Spectroscopy of dipolar fermions in layered two-dimensional and three-dimensional lattices

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Motivated by ongoing measurements at JILA, we calculate the recoil-free spectra of dipolar interacting fermions, for example ultracold heteronuclear molecules, in a one-dimensional lattice of two-dimensional layers or “pancakes,” spectroscopically probing transitions between different internal (e.g., rotational) states. We additionally incorporate $p$-wave interactions and losses, which are important for reactive molecules such as KRb. Moreover, we consider other sources of spectral broadening: interaction-induced quasiparticle lifetimes and the different polarizabilities of the rotational states used for the spectroscopy. Although our main focus is molecules, some of the calculations are also useful for optical lattice atomic clocks. For example, understanding the $p$-wave shifts between identical fermions and small dipolar interactions coming from the excited clock state is necessary to reach future precision goals. Finally, we consider the spectra in a deep three-dimensional lattice and show how they give a great deal of information about static correlation functions, including all the moments of the density correlations between nearby sites. The range of correlations measurable depends on spectroscopic resolution and the dipole moment.

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I. INTRODUCTION

Recoil-free spectroscopy is a powerful, pervasive probe of ultracold atomic systems that includes radiofrequency and microwave spectroscopy, as well as Doppler-free optical or Raman spectroscopy. Interatomic interaction effects on the spectra give information about the phase diagram and behavior of many-body systems. For example, these effects on recoil-free spectra were used to first detect Bose-Einstein condensation (BEC) in dilute spin-polarized atomic hydrogen [1], locate and probe the Mott insulator/superfluid quantum phase transition of bosons in optical lattices [2–6], and study Cooper pair binding [7,8], polaron quasiparticle residue [9], and pseudogap behavior of ultracold fermions across the BEC/BCS crossover [10–12]. In precision measurements, e.g., atomic clocks, understanding the effects of interactions is crucial because they can limit [13] or enhance [14] the achieved accuracy.

Here we calculate the recoil-free spectra of dipolar molecules in a one-dimensional lattice of two-dimensional layers or “pancakes” and in a three-dimensional lattice. Such systems have been realized at JILA [15–19]. As we demonstrate, recoil-free spectroscopy gives information complementary to density profile or time-of-flight measurements. Additional probes are especially important for molecules, where direct imaging is extremely difficult due to the absence of cycling transitions [20]. To be concrete, we consider driving the transition between two rotational states of the molecule, but the same calculations apply to transitions between vibrational, hyperfine, and electronic states when the transitions are measured with techniques imparting negligible recoil momentum.

There is great excitement about ultracold molecules: they offer additional internal degrees of freedom (rotational, vibrational) compared to atoms, and the long-range character of their interactions can lead to interesting many-body physics [21–24]. Examples are quantum liquid crystals [25–27], Wigner crystals [28,29], exotic superfluids [30–32], supersolids [33], topological phases [34–37], and the physics resulting from quantum phase transitions among this plethora of phases [38,39]. We expect recoil-free spectroscopy to play a major role in observing and characterizing these systems.

The outline of our paper is as follows. Section II introduces these systems and derives the Hamiltonian describing them. Section III A gives general expressions, valid even for strongly correlated systems, relating the average spectral shifts to simple static correlation functions. Section III B evaluates these expressions explicitly for weakly interacting homogeneous systems. Section III C applies these expressions to obtain the trapped system’s spectra, while Sec. III D presents examples of results. These calculations are directly relevant for the ongoing experiments at JILA [15–19] and should be useful in a number of related ongoing efforts to make ultracold molecules, e.g., [40].

We quantitatively calculate the spectral shifts and, in Sec. IV B, line shapes due to the dipolar interactions, $p$-wave interactions—including $p$-wave losses that are important for reactive molecules such as KRb—and the differing polarizabilities of the different molecular rotational states that may be used in the spectroscopy. We also qualitatively consider and bound other effects, such as higher lattice band occupation and the validity of treating the atoms in the cold collision regime, i.e., treating interactions via a pseudopotential.

To briefly motivate the utility of these spectra, we point out that measuring them in the dilute thermal gas allows one to ensure that these various nontrivial contributions—many of which are just beginning to be explored in the context of cold atoms—behave as expected. Furthermore, deviations from the simple dilute thermal gas values allow one to diagnose more interesting behavior as the temperature is lowered, the simplest example probably being the degree of quantum degeneracy of the gas. The onset of strongly correlated quantum phases will also have signatures.

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As another application of recoil-free spectra to dipolar interacting systems, Sec. V considers a deep three-dimensional (3D) lattice. We show that one can measure the joint probability distribution $P(1,n_1,n_2,\ldots)$ of a particle occupying a site while having $n_1$ molecules in the nearest-neighbor sites, $n_2$ in the next-nearest-neighbor sites, and so on. This includes all moments $\langle n_1^{a_1}n_2^{a_2}\cdots \rangle$ for arbitrary $a_j$. The spatial range of correlations that can be measured is set by the spectral resolution and interaction strength, and is typically a few sites for the molecules presently considered. This provides a powerful way to characterize the system, measuring a much more complete set of correlations and with greater spatial range than other techniques, such as modulation spectroscopy [41].

While our quantitative estimates of parameters are for dipolar molecules, many of the calculations are also relevant for Rydberg atoms and alkaline-earth-metal atomic clock experiments. As one example, Rydberg atoms have long-range dipolar interactions. As another, $p$-wave interactions can be important in optical atomic clocks, as recently revealed by Ramsey spectroscopy of Yb lattice clocks with the atoms trapped in 1D arrays of 2D pancakes or 2D arrays of 1D tubes [42]. Moreover, as experiments in these clocks expand their capabilities to approach expected $10^{-18}$ accuracy and mHz frequency resolution, small dipolar interactions between the atoms become relevant. We expect that our calculations may provide insight into these experiments. However, we mention that while our theory assumes that few atoms are transferred to the excited state (i.e., linear response), many clock interrogation techniques transfer a substantial fraction of the atoms [43,44].

II. THEORETICAL AND EXPERIMENTAL BACKGROUND

We are interested in heteronuclear molecules in a one-dimensional optical lattice of two-dimensional pancakes in the presence of an electric field. We assume that the energy scale set by this field is large compared to the hyperfine splitting; we explain this in more detail below. Figure 1 illustrates this system and defines notation. Section V considers the effects of adding a lattice to the transverse directions. This system displays a hierarchy of energy scales, which dictates the physical behavior. We begin by describing this hierarchy.

![Figure 1](image1.png)

**FIG. 1.** (Color online) Schematic of the experimental geometry considered. Heteronuclear dipolar molecules are trapped in a one-dimensional lattice of two-dimensional pancakes, with the lattice periodic in the $z$ direction. The angle $\theta$ is the angle between the intermolecule separation $r-r'$ and the molecular dipole (applied electric field axis), here taken to be along the $z$ axis.

![Figure 2](image2.png)

**FIG. 2.** (Color online) Lowest rotational energy levels of a single molecule in an electric field. These are described by the rigid rotor Hamiltonian in an electric field $H = B N^2 - d \cdot E$, with $N$ the angular momentum operator, $B$ the rotational constant, and $d$ the dipole moment along $z$. The label $(M,m_z)$ indicates the state that is adiabatically connected to the $E=0$ state with angular momentum $M$ and angular momentum $z$-component projection $m_z$. However, the $E$ field mixes states with the same $m_z$ but with different $M$, so $M$ is not a good quantum number. The energy splitting $\nu$ increases with increasing electric field and is a large energy scale even for small $E$: interactions, the trapping potential, and the kinetic energy cannot mix in states that are off-resonant by this energy scale.

Figure 2 shows the lowest energy rotational levels of a molecule in a dc electric field. This structure is well-described by the rigid rotor Hamiltonian in an electric field, $H = B N^2 - d \cdot E$, with $N$ the angular momentum operator, $B$ the rotational constant, and $d$ the dipole moment operator. This single-molecule physics sets the largest energy scales in the problem. In the absence of an electric field, there is the usual rotational level structure: a ground state, a threefold degenerate spin-1 first excited state manifold, a fivefold degenerate spin-2 second excited state manifold, etc. These are labeled by their total angular momentum $M$ and $z$-axis projection $m_z$. In the presence of an electric field $E$, the rotational symmetry is broken. Here we take $E$ to be along the $z$ direction, perpendicular to the pancakes, as is used in the ongoing KRb experiments to suppress the reactive losses [15]. The electric field mixes states with different $M$ but the same $m_z$.

