Magnetic Field Induced Two-Body Phenomena in Atoms

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Abstract

We report on a number of recently discovered phenomena which arise due to the interaction of the collective (CM) and internal motion of atoms moving in magnetic fields. For neutral atoms the properties of the so-called giant dipole states are discussed and their experimental preparation is outlined. For ions moving in magnetic fields the energy flow between the collective and electronic motions is studied. For low CM velocities this results in a crossover from localization to delocalization with respect to the spreading of the mixing of the CM and internal motion in quantum number space. Large CM velocities induce for both the classical as well as quantum ion the self-ionization process. Future perspectives for both multi-electron atoms as well as molecules are drawn.

1 Introduction

External magnetic fields are well-known to have a strong impact on the properties of particle systems. A number of intriguing phenomena in different areas of physics like, for example, the quantum Hall effect in solid state physics or the interplay of regularity and chaos in few-body atomic systems have their origin in the combination of the magnetic and Coulomb forces. By nature the Coulomb potential together with the paramagnetic and diamagnetic interaction represents a nonseparable and nonlinear problem already on the one-particle level. As a consequence both the dynamics as well as the electronic structure of atoms or molecules are subject to severe changes in the presence of a strong magnetic field.

In the present article we review on selected recent developments for Rydberg atoms in strong laboratory magnetic fields and outline future
perspectives. We also provide concrete suggestions how to experimentally access and observe the corresponding atomic states and processes. Of course, similar phenomena are to be expected also for astrophysical field strengths but the focus in this work is on highly excited atoms exposed to laboratory field strengths. Our central issue are phenomena and effects arising due to the nonseparability of the center of mass and internal motion for two-body systems in external magnetic fields and/or crossed electric and magnetic fields \[3\]. Due to the smallness of the ratio of the electron to nuclear mass one might be, at first glance, tempted to believe that the effects of the coupling of the collective and internal motion provide only tiny corrections to the overall dynamics. This is however in general not correct and indeed we will provide a number of different physical situation in which even the correct qualitative behaviour cannot be obtained without including the interaction of the center of mass and relative motion. Prominent effects due to this interaction are the chaotic classical diffusion of the center of mass \[4, 5, 6, 7\], the existence of giant dipole states for neutral atoms and dynamical phenomena like the self-ionization process for rapidly moving ions in strong magnetic fields. As we shall see in the following the ongoing research in this area reveals more and more of the beautiful phenomena created by the competition of magnetic and Coulomb forces.

In detail we proceed as follows. Section I gives first a brief account of the separation of the CM motion for neutral particle systems in magnetic fields and subsequently investigates a class of decentred magnetically dressed Rydberg states possessing a huge electric dipole moment. A possible experimental set-up for the preparation of these states as well as applications to the positronium atom are discussed. Section II reviews on recent developments for ionic systems like the classical self-ionization process for rapidly moving ions in magnetic fields. A number of most recently discovered quantum properties of ions in fields are outlined and clearly indicate that quantization introduces a number of new features.

2 Neutral atoms in magnetic fields

2.1 Giant dipole states in crossed fields

In the absence of an external magnetic field the CM and electronic motions of an atom separate due to the conservation of the total canonical momentum \( P \). \( P \) is the generator for coordinate space translations which represent a symmetry for any system of particles interacting through a
potential which depends only on the distances of the particles. In a homogeneous magnetic field this symmetry is lost since the Hamiltonian depends explicitly on the vector potential associated with the external field. Nevertheless, there exists a so-called phase space translation group which represents a symmetry in the presence of the external field. This group is generated by the so-called pseudomomentum. Historically the pseudomomentum was implicitly used by Lamb [8] in order to perform a so-called pseudoseparation of the CM motion for the hydrogen atom in a homogeneous magnetic field. In the late seventies a more profound mathematical treatment of the pseudoseparation for two-body systems was achieved [9]. In the eighties and nineties the pseudoseparation for many-body systems has been reviewed (see ref. [10] and references therein) and it was adapted to the needs of molecular physics [11].

