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Observation of Cold Collisions between Trapped Ions and Trapped Atoms

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We study cold collisions between trapped ions and trapped atoms in the semiclassical (Langevin) regime. Using Yb⁺ ions confined in a Paul trap and Yb atoms in a magneto-optical trap, we investigate charge-exchange collisions of several isotopes over three decades of collision energies down to 3 μ eV ($k_B \times 35$ mK). The minimum measured rate coefficient of 6×10^{-10} cm³ s⁻¹ is in good agreement with that derived from a Langevin model for an atomic polarizability of 143 a.u.

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Studies of cold collisions between trapped neutral atoms have revealed a plethora of fascinating quantum phenomena, including Wigner threshold laws [1], magnetically tunable Feshbach resonances [2], controlled molecule formation [3], and the suppression of individual scattering channels [4]. Collisions between trapped ions, on the other hand, are featureless since the strong long-range repulsive Coulomb interaction prevents the ions from approaching each other. Collisions between ions and neutral atoms [5-8] fall into an intermediate regime where an attractive long-range r^{-4} potential leads to semiclassical behavior for a wide range of collision energies, but where quantum phenomena dominate at very low energies. Cold ion-atom collisions have been proposed as a means to implement quantum gates [9], to cool atoms [7,10] or molecules [11,12] lacking closed optical transitions, to bind small Bose-Einstein condensates to an ion [13], or to demonstrate novel charge-transport dynamics [14].

As a function of collision energy, charge-exchange and momentum-transfer ion-atom collisions exhibit three distinct regimes [6,7]: a high-energy classical (hot) regime with a logarithmic dependence of cross section σ on energy E, a wide semiclassical Langevin (cold) regime with a power-law dependence $\sigma(E) \propto E^{-1/2}$ [5,6], and a quantum (ultracold) regime where contributions from individual partial waves can be distinguished. However, given the large forces on ions produced by small stray electric fields, it has been difficult to reach experimentally even the semiclassical regime. References [15,16], studying charge exchange at $E \sim 100$ meV, report the only observations of Langevin-type ion-atom collisions. In ion-molecule systems, experimental signatures of Langevin collisions have been seen at high temperature [17–19], and recently also at 1 K (80 μ eV) [20]. In all previous work, at most one of the collision partners was trapped.

In this Letter, we study collisions between independently trapped, laser-cooled ions and atoms down to unprecedented low energy $(3 \ \mu eV)$ in the semiclassical collision regime. Using a double-trap system [21], we investigate resonant charge-exchange collisions for different Yb⁺ + Yb isotope combinations and find agreement with the Langevin model to within a factor of 2 over three decades of energy [5,6]. The highest energy, 4 meV = $k_B \times 45$ K, corresponds to the transition to the classical regime [22,23], while at the lowest energy, 3 μ eV = $k_B \times$ 35 mK, where approximately 40 partial waves contribute to the cross section, isotope shifts should become relevant. The lower limit on collision energy is set by our ability to detect and minimize the micromotion of a single ion in the Paul trap.

The long-range interaction potential between a singly charged ion and a neutral atom is the energy of the induced atomic dipole in the ion's electric field, given by V(r) = $-C_4/(2r^4)$, where $C_4 = \alpha q^2/(4\pi\epsilon_0)^2$ is proportional to the atomic polarizability α , and q is the electron charge. For a given collision energy E in the center-of-mass frame, there exists a critical impact parameter $b_c = (2C_4/E)^{1/4}$ that separates two types of collisions: those with impact parameter $b < b_c$ that result in inward-spiraling orbits of radius approaching zero, and those with $b > b_c$ that never cross the angular-momentum barrier [5,24].

For collisions of an ion with its parent atom, a semiclassical resonant charge-exchange cross section σ_{ce} can be simply derived, provided that b_c is large compared to the range of the molecular potential. For collisions with $b > b_c$, the electron should remain bound to the incoming atom. For close-range collisions with $b < b_c$, the electron in the resonant process $A^+ + A \rightarrow A + A^+$ is equally likely to exit attached to either nucleus, resulting in either a chargeexchange collision or an elastic collision. It follows that the resonant charge-exchange cross section is $\sigma_{ce} = \sigma_L/2$, where $\sigma_L = \pi b_c^2$ is the Langevin cross section. The corresponding semiclassical rate coefficient $K_{ce} = \sigma_{ce}v = \pi \sqrt{C_4/\mu}$, where v is the relative velocity and μ the reduced mass, is independent of energy.