Figure 2 shows the resulting level structure for a nonzero value of $E$, where we label by $|M,m_z\rangle$ the state that is adiabatically connected to the $E=0$ state with angular momentum $M$ and $z$ projection $m_z$. An electric field induces a level splitting $\nu$ in the previously degenerate excited state manifold, and this energy scale is large compared to the other energy scales that will be of interest. As a consequence, when we treat the interactions, lattice, and trap, it is quantitatively accurate to restrict oneself to processes occurring in the resonant subspace where we neglect transitions that require energy changes on the order of $\nu$.

We now present the Hamiltonian describing the period lattice, the harmonic confining potential, and intermolecular interactions for molecules in an electric field. We consider two molecular states, choosing $|0,0\rangle$ and $|1,0\rangle$ to be concrete, but the vast majority of our calculation is general and
straightforwardly extended to arbitrary internal states. We will denote these states as $|\alpha\rangle$ with $\alpha = 1, 2$ henceforth. Only the quantitative numbers presented depend on the choice of these states.

The hyperfine coupling is the next largest energy scale after the rotational and electric field, and can couple the $\alpha = 1, 2$ and other rotational states, shown in Fig. 2. However, even a small electric field—one that generates dipole moments more than an order of magnitude smaller than the permanent dipole moment—suffices to generate level splittings $\sim\hbar$ that are larger than the hyperfine coupling. Thus for even such small electric fields, if the initial populations are confined to these two rotational states, we can neglect the other states, which are then far off-resonant compared to the hyperfine energy.

The lattice energy scale is the largest after the rotational, electric field, and hyperfine structure of the molecules, all set by single-molecule physics. The lattice separates the energy levels into bands, and because the other energy scales—trap, interactions, and temperature—are small relative to the band splittings for the deep lattices of present interest, we can thus project the Hamiltonian to the lowest lattice band for states $\alpha = 1, 2$. We neglect terms that are exponentially small in the lattice depth (e.g., tunneling terms). The molecular system is described by the Hamiltonian

$$H = H_0 + H_{\text{sw}} + H_{\text{dipole}} + H_{\text{pw}} + H_{\text{Isap}}$$

with the noninteracting part

$$H_0 = \sum_{i,\alpha} \int d\rho \, \psi_{i,\alpha}^\dagger(\mathbf{r}) \left(-\frac{\nabla^2}{2m} - \mu\right) \psi_{i,\alpha}(\mathbf{r}),$$

where $\psi_{i,\alpha}(\mathbf{r})$ and $\psi_{i,\alpha}^\dagger(\mathbf{r})$ are fermionic annihilation and creation operators acting on atoms in rotational state $\alpha$ at site $i$ in the lowest band at the transverse two-dimensional position $\mathbf{r}$. They satisfy the anticommutation relationships $\{\psi_{i,\alpha}(\mathbf{r}), \psi_{j,\beta}^\dagger(\mathbf{r}')\} = \delta_{ij} \delta_{\alpha\beta} \delta(\mathbf{r} - \mathbf{r}')$. Here $m$ is the molecular mass and $\mu$ is the chemical potential.

The projected s-wave interaction Hamiltonian is

$$H_{\text{sw}} = \sum_{i,\alpha,\beta} \frac{g_{\text{sw}}}{2} \int d\rho \, \psi_{i,\alpha}^\dagger(\mathbf{r}) \psi_{i,\alpha}^\dagger(\mathbf{r}) \psi_{i,\alpha}(\mathbf{r}) \psi_{i,\alpha}(\mathbf{r})$$

with $g_{\text{sw}} = (4\pi a_{\text{sw}}/m) \int dz \, [w_{\alpha}(z)^2 |w_{\alpha}(z)^2]$ (we set $\hbar = k_B = 1$ throughout), where $a_{\text{sw}}$ is the s-wave scattering length for scattering between molecules in states $\alpha$ and $\beta$, and $w_{\alpha}(z)$ is the lowest band Wannier function for particles in state $\alpha$.

This depends on $\alpha$ because the lattice depths felt by each species need not be equal since the molecular polarizabilities differ. The interaction $g_{11}$ is undefined for identical fermions, which do not scatter in the s-wave channel, and so we can set it to zero, but $g_{12}$ is nonzero. When the Wannier functions for the two states are not identical, the three-dimensional p-wave interaction projected onto the lowest band gives a renormalization of the two-dimensional 1-2 s-wave interaction (not included above) in addition to the contribution coming from the three-dimensional s-wave scattering, because the particles effectively become distinguishable. However, the s-wave interaction will turn out to be irrelevant for the recoil-free spectra in the linear-response regime with all molecules initially in the ground internal state, as a consequence of the Pauli exclusion principle. Hence we can ignore the s-wave contact interaction.

The dipolar interaction Hamiltonian in 3D is $[45, 46]$

$$V_{\text{dip}}^{(3D)} = \int d\mathbf{r} d\mathbf{r}' \frac{1 - 3 \cos^2 \theta}{4\pi e_0 |\mathbf{r} - \mathbf{r}'|^3}$$

$$\times d_{12}d_{21} \psi_{i,\alpha}^\dagger(\mathbf{r}) \psi_{i,\beta}^\dagger(\mathbf{r}) \psi_{i,\alpha}(\mathbf{r}) \psi_{i,\beta}(\mathbf{r}) + \text{H.c.}$$

$$+ \sum_{m,n} d_{mn} d_{nm} \psi_{m}^\dagger(\mathbf{r}) \psi_{n}^\dagger(\mathbf{r}) \psi_{m}(\mathbf{r}) \psi_{n}(\mathbf{r}).$$

where $\theta$ is the angle between the dipole orientation direction and $\mathbf{r} - \mathbf{r}'$, $d_{mn} = (m|d^{(0)}|n)$, where $d^{(0)}$ is the $m_z = 0$ spherical component of the dipole operator in the basis aligned with the electric field, and $\psi_{m}(\mathbf{r})$ and $\psi_{m}^\dagger(\mathbf{r})$ are fermionic annihilation and creation operators satisfying the anticommutation relations $\{\psi_{m}(\mathbf{r}), \psi_{n}^\dagger(\mathbf{r}')\} = \delta_{mn} \delta(\mathbf{r} - \mathbf{r}')$. The reason why only the $m = 0$ spherical component of the dipole operator is retained is that we are considering the transition between $m_z = 0$ levels and even a small electric field drives the $m = \pm 1$ levels off-resonance, as shown in Fig. 2. For other transitions, the coefficients would be more complicated $[46]$. To be sufficiently off-resonant compared to the other energy scales—interactions, lattice, and hyperfine coupling (to allow one to neglect the nuclear degrees of freedom)—requires only a small field, where the induced dipole moment is still an order of magnitude or more smaller than the permanent dipole moment. The same reason justifies why only the shown matrix elements of the dipole interaction are kept—the other matrix elements leave the resonant subspace. Note that the permanent dipole moments of these levels are nonvanishing, in contrast to the zero-field rotational levels, because the electric field induces a dipole moment in the eigenstates.