In order to perform the pseudoseparation transformation as mentioned above one starts with the Hamiltonian of the atom in the laboratory frame which reads for a two-body system

\[
\mathcal{H}_L = \sum_{i=1}^{2} \left[ \frac{1}{2m_i} (p_i - e_i A_i)^2 - e_i E r_i \right] + V(|r_1 - r_2|) \tag{1}
\]

where \(e_i, m_i, A_i\) and \(E\) denote the charges, masses, vector potential and electric field vector, respectively. \(\{r_i, p_i\}\) are the Cartesian coordinates and momenta in the laboratory coordinate system. Subsequently one chooses a certain gauge for the vector potential \(A_i\) belonging to the magnetic field \(B\). Most commonly this is done by introducing the symmetric gauge \(A_i = \frac{1}{2} B \times r_i\). As a next step a coordinate transformation from laboratory to relative and center of mass coordinates follows. Finally the pseudomomentum is introduced as a canonical momentum of the center of mass coordinate by performing a corresponding transformation in momentum space which gives the final transformed Hamiltonian. Choosing, according to the above, a specific gauge in the laboratory Hamiltonian has, however, a serious drawback: It is not possible to discern between gauge dependent and gauge invariant terms in the transformed Hamiltonian. In order to extract the physics, which is inherently gauge-independent, from the transformed Hamiltonian it is a central issue to identify gauge invariant parts of it. The necessity of a gauge-invariant approach has been realized in the nineties by different authors [12, 13, 14] which have chosen different approaches in order to obtain a gauge-independent formalism and results for the problem of two interacting particles. Most importantly it was finally shown [14, 15]...
that there exists a potential picture (see below) due to a number of gauge-invariant terms emerging from the generalized pseudoseparation.

Let us consider the final Hamiltonian resulting from the afore-mentioned gauge independent pseudoseparation

$$\mathcal{H} = \mathcal{T} + \mathcal{V}$$

where

$$\mathcal{T} = \frac{1}{2\mu} (p - qA(r))^2$$

and

$$\mathcal{V} = \frac{1}{2M} (K - eB \times r)^2 + V(r) + \frac{M}{2}v_D^2 + Kv_D$$

with the charge

$$q = \frac{m}{\mu}$$

where

$$\mu = \frac{mM_0}{M}$$

and

$$\hat{\mu} = \frac{mM_0}{M_0 - m}$$

are different reduced masses. K is now the constant vector of the pseudomomentum which contains a term linear in the external electric field. \{r, p\} denote the canonical pair of variables for the internal relative motion.

The explicit form of the kinetic energy \(\mathcal{T}\) depends on the chosen gauge via the vector potential \(A\). The important novelty with respect to our Hamiltonian \(\mathcal{H}\) is the appearance of the potential term \(\mathcal{V}\). Apart from the Coulomb potential \(V\) and the constant term \(\frac{M}{2}v_D^2 + Kv_D\) \(\mathcal{V}\) contains an additional potential term

$$V_o = \frac{1}{2M} (K - eB \times r)^2$$

The latter is gauge independent, i.e. does not contain the vector potential, and, therefore, fully deserves the interpretation of an additional potential for the internal motion with the kinetic energy (2). Apart from the constant \(\frac{K^2}{2M}\) the potential \(V_o\) contains two coordinate dependent parts. The term linear in the coordinates \(-\frac{e}{\mu}(K \times B)r\) consists of two Stark terms: one which is due to the external electric field \(E\) and a second one which is a motional Stark term with an induced constant electric field \(\frac{1}{\mu}((K + Mv_D) \times B)\). The latter electric field is always oriented perpendicular to the magnetic one and arises due to the collective motion of the atom through the homogeneous magnetic field. Besides the linear terms there exists a quadratic, i.e. diamagnetic, term \(\frac{e^2}{2M}(B \times r)^2\)
in the potential $V_o$ which is of great importance for the main subject of the present section, i.e. the existence of giant dipole states of two-body system in external fields.

In the following we discuss the qualitative properties of the potential $V$. Figure 1 shows for the choice $B = (0,0,B)$, $K = (0,K,0)$ and a vanishing external electric field a two-dimensional intersection of the potential $V$ for $z = 0$. In the neighbourhood of the coordinate origin the Coulomb potential dominates. This regime corresponds in figure 1 to the tube around the origin of the coordinate plane $(x,y)$. With increasing distance from the origin along the $x-$axis ($y = 0$) the Stark term $\left(\frac{e}{M}B K x\right)$ increases and becomes eventually comparable with the Coulomb potential. This means that we encounter an approximately linear raise of the potential for positive values of the $x-$coordinate. For negative values of the $x-$coordinate eventually a saddle point emerges. In this coordinate regime the diamagnetic term of eq.(4) represents a small correction. For even larger distances the Coulomb potential becomes small and the shape of the potential $V$ is more and more determined by the diamagnetic term $\left(\frac{e^2}{2M}B^2(x^2 + y^2)\right)$. Due to the competition of the Stark and diamagnetic terms our potential $V$ develops an outer minimum and a corresponding potential well. The existence of both the saddle point and the outer minimum/well depends, of course, on the values of the magnetic field strength and the motional/external electric field strength. For a derivation and discussion of the corresponding conditions we refer the reader to the literature [14, 15]. We emphasize that the potential $V_o$ is inseparably connected with the finite nuclear mass. Assuming an infinite nuclear mass simply yields $V_o \equiv 0$ and the described features of the total potential $V$ disappear.