At high energies this model becomes invalid when b_c becomes so small that charge exchange outside the centrifugal barrier starts to play a role; for Yb, this occurs at energies $E \gtrsim 10 \text{ meV} = k_B \times 120 \text{ K}$ [6,22,23]. At very

low energies, the semiclassical Langevin model ceases to be valid when the *s*-wave scattering limit is reached, which happens near $E_s \sim \hbar^4/(2\mu^2 C_4) = 4$ peV = $k_B \times 50$ nK. Quantum simulations [6] of this system [22] confirm the semiclassical Langevin model in the indicated energy range. For collisions between different isotopes, the Langevin expression should be modified at low collision energies comparable to the small difference in binding energy of the electron to the two nuclei, i.e., the isotope shift of the ionization potential. In this case, we expect endoenergetic charge-exchange collisions to be suppressed [25,26].

We use a magneto-optical trap (MOT) in combination with a Paul trap, as originally proposed by Smith [11]. The setup, shown in Fig. 1, is described in detail elsewhere [21]. In the present work, ¹⁷²Yb, ¹⁷⁴Yb, or ¹⁷¹Yb atoms are selectively loaded from an atomic beam into a MOT by tuning the laser frequency near the ${}^{1}S_{0} \rightarrow {}^{1}P_{1}$ transition at $\lambda_{a} = 398.8$ nm. Typically, the MOT is operated at a 75 G/cm magnetic-field gradient and contains 3×10^{5} atoms at a peak density of 2×10^{8} cm⁻³ and a temperature of 700 μ K as determined by a time-of-flight measurement.

Cold ions are produced by nonresonant photoionization from the excited ${}^{1}P_{1}$ state of the MOT with 370-nm light from a semiconductor laser [21]. The ion trap is a surfaceelectrode Paul trap printed on a vacuum-compatible substrate [21], and is typically operated at 1.4 MHz to create a 0.3 eV deep pseudopotential trap 3.6 mm above the trap surface with a secular frequency of 67 kHz. Ion trap populations can be adjusted between a single and 10⁴ ions by varying the trap loading time.

Two beams from the same 370 nm laser used for photoionization provide Doppler cooling of the ions on the ${}^{2}S_{1/2} \rightarrow {}^{2}P_{1/2}$ transition along all three principal trap axes. Ions that decay to a metastable *D* state are repumped [27] using a laser operating at 935 nm [21]. We detect the ion population by monitoring trap fluorescence at 370 nm with a photomultiplier tube. A single cold, trapped ion produces 5 kcounts/s.

Collisions that change only the particles' energy and momentum are difficult to observe in our setup due to the continuous laser cooling. On the other hand, chargeexchange collisions between different isotopes ${}^{\alpha}Yb^{+}$ + ${}^{\beta}Yb \rightarrow {}^{\alpha}Yb + {}^{\beta}Yb^{+}$ are easy to observe using isotopeselective ion fluorescence: we first load the ion trap from the MOT with isotope ${}^{\alpha}Yb^+$, change the MOT isotope to $^{\beta}$ Yb by adjusting the frequency of the 399-nm laser, and then monitor the decay of the ${}^{\alpha}Yb^{+}$ ion population through the decay of the 370-nm fluorescence. In order to prevent photoionization and direct loading of the ion trap with ${}^{\beta}Yb^{+}$, we modulate the MOT and ion light out of phase to ensure that the MOT contains no excited atoms when the 370-nm light is present. Without the MOT, the ion trap loss is exponential with a typical lifetime $\tau_0 = 400$ s for ion crystals (Fig. 2, circles), presumably due to collisions with background gas atoms. In the presence of the MOT, τ is substantially shortened, with a shorter lifetime for higher MOT density. