The Schrödinger equation for quantum dot lattices with non-cubic, non-Bravais lattices built up from elliptical dots is investigated. The Coulomb interaction between the dots is considered in dipole approximation. Then only the center of mass (c.m.) coordinates of different dots couple with each other. This c.m. subsystem can be solved exactly and provides magneto-phonon like collective excitations. The inter-dot interaction is involved only through a single interaction parameter. The relative coordinates of individual dots form decoupled subsystems giving rise to intra-dot excitation modes. As an example, the latter are calculated exactly for two-electron dots.

Emphasis is layed on qualitative effects like: i) Influence of the magnetic field on the lattice instability due to inter-dot interaction, ii) Closing of the gap between the lower and the upper c.m. mode at B=0 for elliptical dots due to dot interaction, and iii) Kinks in the intra dot excitation energies (versus magnetic field) due to change of ground state angular momentum. It is shown that for obtaining striking qualitative effects one should go beyond simple cubic lattices with circular dots. In particular, for observing effects of electron–electron interaction between the dots in FIR spectra (breaking Kohn’s Theorem) one has to consider dot lattices with at least two dot species with different confinement tensors.

PACS: 73.20.D (Quantum dots), 73.20.Mf (Collective Excitations)

I. INTRODUCTION

Quantum dots have been in the focus of intensive research already for at least a decade which lead to a countless number of publications (for a recent book see Ref. 1). Although almost all experiments are performed at dot lattices, in the vast majority of theoretical investigations the interaction between dots is neglected. This is for the following reasons: i) Because the confinement frequency $\omega_0$ is a parameter, which is mainly extracted from optical properties, it is difficult to tell the influence of dot interaction apart from the intrinsic single–dot value. (Possibilities to overcome this problem are discussed in the present work.) ii) The theory of Raman spectra, which can in principle monitor the dispersion (wave number dependence) of excitation energies as a direct consequence of interdot–interaction, is not yet advanced enough to extract the dispersion. iii) The lattice constant of dot arrays produced with current technologies is so large ($> 2000$Å) that large electron numbers $N$ per dot are necessary to obtain a seizable amount of shift. For these $N$, however, reliable first principle calculations are not possible. With the advent of self–assembled dot arrays the last item might change.

The scope of this paper is to investigate conditions, which lead to qualitative and observable effects of interdot–interaction on excitation spectra and the phase transition found in Ref. 2. Unlike in Ref. 2, a magnetic field $B$ is explicitly taken into account and a microscopic theory is applied. Our approach is purely microscopic, i.e. we solve the Schrödinger equation of a model system exactly. Our model comprises the following approximations: i) The dot confinement is strictly parabolic in radial direction, but with anisotropic confinement frequencies $\omega_i$ ($i = 1, 2$) and independent of $N$ and $B$. ii) Overlap of wave functions between different dots is neglected (no hopping). iii) The Coulomb interaction of the electrons in different dots is treated in dipole approximation (second order in dot diameter over lattice constant). Our model is similar to that in Ref. 3, but allows more complicated dots and lattice structures. Besides we calculate also the intra dot excitations (apart from the collective center–of–mass excitations) for $N = 2$ explicitly and discuss the instability in this microscopic model. Our results on the lateral dot dimer are compared with a former paper, which uses a high magnetic field approach, in Sect. III.

The plan of this paper is as follows. For further reference, we briefly summarize in Sect. I some relevant results for one single dot, or for dot lattices, where the distance between the dots is very large. This is important, because all

\[1\text{Therefore we will refer here only to papers which are directly connected to the scope of this work} \]
exact solutions in the center–of–mass subsystem are traced back (by special transformations) to the solution of this one–electron Hamiltonian. This is analogous to ordinary molecular and lattice dynamics. After this, we consider a dot dimer, which mimics a lattice, where the dots are pairwise close to each other. This model can give an idea of the effects expected in dot lattices with a basis. Next we consider a rectangular, but primitive lattice in order to obtain the dispersion in the spectra. Finally, the intra-dot excitations of the Hamiltonian in the relative coordinates are calculated numerically. The paper ends with a summary. In the Appendix we give a short and elementary proof for the fact that the Generalized Kohn Theorem holds even for arbitrary arrays of identical non-circular quantum dots with Coulomb interaction (between the dots) in an homogeneous magnetic field.

II. SINGLE DOT

The Hamiltonian considered here reads (in atomic units $\hbar = m = e = 1$)

$$H = \sum_{i=1}^{N} \left\{ \frac{1}{2m^*} \left[ p_i + \frac{1}{c} A(r_i) \right]^2 + \frac{1}{2} r_i \cdot C \cdot r_i \right\} + \frac{1}{2} \sum_{i \neq k}^{N} \frac{\beta}{|r_i - r_k|}$$  

(1)

where $m^*$ is the effective mass (in units of the bare electron mass $m$), $\beta$ the inverse dielectric constant of the background, and $C$ a symmetric tensor. In case of a single dot, $C$ is given by the confinement potential and we define $C = \Omega$. It is always possible to find a coordinate system where $\Omega_{12} = \Omega_{21} = 0$ and $\Omega_{ii} = \omega_{c}^2 = m^* \omega_{c}^2$. We use the symmetric gauge $A = \frac{1}{2} B \times r$ throughout. The Zeeman term in $H$ is disregarded at the moment. For $N = 1$, the Hamiltonian

$$H = \frac{1}{2m^*} \left[ p + \frac{1}{c} A(r) \right]^2 + \frac{1}{2} r \cdot C \cdot r$$  

(2)

can be diagonalized exactly. Later on we will see that also the case of interacting dots can be traced back to the solution of type (2). (Therefore, we kept the off diagonal elements of $C$ in the results given below because the dynamical matrix, which also contributes to $C$, is generally non–diagonal and we want to use the same coordinate system for all $q$ values.) After transforming the operators $r_i$ and $p_i$ to creation– annihilation operators (see e.g. Ref. [1] and using the procedure described by Tsallis [1]), we obtain for the eigenvalues

$$E(n_+, n_-) = (n_+ + \frac{1}{2}) \omega_+ + (n_- + \frac{1}{2}) \omega_- ; \quad n_{\pm} = 0, 1, 2, ...$$  

(3)

where

$$\omega_{\pm} = \sqrt{\frac{\omega_{c}^2}{2} + \tilde{\omega}_{0}^2 \pm \sqrt{\frac{\omega_{c}^2}{4} + \omega_{c}^2 \tilde{\omega}_{0}^2 + \frac{\Delta^2}{4} + C_{12}^2}}$$  

(4)

and

$$\tilde{\omega}_{0}^2 = \frac{1}{2}(C_{11} + C_{22}) ; \quad \Delta = C_{11} - C_{22}$$  

(5)

and $\omega_{c} = \frac{B}{m^* c}$ is the cyclotron frequency with the effective mass. (The results for the special case $C_{12} = 0$ can also be found in Ref. [1]). The optical selection rules are the same as in the circular case, i.e., there are two possible types of excitations

$$(\Delta n_+ = \pm 1 \quad \text{and} \quad \Delta n_- = 0) \quad \text{or} \quad (\Delta n_- = \pm 1 \quad \text{and} \quad \Delta n_+ = 0)$$  

(7)

leading to the excitation energies $\Delta E = \omega_+ + \omega_-$. It is easily seen that the form (3) reduces to the familiar formula in the circular case, where $\Delta = 0$ and $C_{12} = 0$. By inspection of (4) we find that a soft mode $\omega_-(B) = 0$ can only occur if $C_{11} \cdot C_{22} = C_{12}^2$. For a diagonal $C$ this means that $\min(C_{11}, C_{22}) = 0$. The last condition is of importance for interacting dots considered in the next Sections.

In the limiting case $B = 0$ we obtain from (4)

$$E(0, n_-) = (n_- + \frac{1}{2}) \omega_+$$  

$$E(n_+, 0) = (n_+ + \frac{1}{2}) \omega_-$$  

$$E(n_+, n_-) = (n_+ + \frac{1}{2}) \omega_+ + (n_- + \frac{1}{2}) \omega_-$$  

for interacting dots considered in the next Sections.
\[ \omega_{\pm}(B = 0) = \sqrt{\frac{(C_{11} + C_{22})}{2}} \pm \sqrt{\frac{(C_{11} - C_{22})^2}{4} + C_{12}^2} \]  

We see that degeneracy \( \omega_{+}(B = 0) = \omega_{-}(B = 0) \) can only happen if \( C_{12} = 0 \) and \( C_{11} = C_{22} \). For a diagonal confinement tensor with \( C_{12} = 0 \) we obtain \( \omega_{+}(B = 0) = \max(\omega_1, \omega_2) \) and \( \omega_{-}(B = 0) = \min(\omega_1, \omega_2) \). As to be expected, we observe a gap between the two excitation curves \( \omega_{+}(B) \) and \( \omega_{-}(B) \) at \( B = 0 \), if the two confinement frequencies do not agree.

Alternatively we can introduce the quantum numbers

\[ k = \frac{(n_+ + n_-) - |n_+ + n_-|}{2}; \quad m_z = n_+ - n_- \]  

where \( k \) is the node number and \( m_z \) turns in the circular limit into the angular momentum quantum number.

For arbitrary \( N \), the center of mass (c.m.) \( R = \frac{1}{N} \sum_i r_i \) can be separated \( H = H_{c.m.} + H_{rel.} \) with

\[ H_{c.m.} = \frac{1}{N} \left\{ \frac{1}{2m^*} \left[ P + \frac{N}{c} A(R) \right]^2 + \frac{N^2}{2} R \cdot C \cdot R \right\} \]  

where \( P = -i \nabla_R \) (see Appendix). As well known, \( H_{c.m.} \) does not contain the electron–electron–interaction. \( H_{c.m.} \) can be obtained from the one–electron Hamiltonian (2) by the substitution: \( B \rightarrow NB, \ C \rightarrow N^2C \) and \( H \rightarrow \frac{1}{N} H \). If we make the same substitution in the eigenvalues (3), we obtain

\[ E_{c.m.}(n_+, n_-) = E_{N=1}(n_+, n_-) \]  
i.e., the eigenvalues of the c.m. Hamiltonian are independent of \( N \). In other words, in \( H \) there are excitations, in which the pair correlation function is not changed, or classically speaking, where the charge distribution oscillates rigidly. Because FIR radiation (in the limit \( \lambda \rightarrow \infty \)) can excite only the c.m. subspace, all we see in FIR spectra is the c.m. modes.

### III. DOT DIMER

We consider two identical elliptical dots centered at \( a_1 = (-a/2, 0) \) and \( a_2 = (+a/2, 0) \). We expand the Coulomb interaction between electrons in different dots in a multi-pole series and restrict ourselves to the dipole approximation. By introduction of c.m. and relative coordinates within each dot, the c.m. coordinates and the relative coordinates decouple:

\[ \tilde{H} = \tilde{H}_{c.m.}(R_1, R_2) + \sum_{\alpha} \tilde{H}_{\alpha} \{r\}^{(N-1)}_{\alpha} \]  

\( \{r\}^{(N-1)}_{\alpha} \) symbolizes \((N-1)\) relative coordinates in the \( \alpha^{th} \) dot. This means, we have 3 decoupled Hamiltonians: the c.m. Hamiltonian and two Hamiltonians in the relative coordinates of either dot. This leads to two types of excitations:

i) **Collective excitations** from \( \tilde{H}_{c.m.} \) which involve the c.m. coordinates of both dots simultaneously. Because of the harmonic form (in the dipole approximation), there are exactly two modes per dot, thus a total of four. Each excitation can be classically visualized as vibrations of rigidly moving charge distributions of both dots.

ii) **Intra dot excitations** which are doubly degenerate for two identical dots. Because \( \tilde{H}_{\alpha}(\{r\}^{(N-1)}_{\alpha}) \) is not harmonic (it includes the exact Coulomb interaction between the electrons within each dot, which is not harmonic), this spectrum is very complex. It is the excitation spectrum of a single dot in a modified confinement potential where the c.m. coordinate is fixed. The extra term in the modified confinement potential comes from the dipole contribution of the interdot Coulomb interaction.

In this Section we consider only the c.m. Hamiltonian and focus our attention to the the effects of ellipticity in the

---

2 The tilde indicates that in this preliminary Hamiltonian a common gauge center for both dots is used.
dot confinement potential. The relative Hamiltonian for \( N = 2 \) is explicitly given in the last Section and solved for circular dots. For the elliptical confinement potential considered in this Section, the relative Hamiltonian cannot be solved easily, even if we restrict ourselves to \( N = 2 \), because the elliptic confinement potential breaks the circular symmetry of the rest of the relative Hamiltonian.

A. Center of Mass Hamiltonian of the Dimer

The c.m. Hamiltonian in the dipole approximation reads

\[
\tilde{H}_{\text{c.m.}} = \frac{1}{N} \left\{ \sum_{\alpha}^{1,2} \frac{1}{2m^*} \left[ \mathbf{p}_\alpha + \frac{N}{c} \mathbf{A} (\mathbf{U}_\alpha + \mathbf{a}_\alpha) \right]^2 + \frac{N^2}{2} \sum_{\alpha,\alpha'} \mathbf{U}_\alpha \cdot \mathbf{C}_{\alpha,\alpha'} \cdot \mathbf{U}_{\alpha'} \right\}
\]

where the small elongation \( \mathbf{U}_\alpha \) is defined by \( \mathbf{R}_\alpha = \mathbf{a}_\alpha + \mathbf{U}_\alpha \) and \( \mathbf{P} = -i \nabla = -i \nabla \). The tensor \( \mathbf{C} \) is

\[
\mathbf{C}_{\alpha,\alpha'} = \Omega + \beta N \sum_{\alpha''(\neq \alpha)} \mathbf{T}(\mathbf{a}_{\alpha,\alpha''})
\]

for \( \alpha \neq \alpha' \)

\[
\mathbf{T}(\mathbf{a}) = \frac{1}{a^3} \left[ 3 \mathbf{a} \circ \mathbf{a} - a^2 \mathbf{I} \right]
\]

containing a dyad product (\( \circ \)) and the unit tensor \( \mathbf{I} \). As in the c.m. system of a single dot, the explicit \( N \)-dependence in (12) cancels in the eigenvalues. What is left is only the \( N \)-dependence in the dipole contribution of the dot interaction appearing in (13) and (14). This means, that the c.m. spectrum of interacting dots is no longer independent of \( N \).

The term \( \mathbf{a}_\alpha \) in the argument of the vector potential in (12) causes trouble in finding the eigenvalues. This shift is a consequence of the fact that we have to adopt a common gauge center for both dots (we chose the middle between both dots). This problem can be solved by applying the following unitary transformation

\[
H_{\text{c.m.}} = Q^{-1} \tilde{H}_{\text{c.m.}} Q; \quad Q = \prod_{\alpha}^{1,2} e^{-i \frac{\mathbf{a} \times (\mathbf{B} \times \mathbf{a}_\alpha)}{a^2}} \mathbf{U}_\alpha
\]

In other words, \( H_{\text{c.m.}} \) agrees with \( \tilde{H}_{\text{c.m.}} \) except for the missing shift in the argument of the vector potential.