The above-discussed properties of our potential $V$ have important implications on the dynamical behaviour of the atom. First of all, we observe that the ionization of the atom can take place only in the direction of the magnetic field axis: In the direction perpendicular to the magnetic field vector the diamagnetic term $\left(\frac{e^2}{2M}(B \times r)^2\right)$ is dominating for large distances $\rho = (x^2 + y^2)^{1/2}$ and causes a confining behaviour. The second important observation is the fact that the existence of the outer well leads to new weakly bound states in this well. Let us provide some main characteristics of these states. In refs.[14, 15] an explicit approximation formula has been given for the position of the outer minimum: $x_0 \simeq -\frac{K}{B} + \frac{K M}{2 MB} y = z = 0$. Hence, for a laboratory field strength $B \sim 2.35$ Tesla and a motional/external electric field of the order of $E \sim 2.8 \times 10^4 \sqrt{V/m}$ the minimum is located at a distance of about
5.3 \times 10^{-6} m from the Coulomb singularity. For states in the outer well the electron and proton are therefore separated about 100,000 times as much as they are in the ground state of the hydrogen atom without external fields, i.e. we encounter a strongly delocalized atom of mesoscopic size. Since the well exists only on the half-hyperplane with a negative Stark term (see fig.1) these states possess a huge permanent electric dipole moment of the atom. This is in contrast to the well-known Rydberg states in a pure magnetic field which do not exhibit a permanent dipole moment. For energies close to the minimum the outer well is approximately an anisotropic harmonic potential. Low-lying quantum states can therefore be described by an anharmonic oscillator in a magnetic field \[14\]. The field-dependent kinetic energy (see eq.(2)) hereby determines the extension of the wave function in the plane perpendicular to the magnetic field. The deviation of the exact energies from those of the harmonic approximation changes, as expected, significantly with increasing degree of excitation. It grows stepwise and a major contribution to the anharmonicity comes from the quantum number \( n_z \) i.e. the excitation in the direction parallel to the magnetic field. This can also be seen in perturbation theory for higher terms of the expansion of the Coulomb potential where the major contributions to the energy corrections are due to those terms containing high powers of \( z \). For the computational techniques to obtain energies and wavefunctions for a large number of states in the outer well as well as a detailed discussion of their properties we refer the reader to ref.\[14\].

2.2 Experimental preparation

At this point it is natural to pose the question how one can experimentally prepare hydrogen atoms in crossed fields in their giant dipole states. This question has been investigated and discussed in detail in ref.\[16\]. We provide here the key ideas of the approach developed in this work. The important parameter which controls the formation of the outer well and is at disposal to the experimentalist is the external electric field strength. The preparation scheme consists of a sequence of steps which correspond to different electric field configurations. The first step excites the atom in the presence of a magnetic field but no electric field from the ground state with a laser pulse to a highly excited Rydberg state. In the second step an electric field is switched on within a time period of a few \( ns \) to a value \( E = E_c \) which corresponds to the existence of a shallow outer well. Subsequently, during the third step the
The electric field is kept constant for a time period $\Delta t$. The Rydberg states prepared in step one are localized in the Coulomb well. After the switching of the electric field to $E_c$ (step two) their energy is above the saddle point. Keeping the electric field constant during the third time step has the reason to achieve a significant spreading of the prepared state over the shallow outer well. Thereafter, in a third step, the electric field is switched to its final value $E_f$ which corresponds to a deep outer well.

This last step of the preparation procedure captures the wave packet in the outer well and a significant portion of the hydrogen atoms therefore ends up in low-energy states of the outer well. The second switching of the electric field to its final value $E_f$ is much slower than the first fast switching to the value $E_c$. The reason herefore is the broadening of the final energy distribution during the trapping of the hydrogen atoms in the outer well. An adiabatic second switching is therefore obligatory in order to end up with a narrow final energy distribution. Figure 2 shows the final energy distribution in the outer well resulting from calculations for an ensemble of classical trajectories which simulate the behaviour of the highly excited Rydberg states during the above-discussed steps of preparation. It can clearly be seen that the maximum of the energy distribution is below the ionization threshold (vertical dashed line in figure 2) and therefore a significant part of the prepared outer well states is strictly bound. For further details of the experimental setup and preparation like, for example, the specific switching procedures of the electric field or the influence of electric stray fields on the preparation scheme we refer the reader to ref.[16].

