Non-local state-swapping of polar molecules in bilayers

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The observation of significant dipolar effects in gases of ultra-cold polar molecules typically demands a strong external electric field to polarize the molecules. We show that even in the absence of a significant polarization, dipolar effects may play a crucial role in the physics of polar molecules in bilayers, provided that the molecules in each layer are initially prepared in a different rotational state. Then, inter-layer dipolar interactions result in a non-local swap of the rotational state between molecules in different layers, even for weak applied electric fields. The inter-layer scattering due to the dipole-dipole interaction leads to a non-trivial dependence of the swapping rate on density, temperature, inter-layer spacing, and population imbalance. For reactive molecules like KRb, chemical recombination immediately follows a non-local swap and dominates the losses even for temperatures well above quantum degeneracy, and could be hence observed under current experimental conditions.

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A new generation of experiments has started to explore the remarkable novel physics of dipolar gases, in which dipole-dipole interactions play a key role \textsuperscript{1}. These interactions, being long-range and anisotropic, differ from the short-range isotropic interactions which have dominated up to now ultra-cold atomic physics. Polar molecules are expected to provide fascinating new scenarios for quantum gases due to their large electric dipole moments. Recent experiments on preparation and control of ro-vibrational and hyperfine states of KRb at JILA \textsuperscript{2} open new perspectives towards a degenerate quantum gas of polar molecules. Unfortunately chemical recombination due to the reactive character of KRb has up to now prevented to reach quantum degeneracy \textsuperscript{3}. However, dipolar interactions between partially polarized KRb molecules have been recently shown to significantly reduce chemical recombination in constrained quasi-2D geometries \textsuperscript{4}.

Although a polar molecule in its lowest ro-vibrational state may have a large dipole moment in the molecular frame, in the absence of an external electric field the dipole moment in the laboratory frame averages to zero. As a consequence, the observation of dipolar effects in gases of polar molecules typically demands the polarization of the molecules in an external electric field \textsuperscript{4}. For example, a KRb molecule in its singlet ground state has a permanent dipole moment of $d = 0.566$ Debye in the molecular frame \textsuperscript{2}, but rather large fields of several kV/cm must be employed to reach effective dipoles of $\sim 0.2$ Debye in the lab frame \textsuperscript{5}.

However, remarkably, dipolar interactions may play a significant role even in the absence of an external electric field, if the molecules are prepared in different rotational states \textsuperscript{6}. In that case, the interaction between molecules in different rotational states is produced by exchanging a quantum of angular momentum (state swap), resembling the resonant interaction of an electronically excited atom with a resonant interaction of an electronically excited atom.

In this Letter, we show that bilayer systems may provide an excellent environment for observing the non-trivial physics associated with dipole-induced state-swaps in current experiments. We consider polar molecules in a bilayer geometry, without inter-layer tunneling, prepared such that the molecules in layer A (B) are in the ground (first-excited) rotational state (Fig. 1). The dipole-dipole interaction induces a molecule in one layer to interchange its rotational state with a molecule in a different state in the other layer, as sketched in Fig. 1. Such a process resembles spin-exchange in the collision of polarized atomic beams \textsuperscript{7} and spin-changing collisions in spinor gases \textsuperscript{8}. We show that the special features of the inter-layer scattering lead, especially for quantum degenerate samples, to a non-trivial dependence of the swap rate on temperature, molecular density, inter-layer spacing, and population imbalance. Interestingly, for reactive molecules like KRb chemical recombinations following a state-swap dominate losses even well inside the non-degenerate regime and could be observable under current experimental conditions.