The 4 modes inherent in \( H_{\text{c.m.}} \) are not yet explicitly known, because the 4 degrees of freedom are coupled. Decoupling into two oscillator problems of type (2) can be achieved by the following transformation:

\[
\mathbf{U}^{(+)} = \frac{1}{2} (\mathbf{U}_2 + \mathbf{U}_1) ; \quad \mathbf{U}^{(-)} = \mathbf{U}_2 - \mathbf{U}_1
\]

This results in

\[
H_{\text{c.m.}} = \frac{1}{2} \tilde{H}^{(+)} + 2 \tilde{H}^{(-)}
\]

where

\[
\tilde{H}^{(+)} = \frac{1}{N} \left\{ \frac{1}{2m^*} \left[ \mathbf{P}^{(+)} + \frac{2N}{c} \mathbf{A} \left( \mathbf{U}^{(+)} \right) \right]^2 + \frac{N^2}{2} \mathbf{U}^{(+)} \cdot \left( 4\mathbf{\Omega} \right) \cdot \mathbf{U}^{(+)} \right\}
\]

\[
\tilde{H}^{(-)} = \frac{1}{N} \left\{ \frac{1}{2m^*} \left[ \mathbf{P}^{(-)} + \frac{N}{2c} \mathbf{A} \left( \mathbf{U}^{(-)} \right) \right]^2 + \frac{N^2}{2} \mathbf{U}^{(-)} \cdot \left( \frac{1}{4} \mathbf{\Omega} + \frac{N}{2} \beta \mathbf{T}(\mathbf{a}) \right) \cdot \mathbf{U}^{(-)} \right\}
\]

and \( \mathbf{a} \) is a vector pointing from one dot center to the other. Then \( \mathbf{T}(\mathbf{a}) \) has the following components

\[
T_{11} = \frac{2}{a^3} ; \quad T_{22} = -\frac{1}{a^3} ; \quad T_{12} = T_{21} = 0
\]
Now we assume that the principle axes of the confinement potentials are in x-y-direction. This means
\[ \Omega_{11} = \omega_1^2 ; \quad \Omega_{22} = \omega_2^2 ; \quad \Omega_{12} = \Omega_{21} = 0 \] (22)
The eigenvalues of \( H^{(+)} \) can be obtained from (3) and (4) with
\[ \omega_0^2 = \frac{1}{2} (\omega_1^2 + \omega_2^2) ; \quad \Delta = (\omega_1^2 - \omega_2^2) \] (23) and for \( H^{(-)} \) with
\[ \omega_0^2 = \frac{1}{2} (\omega_1^2 + \omega_2^2) + \frac{1}{2} p ; \quad \Delta = (\omega_1^2 - \omega_2^2) + 3 p \] (24)
where the interaction parameter is defined by
\[ p = \frac{2N\beta}{a^3} \] (25)
Observe that the dependence on \( N \) cancels, except that included in \( p \) (see discussion following (11)).
It is important that the dot interaction influences the result only through a single parameter. This conclusion agrees with the semi–phenomenological theory in Ref. 3.

In all our figures we express frequencies in units of the average confinement frequency \( \omega_0 = \frac{1}{2} (\omega_1 + \omega_2) \), and \( \Delta \) and \( p \) in units \( \omega_0^2 \). Then, all systems can be characterized by the two parameters: \( \omega_1/\omega_2 \) and \( p \). In other words, all systems having the \( \omega_1/\omega_2 \) ratio indicated in the figures are represented by the family of curves with the \( p \) values shown. The only exception we made is the cyclotron frequency \( \omega_c^\ast \). \( \omega_c^\ast/\omega_0 \) would be a good parameter in this sense, but we chose to use the magnetic field in Tesla instead for better physical intuition. The conversion between both scales is given by \( \omega_c^\ast [\text{a.u.}^\ast] = \frac{0.9134\cdot10^{-2}}{m^\ast} B[\text{Tesla}] \) or, \( \omega_c^\ast [\omega_0] = \frac{0.9134\cdot10^{-2}}{m^\ast/\omega_0[\text{a.u.}^\ast]} B[\text{Tesla}] \) In this paper we used \( \omega_0 = 0.2 \text{ a.u.}^\ast = 2.53 \text{ meV} \) and \( m^\ast \) of GaAs. (We want to stress that this choice effects only the magnetic field scale and not the qualitative features of the figures.) For easy comparison with experimental parameters we add the definition of effective atomic units (a.u.^\ast) in GaAs (\( m^\ast = 0.067, \beta = 1/12 \)) for the energy: \( 1 \text{ a.u.}^\ast = 4.65 \cdot 10^{-4} \text{ double Rydberg} = 12.64 \text{ meV} \), and for the length: \( 1 \text{ a.u.}^\ast = 1.791 \cdot 10^2 \text{ Bohr radii} = 0.9477 \cdot 10^2 \text{ Å} \).

Because \( U^{(+)} \) agrees with the total c.m. \( \mathbf{R} = \frac{1}{2}(\mathbf{R}_1 + \mathbf{R}_2) \) of the system, \( H^{(+)} \) is the total c.m. Hamiltonian. For \( B = 0 \), the eigenmodes can be visualized by classical oscillations. The two eigenmodes of \( H^{(+)} \) are (rigid) in–phase oscillations of the dots in x and y direction, respectively. Because of the Kohn theorem (see Appendix), the independence of \( H^{(+)} \) on the Coulomb interaction does not only hold in the dipole approximation, but it is rigorous. This shows also that the dipole approximation is consistent with the Kohn theorem, which is not guaranteed for single particle approaches. Because FIR radiation excites (in the dipole approximation) only the c.m. modes, it is only the \( p \)-independent eigenmodes of \( H^{(+)} \) which are seen in FIR absorption experiments.
This statement is in contradiction to Ref. 3. They performed numerical diagonalizations for a lateral pair of circular dots confining the set of basis functions to the lowest Landau level and considering parallel spin configurations only. This is justified in the limit of high magnetic fields. They found a splitting of the two dipole allowed modes at \( B = 0 \) due to dot interaction and some anti-crossing structures in the upper mode, whereas the lower mode is always close to the single particle mode. This fact is already a strong indication that the missing higher Landau levels cause both spurious effects. (Observe that the lifting of the degeneracy at \( B = 0 \) in the dipole allowed excitations in Fig.1 is due to the ellipticity of the intrinsic confinement and not due to dot interaction.)
The eigenvalues of \( H^{(-)} \) do depend on \( p \) because the dots oscillate (rigidly) in its two eigenmodes in opposite phase, one mode in x and one mode in y direction. This leads to a change in the Coulomb energy. The two eigenmodes of \( H^{(-)} \) can also be described as a breathing mode (in x direction) and a shear mode (in y direction).
FIG. 1. Excitation energies $\Delta E^{(\pm)} = \omega^{(\pm)}$ for a dimer of elliptical dots as a function of the magnetic field for some discrete values of the interaction parameter $p$ ($p$ values in the inset are in units $\omega_0^2$). The ratio of the oscillator frequencies in the direction of the capital axes $\omega_1/\omega_2$ is a) $3/2$ and b) $2/3$. The dipole allowed excitations $\Delta E^{(+)}$ of $H^{(+)}$ (thick full line) are not influenced by the dot interaction and therefore independent of $p$.