Let us provide some remarks with respect to the experimental detection of the giant dipole states of the hydrogen atom in crossed electric and magnetic fields. Direct state-to-state transitions for bound states in the outer well should be observable in the radio-frequency regime. The energy gap between the ground and first excited state corresponds, for the above-given field strengths, to a frequency of the order of magnitude of a few tens of MHz. Alternatively the large electric dipole moment suggests itself for detection which could be achieved through deflection of the atoms by a slightly inhomogeneous electric field [17]. Even though the binding energies of the states in the outer well are relatively small they should be stable as long as collisional interaction is prevented.

We have discussed the existence and properties as well as the experimental preparation and detection of giant dipole states for the hydrogen atom in crossed fields. The electric field can hereby be either an external one or due to the collective motion of the hydrogen atom through the
magnetic field. For these states the magnetic interaction is dominating the Coulomb attraction in the plane perpendicular to the magnetic field whereas parallel to the magnetic field we exclusively encounter Coulomb forces. The electric field causes the decentred character of these states \[18, 19, 20\]. Of course this kind of states does not only exist for one-electron atoms but should occur also for more-electron systems. Indeed it is a challenging question to ask for the existence and properties of decentred multielectron atoms in crossed external fields. Moreover one can imagine giant dipole molecules which, due to the presence of several heavy particles, might possess completely different features compared to atoms.

### 2.3 Application to Positronium

Besides the above-drawn general perspective there exists an intriguing application of the above results to exotic two-body systems, namely the positronium atom \[21, 22\]. Since the distance of the minimum of the outer well from the Coulomb singularity is approximately proportional to the total mass of the two-body system (\( \propto \frac{EM}{B^2} \)) we expect that the giant dipole states for positronium are significantly less extended than those of hydrogen. At the same time the critical electric field strength necessary for the existence of the outer well scales with \( M^{-\frac{2}{3}} \). As a consequence the typical reduction of size for fixed field strength is of the order of 10, i.e. for \( B \sim 2.35 \, T \) the extension of the Rydberg state is several thousand Bohr radii. The giant dipole states located in the outer well and the traditional Rydberg states located in the Coulomb well are separated by a wide and high potential barrier. For the positronium atom this has important consequences: the potential barrier prevents the particles from contact and therefore decreases the annihilation rate by many orders of magnitude. Indeed, for typical laboratory field strengths the lifetime can become many years and low-lying outer well states of positronium can, for all practical purposes, be considered as stable \[21, 22\]. Crossed fields offer therefore a unique opportunity for the stabilization of matter-antimatter two-body systems. For a detailed investigation of the positronium atom including dipole transition rates, tunneling probabilities as well as spectra and wavefunctions resulting from both perturbation theoretical and finite element calculations we refer the reader to refs.\[21, 22\]. We conclude with an important result of ref.\[22\] which should be of relevance to astrophysical situations: For sufficiently strong fields the energetically lowest decentred outer well state
becomes the global ground state of the atom. This statement holds for both the hydrogen atom as well as the Positronium atom. As a consequence the ground state of isolated positronium in strong crossed external fields is prevented from annihilation and represents a long-lived state.

3 Atomic ions in magnetic fields

3.1 Basic properties

For a charged atom in a magnetic field the interaction of its collective motion with the electronic motion is more intricate than for neutral species. From a physical point of view it is evident that the crudest picture for the CM motion describes the ion as an entity in the magnetic field with a charge and mass identical to the net charge and total mass of the ion. For neutral systems the net charge of the system is zero and the crudest picture for the behaviour of the CM is a free straightlined motion through the magnetic field. However, as we shall see below, we encounter coupling terms for the ions CM and electronic motions which can mix them up heavily thereby causing a number of interesting energy transfer processes between the degrees of freedom.

From a formal point of view the maximum number of commuting constants of motion is the same for neutral as well as charged systems. For neutral species, however, one can choose the three components of the total pseudomomentum $\mathbf{K}$ which are exclusively associated with the CM motion of the system. As a consequence the CM coordinates are cyclic and can be completely eliminated by the so-called pseudoseparation from the Hamiltonian (see section 2.1). For charged systems the only maximal set of commuting constants of motion is $(\mathbf{K}_\perp^2, \mathbf{L}_\parallel, \mathbf{K}_\parallel)$ where $\mathbf{K}_\perp$ is the component of the pseudomomentum perpendicular to the magnetic field and $\mathbf{L}_\parallel$ is the component of the total angular momentum parallel to the field. This set of quantities is not exclusively associated with the CM motion but involves both the CM and internal degrees of freedom. Unlike the neutral system the CM coordinates cannot completely be eliminated from the Hamiltonian. To simplify the Hamiltonian one therefore uses a transformation \[ \mathcal{H} = \mathcal{H}_1 + \mathcal{H}_2 + \mathcal{H}_3 \] (6)
where

