the contribution of a possible spin orbit term.

For the $N = 6$ molecule discussed in Figs. 2 and 3, the $C_{6v}$, symmetry classification of the normal modes is (in order of increasing vibrational frequency) $E_1$ (2-fold degenerate), $E_2$ (2 fold degenerate) and $B_1$ (non-degenerate). The rotational states carry symmetry for $M = 0$ ($A_1$ or $A_2$), $M = 1$ or 5 ($E_1$), $M = 2$ or 4 ($E_2$), $M = 3$ ($B_1$ or $B_2$), where the congruent sign in all these terms refers to congruent (mod 6), and the total wave function must have $C_{6v}$ symmetry $B_2$ in order to fulfill the requirement of anti-symmetry between all the electrons.

As we learned from the above analysis, the lowest band is of rotational nature, while the bands above the energy gap represent vibration in addition to rotation. The three vibrational modes of such a system have energy ratios $\omega_1 : \omega_2 : \omega_3 = 1 : \sqrt{7} / 3 : \sqrt{3}$, in excellent agreement with the three lowest excitations for $M = 0$ shown in Fig. 3. The three highest states have then energies $2\omega_1$, $\omega_1 + \omega_2$ and $\omega_1 + \omega_3$.

A generic model for localized electrons is the Hubbard model which can be solved exactly for a small number of lattice points. The real electrons in the ring interact with the long-range repulsive Coulomb force. Classically the electrons would form a ring of equidistantly localized electrons. In the Hubbard model this case corresponds to the so-called half-filled model, where the number of electrons equals the number of lattice points. In the strong interaction limit it can be transformed to the anti-ferromagnetic Heisenberg model leading to the last term of Eq. (2).

On the basis of the results from exact diagonalizations and their group-theoretical analysis described above it is now interesting to see how well such simple models of localized electrons can describe the many-body spectra in these finite, not truly one-dimensional quantum rings. Fig. 5 compares the energy spectra of the Heisenberg model to the CI spectra described above. In the latter case the center-of-mass motion associated to the orbital angular momentum ($AM^2$-term) is subtracted from each level. The coefficient $A$ is determined to make the lowest energy eigenvalue corresponding to $M = N$ equal to that of $M = 0$. The resulting values of $A$ are very close to $A = \pi^2 / (2N^3 r^2)$ obtained for strictly localized electrons. The scale of the Heisenberg result is determined to get the same band width as in the case of real electrons. Fig. 5 displays for electron numbers $N = 5, 6$ and 7 at $r_s = 2a_B^*$ and $C_F = 10$ the spectra of the states below the first gap which (as analyzed above) are of rotational nature. For $N = 7$, corresponding to the larger particle number, there are now more states as we have more possibilities for the different spin configurations. The above comparison shows that indeed, the electrons in a ring can be rather accurately described as localized electrons: The whole rotational spectrum close to the yrast line can be determined by a spin model combined with a rigid center-of-mass rotation.

It should be noted that the results shown in Fig. 5 are calculated for a ring with a finite width. Making the ring narrower the small disagreement between the Heisenberg model and exact calculation vanishes. We finally mention that the pair correlation function does indicate anti-ferromagnetic coupling of the electrons in the

\begin{figure}
\centering
\includegraphics[width=\textwidth]{fig4.png}
\caption{Schematic view of the spin configurations in a six-electron ring for different values of $S$.}
\end{figure}
ground state. (See configuration schematically shown in the top left corner of Fig. 4). However, the correlation is not as clear as expected from the rotational structure in the energy spectrum. In the limit of an infinitely long one-dimensional Heisenberg model the spin-spin correlation decreases as $1/r_{ij}$. Consequently, even in the more narrow rings, where the rotational and vibrational spectra show a clear localization of electrons, the pair correlation function shows only rather weak spin-spin correlation.

![Figure 5](image.png)

**FIG. 5.** Low-lying (rotational) spectra of quantum rings with $N = 5, 6$ and 7 electrons: Comparison between the Heisenberg model (left columns in each of the spectra for given $M$) and exact diagonalization (right). The spins of the levels are marked by different colors

In conclusion, we reported rotational and vibrational many-body spectra of quantum rings confining up to seven electrons. We saw that the results obtained by configuration interaction calculations are best described by assuming localization in the internal structure of the ground-state many-body wave function. For even electron numbers, antiferromagnetic ordering was found for the ground state. Such so-called spin density waves in quantum rings actually were predicted by density functional theory \[9\]. The many-body spectra of the reported exact diagonalization calculations, their analysis by group-theoretical methods and further the comparison to the Heisenberg model confirm that indeed the simple mean-field picture (as it is provided by density functional theory) can to a rather large degree correctly map out the internal symmetry of the many-body wave function.

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