The projected dipolar interaction Hamiltonian is

$$H_{\text{dipole}} = \frac{1}{2} \sum_{i,\alpha,\beta} \int d\rho d\rho' V_{\text{dip}}(i - j, \rho - \rho')$$

$$\times \psi_{i,\alpha}(\rho) \psi_{j,\beta}^\dagger(\rho') \psi_{j,\beta}(\rho') \psi_{i,\alpha}(\rho)$$

$$+ \frac{1}{2} \sum_{ij} \int d\rho d\rho' V_{\text{dip}}(i - j, \rho - \rho')$$

$$\times [\psi_{i,\alpha}(\rho) \psi_{j,\beta}^\dagger(\rho') \psi_{j,\beta}(\rho') \psi_{i,\alpha}(\rho) + \text{H.c.}]$$

with

$$V_{\text{dip}}(i - j, \rho - \rho') = \int d\mathbf{z} d\mathbf{z}'$$

$$\times d_{12}d_{21} \frac{(1 - 3 \cos^2 \theta)}{4\pi e_0 |(\rho - \rho')^2 + (z - z')^2|^{3/2}}$$

$$\times [w_{\alpha}(z - z_i)^2 |w_{\beta}(z' - z_j)^2]^{1/2}$$

and

$$V_{\text{dip}}(i - j, \rho - \rho')$$

$$= \int d\mathbf{z} d\mathbf{z}'$$

$$\times d_{12}d_{21} \frac{(1 - 3 \cos^2 \theta)}{4\pi e_0 |(\rho - \rho')^2 + (z - z')^2|^{3/2}}$$

$$\times w_{\alpha}(z - z_i) w_{\beta}(z - z_i) w_{\alpha}(z' - z_j) w_{\beta}(z' - z_j).$$

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We refer to the former term as the “direct” term and the second as the “spin-flip” term. We approximate \( V_{af}(j, \rho) \) by
\[
V_{af}(j, \rho) \approx \gamma_{af} U(j, \rho)
\]
and \( V_d \) by
\[
V_d(j, \rho) \approx \eta U(j, \rho),
\]
with
\[
U(j, \rho) = \delta_{0j} \frac{1}{\rho^3 + A} + (1 - \delta_{0j}) \left( 1 - \frac{3 (i |d\rho|^2)^2}{|\rho^2 + (j d\rho)^2|^{3/2}} \right),
\]
where
\[
\gamma_{af} = \frac{d_{\alpha\beta} d_{\beta\rho}}{4 \pi \epsilon_0}, \quad \eta = \frac{|\int dz w_1^*(z) w_2(z) d_j(z)/(4 \pi \epsilon_0)|}{d_j}, \quad \text{and} \quad d_j = \text{the lattice spacing},
\]
and \( d_j \) is the lattice spacing, with \( \epsilon \) the width of the Wannier functions, governs the short-range physics. More details on this approximation follow shortly. In principle, we should let \( \Lambda \) be different for the \( V_d \) and \( W_{af} \) terms, but we will argue that we can set \( \Lambda = 0 \) and maintain quantitative accuracy in the calculated spectra.

We also note that the factor of \( F = |\int dz w_1^*(z) w_2(z) d_j(z)\rangle^2 \) appearing in \( \eta \) is almost always nearly unity in a deep lattice. We denote the polarizability of state 2 by \( (1 + \xi) \) times the polarizability of state 1. In a deep lattice, \( w_{\alpha}(z) \) can be approximated by a Gaussian. Then one finds \( F = \sqrt{\frac{\pi}{2\xi}}(1 + \xi)^{1/4}. \) For \( \xi = 0.4 \) polarization difference, one finds \( F = 0.99. \)

At long distances, these approximate interaction potentials are exact for dipoles aligned by a (perhaps very small) electric field, projecting the dipolar interaction into the lowest band, but at short distances they are just a convenient approximation. The form is chosen so that at short distances the potential goes to a constant, as does the true projected potential, and approximately interpolates between the exact long-distance and zero-distance forms. In fact, we will set \( \Lambda = 0. \) Although for some calculations this can lead to divergent results, everything in this paper remains finite in the \( \Lambda \to 0 \) limit, and comparing the \( \Lambda \to 0 \) limit with alternative calculations using finite realistic values of \( \Lambda \) (not presented) shows excellent agreement. The fundamental reason for this is that at low temperature and for a dilute gas, the physics is fairly insensitive to the short-range physics governed by \( \Lambda. \)

The p-wave interaction Hamiltonian is (the multi-internal state generalization of [47], as given in Ref. [48])
\[
H_{pm} = H_{(11)}^\alpha + H_{(12)}^\alpha + H_{(21)}^\alpha,
\]
where
\[
H_{(11)}^\alpha = u_{af} W_{af} \sum_i \int d\rho \left[ \nabla_\rho \psi^\dagger_\alpha(i, \rho) \right] \psi^\dagger_\alpha(i, \rho) - \psi^\dagger_\alpha(i, \rho) \nabla_\rho \psi^\dagger_\alpha(i, \rho) - \psi_\alpha(i, \rho) \nabla_\rho \psi^\dagger_\alpha(i, \rho)
\]
with \( u_{af} = 6 \pi \hbar^2 \tilde{b}_d^3 / m, \) \( W_{af} = \int dz |w_{\alpha}(z)|^2 |w_{\beta}(z)|^2, \) and \( \tilde{b}_d \) is the p-wave scattering length between states \( \alpha \) and \( \beta. \) In asymptotically deep lattices, the Wannier functions are approximately Gaussian, and \( W_{af} \) simplifies to
\[
W_{af} \approx \frac{4 \pi}{\sqrt{3}} \left( \frac{\sqrt{3}}{\sqrt{\gamma_1}} + \frac{\sqrt{3}}{\sqrt{\gamma_2}} \right)^{-1/2} \langle \tilde{V}_d \rangle \text{ the lattice depth in recoil units for state } \alpha. \] 
If the lattice potentials for \( \alpha \) and \( \beta \) are the same, then \( W_{af} \approx (1/d) \sqrt{\pi/2} \). Alternatively, the p-wave interaction could be written by adding a term to the intermolecular potential \( V_{af} \) in Eq. (5) of the form
\[
(2\pi \hbar^2 / m) \nabla_\rho \delta(\rho) \frac{1}{\pi} \overline{V_d} \bar{V}_d \int dz |w_{\alpha}(z)|^2 |w_{\beta}(z)|^2 \] [47], where an arrow to the left (right) indicates that the gradient operates to functions on the left (right).

There is also a confining potential
\[
H_{\text{trap}} = \sum_i \int d\rho V_\alpha(i, \rho) \psi^\dagger_\alpha(i, \rho) \psi_\alpha(i, \rho),
\]
which we neglect in our first computation of the average spectral shifts for the homogeneous system in Secs. III A and III B. Its effects will be included with a local density approximation in Sec. III C.

One measures the recoil-free spectrum by applying a probe pulse for a short time, where the probe pulse is described by the Hamiltonian
\[
H_\rho = \Omega \sum_j \int d\rho e^{-i\omega t} \psi_{\rho 1}(j, \rho) \psi_{\rho 2}(j, \rho) + \text{h.c.}
\]
in the rotating-wave approximation, with \( \omega \) the probe frequency and \( \Omega \) proportional to the amplitude of the probe field. The spectrum is given by counting the number of atoms transferred to state 2 as a function of \( \omega \) for a fixed probe time. As an aside, we note that in atomic clock experiments, the small variations of \( \Omega \) with the trap quantum number give rise to a shift even for s-wave interacting—initially identical—fermions [43,44]. This effect is quite tiny, coming from the differences in Rabi frequencies associated with each trap mode: they only manifest in clocks because of the clock experiments’ extraordinary frequency resolution (~1 Hz). The effect is even smaller for current experiments: the variation of this Rabi frequency for radiofrequency or microwave pulses is orders of magnitude smaller, and so the resulting shift will be much smaller than a Hertz. This is completely negligible in the current experiments.

Finally, we comment that we calculate all frequency shifts relative to the level splitting of the internal states (e.g., rotational levels) 1 and 2 of a single molecule, \( \delta_v. \) The experimentally measured spectra will then all be shifted by this value.

III. SPECTRAL SHIFT: SUM RULES

A. General expressions for the homogeneous gas

The spectral weight for the recoil-free spectrum is the number of particles transferred from the initial internal state in a time \( t, \) as defined above. Fermi’s golden rule gives
\[
\mathcal{I}(\omega) = \frac{2\pi t}{h} \sum_{i,f} p_i \langle f | H_\rho | i \rangle^2 \delta(\omega - E_f + E_i),
\]
where the sum over \( i \) and \( f \) runs over all possible energy eigenstates of \( H, \) \( p_i \) is the initial probability of being in state \( i, \) and \( E_j \) is the energy of state \( j \) for the Hamiltonian \( H. \)

The average spectral frequency
\[
\langle \omega \rangle = \frac{\int d\omega \omega \mathcal{I}(\omega)}{\int d\omega \mathcal{I}(\omega)}
\]

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is [49,50]

\[ \langle \omega \rangle = \frac{\langle [H_p, H] H_p \rangle}{|H_p|^2}, \]  

(17)

where the expectation includes both the thermal and quantum averages. This is an exact result (within linear response) relating \( \langle \omega \rangle \) to the expectation value of an operator that can simply be evaluated from the definitions of \( H_p \) and \( H \), for any Hamiltonian, and holds even for strongly correlated systems.