$$H_1 = \frac{1}{2M} \left( p - \frac{Q}{2} B \times R \right)^2$$  \hspace{1cm} (7)

$$H_2 = e \alpha \left( B \times \left( p - \frac{Q}{2} B \times R \right) \right) r$$  \hspace{1cm} (8)

$$H_3 = \frac{1}{2m} \left( p - \frac{e}{2} B \times r + \frac{Q m^2}{2 M^2} B \times r \right)^2 + \frac{1}{2M_0} \left( p + \left( \frac{e}{2} - \frac{Q}{2M M_0} (M + M_0) \right) B \times r \right)^2 + V$$  \hspace{1cm} (9)

where $m$, $M_0$ and $M$ are the electron, nuclear and total mass, respectively. $\alpha = (M_0 + Zm)/M$ and $V$ is the Coulomb potential. For the vector potential $A$ we have adopted the symmetric gauge. The magnetic field vector $B$ again is assumed to point along the $z$-axis. $(R, P)$ and $(r, p)$ are the canonical pairs of variables for the CM and relative motion, respectively.

The Hamiltonian $H$ consists of three parts which introduce different types of interaction. The part $H_1$ in eq.(6), which involves solely the CM degrees of freedom, describes the cyclotron motion of a free pseudoparticle with mass $M$ and charge $Q$ in a homogeneous magnetic field (see comments at the beginning of this subsection). This zeroth order picture is, in general, not sufficient to describe the CM motion of the ion. In fact the behaviour of the CM can deviate strongly from the motion given by the Hamiltonian $H_1$ and exhibits a variety of different phenomena depending on the parameter values (energy, field strength, CM velocity) [23, 24, 25, 26, 27, 28]. The origin of this rich dynamics lies in particular in the Hamiltonian $H_2$ in eq.(7) which describes the coupling of the CM and electronic degrees of freedom. It represents a motional Stark term with a rapidly oscillating electric field which is determined by the dynamics of the system. Because of this "dynamical" electric field the collective and internal motion will, in general, mix up heavily. $H_3$ in eq.(8) contains only the electronic degrees of freedom and describes, to zeroth order, the relative motion of the electron with respect to the nucleus.
3.2 Mixing and localization properties for the collective and electronic motions

The Hamiltonian $H_2$ in eq.(7) destroys the picture of a decoupled CM and electronic motion and causes an interaction of these motions whose strength depends on several parameters (CM and internal energy, field strength). In the following we outline both classical and quantum effects and phenomena due to this interaction [23, 24, 25, 26, 27, 28]. We hereby pursue the track of an increasing strength of the coupling of the collective and internal motion which can in particular be achieved by increasing the energy of the CM motion.

In order to investigate the significance and effects of the coupling Hamiltonian $H_2$ one can use the following method for formally solving the total Schrödinger equation $H\Psi = E\Psi$. The most natural way is to expand the total wave function $\Psi$ in a series of products

$$\Psi(R,r) = \sum_{p,q} c_{pq} \Phi^L_p(R) \psi_q(r)$$

where $c_{pq}$ are the coefficients of the product expansion. The functions $\{\Phi^L_p\}$ obey the Schrödinger equation $H_1 \Phi^L_p = E^L_p \Phi^L_p$ for a free particle with charge $Q$ and mass $M$ in a homogeneous magnetic field, i.e. they are the corresponding Landau orbitals. One proper choice for the functions $\{\psi_q\}$ in the expansion (9) (see also below) are the eigenfunctions of the electronic Hamiltonian, i.e. $H_3 \psi_q = E^I_q \psi_q$ (q stands collectively for all electronic quantum numbers). If we insert the product expansion (9) for the total wave function $\Psi$ in the total Schrödinger equation and project on a simple product $\Phi^L_p \psi_q'$ we arrive at the following set of coupled equations for the coefficients $\{c_{pq}\}$:

$$(H_2 + E^L_p + E^I_q)c = Ec$$

where $c$ is the column vector with components $\{c_{pq}\}$. $E^L_p$ and $E^I_q$ are the diagonal matrices which contain the Landau energies $\{E^L_p\}$ and internal energies $\{E^I_q\}$, respectively. The original problem of the investigation of the significance and effects of the couplings among the CM and electronic degrees of freedom is now reduced to that of the solution of the eigenvalue problem (10). $H_2$ contains the matrix elements of the coupling Hamiltonian $H_2$. These elements can, via certain commutation relations, be transformed into pure dipole transition matrix elements of the CM as well as internal degrees of freedom (see ref.[23] for details)

$$H_2 = -i\epsilon(\epsilon^L_{p'} - \epsilon^L_p) \cdot (\Phi^L_{p'} | R | \Phi^L_p) \cdot |B \times (\psi_{q'} | r | \psi_q)\rangle$$

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Obviously the coupling matrix elements \( \langle H_2 \rangle_{\{p'q'\}\{pq\}} \) vanish if the two Landau states \( \Phi_{p'}^L \) and \( \Phi_p^L \) possess the same energy \( E_{p'}^L = E_p^L \). Transitions, therefore, do occur only for total states which involve different states of the collective motion with respect to their energy.

The expansion (9) with \( \{\psi_q\} \) being the eigenfunctions of the electronic Hamiltonian \( H_3 \) is particularly adequate if the coupling is sufficiently weak which means that the series contains only a few dominant terms. Otherwise, i.e.

in the case that many terms contribute significantly to the sum, the mixing of the CM and electronic motion is strong and may require for its proper and efficient description either a different choice for the functions \( \{\psi_q\} \) or an alternative approach (see below) to the expansion (9). Let us evaluate the importance of the coupling terms for different physical situations at laboratory field strengths. We hereby concentrate on the \( He^+ \)-ion moving in a magnetic field. The simplest way to treat the couplings is to assume that the electronic wave functions are to zeroth order well-described by the field-free hydrogenic wave functions and to take into account the electronic diamagnetic interaction via perturbation theory. This description is appropriate for laboratory field strengths for not too high electronic excitations, i.e.

for a field strength of a few Tesla typically up to \( n = 10 - 20 \). The relevant indicator for the mixing of CM and electronic motions is the quotient of the coupling \( \kappa \) and the energy spacing \( \Delta \) of the corresponding diagonal matrix elements in \( (E^L + E^I) \). For electronic states belonging to the same principal quantum number \( n \) (the latter being not too large) i.e.

for the case of dominant intramanifold coupling, we obtain \( 23 \)

\[
(\kappa/\Delta) \propto \frac{\sqrt{N B}}{M} n^2
\]

where \( N \) is the principal quantum number of the Landau orbitals. For \( n = 10 \) and a typical laboratory field strength \( B = 2.35 T \) \( N \) has to be of the order of magnitude \( N \approx 10^8 \) to make the coupling \( \kappa \) as large as the energy spacing \( \Delta \). The corresponding energy of the CM motion is some 10eV. For the case of intermanifold coupling, i.e. the coupling of states belonging to different \( n \)-manifolds we obtain

\[
(\kappa/\Delta) \propto \frac{\sqrt{N B}}{M} B n^5
\]

Choosing \( B = 10 T \) and \( n = 10 \) the requirement that \( \kappa \) should be of the order of magnitude of \( \Delta \) yields \( N \approx 10^{10} \), i.e.

a CM energy of the ion of a few keV. This means that at these energies the couplings become not
only dominant for states within the same $n$ manifold but also important for states belonging to adjacent $n$ manifolds.

The above considerations provide an idea how important the coupling of the CM and electronic motion is for different values of the parameters $n$ (internal energy), $N$ (CM energy), $B$ (field strength). In the case $\frac{B}{n} > 1$ we encounter a strong mixing of the Landau orbitals $\Phi_L^p$ and the electronic functions $\psi_q$. The expansion (9) of an eigenfunction $\Psi$ of $\mathcal{H}$ involves therefore a large number of wave functions $\{\psi_q\}$ which means that the typical spatial extension of $\Psi$ with respect to the electronic coordinates is much larger than that of the individual functions $\psi_q$. A perturbation theoretical approach with respect to the coupling Hamiltonian $\mathcal{H}_2$ is not appropriate in this case. Instead, as already mentioned above, an efficient description of the wave function might either be achieved by a better (intuitive) choice for the functions $\{\psi_q\}$ in the expansion (9) or by pursuing a conceptually different approach (see below).