B. Special Features of the Excitation Spectrum

In Fig.1a and 1b, the four excitation frequencies of the dimer are shown with $p$ as a parameter.
For \( p = 0 \), the two modes \( \omega_\pm^{(-)} \) agree with the two modes \( \omega_\pm^{(+)} \). In all symbols, the superscript sign refers to the system \( H^{(+)} \) and \( H^{(-)} \) (c.m. or relative coordinate), and the subscript sign discriminates the two modes of the same system. The two modes \( \omega_\pm^{(\pm)} \) are independent of \( p \). There are two qualitatively different cases. (Consider that \( \omega_1 \) is the oscillator frequency parallel to the line, which connects the two dot centers, and \( \omega_2 \) is the oscillator frequency perpendicular to this line.) If \( \omega_1 \geq \omega_2 \) (Fig.1a), the gap between \( \omega_\pm^{(-)} \) and \( \omega_\pm^{(-)} \) at \( B = 0 \) increases steadily with increasing \( p \) until, for a critical \( p_{cr} = \omega_2^2 \) (in our numerical case: \( p_{cr}[\omega_0^2] = 16/25 = 0.64 \)) the lower mode \( \omega_-^{(-)} \) becomes soft. This transition is independent of \( B \). For \( \omega_1 \leq \omega_2 \) the gap between \( \omega_\pm^{(-)} \) and \( \omega_\pm^{(-)} \) at \( B = 0 \) first decreases with increasing \( p \) until it vanishes for \( p = \frac{1}{4}(\omega_2^2 - \omega_1^2) \) (in our numerical case: \( p[\omega_0^2] = 4/15 = 0.27 \)). Afterwards, it increases until the lattice becomes soft at \( p_{cr} = \omega_2^2 \) (in our numerical case: \( p_{cr}[\omega_0^2] = 36/25 = 1.44 \)). The dependence of the two excitation energies \( \omega_\pm^{(-)} \) and \( \omega_\pm^{(-)} \) on \( p \) for \( B = 0 \) in the second case is shown in Fig.2. Comparison of Fig.s 2a and 2b demonstrates that the dot architecture in Fig.2a is much more sensitive to interdot interaction than that in Fig.2b. Thus, if we want to observe or use the instability, this event happens in case 2a for for smaller \( p \) (or larger lattice constants) than in case 2b. Additionally, the assumption of non–overlapping dot wave functions (for a given lattice constant) is better fulfilled in case 2a than in case 2b.

![Dot dimer, \( \omega_1/\omega_2=2/3 \), \( B=0 \)](image)

**FIG. 2.** Excitation energies \( \Delta E_\pm^{(-)} = \omega_\pm^{(-)} \) of the Hamiltonian \( H^{(-)} \) for \( B = 0 \) and \( \omega_1/\omega_2 = 2/3 \) as a function of the interaction parameter \( p \).

For GaAs as a typical substance, \( (\omega_0^2) \) can be rewritten in more convenient units as \[ p[\omega_0^2] = \frac{2.26 \cdot 10^7 N}{(a[A])^3 (\omega_0[meV])^2} \] (26)

Obviously, we need large dots (large \( N \), small \( \omega_0 \)– which means large polarizability), and a small dot distance \( a \) for a seizable interaction effect. On the other hand, the dot radius for \( N = 1 \) is of order of the effective magnetic length \( l_0 = \left( (2\omega_0)^2 + (\omega_c^*)^2 \right)^{-1/4} \), which reads for GaAs

\[ l_0[A] = \frac{238}{\left( (\omega_0[meV])^2 + 0.739 (B[\text{Tesla}])^2 \right)^{1/4}} \] (27)

7
and we need small dots and high magnetic fields for small overlap. Consequently, a magnetic field helps avoiding overlap of the dots, although e.g. the critical $p$ for soft modes is independent of $B$. The question is, if there exists a window between these two (partly) conflicting demands. For an order–of–magnitude estimate, let us consider GaAs with $\omega_0$ as chosen above and the worst case $N = 1$. Then (22) with a typical $p[\omega_0^2] = 0.1$ (which seems to be the minimum for any observable effect) provides a dot distance of $a[A] = 327$ and (23) gives for $B = 0$ a radius of $l_0[A] = 150$ and for $B[\text{Tesla}] = 10$ a radius of $l_0[A] = 80$. Consequently, the constraint $l_0 < a/2$ for our model can be fulfilled. For obtaining larger interaction effects the parameters have to be optimized.

The next question is what happens in mode softening physically? Firstly, it is the antisymmetric shear mode $\omega^{(-)}$ which has the lowest frequency and which becomes soft. If the interaction parameter is strong enough ($p > p_{\text{cr}}$), the decrease in interdot–Coulomb energy with increasing elongation of the dots becomes larger than the increase of confinement potential energy. Because in the harmonic model both energies depend quadratically on elongation, the dimer would be ionized, i.e. stripped of the electrons. Clearly, in this case we have to go beyond the dipole approximation for the interdot interaction and beyond the harmonic approximation for the confinement potential.

In order to obtain a hand-waving picture of what happens, the confinement potential of the system for shear mode oscillations is supplemented by a 4th order term in the following way: $V_{\text{conf.}} = 2N \left[\frac{1}{2} \omega_2^2 U^2 - A U^4\right]$ with $(A > 0)$, and the Coulomb interaction in 4th order reads: $V_{\text{int}} = -pN U^2 + (3pN/a^2) U^4$ where $p = 2N\beta/a^3$ as above. Then, the stability condition reads

$$\frac{V_{\text{tot}}}{N} = (\omega_2^2 - p) \frac{U^2}{2} + \frac{3p}{a^2} - 2A \frac{U^4}{2} \geq 0$$

(28)

The condition for the existence of a bound state is that the $U^4$ term is positive: $3p/a^2 > 2A$. For a positive $U^2$-term ($p < \omega_2^2$), the equilibrium position is $U_0 = 0$. If the $U^2$-term becomes negative ($p > \omega_2^2$), the system finds a new equilibrium at a finite elongation

$$U_0 = \pm \sqrt{(p - \omega_2^2)/(2(3p/a^2 - 2A))}$$

(29)

This new ground state is doubly degenerate: $U_1 = (\pm a, U_0)$, $U_2 = (\mp a, -U_0)$ and $U_{1} = (\mp a, -U_0)$, $U_2 = (+a, +U_0)$ have the same energy. In short, at $p_{\text{cr}} = \omega_2^2$ there is an electronic phase transition to a polarised state, where the equilibrium position of the c.m. is no more in the middle of the dots. At the end we want to stress that all these stability considerations are only valid if the confinement potential is not changed under elongation of the c.m. of the dots. Secondly, it is not rigorous to include the fourth order terms after separation of c.m. and relative coordinates, because in fourth order these two coordinates do not decouple exactly.

IV. DOT LATTICE

We consider a periodic lattice of equal quantum dots at lattice sites $R^{(0)}_{n,\alpha} = R^{(0)}_{\alpha} + a_\alpha$. The vectors $R^{(0)}_{\alpha}$ form a Bravais lattice and $a_\alpha$ runs over all sites within an unit cell. In developing a theory for these lattices we have to repeat all steps in Sect. II from [11] to [14] just by supplementing the index $\alpha$ by the index $n$ for the unit cell.

A. Center of Mass Hamiltonian of the Dot Lattice

The c.m. Hamiltonian in the dipole approximation then reads

$$H_{\text{c.m.}} = \frac{1}{N} \left\{ \sum_{n,\alpha} \frac{1}{2m^*} \left[ P_{n,\alpha} + \frac{N}{c} A (U_{n,\alpha}) \right]^2 + \frac{N^2}{2} \sum_{n,\alpha} U_{n,\alpha} \cdot C_{n,\alpha; n',\alpha'} \cdot U_{n',\alpha'} \right\}$$

(30)

where $U_{n,\alpha} = R_{n,\alpha} - R^{(0)}_{n,\alpha}$ is the elongation of the c.m. at lattice site $(n, \alpha)$ and the force constant tensor $C$ is defined in analogy to (13) and (14).