We are interested in fermionic polar molecules confined in two thin parallel layers, $A$ and $B$, separated by a distance $\lambda$ along $z$, and with negligible hopping between them (Fig. 1). We consider diatomic molecules...
with a permanent electric dipole moment \( d \) in the electronic singlet \((1\Sigma)\) and in the vibrational ground state. A weak external electric field \( \mathbf{E} = \mathbf{E} \mathbf{z} \) is applied. The rotational states of the molecules are described by the Hamiltonian

\[
H_{\text{mol}} = BJ^2 - d \cdot \mathbf{E},
\]

where \( B \) is the rotational constant, \( J \) is the angular momentum operator, and \( d = d(\sin \theta \cos \varphi, \sin \theta \sin \varphi, \cos \theta) \) is the dipole moment operator, where the angles \( \theta \) and \( \varphi \) describe the orientation of the molecule in the laboratory frame. For a weak electric field, such that \( \beta = dE/B \ll 1 \), the eigenstates \( \varphi_{JM} \) of the Hamiltonian are still well described by the quantum numbers \( J \) and \( M \) associated with the angular momentum \( J \), and they can be found using perturbation theory [1,2]. The electric field lifts the degeneracy of the three \( J = 1 \) states, which become separated by an energy \( \Delta E = E_{1,0} - E_{1,\pm 1} = 3\beta^2 B/20 \).

The effective dipole moment in the laboratory frame, \( \langle \mathbf{d} \rangle \), is very small for a weak field. For example, the effective dipole in the rotational ground state is \( \langle \mathbf{d} \rangle \varphi_{0,0} = 3\beta d/3 \mathbf{z} \), giving 0.019 Debye for KrB and \( \beta = 0.1 \), corresponding to \( E = 0.4 \text{kV/cm} \); this results in a dipolar interaction five times smaller than that for atomic magnetic dipoles in Chromium [1].

We suppose that, initially, the molecules in layer \( A \) are prepared in the state \( \varphi_{0,0} \), while those in \( B \) are prepared in \( \varphi_{1,0} \), as illustrated in Fig. 1. Such selective preparation may be achieved by applying a spatially varying electric field, and then transferring the molecules of layer \( B \) into the desired state with a microwave transition [1].

The dipole-dipole interaction between two molecules with dipole \( \mathbf{d}_{1,2} \) at positions \( \mathbf{r}_{1,2} \) is

\[
H_{DD} = \frac{1}{4\pi \varepsilon_0} \left[ \frac{\mathbf{d}_1 \cdot \mathbf{d}_2}{|\mathbf{r}|^3} - \frac{3(\mathbf{d}_1 \cdot \mathbf{r})(\mathbf{d}_2 \cdot \mathbf{r})}{|\mathbf{r}|^5} \right],
\]

with \( \mathbf{r} = \mathbf{r}_1 - \mathbf{r}_2 \). Since the anisotropic dipole-dipole interaction does not preserve the rotational quantum number, a collision may lead to an interchange of the rotational state of molecules in different layers:

\[
\varphi_{0,0}^{A} \varphi_{1,0}^{B} \rightarrow \varphi_{1,0}^{A} \varphi_{0,0}^{B}.
\]

The matrix elements of \( H_{DD} \) involving the states \( \varphi_{0,0} \) and \( \varphi_{1,0} \) are given by, neglecting terms \( O(\beta^2) \),

\[
\langle \varphi_{1,0}^{A} \varphi_{0,0}^{B} | H_{DD} | \varphi_{0,0}^{A} \varphi_{1,0}^{B} \rangle = 0,
\]

\[
\langle \varphi_{1,0}^{A} \varphi_{1,0}^{B} | H_{DD} | \varphi_{0,0}^{A} \varphi_{1,0}^{B} \rangle = \frac{d^2}{3\varepsilon_0} \left[ 1 - \frac{3z^2}{r^2} \right] = V_{dd}(\mathbf{r}).
\]

Transitions into rotational states other than \( \varphi_{0,0} \) and \( \varphi_{1,0} \) can be neglected. They involve an energy change of \( \Delta E \), which is much larger than other energy scales in the system (the rotational constant \( B \) is typically very large), such as the dipolar energy \( E_D = d^2/(4\pi \varepsilon_0 \lambda^3) \). For KrB, for example, \( B/h = 1.11 \text{ GHz} \), thus \( \Delta E/k_B = 80 \mu\text{K} \) is much larger than \( E_D/k_B = 15 \text{nK} \) for \( \lambda = 532 \text{nm} \).