On the other hand side it is interesting to consider the situation of increasing internal energy $\mathcal{H}_3$ for fixed field strength. It is well-known that the fixed nucleus one-electron atom undergoes a classical transition from regularity to chaos with increasing excitation energy and the quantized atom shows the corresponding quantum signatures of chaos [1]. Perturbation theory with respect to the magnetic interaction terms is only applicable for sufficiently small field strengths/internal energies which corresponds, in a classical language, to the regime for which phase space is dominated by regular structures. Treating the ion in the intermediate case of mixed regular and chaotic classical phase space is a difficult task. However for another limiting case, namely the situation of a completely chaotic phase space (Coulomb- and diamagnetic interaction are of equal strength) a statistical approach to the matrix elements and spectrum of the Hamiltonian $\mathcal{H}_3$ seems appropriate. Such an approach has been developed in ref. [26] and allows to extract important statistical properties for the mixing of the electronic (due to $\mathcal{H}_3$) and CM (due to $\mathcal{H}_1$) wave functions. The Hamiltonian $\mathcal{H}_3$ is hereby represented by a random matrix ensemble, i.e. the Gaussian orthogonal ensemble (GOE), which is the appropriate semiclassical description of a completely chaotic system [29]. The GOE provides the fluctuations of the chaotic levels. The mean level density (MLD) as a function of energy, field strength and in particular electronic angular momentum $L_z$ is a key ingredient for the specification of the random matrix ensemble and can in our case be obtained via the semiclassical Thomas-Fermi formula [26]. It represents the density of irregular states. The size of the matrix
elements of $\mathcal{H}_2$ can be determined from a semiclassical relation between off-diagonal matrix elements of an operator, and the Fourier transform of its classical autocorrelation function \[30\]. For more details on the concrete appearance of the statistical-semiclassical model for the moving ion we refer the reader to ref.\[26\] and report here only on some major results. Of particular interest are the properties of the eigenvectors which are obtained through diagonalization of the total matrix Hamiltonian consisting of the parts $\mathcal{H}_1$, $\mathcal{H}_2$ and $\mathcal{H}_3$. For sufficiently low CM energies we encounter an exponential localization of the components of the eigenvectors around some maximum component. With increasing CM energy the localization length $\lambda$, which reflects the typical length of the mixing process in the space of the quantum states, increases. In ref.\[26\] it was, however, shown that the eigenvectors of this statistical model possess a finite length $L_c$. At some critical CM energy the localization length becomes therefore larger than $L_c$, i.e. larger than the size of the system in the space of the quantum states, and we encounter a crossover from localization to delocalization. We conclude with the remark that there exists an intriguing analogy of a simplified version of the above statistical model with models for transport in disordered finite-size wires whose localization lengths and related properties are known exactly.

3.3 Energy transfer processes for rapidly moving atomic ions

In the present subsection we focus on the situation of a rapidly moving highly excited ion. The corresponding CM energy is much larger than the initial electronic binding energy. Since the coupling of the collective and electronic motion will be large, energy exchange processes between the CM and electronic motion are extremely relevant and provide some intriguing new phenomena \[24, 25, 27, 28\]. The study of both the classical as well as quantum properties provides additional insight into our understanding of the quantization on the classical energy flow.

The classical energy exchange equation for the CM energy $E_{cm}$ and for the internal energy $E_{int} = \frac{\mu}{2} \dot{r}^2 + V$ reads as follows \[24, 25\]

\[
\frac{d}{dt} E_{cm} = -\frac{d}{dt} E_{int} = e\alpha \left( \mathbf{B} \times \mathbf{R} \right) \dot{\mathbf{r}} \quad (15)
\]

This equation shows that a permanent flow of energy from the CM to the electronic degrees of freedom and vice versa has to be expected. Let us consider a typical classical trajectory corresponding to the above-
described situation of a rapidly moving highly excited Rydberg atom. After a transient time of bound oscillations in the internal motion (energy) a strong flow of energy from the CM to the internal motion takes place. The internal energy is hereby increased above the threshold for ionization, $E_{th} = 0$, and the ion eventually ionizes, i.e. the electron escapes in the direction parallel to the magnetic field. Note that the motion of the electron is confined in the direction perpendicular to the magnetic field.

Figure 3 provides a prototype example for such an ionizing trajectory. The subfigures 3a and 3b illustrate the time-dependencies of the CM-energy and the z-component of the internal relative coordinate, respectively. After the above-mentioned initial phase of oscillations there occurs at approximately $T = 7 \times 10^6\text{a.u.}(1.7 \times 10^{-10}\text{s})$ a sudden loss of CM kinetic energy simultaneously accompanied by an increase in the internal energy which causes the electron to move away from the nucleus in the positive z-direction. The transferred energy, which is in our case of figure 3 approximately $6 \times 10^{-3}\text{a.u.}(0.2\text{eV})$, corresponds to a small fraction of the total initial CM energy which is for our example about $12.2677\text{a.u.} (333.8\text{eV})$. This self-ionization process is only possible due to the presence of the coupling term $H_2$ in the Hamiltonian (7) which involves both the internal and CM degrees of freedom. The ionization time for an individual trajectory depends, apart from its intrinsic dynamics, on the field strength and in particular on the CM kinetic energy of the ion.