The Hamiltonian (30) is a phonon Hamiltonian in an additional homogeneous magnetic field. The first stage of decoupling can be achieved by the usual phonon transformation.
\[ U_{n,\alpha} = \frac{1}{\sqrt{N_c}} \sum_{q}^{BZ} e^{-i q \cdot R_{n}^{(0)}} U_{q,\alpha} \]  
\[ P_{n,\alpha} = \frac{1}{\sqrt{N_c}} \sum_{q}^{BZ} e^{i q \cdot R_{n}^{(0)}} P_{q,\alpha} \]

where \( N_c \) is the number of unit cells and the transformed coordinates have the following properties \( U_{-q,\alpha} = U_{q,\alpha}^* = U_{q,\alpha}^\dagger \) and \( P_{-q,\alpha} = P_{q,\alpha}^\dagger \). The Hamiltonian in the new coordinates is a sum of \( N_c \) subsystems of dimension \( 2 \times \) number of dots per unit cell: \( H_{c.m.} = \sum_q H_q \), where

\[ H_q = \frac{1}{N} \left( \sum_a \frac{1}{2m^*} \left[ P_{q,\alpha} + \frac{N}{c} A(U_{q,\alpha}^*) \right]^\dagger \right) \cdot \left[ P_{q,\alpha} + \frac{N}{c} A(U_{q,\alpha}) \right] + \frac{N^2}{2} \sum_{a,a'} U_{q,\alpha}^* \cdot C_{q\alpha,a,a'} \cdot U_{q,\alpha'} \]  

(33)

The dynamical matrix is defined by

\[ C_{q\alpha,a,a'} = \sum_n e^{i q \cdot R_{n}^{(0)}} C_{a,a'} \left( R_{n}^{(0)} \right) ; \quad C_{a,a'} \left( R_{n}^{(0)} \right) = C_{n,\alpha; 0,\alpha'} \]  

(34)

and it is hermitean \( C_{q\alpha';\alpha} = C_{q\alpha,a,a'}^* = C_{-q\alpha,a,a'} \).

Next we want to recover the limiting case considered in Sect. III. If the dots in a given unit cell are far away from those in neighboring cells, then in (34) only the term with \( R_{1}^{(0)} = 0 \) contributes, \( C \) does not depend on \( q \), consequently the index \( q \) is redundant, and (33) agrees with (12).

Our preliminary result (33) is not yet diagonal in \( \alpha,\alpha' \). In some special cases (see e.g. two identical dots per unit cell considered in Sect. III) this can be achieved by an unitary transformation

\[ U_{q,\alpha} = \sum_{\alpha'} Q_{q,\alpha,\alpha'} \cdot \hat{U}_{q,\alpha'} ; \quad Q_{q,\alpha',\alpha} = Q_{q,\alpha,\alpha'}^{-1} \]  

(35)

under which the one– particle term in (33) is invariant and the transformed interaction term

\[ \frac{1}{2} \sum_{\alpha,\alpha'} \hat{U}_{q,\alpha}^* \cdot \hat{C}_{q\alpha,\alpha',\alpha} \cdot \hat{U}_{q,\alpha'} \]  
\[ \hat{C}_{q\alpha,\alpha',\alpha} = \sum_{\alpha_1,\alpha_2} Q_{\alpha,\alpha_1}^{-1} C_{q\alpha_1,\alpha_2} Q_{\alpha_2,\alpha'} \]  

(36)

can be made diagonal \( \hat{C}_{q\alpha,\alpha',\alpha} = \delta_{\alpha,\alpha'} \) by a proper choice of \( Q_{\alpha,\alpha'} \). Now, (33) reads \( H_q = \sum_\alpha H_{q,\alpha} \), where

\[ H_{q,\alpha} = \frac{1}{N} \left( \frac{1}{2m^*} \left[ \hat{P}_{q,\alpha} + \frac{N}{c} A(\hat{U}_{q,\alpha}^*) \right]^\dagger \right) \cdot \left[ \hat{P}_{q,\alpha} + \frac{N}{c} A(\hat{U}_{q,\alpha}) \right] + \frac{N^2}{2} \sum_{a,a'} \hat{U}_{q,\alpha}^* \cdot \hat{C}_{q\alpha,a,a'} \cdot \hat{U}_{q,\alpha} \]  

(37)

The eigenvalues of (37) can be obtained from those of (33) because corresponding quantities have the same commutation rules. Such an unitary transformation does not exist, e.g., for two different dots per cell. Then (33) has to be solved directly using the method described in Ref. 3.

B. Dynamical Matrix for Bravais lattices

From now on we consider Bravais lattices what means that we can forget the indices \( \alpha \) in the first part of this Section. Then the dynamical matrix

\[ C_q = \Omega + \beta N \sum_{R_n^{(0)} \neq 0} \left( 1 - e^{i q \cdot R_n^{(0)}} \right) T \left( R_n^{(0)} \right) \]  

(38)
is real and symmetric, but generally not diagonal, even if $\Omega$ is diagonal. A very important conclusion is apparent in (38). In the limit $q \to 0$, the inter-dot interaction (represented by $\beta$) has no influence on $C_q$ and therefore on the spectrum. This means, that the excitation spectrum observed by FIR spectroscopy is not influenced by inter-dot interaction and agrees with the one-electron result (as in the single dot). This statement is rigorous for parabolic confinement (see Appendix). It can also be understood intuitively, because a $q=0$ excitation is connected with homogeneous in-phase elongations of the dots which do not change the distance between the electrons. We want to mention that this conclusion seems to be in contradiction with the experimental work in Ref. [11]. They found a splitting of the upper and lower excitation branch at $B=0$ and $q=0$ for circular dots in a rectangular lattice, which they interpreted within a phenomenological model of interacting dipoles as a consequence of lattice interaction. However, they use mesoscopic dots with a diameter of 370000 Å and lattice periods of 400000 and 800000 Å. These dots are clearly beyond our microscopic quantum mechanical model, which rests on a parabolic confinement.

For the rectangular lattices considered in our numerical examples we define $R^{(0)} = N_1 a_1 e_1 + N_2 a_2 e_2$ and $q = q_1 \frac{2\pi}{a_1} e_1 + q_2 \frac{2\pi}{a_2} e_2$ with the lattice constants $a_1$ and $a_2$ and integers $N_1$ and $N_2$ characterizing the lattice sites. The components of $q$ vary in the Brillouin zone (BZ) in the range $[-1/2, +1/2]$. The dipole tensor (15) reads

$$T(N_1, N_2) = \frac{1}{(N_1^2 a_1^2 + N_2^2 a_2^2)^{1/2}} \begin{pmatrix} 2N_1^2 a_1^2 - N_2^2 a_2^2 & 3N_1 N_2 a_1 a_2 \\ 3N_1 N_2 a_1 a_2 & 2N_2^2 a_2^2 - N_1^2 a_1^2 \end{pmatrix}$$

(39)

Although for all figures the exact dynamical matrix is used, it is useful to consider the results with nearest neighbor (n.n.) lattice sums in (38) separately. This provides simple formulas for order-of-magnitude estimates.

$$C_{11} = \omega_1^2 + 2p_1 \left[1 - \cos(2\pi q_1)\right] - p_2 \left[1 - \cos(2\pi q_2)\right]$$

$$C_{22} = \omega_2^2 + 2p_2 \left[1 - \cos(2\pi q_2)\right] - p_1 \left[1 - \cos(2\pi q_1)\right]$$

$$C_{12} = \Omega_{12}$$

(40)

where we introduced the interaction parameters $p_i = \frac{2\beta N_{i}}{a_i^2}$.