We evaluate the commutators in Eq. (17) for each interaction term in \( H \) in turn. The spectral shift from \( s \)-wave interactions vanishes.

The spectral shift from the direct and spin-flip terms of the dipolar potential is

\[ \langle \omega_d \rangle = n \sum_j \int d\rho [V_{12}(j, \rho) - V_{11}(j, \rho) + V_{s2}(j, \rho)] g_2(j, \rho), \]  

(18)

with

\[ n = \langle \psi_i^\dagger(0,0) \psi_i(0,0) \rangle \]  

(19)

the two-dimensional density and

\[ g_2(i, \rho) = \frac{1}{n^2} \langle \psi_i^\dagger(i, \rho) \psi_i^\dagger(0,0) \psi_i(0,0) \psi_i(i, \rho) \rangle. \]  

(20)

This holds for any two-body potential, and one sees that it vanishes for an \( s \)-wave contact interaction because \( \langle \psi_i^\dagger(0) \psi_i^\dagger(0) \psi_i(0) \psi_i(0) \rangle = 0 \). The key qualitative features of this result are understood by mean-field theory: the expression, up to constant factors in each term, is the mean-field interaction energy of the state with one molecular state changed at position \( \rho \), summed over \( \rho \), giving \( V_{12} \) and \( V_{s2} \) terms minus the state with all the molecules in the same initial state (\( V_{11} \) term).

The spectral shift for the \( p \)-wave potential is

\[ \langle \omega_{pw} \rangle = \frac{8(u_{12}W_{12} - u_{11}W_{11})}{n} \times \langle [\nabla \psi_i^\dagger(0, \rho)] n_1(0, \rho) \cdot [\nabla \psi_i(0, \rho)] \rangle \]  

(21)

with \( n_1(j, \rho) = \psi_i^\dagger(j, \rho) \psi_i(j, \rho) \). The total average spectral shift is \( \langle \omega \rangle = \langle \omega_d \rangle + \langle \omega_{pw} \rangle \).

### B. Weakly interacting homogeneous gas

Equations (18) and (21) give exact, general relations for the spectral shift, valid in any system. Here we simplify these for weakly interacting systems. In the limit in which the interactions induce negligible correlations, the system’s correlations are that of a free Fermi gas. In other words, since we want to know the shifts to lowest order in the interactions, and the interaction already appears multiplying the operator expectation values, we may evaluate these expectation values to zeroth order in the interaction. Consequently, Wick’s theorem determines the expectation values.

The spectral width of a homogeneous gas narrows as the interactions decrease, as we show in Sec. IV B. For experiments considered in the present paper, the spectral width is very small and the spectrum is nearly a perfect \( \delta \) function.

In this case, the sum rule gives the exact location of the well-defined spectral peak.

#### 1. Expressions at general temperature

For the dipolar interaction, we first compute \( g_2 \). Wick’s theorem gives

\[ g_2(i, \rho) = 1 - \left( \frac{G(i, \rho)}{n} \right)^2, \]  

(22)

where \( G(i, \rho) = \langle \psi_i^\dagger(i, \rho) \psi_i(0, 0) \rangle \). Because there is no tunneling, the Green’s function has the form

\[ G(i, \rho) = \delta_{i,0} \tilde{G}(\rho) \]  

(23)

with

\[ \tilde{G}(\rho) = \frac{1}{(2\pi)^2} \int dk e^{ik \cdot \rho} \langle \tilde{n}_k \rangle, \]  

(24)

where the integral runs over all the transverse momentum states of the system (which, recall, is assumed in this section to be homogeneous).

The expectation values are, from Eq. (19),

\[ n = \frac{1}{2\pi} \int dk dk' \frac{1}{e^{(k^2/2m) - \mu} + 1} = \frac{m}{2\pi \beta} \ln \left( 1 + e^{\beta \mu} \right) \]  

(25)

and, from Eq. (24),

\[ \tilde{G}(\rho) = \frac{1}{2\pi} \int dk \frac{J_0(k \rho)}{e^{(k^2/2m) - \mu} + 1} \]  

(26)

with \( J_0 \) the zeroth-order Bessel function of the first kind. Equations (18), (22), and (26) give the dipolar shift.

For the \( p \)-wave interaction, Wick’s theorem gives

\[ \langle \omega_{pw} \rangle = \frac{8}{n} \left( u_{12}W_{12} - u_{11}W_{11} \right) \times \langle [\nabla \psi_i^\dagger(0, \rho)] n_1(0, \rho) \cdot [\nabla \psi_i(0, \rho)] \rangle \]  

\[ = \frac{8}{2\pi} \left( u_{12}W_{12} - u_{11}W_{11} \right) \times \int_0^\infty dk \frac{k^3}{e^{(k^2/2m) - \mu} + 1} \]  

\[ = -\frac{8m^2 T^2}{\pi} \left( u_{12}W_{12} - u_{11}W_{11} \right) L_2(-e^{\beta \mu}), \]  

(27)

where \( L_2 \) is the polylog function.

#### 2. Thermal (Boltzmann) and degenerate gas limits

In this section, we evaluate the shifts, given above for a weakly interacting system at arbitrary temperature, in the thermal/Boltzmann \( (T \gg \mu) \) and degenerate \( (T \ll \mu) \) limits.

In the thermal limit \( T \gg \mu \), the dipolar shift, given by Eqs. (18), (22), and (26), taking \( A = 0 \), is

\[ \langle \omega_d \rangle = n (\gamma_{12} - \gamma_{11} + \eta) \left[ \int d\rho \frac{1 - e^{-mp^2/\beta}}{\rho^3} \right. \right. \]  

\[ + 2 \sum_{j=1}^{\infty} \int d\rho \frac{1 - 3(jd_j^2)^2}{\rho^3 + (jd_j^2)^{1/2}} \]  

\[ = 2\pi^{3/2} \frac{m}{\beta^{3/2}} (\gamma_{12} - \gamma_{11} + \eta) \]  

(thermal),

(28)
where \( n = [m/(2\pi \beta)]e^{\beta \mu} \). Note that the integrals in the sum over \( j \) vanish. The physical reason for this is that these integrate the electric field generated by the zeroth pancake over the plane of pancake \( j \). Closing this planar surface at infinity, the total electric flux penetrating the surface must be zero. Since the dipolar field decays as \( 1/r^3 \), the contribution from the surface at infinity is negligible, the electric flux through the plane must be zero, and hence the integral vanishes.

In the degenerate limit \( T \ll \mu \), the dipolar shift is

\[
\langle \omega_d \rangle = n(\gamma_{12} - \gamma_{11} + \eta) \left[ \int d\rho \left( 1 - 4 \frac{[(k_x \rho)]^2}{\rho^3} \right) \right]
\]

\[
= \frac{256}{45} n k_F (\gamma_{12} - \gamma_{11} + \eta) \quad \text{(degenerate)} \quad \text{(29)}
\]

with \( k_F \equiv \sqrt{2m\mu} \) the Fermi wave vector and \( n = k_F^2/(4\pi) \) in this limit.

The \( p \)-wave shift, Eq. (27), simplifies in the thermal limit \( T \gg \mu \) gas to

\[
\langle \omega_{pw} \rangle = \frac{8m\pi}{T} \int n(u_{12} W_{12} - u_{11} W_{11}) \quad \text{(thermal)} \quad \text{(30)}
\]

and in the degenerate limit \( T \ll \mu \) gas to

\[
\langle \omega_{pw} \rangle = \frac{k_F^4}{\pi} (u_{12} W_{12} - u_{11} W_{11}) \quad \text{(degenerate)} \quad \text{(31)}
\]

### C. Spectra in a trap

In this section, we calculate the spectra of systems in a harmonic trap. We use the local density approximation (LDA): we approximate the properties at a position \((i,\rho)\) in the trap by those of a homogeneous system at an effective chemical potential \( \mu_{\text{eff}}(j,\rho) = \mu_0 - (m\omega^2/2)(jdj)^2 + \rho^2 \) with \( \mu_0 \) the global chemical potential of the system and \( \omega \) the harmonic-oscillator potential’s trapping frequency, assuming an isotropic trap. As usual, the LDA is accurate only when the local chemical potential variation over the correlation lengths is sufficiently small [51]. This condition is met in the experiments of present interest. There is an additional requirement for the accuracy of the LDA, because we are considering a rather nonlocal quantity, the shifts from the dipolar interaction energy. For this quantity, the accuracy of the LDA requires that the spatial variation of the local chemical potential is sufficiently slow compared to the length over which the dipolar interaction becomes negligible. Here, the accuracy of local quantities in a spatially varying system will not be expected to converge exponentially, but rather the error \( \epsilon \) scales as a power law \( \epsilon \sim n \int L^2(2\pi \rho) d\rho \ll n/L \) in the scale \( L \) over which \( \mu \) varies (\( L \) is on the order of the cloud width). Meanwhile, as seen in Eq. (18), the contributions to the shift scale as \( n/\ell \) with \( \ell \) a characteristic microscopic scale (thermal de Broglie wavelength in the thermal gas or Fermi wavelength in the degenerate gas). Since (for our trap parameters) \( L \) is typically much larger than tens of micrometers, while \( \ell \) is typically a tenth of a micrometer, we expect this to be quite accurate. However, it will lead to quantitative corrections at the \( \lesssim 1\% \) level. Consequently the LDA is a good approximation for these systems.

Within the LDA, in the limit that the homogeneous system’s spectrum is a \( \delta \) function (validated in Sec. IV for current, weakly interacting systems), the trap summed spectrum is

\[
I(\omega) = 2\pi \Omega^2 \int \left[ \int d\rho n[\mu_{\text{eff}}(i,\rho)] \delta[\omega - \omega_{\mu_{\text{eff}}}(i,\rho)] \right].
\]

(32)

We can generally numerically sample the integral, approximating the \( \delta \) function as a sharply peaked Lorentzian of width \( \Gamma \), \( \delta(x) \to \delta_{\Gamma}(x) \), with \( \delta_{\Gamma}(x) \equiv (1/\pi)^{1/2}(x^2 + \Gamma^2) \). In some cases, however, an analytic formula can be derived, as we now discuss.

The spectral weight in the frequency window \((\omega,\omega + d\omega)\), given by \( I(\omega) d\omega \), is \( 2\pi \Omega^2 \Gamma \) times the number of atoms with spectral shifts in that window:

\[
I(\omega) = (2\pi \Omega^2 \Gamma) 4\pi [\rho(\omega)]^2 n(\omega) |d\rho/d\omega| \quad \text{(33)}
\]

where \( \rho(\omega) \) is the distance to the trap center of atoms that have spectral shift equal to \( \omega \). Within the LDA, \( \omega \) as a function of \( r \) is inverted to give \( \rho(\omega) \) and one similarly can calculate \( d\rho/dr \).

In the thermal gas, we obtain analytic formulas. Note that in this regime, the sole chemical potential dependence of both the \( p \)-wave and dipolar interactions comes from the density, which multiplies a term independent of the chemical potential: \( \langle \omega \rangle = Cn \) for some constant \( C \). Moreover, in this limit, \( n(r) = [m/(2\pi \beta)]e^{\beta \mu(r)} \). Then \( \rho(\omega) = (1/\omega_0)^{1/2} [(m/\beta)]^{1/2} [\beta \mu_0 - \ln(2\pi \beta \mu_0/(Cm))] \) \( |d\rho/d\omega| = \{2m[\beta \mu_0 - \ln(2\pi \beta \mu_0/(Cm))] \}^{-1} \). Thus the trap summed spectrum of the thermal gas is

\[
I(\omega) \propto \frac{\beta \mu_0 - \ln(2\pi \beta \mu_0/(Cm))}{\beta \mu_0} \quad \text{(34)}
\]

for \( \omega \) such that the term under the square root is real and positive, and zero otherwise.

No simple expression exists for the case of general temperature, nor does one exist even if the gas is deeply degenerate, although we mention that such formulas do exist for the deeply degenerate gas if either the \( p \)-wave or dipolar shift is included alone. For a purely dipolar gas in the deeply degenerate regime \((T = 0)\), one finds

\[
I(\omega) \propto \omega^{1/3} \left[ \frac{\mu_0}{\omega_{\text{deg}}} \left( \frac{\sqrt{2}\pi \omega}{\omega_{\text{deg}}} \right)^{-2/3} \right] \quad \text{(35)}
\]

with \( \omega_{\text{deg}} \equiv \{m^3/45(\gamma_{12} - \gamma_{11} + \eta)^2\}^{-1} \).

### D. Example spectra

Figure 3(a) shows an example spectrum for a thermal gas calculated by numerically integrating the LDA expression Eq. (32) with the dipolar shifts Eq. (28), for a finite-size system of 200 pancakes (although only \( \approx 30 \) of these are significantly occupied) and trapping frequencies and temperatures roughly corresponding to ongoing JILA experiments: \( \omega_i = 2\pi \times 100 \text{ Hz} \) and \( T = 800 \text{ nK} \), with 2200 particles in the central tube. In fact, the JILA traps are quite anisotropic, but, since our goal here is not to construct detailed models...
SPECTROSCOPY OF DIPOLEAR FERMIONS IN LAYERED...

FIG. 3. (Color online) (a) Recoil-free spectrum with a rough modeling of the JILA experimental parameters (see text). Here \( N_e(\omega)/N_e(\omega_{\text{max}}) \) is the number of excited particles after a probe pulse of frequency \( \omega \) and duration \( t \) rescaled by the peak’s maximum value. The overall frequency scale, related to the dipolar interaction strengths, is \( \omega_{\text{scale}} = (\gamma_{12} - \gamma_1 + \eta) \sqrt{\frac{\pi m}{2\hbar^2}} \). The shape of this spectrum is entirely determined by the density profile. The infinite system has a logarithmically divergent peak at \( \omega = 0 \), but this is rounded off by a small spectral broadening \( \Gamma = 7 \times 10^{-4} \omega_{\text{scale}} \) added to the calculation here and, less so, by the finiteness of the present system, assumed to be 200 lattice sites wide. (b) Recoil-free spectrum at \( T = 0 \) (deeply degenerate) with \( \omega_{\text{deg}} \equiv (m^{3/2} \frac{\hbar}{\omega_{\text{scale}}})^{1/5} \), where we have chosen \( \hbar = 2.6\omega_{\text{deg}} \); different \( \hbar = \hbar_{\text{deg}} \)'s just rescale the plot.

of particular experiments, an isotropic trap gives an adequate caricature. Figure 3(b) shows the spectrum at \( T = 0 \), showing how the spectral shape changes when the gas becomes deeply degenerate. In contrast to the asymmetric thermal cloud with its small-\( \omega \) logarithmic divergence, a more symmetric, broad peak forms. At intermediate temperatures, the spectrum looks roughly like a sum of these two spectra: there is a broad spectral peak from the degenerate portion of the trap and a small-\( \omega \) logarithmic divergence. This shows how the spectra may be used to assess the degeneracy of the gas.

The structure shown in Fig. 3 is quite general. A trapped system’s spectra generically show two characteristic features: a logarithmic divergence at low frequency due to thermal tails and a broad structure when the gas becomes strongly interacting and/or degenerate [52].

Such a structure has also been seen in other systems, such as dilute spin-polarized atomic hydrogen [1].

The dipolar energy scale \( \omega_{\text{scale}} = \omega_{\text{dir}} + \omega_{\text{sf}} \), with \( \omega_{\text{dir}} \equiv (\gamma_{12} - \gamma_1) \sqrt{\frac{\pi m}{2\hbar^2}} \) and \( \omega_{\text{sf}} \equiv \eta \sqrt{\frac{\pi m}{2\hbar^2}} \), depends on the rotational states being used and the external electric field applied. For concreteness, we estimate the shift in the transition from the ground rotational state \( |0,0\rangle \) to the excited rotational state \( |1,0\rangle \). The magnitudes of shifts for other low-lying rotational transitions are generically the same order of magnitude.

Figure 4 plots the energy scale \( \omega_{\text{scale}} \) versus the dipole moment in units of the permanent dipole moment. The permanent dipole moment for KRb is about 0.5 D. We calculate the dipolar interaction and \( \omega_{\text{scale}} \) by computing the dipole moments of these rotational states using a quantum rigid rotor model in an electric field with parameters given in Ref. [53]. We observe that this energy scale can be greater than 50 kHz. Then we see from Fig. 3(a) that even for the low densities of the JILA experiments, the width of the spectra is as large as about 5 kHz. Experimentally resolving such scales is standard practice.