In order to gain an idea of the statistical measure for the ionization process it is instructive to consider for an ensemble of trajectories the fraction of ionized orbits as a function of time. The initial internal energy is chosen to correspond to a completely chaotic phase space of the $\text{He}^+-$ion if the nuclear mass were infinite. In figure 4 we have illustrated the fraction of ionized orbits as a function of time up to $T = 10^{10}\text{a.u.}$ for a series of different CM energies and for a strong laboratory field strength $B = 23.5T$. For an initial CM energy of $E_{cm} = 0.053\text{a.u.}$ which corresponds to an initial CM velocity of $V_{cm} = 8.4 \times 10^3\text{m/s}$ about 70% of the trajectories are ionized within a time of $T = 10^9\text{a.u.} (2.4 \times 10^{-8}\text{s})$ which is the tenth part of the integration time. In contrast to this we have for $E_{cm} = 0.01\text{a.u.}$ only about 30% of ionized orbits within the total integration time of $T = 10^{10}\text{a.u.} (2.4 \times 10^{-7}\text{s})$. The ionization process depends, therefore, very sensitively on the initial CM kinetic energy of the ion.

The above investigations have shown the existence of the self-ionization
process through energy transfer from the CM to the electronic motion for the classically moving ion in a magnetic field. For the limit of very highly excited electronic states and a large CM energies it is expected that the above described behaviour reflects the true dynamics of the ion. The natural question arises now how quantization changes or enriches this picture for the typical energies accessible experimentally. It is well-known that quantization can alter the effects observed in classical dynamics (see ref. [32] and references therein). Very recently an approach which consists of a combination of semiclassical and wave packet propagation techniques has been developed which is appropriate for the description of the quantum dynamics of the highly excited and rapidly moving quantum ion in a magnetic field. The results of the corresponding investigations [27] show that the quantum self-ionization obeys a time scale which is by orders of magnitude larger than the corresponding classical process. The typical ionization times are for both the classical as well as quantum processes much smaller than the life times of the highly-excited Rydberg states. In addition the ionization signal is seemingly affected by quantum coherence phenomena which yield strong fluctuations for the time-dependence of the ionization rate. More studies have to be performed in order to elucidate the dynamics in the semiclassical and deep quantum regime both theoretically as well as experimentally [31].

Let us comment on a very recent application [28] of the effects of the coupling and mixing of the collective and internal motion in exotic atoms. It was shown that one can stimulate, by using present-day laboratory magnetic fields, transitions between the $lm$ sub-levels of fast $\mu He^+$ ions forming in muon catalyzed fusion. This gives a possibility to drive the population of the $lm$ sub-levels by applying a field of a few Tesla, which affects the reactivation rate of the fusion process and is especially important to its $K\alpha X$-ray production.

To conclude, we have discussed a number of intriguing phenomena for two-body atoms in magnetic fields which have their origin in the nonseparability of the CM and internal motion causing an interaction of the collective and internal motion. It has to be expected that this interaction will induce even more phenomena for multi-electron atoms or molecules. For the case of heteronuclear molecules the relevant coupling Hamiltonian contains both the vibrational and rotational degrees of freedom and new vibrational or rotational structures as well as dynamical processes might therefore arise once the coupling becomes significant i.e. comparable to the spacing of the vibrational or rotational energy levels.
Figure captions

Figure 1: A two-dimensional intersection of the potential $V$ in the plane perpendicular of the magnetic field.

Figure 2: Distribution over energy in the outer well after the preparation is completed. In the calculation a magnetic field strength $B = 14T$ was used. The ionization threshold is shown by a vertical dashed line. The inset shows the spreading of the trajectories over the saddle point into the outer well during the intermediate time step of a constant electric field $E_c$.

Figure 3: (a) The CM energy as function of time. (b) The $z$-component of the internal relative coordinate as a function of time. The total and initial internal energy of the ionizing trajectory are $E = 333.68\text{eV}$ and $E_{int} = -8.16 \times 10^{-2}\text{eV}$, respectively. The field strength is $B = 23.5T$.

Figure 4: The ionized fraction for an ensemble of 250 trajectories as a function of time. From top to bottom the CM-energies belonging to the ionization curves are $E_{CM} = 1.45, 0.63, 0.47, 0.34, 0.27\text{eV}$, respectively. The initial internal energy is always $E_{int} = -9.25 \times 10^{-3}\text{eV}$. The field strength is $B = 23.5T$.

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