Convergency of Dynamical matrix

(for $a=2a_1$ in the middle of the irreducible Brillouin zone)

![Convergency of the lattice sums $S_{ik}$ as defined in (41) with increasing number of cubic shells $N_{max}$ for $q$ in the middle of the Brillouin zone.](image)

The convergence of the lattice sums $S_{ik}$ in the dynamical matrix is shown Fig.3. $S_{ik}$ is defined by

$$C_{ik} = \Omega_{ik} + p_2 S_{ik}$$

(41)
and depends only on $\mathbf{q}$ and the ratio $a_1/a_2$. Apart from the off-diagonal elements, which vanish in n.n. approximation, the error of the n.n. approximation is less than 30%.

C. Special Features of the Magneto– Phonon Spectrum

Fig.s 4-6 show the magneto– phonon spectrum for circular dots on a rectangular lattice with $a_1 = 2a_2$. Because the two interaction parameters have a fixed ratio, it suffices to use one of them for characterizing the interaction strength. We chose the larger one $p_2 = p$. For $B = 0$ and isolated dots ($p = 0$), the two excitation modes are degenerate. If we tune up the interaction strength represented by $p$, a $\mathbf{q}$ dependent splitting appears (see Fig. 4).

![Magneto– phonon dispersion at $B = 0$ for several interaction parameter values and $\mathbf{q}$ on symmetry lines of the Brillouin zone. ($p = p_2$ values in the inset are in units $\omega_0^2$.)](image)

This splitting is a manifestation of the dot interaction. For a certain critical $p_{cr}$ the lower mode becomes soft. This feature will be discussed below. There are points in the BZ, however, where the degeneracy for finite $p$ remains. These points will be investigated now. We demonstrated in Sect.II after formula (8) that necessary for degeneracy is $C_{12} = 0$, i.e., the dynamical matrix must be diagonal. Then the points with degeneracy are defined by the condition $C_{11} = C_{22}$. As seen in (38), for circular dots $\omega_1 = \omega_2 = \omega_0$ this happens in the center of the BZ $\mathbf{q} = 0$. The next question to be discussed is if there are other points with degeneracy. The first condition $C_{12} = 0$, is fulfilled for all points on the surface of the BZ. The second condition must be investigated for special cases. We find, that for quadratic lattices $a_1 = a_2$ with circular dots $\omega_1 = \omega_2$ both modes are degenerate at the point $\mathbf{q} = (1/2, 1/2)$. In the case shown in Fig.4 this point is somewhere between $(1/2, 1/2)$ and $(1/2, 0)$. In n.n. approximation (40), however, this equation is even fulfilled on full curves in the BZ defined by $p_1 [1 - \cos(2\pi q_1)] = p_2 [1 - \cos(2\pi q_2)]$. In a cubic lattice, this is the straight lines $q_2 = \pm q_1$. The contributions beyond n.n.s remove the exact degeneracy on this curve in the interior of the BZ, but leave a kind of anti-crossing.

3 The term magneto– phonon is attributed to the fact that there is no exchange and there are harmonic forces between the oscillating individuals. One could also call them magneto– plasmons, if one wants to emphasize that it is only electrons which oscillate, and no nuclei.
behavior of the two branches.

An important parameter, which characterizes the influence of the dot interaction in circular dots, is the *band width* at $B=0$, i.e., the maximum splitting of the two branches due to dot interaction. (Remember that this splitting vanishes for noninteracting circular dots.) Assume $a_1 > a_2$. Then the largest splitting for circular dots in n.n. approximation appears at $q = (0, 1/2)$ and has the amount

$$W = \max(\Delta E_+ - \Delta E_-) = \max(\omega_+ - \omega_-) = \sqrt{\omega_0^2 + 4p_2} - \sqrt{\omega_0^2 - 2p_2}$$

(42)

For small dot interaction and in units $\omega_0$, this is proportional to the interaction parameter $W/\omega_0 \rightarrow 3p_2$.

We next discuss the appearance of *soft modes*. The question is, for which $q$, $B$ and interaction parameter $p$ this happens. The general condition for vanishing of the lowest mode is $C_{11} \cdot C_{22} = C_{12}^2$ (see Sect. II). In this condition the magnetic field does not appear. For circular dots and with the definition (41) this equation reads

$$[\omega_0^2 + p_2 S_{11}][\omega_0^2 + p_2 S_{11}] = p_2^2 S_{12}^2$$

(43)

After introducing a dimensionless critical interaction parameter $P_{cr} = p_2/\omega_0^2$, we obtain a quadratic equation for $P_{cr}$ which has the solution

$$P_{cr} = -\frac{1}{2} \frac{tr}{det} \pm \sqrt{\left(\frac{1}{2} \frac{tr}{det}\right)^2 - \frac{1}{det}}$$

(44)

where $det = S_{11}S_{22} - S_{12}^2$ and $tr = S_{11} + S_{22}$. For our numerical case $a_1 = 2a_2$ and n.n. interaction for $S_{ik}$ the lowest mode becomes soft at $q = (0, 1/2)$ and the critical interaction parameter is $P_{cr} = 1/2$. Inclusion of lattice contributions beyond n.n. shifts $P_{cr}$ to 0.7543. The most important result of this paragraph is that lattice softening is *independent* of $B$ (see also Fig.s 5 and 6). The latter conclusion is *exact* within the range of validity of the Hamiltonian (30) and no consequence of any subsequent approximation or specialization.

![Diagram](image)

FIG. 5. The same as Fig.4, but for $p = p_{cr}$ and several magnetic fields.
FIG. 6. Magneto–phonon excitations as a function of $B$ for the symmetry points in the Brillouin zone. The upper abscissa is independent of the effective mass, the lower one applies to GaAs.

V. INTRA-DOT–EXCITATIONS FOR $N=2$

Intra-dot excitations for circular dots in a cubic lattice and for $N = 2$ can be calculated easily. We define the relative coordinate $r = r_2 - r_1$, and assume that all dots are equivalent (also with respect to their environment). Then the indexes $(n, \alpha)$ can be chosen as $(0, 0)$ and omitted. The relative Hamiltonian reads

$$H_{rel} = 2 \left\{ \frac{1}{2m^*} \left[ p + \frac{1}{2c} A(r) \right]^2 + \frac{1}{2} r \cdot D \cdot r + \frac{\beta}{2r} \right\}$$

where $p = -i \nabla_r$ and

$$D = \frac{1}{4} \Omega + \frac{\beta}{2} T_0; \quad T_0 = \sum_{n, \alpha(\neq 0, 0)} T \left( R^{(0)}_{n,\alpha} \right)$$

It is worth emphasizing that $H_{rel}$ contains a contribution from neighboring dots, originating from the interdot Coulomb interaction. A trivial angular dependent part can only be decoupled from $H_{rel}$, or, the 2-dimensional Schrödinger equation can be traced back to an ordinary radial Schrödinger equation, if the term $r \cdot D \cdot r$ has the same circular symmetry as the intra-dot Coulomb term $\beta/(2r)$. Therefore we confine ourselves to circular dots on a cubic lattice, and we have

$$T_0 = \frac{1}{a^3} \sum_{N_1,N_2 \neq 0,0} \frac{1}{(N_1^2 + N_2^2)^{3/2}} I \approx \frac{4}{a^3} I$$

where the simple result is in n.n. approximation. Using the interaction parameter $p = 2N\beta/a^3$ (with $N = 2$) defined above, we obtain

$$D = \frac{1}{4} (\omega_0^2 + 2p) I$$
In this way, dot interaction defines an effective confinement frequency \( \omega_{0,\text{eff}}^2 = \omega_0^2 + 2p \). This means that the c.m. excitations have to be calculated (or interpreted) with another confinement potential then the relative excitations. In our figures we present results for \( \omega_{0,\text{eff}} = 0.2 \text{ a.u.}^* \), which agrees with the bare confinement potential used in Sect.IV and the mean value in Sect.III. Because our results are presented in units of \( \omega_0 \), they depend on \( \omega_0 \) only weakly through the differing influence of electron-electron interaction. For the absolute values, however, the influence of the dot interaction can be tremendous.

In the relative motion there is a coupling between orbital and spin parts through the Pauli principle. For \( N = 2 \) and a circular effective confinement, Pauli principle demands that orbital states with even and odd relative angular momentum \( m_i \) must be combined with singlet and triplet spin states, respectively (see e.g. Ref. [7]). For the c.m. motion there is no interrelation between orbital and spin part because the c.m. coordinate is fully symmetric with respect to particle exchange. Consequently, any c.m. wave function can be combined with a given spin eigen function. The only spin dependent term in the total energy considered here is the Zeeman term, which reads in our units

\[
E_B = 0.9134 \cdot 10^{-2} \frac{g_s B[\text{Tesla}]}{\omega_0[\text{a.u.}^*]} \frac{M_s}{2} \tag{49}
\]

where we used \( g_s = -0.44 \) for the gyro-magnetic factor of GaAs from Ref. [8]. The total spin quantum number is \( M_s = 0 \) for the singlet state and \( 0, \pm 1 \) for the triplet state. One of the most interesting points in quantum dot physics is that the total orbital angular momentum of the ground state depends on the magnetic field (see e.g. Refs [8, 9]). This feature is a consequence of electron–electron interaction. For our parameter values, the relative orbital angular momentum of the ground state \( m_i \) changes from 0 to -1, from -1 to -2, and from -2 to -3 at \( B = 1.250, 4.018, \) and 5.005 Tesla. This corresponds to a sequence \( M_s=0, +1, 0, +1 \) for the spin quantum number. Figures 7a-c show the excitation frequencies for three B-values lying within the first three regions. \( m_f \) is the relative orbital angular momentum of the final state. All excitations are included irrespective of selection rules. For dipole transitions only two of them would remain (the lowest excitation with \( m_f = m_i \pm 1 \)). For \( B = 0 \), the lowest excitation energy (in units \( \omega_0 \)) for noninteracting electrons would be 1. As seen in in Fig.7a, electron–electron interaction decreases this value by at least a factor of 1/2. The same holds qualitatively for finite \( B \). This is connected to the fact, that the ground state depends on \( B \). Let’s consider an example. For \( B = 1.250 \text{ Tesla} \) the ground state switches from \( m_i = 0 \) to \( m_i = -1 \). This implies that for \( B \) approaching this transition field from below, the excitation energy for dipole allowed transition from \( m_i = 0 \) to \( m_f = -1 \) converges to 0. In other words, there is a level crossing at the transition field. Therefore, very small transition energies and switching of the ground state are connected.
FIG. 7. Intradot excitation energies (in units of the effective confinement frequency) for $B=0$ (a), $B=3$ Tesla (b) and $B=4.5$ Tesla (c). The corresponding relative orbital angular momenta of the ground state are $m_i=0$ (a), $-1$ (b), and $-2$ (c) and the spin angular momenta $M_s=0$ (a), $+1$ (b), and 0 (c).

For a qualitative understanding, Figures 7a-c can be used together with Figures 1a, 1b, 4, and 5 to investigate the relative position of collective and intra-dot excitations. The conclusion is that for small dot interaction (for $p$ well below $p_{cr}$), the lowest intra-dot excitation energies lie well below the lowest c.m. excitations. Apart from using a different terminology, this conclusion agrees with the experimental findings in Ref. 10.

Fig. 7b and c, which belong to finite $B$, show the Zeeman splitting. All transition energies to final states with odd
$m_f$ are triplets because the corresponding spin state is a triplet state. The thin lines of a triplet belong to spin-flip transitions.

**FIG. 8.** Intradot excitation energies (in units of the effective confinement frequency) as a function of the magnetic field. The small Zeeman splitting is neglected. The B-values, where the angular momenta of the ground state change, are indicated by vertical lines. The absolute value of the final state orbital momentum $m_f$ of the curves at B=0 grows from bottom to top by 1 starting with 0.

In Fig.8 the B– dependence of the lowest excitation energies is shown. It is clearly seen that the curves exhibit a kink at those B– values, where the ground state configuration changes. The size of the kink decreases with increasing $B$. If this kink could be resolved experimentally (e.g. by electronic Raman spectroscopy), it would be a direct indication for the change of the ground state configuration, and thus an experimentally observable consequence of electron–electron interaction.
VI. SUMMARY

We solved the Schrödinger equation for a lattice of identical parabolic (but not necessarily circular) quantum dots with Coulomb interaction (in dipole approximation) between the dots. We provide an overview over the state of art of these systems which includes the results of former publications. References can be found in the text.

- Similar to single dots, the center of mass coordinates of all dots can be separated from the relative coordinates. Only the c.m. coordinates of different dots are coupled to each other. The relative coordinates of different dots are neither coupled to each other nor to the c.m. coordinates.

- This gives rise to two types of excitations: two collective c.m. modes per dot and a complex spectrum of intra-dot excitations. In periodic arrays only the collective c.m. modes show dispersion. Intra-dot excitations are dispersion-less.

- The c.m. system can be solved exactly and analytically providing magneto–phonon excitations characterized by a certain wave number $q$ within the Brillouin zone. For $q = 0$ and one dot per unit cell, interdot interaction does not have any influence on the c.m. excitations.

- All dipole allowed excitations (seen in FIR experiments) are not influenced by the dot interaction.

- Interdot interaction between two dots influences the spectrum through a single parameter $p = 2N\beta/a^3$, where $a$ is the distance between the dots, $N$ the number of electrons per dot and $\beta$ the inverse background dielectric constant.

- If $p$ exceeds a certain critical value $p_{cr}$, the lowest c.m. mode becomes soft leading to an instability. This transition is independent of the magnetic field.

- For $B=0$ and one circular dot per unit cell, the two c.m. modes are not only degenerate in the middle of the Brillouin zone, but also at some points on the surface. If we use the n.n. approximation for the lattice sums in the dynamical matrix, degeneracy is maintained even on full curves in the Brillouin zone.

- Intra-dot excitations have to be calculated from an effective confinement. In circular dots with a cubic environment in nearest neighbor approximation the effective confinement frequency reads $\omega_{0,eff}^2 = \omega_0^2 + 2p$. This effective confinement differs from that for the c.m. motion.

- For $p$ well below $p_{cr}$, the lowest intra-dot excitations are much smaller than the lowest collective excitations.

- The intra-dot excitation energies versus magnetic field exhibit kinks at those fields, where the angular momentum of the ground state changes.

In the Appendix we prove a Kohn Theorem for dot arrays with Coulomb interaction between the dots without the dipole approximation. The individual confinement potentials can be arbitrarily arranged and can carry different electron numbers, but have to be described by identical confinement tensors. This means that for breaking Kohn’s Theorem in dot arrays, we have to have at least two different confinement species.

APPENDIX

We are going to prove that for an arbitrary array\(^4\) of identical parabolic quantum dot potentials\(^5\) in an homogeneous magnetic field: i) the total c.m. degree of freedom can be separated from the rest, ii) the total c.m. Hamiltonian is not influenced by Coulomb interaction, and iii) the eigenvalues of the total c.m. Hamiltonian are independent of the electron number $N$ in each dot\(^6\).

