Discontinuities in the level density of small quantum dots under strong magnetic fields

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Exact diagonalization studies of the level density in a six-electron quantum dot under magnetic fields around 7 T (“filling factor” around $\frac{1}{2}$) are reported. In any spin-polarization channel, two regimes are visible in the dot excitation spectrum: one corresponding to interacting quasiparticles (i.e., composite fermions) for excitation energies below 0.4 meV, and a second one for energies above 0.4 meV, in which the level density exponentially increases at the same rate as in the noninteracting composite-fermion model.

The lowest-energy states of relatively small quantum dots (number of electrons $N_e \geq 3$) in strong magnetic fields have been qualitatively described within the composite-fermion picture. The position of cusps in the ground-state energy as a function of the angular momentum, and the spin quantum numbers of the low-lying excited states are nicely reproduced by this theory. The residual interaction between composite fermions (CF’s) is expected to be a weak, contact interaction.

The present paper is aimed at giving a quantitative characterization of the density of energy levels of small quantum dots in an energy interval below 1–1.5 meV, where dozens or hundreds of states exist. Fully converged exact diagonalization results for a six-electron GaAs dot in magnetic fields corresponding to “filling factors” $\nu \approx \frac{1}{2}$ are presented below. The error in computing energy eigenvalues is estimated to be lower than 0.02 meV for levels with excitation energies below 1 meV. The studied excitation energy range is small as compared to the cyclotronic ($\hbar \omega_c = 12$ meV), Coulomb (8.4 meV), or confinement ($\hbar \omega_0 = 3$ meV) energies of the model dot. Three Landau levels (LL’s) are included in the calculations, and a 25-meV cutoff in the energy of the noninteracting many-electron states used as basis functions allows us to deal with Hamiltonian matrices of dimension less than 850,000, which are diagonalized by means of a Lanczos algorithm. The main result of the paper is the existence of two regimes in the excitation spectra, corresponding to low $(\Delta E<0.4$ meV) and intermediate $(\Delta E>0.4$ meV) excitation energies, in which the level density increases at different rates. Thus, at an energy interval $\delta E$ around 0.4 meV the level density experiences a “discontinuity.”

The model parameters are similar to those used in Ref. 5. The confinement potential is parabolic. The bare Coulomb interaction is weakened by a factor 0.8 to approximately account for quasibidimensionality (instead of exact bidimensionality). The Zeeman energy is written in the form $0.014 \times 32 B s_z$ meV ($B$ in Tesla and $s_z = \pm \frac{1}{2}$), corresponding to a 8-nm-width well in magnetic fields around 7 T.

The lowest-energy states in each angular momentum and spin-polarization tower (the yrast spectrum) for magnetic fields $B=7$ and 8 T are shown in Fig. 1. Energy jumps between adjacent angular momentum states are about 0.6 meV in the spin-polarized case, but roughly three times smaller in any other spin-polarization sector. At this point it is important to stress the role of the higher LL’s in the energy eigenvalues. The absolute contribution of the second and third LL’s is around $\sim 0.4$ meV, a magnitude much greater than the Zeeman splitting (0.1 meV), and than the characteristic energy spacing between states near the absolute minimum. Excitation energies are pushed down 0.1–0.3 meV by the higher LL’s.

We show in Fig. 2 the number of states as a function of the excitation energy, $\Delta E$. In this figure, the reference energy

$$\text{FIG. 1. The lowest-energy levels in each angular momentum and spin-polarization sectors. Lines are guides to the eyes.}$$
The temperature parameter, $Q$, quantities correspond to the low-energy sectors. Notice that the excitation energy is written in the form $tation spectrum starts from a simplified first LL picture in the state, of the effective LL occupation numbers. By definition, then $B = 7$ T, $S_z = 3$

\[
\ln(n(\Delta E)) = n_0 \exp \frac{\Delta E}{\Theta},
\]

i.e., described by the so-called “constant-temperature approximation.” In the polarized case, there are only a few states for low excitation energies, thus we do not fit them. The fitting parameter $n_0$ and $\Theta$ are given in Table I. Primed quantities correspond to the low-energy sectors. Notice also the apparent jump of the temperature parameter at $\delta E$, which means a discontinuity in the level density.

The noninteracting CF (NICF) understanding of the excitation spectrum starts from a simplified first LL picture in which the excitation energy is written in the form $^2$

\[
\Delta E = \hbar \omega_{EF} \Delta n_{LL} + \hbar \omega_0^2/\omega_c \Delta L.
\]

$\Delta n_{LL}$ is the variation, with respect to the quasiparticle ground state, of the effective LL occupation numbers. By definition, $\Delta L = |L| - |L_{gs}|$, where $L_{gs}$ is the ground-state angular momentum. $\hbar \omega_c$ is the electron cyclotronic energy in GaAs, and the effective cyclotronic energy of CF’s is extracted from the polarized yrast spectrum, $S_z = 3$, as $^2$

\[
\hbar \omega_{CF} = E(L_3 + 1) + E(L_3 - 1) - 2E(L_3),
\]

where $L_3$ is the angular momentum of the lowest polarized state. It gives $\hbar \omega_{CF} = 1.15$ and $1.19$ meV at $B = 7$ and 8 T, respectively. The quasiparticles are supposed to occupy states in effective Landau levels, which are separated by $\hbar \omega_{CF}$. The angular momentum of the quasiparticle system is $L_{CF}$ which, due to the special form of the variational CF wave function, is related to $L$ by $^2$

\[
L = -N_e(N_e - 1) + L_{CF}.
\]

The NICF model gives qualitatively correct answers to questions such as the position of cusps in the yrast spectrum, the nature of the first excited states, etc. One may ask for other visible manifestations of the NICF in the spectrum, and indeed, it gives the correct $\Theta$ parameter at intermediate excitation energies, $\Delta E > \delta E$. In other words, the NICF curve grows exponentially with the same $\Theta$, although it is shifted with respect to the actual spectrum. This fact is illustrated in Fig. 2 also, where the NICF curve at a certain level (the sixth in the upper figure, for example) is forced to meet the actual curve. From this point on the two spectra have the same slope on average.

There is a natural interpretation of this behavior. In the ground and first excited states (low $\Delta E$), the quasiparticles form compact clusters $^2$ and the interaction between CF’s plays an important role. The low-energy parameter $\Theta'$ is a

<table>
<thead>
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<th>$S_z$</th>
<th>$B = 7$ T</th>
<th>$B = 8$ T</th>
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</thead>
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<td>3</td>
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<tr>
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<td>$n_0$</td>
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<tr>
<td></td>
<td>$\Theta$</td>
<td>0.423</td>
</tr>
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</table>

$\Delta E = |L| - |L_{gs}|$, where $L_{gs}$ is the ground-state angular momentum. $\hbar \omega_c$ is the electron cyclotronic energy in GaAs, and the effective cyclotronic energy of CF’s is extracted from the polarized yrast spectrum, $S_z = 3$, as $^2$

$\hbar \omega_{CF} = E(L_3 + 1) + E(L_3 - 1) - 2E(L_3)$.

$\Delta L = |L| - |L_{gs}|$, where $L_{gs}$ is the ground-state angular momentum. $\hbar \omega_c$ is the electron cyclotronic energy in GaAs, and the effective cyclotronic energy of CF’s is extracted from the polarized yrast spectrum, $S_z = 3$, as $^2$

$\hbar \omega_{CF} = E(L_3 + 1) + E(L_3 - 1) - 2E(L_3)$.

$\hbar \omega_{CF} = E(L_3 + 1) + E(L_3 - 1) - 2E(L_3)$.
consequence of this interaction. On the other hand, at intermediate energies and assuming that the interaction is weak, one expects the level density to be dominated by the relevant energy scales, i.e., $\hbar \omega_{CF}$ and $\hbar \omega_0^2/\omega_c$. One may visualize this regime in terms of Rydberg-like excitations, in which a few noninteracting CF’s orbit around a core of weakly interacting CF’s. Energy differences will, of course, follow the NICF rules.

The conclusion to be extracted from this figure is that traces of the effective LL structure of CF’s may also be looked for at intermediate excitation energies, $D E \approx 0.4$ meV. For $D E > 0.4$ meV the transition to a regime of weakly interacting quasiparticles takes place. Temperature parameters are different from both sides of $\delta E$, thus a discontinuity in the level density is expected.

The level density may be directly measured in Raman-scattering experiments under extreme resonance, where the Raman amplitude depends on the density of energy levels in final states. Magnetoconductance measurements under equilibrium or nonequilibrium conditions could also give evidence about the level-density behavior at intermediate excitation energies.

Finally, we show in Fig. 3 the magnitudes $\hbar \omega_{CF}$ and $\Theta_{S'=3}$ as a function of magnetic-field intensity $B$. Part of this work was carried out at the Abdus Salam ICTP. A. G. acknowledges the ICTP Associate and Federation Schemes for support. R. C. acknowledges support from the Ministerio de Educación, Deportes y Cultura de España, Secretaría de Estado de Educación y Universidades.

FIG. 3. The magnitudes $\hbar \omega_{CF}$ (squares) and $\Theta_{S'=3}$ (circles) as a function of magnetic-field intensity $B$.