The real part of the \( p \)-wave interaction is unknown, but it should be quite small at these low temperatures. On the other hand, the imaginary part of the \( p \)-wave scattering, due to the reactive loss when molecules collide, can be much faster, and limits the cloud lifetime for KRb, even when it is suppressed by using a two-dimensional “pancake” geometry [15]. It gives an imaginary shift to the average frequency from the sum rule, and thus a Lorentzian broadening. To determine the value of this broadening, note that Eq. (30) with \( b_{12} = 0 \) is the expectation value of the \( p \)-wave interaction energy. The imaginary part is the particle lifetime, since at the level of calculation of this equation the imaginary part of this energy is the quasiparticle lifetime, and the only source of quasiparticle decay is the \( p \)-wave particle loss. Thus the cloud lifetime is simply this quasiparticle lifetime. Using cloud lifetimes measured in Ref. [15] that decay is around 4/s, or a little less than a Hertz. Since the shift is the difference of two such rates—the 1-2 channel and the 1-1 channel—we expect it to be at most on this order, and likely considerably less. Of course, we cannot rule out the possibility that the 1-2 interaction is much larger, but it is highly unlikely to be the orders of magnitude larger required to contribute substantially to the spectral broadening, compared to the dipolar shifts. Thus the \( p \)-wave broadening will be negligible under the current conditions of the JILA experiments.

FIG. 4. (Color online) The characteristic frequency of the collisional shifts in the thermal gas, \( \omega_{\text{scale}} \), for transitions between KRb’s two lowest rotational states vs the dipole moment \( d_{11} \) in units of the permanent dipole moment \( d_p \) (roughly 0.5 D for KRb). Middle curve: total characteristic frequency \( \omega_{\text{scale}} = \omega_{\text{dir}} + \omega_{\text{sf}} \). Top curve: \( \omega_{\text{dir}} \), the contribution from the spin-flip term. Bottom curve: \( \omega_{\text{sd}} \), the contribution from the direct term.
IV. SPECTRAL SHAPE: BROADENING

Previously, we considered the mean spectral shifts. Also, because the \( p \)-wave shifts may have an imaginary part due to the reactive nature of certain molecules, such as K\( \text{Rb} \), this gives a broadening. Here we go beyond treating the spectra as sharp \( \delta \) functions with this simple broadening, and we consider the spectral line shape for the experimentally relevant dilute gas. Broadening arises from several sources: differing trapping potentials felt by the two states due to their differing polarizabilities (coming from vector and tensor light shifts), collisional broadening (related to the quasiparticle lifetime), temperature broadening from going beyond the pseudopotential approximation, and higher band occupation. We will see that in this limit, all sources of broadening are very small, except the differing state polarizabilities. However, even for current conditions, these broaden the spectra by an amount that is comparable to the width of the trap-summed spectra. Consequently, some key features survive, such as the maximal shift. Moreover, by applying the probe after aligning the dipoles at the “magic angle” or “magic field strength” \([53]\), even this broadening can be eliminated.

There is also the natural linewidth of the excited rotational state, given by \( \Gamma_{\text{sat}} = (\delta c/\epsilon_0) d_{12}^2/(3\pi \epsilon_0 \hbar) \), where \( \delta c \) is the transition frequency between the excited state \([1, 0]\) and the ground state \([0, 0]\), and \( d_{12} \) is the dipole matrix element between these states. Even for the strongest dipole moments, this is \( \sim 10^{-10} \) Hz for K\( \text{Rb} \), calculating \( d_{12} \) and \( \delta c \) in the same manner we calculated the energy scales above, and thus completely negligible.

A. Differing polarizabilities

The polarizabilities of rotational states can differ substantially for different states due to vector and tensor light shifts. For example, the polarizability difference of K\( \text{Rb} \) between the lowest and first excited rotational states is about 30\% for lattice wavelengths giving a lattice spacing of 545 nm \([53]\). In systems using optical potentials to confine the molecules, this leads to the molecules experiencing different external potentials. In the presently considered case, this means the trapping frequencies differ by about 15\% and the lattice depths differ by about 30\%. Since we are taking the lattice to be sufficiently deep to suppress tunneling, its only effect on physics along the lattice direction is to change the Wannier functions for each state and thus renormalize the interactions. This was accounted for in our derivation of the lattice Hamiltonian. However, the differing trapping frequencies in the transverse direction leads to broadening.

Within the LDA, we note that if the trapping potentials differ by \( \Delta r \) at location \( r \), then this shifts the spectral line of atoms at that position by \( s(r) = \Delta(r) \) [this intuitive result follows formally from calculating the sum-rule commutators of Eq. (17)]. For harmonic traps, this shift is \( s(r) = -A_2 + A_1 + \frac{m \omega_0^2 - \omega_1^2}{2 \omega_0} r^2 \), where \( \omega_0 \) is the resonant frequency in the transverse direction for state \( s \), and \( A_0 \) and \( A_1 \) are constants. Note that \( A_0 \) are positive since the optical intensity is maximal at the center of the trap and that there is a frequency shift there as well. The magnitude of the shift \( A_1 - A_2 \) depends on the exact trapping laser used and just gives a constant shift to the spectrum. What is more important for our present concerns is the broadening coming from the second, spatially dependent term. At finite temperatures, there is (exponentially small) occupation arbitrarily far from the trap center, and the shift from the second term increases indefinitely as one moves outward in the trap. However, because the spectral density for a given value of this shift is proportional to density, the result of the polarizability difference is an exponentially decaying tail, in frequency, in the spectrum. The spectrum for locations at distance \( r \) from the center of the trap has a polarizability difference shift

\[
\omega = \frac{m (\omega_1^2 - \omega_2^2)}{2} \frac{r^2}{\omega_1^2 - \omega_2^2}, \tag{36}
\]

and the spectral weight is proportional to \( n(r) \). In the thermal gas, Eq. (33) with the frequency position relation Eq. (36) thus gives the spectrum, at least in the tails where the polarizability difference shift is the only relevant one and the cloud is thermal.

Thus \( I(\omega) \propto r^2 (dr/d\omega) e^{-\frac{m (\omega_1^2 - \omega_2^2)}{2T(\omega_1^2 - \omega_2^2)} \frac{r^2}{\Lambda}} \) yielding

\[
I(\omega) \propto \sqrt{\frac{\omega}{\omega_c}} e^{-\frac{\omega}{\omega_c} - \frac{\omega}{\omega} - \frac{\omega}{\omega}} \Theta((\omega_2^2 - \omega_1^2)\omega), \tag{37}
\]

where \( \Theta \) is the Heaviside step function. The spectral density decays exponentially with frequency, with the characteristic decay frequency \( |(\omega_2^2 - \omega_1^2)\omega| \). Note that this decreases with decreasing temperature, by virtue of the cloud shrinking, so that fewer particles occupy the wings, where the potential difference is largest. Also in the tail, this is the only source of shifts since densities are low, so this always gives the complete spectral shape of the spectral tail.

For an 800 nK gas with polarizabilities of the two states differing by 30\%, this characteristic frequency scale over which the spectral density decays is about 4.9 kHz, comparable to the spectral width of the trap-summed spectra obtained without this source of broadening. However, we emphasize that these shifts can be eliminated by performing the recoil-free spectroscopy while aligning the molecules at the magic angle or at the magic field strength, using states with nearly identical polarizabilities, or using other techniques to compensate the polarizability difference. As such, the spectral tails can be a useful diagnostic of how close the trapping potentials of the two species are, and can help, for example, to experimentally pinpoint the magic angle.

Figure 5(a) illustrates the spectral shape due to polarizability difference broadening, with a sufficiently dilute system such that there is no interaction shift. In this limit, the spectrum without the polarizability difference broadening would simply be a \( \delta \) function. For illustration, we assumed that \( \omega_2^2 < \omega_1^2 \). For \( \omega_2^2 > \omega_1^2 \), the spectrum will be reflected across the \( y \) axis. Figures 5(b) and 5(c) illustrate the effect of this broadening on the spectrum in the presence of dipolar interaction shifts, analogous to Fig. 3.