The Hamiltonian $H = H^{(0)} + V$ consists of an one–particle term $H = H^{(0)}$ and the Coulomb interaction between all dots.

---

\(^4\) The dot centers can be arranged arbitrarily.

\(^5\) The confinement tensors $\Omega$ of all dots must be equal.

\(^6\) The electron number in different dots can be different.
electrons $V$. The dot centers are located at $R^{(0)}_\alpha$ and the electron coordinates are denoted by $r_{i\alpha} = R^{(0)}_\alpha + u_{i\alpha}$. Then we have

$$H^{(0)} = \sum_{i\alpha} \left\{ \frac{1}{2m^*} \left[ p_{i\alpha} + \frac{1}{c} A(R^{(0)}_\alpha + u_{i\alpha}) \right]^2 + \frac{1}{2} u_{i\alpha} \cdot C \cdot u_{i\alpha} \right\}$$

(50)

First of all we shift the gauge center for each electron into the middle of the corresponding dot using an unitary transformation similar to (16). This transforms the shift $R^{(0)}_\alpha$ in the argument of the vector potential away. Next we drop the index $\alpha$ in (50) so that the index 'i' runs over all electrons in all dots. Now we perform a transformation to new coordinates $\tilde{u}_i$

$$u_i = \sum_k Q_{ik} (\sqrt{N} \tilde{u}_k) ; \quad (\sqrt{N} \tilde{u}_i) = \sum_k Q_{ki} u_k$$

(51)

where $Q_{ik}$ is an unitary matrix. This implies

$$p_i = \sum_k Q_{ik} (\sqrt{N} \tilde{p}_k) ; \quad (\sqrt{N} \tilde{p}_i) = \sum_k Q_{ki} p_k$$

(52)

It is possible to choose for the first column $Q_{k1} = \frac{1}{\sqrt{N}}$. The other columns need not be specified. Then $\tilde{u}_1 = (1/N) \sum_i u_i = U$ is the c.m. of all elongations, or, the c.m. of the electron coordinates with respect to the weighted center of the dot locations $R^{(0)}_\alpha = (1/N) \sum_\alpha N_\alpha R^{(0)}_\alpha$, where $N_\alpha$ is the number of electrons in dot $\alpha$. The corresponding canonical momentum $\tilde{p}_1 = (1/i) \nabla \tilde{u}_1 = \tilde{P}$ is the c.m. momentum. Inserting our transformation into (50) provides

$$H^{(0)} = \sum_i \left\{ \frac{1}{2m^*} \left[ \frac{1}{\sqrt{N}} \tilde{p}_1 + \sqrt{N} c A(\tilde{u}_1) \right]^2 + \frac{N}{2} \tilde{u}_1 \cdot C \cdot \tilde{u}_1 \right\}$$

(53)

The term $i = 1$ in (53) is the (separated) c.m. Hamiltonian

$$H_{c.m.} = \frac{1}{2m^*} \left[ \frac{1}{\sqrt{N}} \tilde{P} + \sqrt{N} c A(\tilde{U}) \right]^2 + \frac{N}{2} \tilde{U} \cdot C \cdot \tilde{U}$$

(54)

which agrees with (10). Clearly, the Coulomb interaction $V$ in $H$ is independent of the c.m., and does not contribute to $H_{c.m.}$. For the independence of the eigenvalues of $N$ see the discussion following (10).

This prove, in particular the step from (50) to (53), is not correct if the dot confinement tensor $C$ depends on $\alpha$ (or 'i' in the changed notation). Therefore, all dots must have the same $C$, but can have different electron numbers $N_\alpha$. In other words, the total c.m. excitations in dot arrays, which are seen in FIR spectra, are not affected by the e− e− interaction, if and only if all confinement tensors $C$ are equal. On the other hand, if we want to observe e− e− interaction in the FIR spectra and break Kohn’s Theorem, we have to use dot lattices with at least two different confinement tensors. The simplest way to implement this is using a lattice with two non-circular dots per cell, which are equal in shape, but rotated relative to each other by 90 degrees.

VII. ACKNOWLEDGMENT

I am indebted to D.Heitmann, H.Eschrig, and E.Zaremba and their groups for very helpful discussion and the Deutsche Forschungs– Gemeinschaft for financial funding.
1. L.Jacak, P.Hawrylak, and A.Wojs, *Quantum Dots*, Springer 1998
2. K.Kempa, D.A.Broido, and P.Bakshi; Phys. Rev. B43, 9343 (1991);
   A.O.Govorov and A.V.Chaplik; Sov.Phys.72, 1037(1991)
   A.O.Govorov and A.V.Chaplik; J.Phys.: C 6, 6507(1994)
   A.V.Chaplik and A.O.Govorov ; J.Phys.: C 8, 4071(1996)
3. J.Dempsey, N.F.Johnson, L.Brey, and B.I.Halperin; Phys.Rev. B 42, 11708 (1990)
4. T.Chakraborty, V.Halone and P.Pietiläinen; Phys.Rev. B 43, 14289 (1991)
5. A.O.Govorov and A.V.Chaplik; J.Phys.: C 6, 6507(1994)
   A.V.Chaplik and A.O.Govorov ; J.Phys.: C 8, 4071(1996)
6. J.Dempsey, N.F.Johnson, L.Brey, and B.I.Halperin; Phys.Rev. B 42, 11708 (1990)
7. T.Chakraborty, V.Halone and P.Pietiläinen; Phys.Rev. B 43, 14289 (1991)
8. M.Taut, J.Phys. A27, 1045 (1994) and erratum J.Phys. A27, 4723 (1994). Additionally, in formula (10) in the term containing
   $\frac{\partial}{\partial \alpha}$ a factor $\frac{1}{2}$ is missing, and on the r.h.s. of (19a) and (20a) $\tilde{\omega}$ must be replaced by $\tilde{\omega}_r$.
9. U.Merkt, J.Huser, M.Wagner, Phys.Rev. B 43, 7320 (1991);
   M.Wagner, U.Merkt, and A.V.Chaplik, Phys. Rev. B 45, 1951 (1992)
10. P.A.Maksym, Physica B 249, 233 (1998); O.Dippel, P.Schmelcher, and L.S.Cederbaum; Phys.Rev.A 49, 4415 (1994);
   B.Schuh; J.Phys. A 18, 803 (1985)
11. C.Tsallis; J.Math.Phys. 19, 277 (1978)
12. Y.Tikosinsky; J.Math.Phys. 20, 406 (1979)
13. M.Taut, Phys. B 27, 1045 (1994) and erratum J.Phys. A27, 4723 (1994). Additionally, in formula (10) in the term containing
   $\frac{\partial}{\partial \alpha}$ a factor $\frac{1}{2}$ is missing, and on the r.h.s. of (19a) and (20a) $\tilde{\omega}$ must be replaced by $\tilde{\omega}_r$.
14. U.Merkt, J.Huser, M.Wagner, Phys.Rev. B 43, 7320 (1991);
   M.Wagner, U.Merkt, and A.V.Chaplik, Phys. Rev. B 45, 1951 (1992)
15. P.A.Maksym, T.Chakraborty, Phys.Rev.Lett. 65, 108 (1990) and Phys.Rev.B 45, 1947 (1992);
   P.A.Maksym, Physica B 184, 385 (1993)
16. C.Schüller, K.Keller, G.Biese, E.Ullrichs, L.Rolf, C.Steinebach, and D.Heitmann, Phys.Rev.Lett. 80, 2673 (1998)
17. C.Dahl, J.P.Kotthaus, H.Nickel and W.Schlapp;
    Phys.Rev. 46, 15590 (1992)