Therefore, we only consider processes between the states \( \varphi_{0,0} \) and \( \varphi_{1,0} \).

At this point the dipolar interaction is easily diagonalized using the symmetric and antisymmetric two-body states \( |S, A\rangle \equiv \langle \varphi_{0,0}^{A} \varphi_{1,0}^{B} \rangle \pm |\varphi_{1,0}^{A} \varphi_{0,0}^{B} \rangle \sqrt{2} \), for which \( \langle A | H_{DD} | A \rangle = -V_{dd}(\mathbf{r}) \), \( \langle S | H_{DD} | S \rangle = V_{dd}(\mathbf{r}) \), and \( \langle S | H_{DD} | A \rangle = 0 \). The inter-layer scattering is hence decomposed into two separate channels, namely the symmetric one, with interaction potential \( V_{dd}(\mathbf{r}) \), and the antisymmetric one, with \( -V_{dd}(\mathbf{r}) \). It may be noted that in analogy with spin-exchange collisions between polarized atomic beams [2], the interaction between two molecules can be written as \( V = V_{dd}(\mathbf{r})P_S - V_{dd}(\mathbf{r})P_A \), where \( P_S, P_A \) are the projection operators for states \( |S, A\rangle \), \( |A\rangle \).

We introduce the scattering amplitudes \( f_+ \) and \( f_- \), which characterize the low-energy inter-layer scattering for the symmetric and antisymmetric channel, respectively. The scattering amplitude for the state-changing process in Eq. (7) is then given by

\[
f_{sc}(k) = \frac{1}{2}(f_+(k) - f_-(k)),
\]

with \( \hbar k \) the relative momentum between the colliding particles. The amplitude for state-preserving collisions, \( \varphi_{0,0}^{A} \varphi_{1,0}^{B} \rightarrow \varphi_{0,0}^{A} \varphi_{1,0}^{B} \), is \( f_{sp} = \frac{1}{2}(f_+ + f_-) \).

Collisions involving particles in different layers can be described here by two-dimensional scattering (s-wave only). The amplitudes \( f_+ \) and \( f_- \) are found by numerically solving the 2D Schrödinger equation with the potential \( \pm V_{dd}(x, y, z = \lambda) \). This may be compared, using Ref. [12], to the result from the Born approximation for \( f_\pm \),

\[
|f_{sc}(q)|^2 = U_0^2 q^2 \left[ 2 + \pi(\mathbf{L}_1(2q) - I_1(2q))^2 \right] + O(U_0^3),
\]

with \( q = k \lambda \) and \( \mathbf{L}_1(x) (I_1(x)) \) the modified Struve (Bessel) function. We use the unit of energy \( E_0 = \hbar^2/(m \lambda^2) \), with \( m \) being the mass of the molecule, and the dimensionless coupling strength \( U_0 = E_D/(3E_0) \).

To be specific, we will take the values for KrB and \( \lambda = 532 \text{nm} \) for the plots, so \( E_0/k_B = 13.5 \text{nK} \) and \( U_0 = 0.38 \). In Fig. 2(a) we show that the expression in Eq. (17) is in good agreement with the numerical solution of the Schrödinger equation, for the regime of interest.

Note that \( |f_{sc}|^2 \) presents a non-monotonical dependence with the relative kinetic energy, and falls off slowly for large energies, see Fig. 2(a). Interestingly, the swapping amplitude in Eq. (7) is 1/3 of the scattering amplitude expected for fully polarized molecules calculated in the same approximation. On the other hand, the scattering amplitude for state-preserving collisions is much smaller. Since there is no direct process (the matrix element vanishes, Eq. (11)), these collisions are of higher order: \( |f_{sp}|^2 = O(U_0^3) \).
State-swapping leads to molecules in \( \varphi_{0,0} \) (\( \varphi_{1,0} \)) in layer \( B \) \( (A) \), see Fig. 1. For short times, the number density \( n_d \) of these “defects” satisfies the rate equation

\[
dn_d/dt = K n_A n_B,
\]

where \( n_A \) and \( n_B \) are the densities in layers \( A \) and \( B \). The rate \( K \) is determined by the cross-section for state-changing collisions \( \sigma_{sc} \), averaged over the velocity distributions \( f_{A,B} \) of particles in layer \( A \) and \( B \) \( \text{(13):} \)

\[
K = \langle v \sigma_{sc} \rangle = \int \mathcal{V} v \sigma(p) f_A(v_A) f_B(v_B) d^2v_A d^2v_B,
\]

where \( v = v_A - v_B \) is the relative velocity for a pair of molecules and \( p = (m/2)v \). The relation to the scattering amplitude in two dimensions is \( v \sigma = 4 \frac{1}{m_2} |f|^2 \).

For short times or for a small defect density, Pauli-blocking effects can be neglected. For reactive molecules, in fact, the defect density is always small because swapped molecules are quickly lost, as discussed below. For equilibrium distributions which depend only on energy, the collision rate \( K \) becomes:

\[
K = 8 \frac{\hbar}{m} \int_0^\infty |f_{sc}(k_{rel})|^2 F(E_{rel}) dE_{rel},
\]

where \( E_{rel} = \hbar^2 k_{rel}^2/m = 1/2 (m/2) v^2 \). Here

\[
F(E_{rel}) = \int_0^{2\pi} \frac{d\gamma}{2\pi} \int_0^\infty f_A(E_A) f_B(E_B) dE_{cm}
\]

is the distribution of relative kinetic energies, where \( \cos \gamma = \frac{v_{rel}}{V}, V = (v_A + v_B)/2, E_{AB} = m v_{AB}^2/2, \) and

\[
E_{cm} = m V^2
\]

is the center-of-mass kinetic energy. The \( f_{A,B} \) are normalized to \( \int f(E)dE = 1 \).

The distribution of relative energies \( F(E) \) can be calculated from Eq. \( \text{(11)} \), assuming a Fermi distribution in each layer \( f_{A,B}(E) = N^{-1}(e^{(E-E/k_B T)} + 1)^{-1} \), with the normalization constant \( N = T \log(1 + e^{\mu/k_B T}) \). The chemical potential \( \mu(T) \) is that of the ideal 2D Fermi gas, i.e. the solution of \( \int f(E)dE = 1 \).

Figure 2 shows \( F(E) \) for different temperatures. For temperatures \( k_B T \gg \epsilon_F \), the relative energies follow a Boltzmann distribution, \( F(E) = 1/k_B T \exp(-E/k_B T) \).

The swap rate \( K \), as given by Eq. \( \text{(12)} \), is the overlap integral between \( F(E_{rel}) \) and \( |f_{sc}(E_{rel})|^2 \). The dependence of the collision rate on the density, inter-layer spacing \( \lambda \), and temperature can be obtained by analyzing this overlap. The scattering amplitude \( |f_{sc}|^2 \) has a maximum around \( E_{rel}/E_0 = (\lambda k_{rel})^2 = 0.5 \), see Fig. 2b, so at \( T = 0 \) the maximal swapping rate occurs for Fermi momenta \( k_F \lambda \sim 1 \). Hence, Fermi degenerate molecular samples show a peculiar non-monotonical behavior of the swapping rate as a function of density, or inter-layer spacing, as shown in Fig. 3a. This effect is visible if \( k_B T \ll E_0 \), and it is smeared for higher temperatures, see Fig. 3a.

The swap rate \( K \) presents as well an interesting dependence on the population imbalance between both layers. At \( T = 0 \) the function \( F(E) \), which may be analytically evaluated, depends on the chemical potentials \( \mu_A \) and \( \mu_B \) of the layers (or, equivalently, particle densities), acquires the form of a smoothed step function, as Fig. 2b). In-
terestingly, as shown in Fig. 3(b), the swap rate $K$ may show a maximum as a function of the imbalance $\mu_A/\mu_B$, if $\mu_B \lesssim E_0$. Again, as can be seen in Fig. 3(b), at finite $T$, the maximum persists although smeared out.

The previous results suggest that the state swapping collisions may lead to very relevant effects even in the high-temperature regime, especially in the context of reactive molecules. Typical experimental values for $E_0/k_B$ and $\epsilon_F/k_B$ are in the range of 10nK, while currently available experimental temperatures are much higher, namely in the range of several 100nK. In the following we discuss the particularly relevant scenario of KRB molecules, assuming a molecular density of $5.6 \times 10^7$ cm$^{-2}$, which gives $\epsilon_F/E_0 = 1.0$. Figure 3 shows the temperature dependence of the swapping rate under these conditions. For $k_B T \gg \epsilon_F$, it follows from Eq. (7) that the swapping rate acquires the form

$$K(T) \approx 2\frac{\hbar}{m} U_0^2 \log\left(\frac{k_B T}{E_0}\right) \frac{1}{k_B T/E_0}.$$  \hfill (11)

Once chemically reactive molecules collide within one layer, they undergo a reaction of the form KRB $\rightarrow$ K$_2$ + Rb$_2$ and leave the trap [4]. For weak electric fields (no significant polarization of the dipoles), this results in a loss rate of $K_p \approx 2 \times 10^{-7}$ cm$^2$/s for $T = 800$ nK due to $p$-wave scattering, scaling linearly with $T$ due to Wigner threshold law [4]. The swapping process discussed here leads to an additional loss mechanism for reactive molecules, since the creation of defect molecules (state $\varphi_{0,0}$ in layer $B$ and state $\varphi_{1,0}$ in layer $A$, see Fig. 1) allows for intra-layer $s$-wave collisions. Their typical rates are $K_s \approx 10^{-5}$ cm$^2$/s [4], i.e. at least one order of magnitude larger than the swapping rate, see Fig. 3. Since $K_s \gg K$, the defect molecules are quickly lost (the instantaneous number of defect molecules at short times remains at any time very small $n_d \approx (K/K_s)n_A$). Hence we may equate the swapping rate with molecular loss rate, $dn_A/dt \approx -(K + K_p)n_A^2$, assuming balanced mixtures with $n_A = n_B$. Note that the swapping mechanism discussed here becomes the dominant loss mechanism at low-enough $T$ characterized by $K_p \ll K$. Since swap-induced losses decrease with $T$ while losses associated to $p$-wave scattering increase linearly with $T$, there is a temperature threshold $T_c$ at which both loss mechanisms are equally relevant. From Fig. 3 and Ref. [4], we estimate this threshold at approximately $T_c \approx 530$ nK$\approx 40 \epsilon_F/k_B$ [13]. Therefore even in the absence of external polarization, non-local dipole-induced swapping dominate over $p$-wave losses even deeply inside the non-degenerate regime.

In summary, polar molecules in bilayer systems in a different rotational state in each layer provide a distinctive setting for the study of non-local state-swaps. We have shown that the special features of the inter-layer dipole-dipole interaction lead to a non-trivial dependence of the swap rate with the parameters of the system, especially within the degenerate regime. Moreover, for chemically reactive molecules, the swapping leads to losses, since a swap is followed by a quick recombination due to an intra-layer $s$-wave collision. As a result of that, non-local state-swapping should lead to significantly enhanced losses below a temperature threshold, which lies well inside the non-degenerate regime. Hence, our results show that strong non-local dipolar effects should be observable in current experiments in the absence of a significant external polarization.

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\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig4.png}
\caption{Swapping rate $K$ as a function of temperature, for $\epsilon_F/E_0 = 1$. The dashed line is the result of Eq. (11).}
\end{figure}

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