B. Collisional lifetime

Interparticle interactions broaden the spectral line by giving the quasiparticle excitations associated with the response a finite lifetime. For weak interactions, the line shape is roughly Lorentzian. We emphasize that the full line shape is captured in Eq. (15). Only our subsequent determination of trapped
We start by calculating the quasiparticle lifetime for particles in the initial rotational state. The standard theory of collisional lifetimes for a quasiparticle of momentum $p$ and frequency $\omega$ gives the decay rate [54]

$$\Gamma(p,\omega) = \frac{1}{2(2\pi)^3} \int dq \left[ \delta(\omega + \epsilon_p - \epsilon_{p+q} - \epsilon_{p-q}) \times [v(-q) - v(p - p' + q)] \times \{f(\epsilon_p)[1 - f(\epsilon_{p+q})][1 - f(\epsilon_{p-q})] - [1 - f(\epsilon_p)]f(\epsilon_{p+q})f(\epsilon_{p-q})\} \right],$$

(38)

where $f(\epsilon) = \frac{1}{e^{\epsilon/k_B T} + 1}$ is the Fermi function, $\epsilon_p \equiv p^2/(2m)$, and $v(q)$ is the Fourier transform of the interaction potential, here the Fourier transform of $V_{11}(p)$. This can be understood by the decay processes occurring in Fermi’s golden rule, perturbing in the interactions. For a thermal gas, this simplifies to

$$\Gamma(p,\omega) = \frac{1}{2(2\pi)^3} \int dq \left[ \delta(\omega + \epsilon_p - \epsilon_{p+q} - \epsilon_{p-q}) \times [v(-q) - v(p - p' + q)]^2 e^{-\beta(\epsilon_p - \mu)} \right].$$

(39)

Rather than worry about the full $p$ and $\omega$ dependence of this broadening, we calculate the average broadening of the line and for each $p$ treat the frequency as being fixed to $\omega = \epsilon_p$. With this, the typical broadening—that is, the broadening averaged over momenta $\Gamma_{\text{typ}} = \langle 1/N \rangle \sum_p \Gamma(p,\epsilon_p)$—is

$$\Gamma_{\text{typ}} = \frac{1}{n} \int \frac{dp}{(2\pi)^3} \Gamma(p,\epsilon_p).$$

(40)

Calculating $\Gamma_{\text{typ}}$ from Eq. (39) requires the Fourier transform of the interaction potential. The Fourier transform, neglecting $A$ as argued to be valid earlier, is simply

$$v(q) = C_0 - 2\pi \gamma_{11} q + O(q^2)$$

(41)

for some constant $C_0$. The constant $C_0$ cancels in Eq. (39). Physically, this is because it describes a renormalization of the contact potential, which affects nothing in a gas of identical fermions. Then, evaluating the integrals, the typical quasi-particle lifetime is $\Gamma_{\text{typ}} = \frac{m^2(\pi + 2)}{2\pi^3} T n$. Reference [55] argued that the recoil-free spectral broadening is simply the expression for the quasiparticle decay rate with $\gamma_{11} \to \gamma_{12} - \gamma_{11}$, giving the collisional spectral broadening

$$\Gamma_{\text{col}} = \frac{m^2(\pi + 2)(\gamma_{12} - \gamma_{11})^2}{2\pi^3} T n.$$  

(42)

It is useful to compare this broadening to the mean dipolar spectral shift:

$$\frac{\Gamma_{\text{col}}}{\langle \omega_d \rangle} = \frac{\pi}{4\pi m^2 T} \frac{m^2(\pi + 2)(\gamma_{12} - \gamma_{11})}{2\pi^3} T n.$$  

(43)

Note that this ratio decreases with temperature, and is proportional to $\gamma_{12} - \gamma_{11}$. Thus as the shifts decrease, the ratio of the broadening to the shift also decreases. Even for the largest experimentally realistic differences $\gamma_{12} - \gamma_{11}$, for example setting $\gamma_{11} = 0$ and taking $\gamma_{12}$ to be that associated with the permanent dipole, this ratio is about 0.02. More typical differences will be a small fraction of this ($\lesssim 10\%$ or smaller). Given the already small shifts, this broadening is negligible, perhaps at most 100 Hz.

FIG. 5. (Color online) Polarizability difference broadening in the spectra. (a) Polarizability difference broadening when other shifts are negligible (as applies in a very dilute gas, or in the tails of the cloud). (b,c) Broadening including interactions and with the polarizability difference for positive (b) and negative (c) $\omega_2 - \omega_1$, with magnitude decreased fivefold from its true value. They compare the thermal spectra without broadening [blue, dashed, exactly as Fig. 3(a)] with the spectra with broadening (red, solid) for dipole moments giving the maximal values of the collisional shifts relevant for the JILA parameters computed previously.
So far, we have neglected the interactions with other pancakes and the spin-flip terms. Neglecting the interpancake interaction is reasonable since the interpancake interactions are somewhat smaller than the intrapancake interactions—by being far away by at least one lattice spacing.

The fact that the shift vanishes for interactions between pancakes suggests that the broadening may be even smaller than this naive argument suggests. Since the effect of the intrapancake interactions is quite small, it is almost certainly an excellent approximation to neglect the intrapancake interaction induced collisional broadening. Regarding the spin-flip terms, their maximal value is less than the maximal direct term’s, and thus the spin-flip term’s effect is likely to be comparably small.

C. Broadening from interaction potential shape

At sufficiently high temperatures, multiple scattering channels will become relevant, and rather than merely shifting the spectral line, interactions will also broaden it. This leads to a broadening as one leaves the cold collisional regime. This occurs when the thermal de Broglie wavelength $\lambda_T$ becomes less than or comparable to the range of the potential. Here, however, the temperature is sufficiently low that $\lambda_T$ is somewhat longer than the characteristic range of the potential, even for the dipolar potential. To see this, observe that presently $\lambda_T \approx 70 \text{ nm}$ while the dipole length is $\sim 50 \text{ nm}$. As experiments achieve colder temperatures, $\lambda_T$ will increase.

D. Higher band occupation

We have so far neglected higher band occupations. These have various effects, including a broadening of the spectra. Such broadening should mostly be, relative to the shifts, on the order of the relative occupation of the higher bands, perhaps $\sim 10$–$20\%$ at most here. A somewhat larger contribution can come from s-wave collisions between molecules in different bands. Regardless, these occupations are exponentially suppressed as the temperature is lowered, and additionally it may be possible to remove them by purification techniques: e.g., higher bands tunnel more rapidly than the lowest band, and it may be possible to remove them from the lattice on a short time scale.

V. 3D LATTICE

This section considers the recoil-free spectra of molecules in a sufficiently deep 3D lattice such that tunneling is negligible. This limit is interesting both for its simplicity and, as we show, its utility as a direct measurement tool for numerous important correlation functions, including for states created when the tunneling is significant. Experiments on KRb molecules in 3D lattices are underway at JILA [56].

In the infinitely deep lattice, the eigenstates of the initial Hamiltonian, prior to application of the spectral probe, are $|i\rangle \equiv \bigotimes_i |n_i\rangle$, where $|n_i\rangle$ is the Fock state with $n$ particles at site $i$. Then the total spectrum is the sum over sites $j$ of $\tilde{\delta}$ functions whose spectral shift is that for exciting particles at site $j$; in the eigenstate $|\{n_j\}\rangle$, the shift associated with exciting site $j$ is

$$\langle \omega \rangle_j = \sum_{i \neq j} [V_{12}(R_{ij}) - V_{11}(R_{ij}) + V_{df}(R_{ij})] n_i,$$

where $R_{ij}$ is the separation of the $i$th and $j$th lattice sites. This is essentially the integrand/summand of Eq. (18), since we are considering the shift associated with changing the rotational state at a single site $j$, projected to a 3D lattice. Each interatomic separation between $i$ and $j$ will have its own unique shift, and this will allow us to spectroscopically measure the correlations at various distances. Broadening can be neglected, as the only source of broadening when the tunneling is zero is the $\sim \text{Hz}$ $p$-wave losses.

Any many-body wave function $|\Psi\rangle$ initially all in the $\alpha = 1$ internal state can be expanded in this basis:

$$|\Psi\rangle = \sum_{|n_1\rangle} A_{\{n_1\}} |\{n_1\}\rangle.$$

The total spectrum is a sum of peaks with shifts given by Eq. (44). In particular, the total spectrum in the present zero-tunneling limit is

$$I_{\{n_1\}}(\omega) = \sum_{\{n_2\}} A_{\{n_1\}} I_{\{n_2\}}(\omega),$$

where $I_{\{n_1\}}(\omega)$ is the spectrum for the eigenstate $|\{n_1\}\rangle$. This spectrum is

$$I_{\{n_1\}}(\omega) = \sum_{j} n_j \delta(\omega - \langle \omega \rangle_j)$$

with $\langle \omega \rangle_j$ given by Eq. (44).

For simplicity, we consider the case in which the 3D lattice is broken into 2D sheets such that the interlayer interactions are negligible. This can be achieved either by depopulating all but one sheet or by increasing the lattice spacing along one axis. The basic idea to read off correlations, discussed in the next paragraph, can be used in more general geometries, but the procedure becomes more complicated.

Figure 6 shows an example of how to read correlation functions from the splitting of spectral peaks, explained in the caption. Figure 6(a) shows this including only nearest-neighbor interactions, and Fig. 6(b) includes next-nearest-neighbor interactions. Contributions from further separated sites’ correlations will continue the pattern of splitting the lines by smaller and smaller values.

Thus we observe that recoil-free spectra can be used to cleanly extract a tremendous amount of information about the static correlation functions of a system in a 3D lattice that have either zero or one particle per site (somewhat higher occupations can be accounted for straightforwardly, but lead to complications where peaks start to overlap). In particular, one can measure the joint probability distribution $P(1, n_1, n_2, \ldots)$, the probability of a site having one particle on it, $n_1$ atoms on all the nearest-neighbor sites, $n_2$ the number of atoms on all the next-nearest-neighbor sites, and so on. Neglecting occupations of more than one molecule per site, the arguments above show that this function $P(1, n_1, n_2, \ldots)$ is simply proportional to the spectral weight of a particular spectral peak.

Measuring $P(1, n_1, \ldots, n_j)$ requires finer spatial resolution as $j$ increases. For a given string of occupancies $(n_1, \ldots, n_{j-1})$, Eq. (44) gives the frequency splitting between having $n_j$ and $n_j + 1$ particles at lattice sites that are $j$th nearest neighbors to be $\delta = V_{12}(r_j) - V_{11}(r_j) + V_{df}(r_j)$, where $r_j$ is the lattice vector corresponding to the $j$th nearest-neighbor separation. The energy scales for KRb for current 532-nm lattice spacing square lattices (neglecting interactions between the 2D sheets
that occur in a 3D lattice, and assuming that the difference in dipole moments for the initial state and final state can be made comparable to half the permanent dipole moment) enable one to resolve $P(1, n_1, n_2)$ if one has 30 Hz spectral resolution and $P(1, n_1, n_2, n_3)$ if one can obtain 10 Hz spectral resolution. For other molecules, the measurements can be even more informative. For example, with RbCs one can measure $P(1, n_1, \ldots, n_4)$ using a 30 Hz spectral resolution and $P(1, n_1, \ldots, n_5)$ using a 10 Hz spectral resolution. Shorter lattice spacings enhance the range of correlations measurable for a given spectral resolution.

Note that this joint probability distribution includes moments, e.g., $\langle n^2 \rangle$ and $\langle n_0 n_2 \rangle$, but is substantially more informative. In particular, one can compute any moment between $n_j$’s that can be resolved: i.e., it gives $\langle n_{i_0}^a n_{i_1}^{a_1} n_{i_2}^{a_2} \cdots n_{i_j}^{a_j} \rangle$ for arbitrary $(a_0, a_1, \ldots, a_j)$.

Importantly, this technique can probe correlations of many-body systems generated by the system with a shallower lattice, where tunneling may be important. In this case, one can rapidly ramp up the lattice depth from the value where the physics is of interest to a value where the tunneling is effectively zero. This projects the initial state onto a basis of the zero-tunneling eigenstates $|n_i\rangle$, and the dynamics during the interrogation then are described by the above considerations. Thus one can measure the joint number distribution function of the initial state. This is analogous to the measurement of on-site number statistics done in collapse-and-revival experiments [37,58]: a shallow lattice equilibrium state is quenched to a deep lattice state for measurement.

Such a technique is straightforwardly extended to measure correlations where the initial state may have several rotational levels populated, and to measure intersite correlation functions between atoms in these various states. This occurs naturally in, and would be particularly useful for, proposals to use molecules’ internal degrees of freedom as “pseudospins” to emulate quantum magnetism [45,46], where the technique would probe the pseudospin correlations.

Here we explicitly considered interpreting these spectra for the 2D lattice in a 2D system. This is straightforwardly extended to a 3D lattice, but the form of the $V_{12} - V_{11} + V_d$ will be slightly more complicated due to interaction anisotropies. However, our discussion does not rely crucially on the form of $V_{12} - V_{11} + V_d$, except for the fact it is long-ranged.

Finally, this measurement technique is not limited to molecules. For example, it may be possible to measure correlations of alkali-metal atoms in optical lattices by exciting all the atoms to a Rydberg state immediately before the spectral interrogation. The spectral pulse would then be chosen to drive the atoms between two Rydberg states, which will experience dipolar effects in a manner analogous to the molecules.

VI. SUMMARY

In this paper, we have calculated the recoil-free spectra for dipolar molecules in one-dimensional lattices of two-dimensional pancakes, and in deep three-dimensional lattices. We particularly focused on the case of driving transitions between different rotational states. We have incorporated both the dipolar interactions and the $p$-wave interactions and losses, the latter of which are important for reactive molecules. We gave general expressions for the shifts, and we evaluated these expressions for the weakly interacting gas as well as their simpler low-temperature ($T \ll \mu$, degenerate) and high-temperature ($T \gg \mu$, thermal) limits. We calculated the broadening of these lines from collisions, losses, differing polarizabilities of rotational states, and other mechanisms. We overviewed typical scales for these various effects for the case of KRb molecules. The overall scale for the trap summed spectrum under current experimental conditions is on the order of $\sim 5$ kHz for strong dipoles due to the dipolar interactions and the inhomogeneous density of the trapped system. This will increase to a constant with decreasing temperature and increase indefinitely with increasing density (until perturbation theory breaks down). Observation of this line shape will confirm that the interactions in this system take the expected form and can be used to diagnose temperature and degeneracy. Deviations from the weakly interacting spectral line shapes can then be used to observe and characterize strong correlation physics.

The $p$-wave losses induce a few Hertz broadening, and the real $p$-wave interactions should be very small. Collisional broadening is also small, probably much less than 100 Hz for current conditions. The largest source of shift and broadening.
other than the dipolar interactions, comes from the differing polarizabilities of rotational states. This is on the order of a few kilohertz, and decreases with temperature. However, it can be eliminated by working, for example, at the “magic angle” or “magic field strength” [53]. Recoil-free spectra offer a useful probe of this physics.

In the three-dimensional lattice, we showed how the spectra give access to a multitude of static correlation functions that are inaccessible by other techniques. The spectral weight in each peak is proportional to the probability of a certain set of spatial configurations of a particular set of lattice occupations. This allows one to measure joint probability distributions for correlations among a few nearest-neighbor sites.

We mention that although we focused on the most immediately relevant case of dipoles aligned perpendicular to the pancakes and in a one-dimensional lattice of two-dimensional pancakes, most of our general formulas extend straightforwardly to other cases, for example three dimensions or other angles of alignment. Only an evaluation of the integrals changes. Presumably, many of the overall magnitudes typically will remain similar.

Finally, although we focused on dipolar molecules in our quantitative estimates, the effects are relevant elsewhere. Rydberg-earth-metal atomic clocks, $p$-wave interactions and reactive losses can be important, and although the dipolar interactions are very small, they may be important as clock accuracies approach their fundamental limits. However, we mention that we have treated the dipole interaction electrostatically. This is valid when the particle separation is much less than the wavelength of the photons mediating the dipole-dipole interaction. For molecules, we expect this to be quantitatively accurate. However, in other contexts such as atomic clocks, the dynamic field becomes important when interparticle separations are comparable to or larger than the wavelength of the photon associated with the dipole transition [59]. In this case, the treatment of dipolar interactions of the excited state is quite different. For example, they scale as $1/r$, where $r$ is the interparticle separation. In this context, careful choice of lattice parameters mitigates these effects [59].

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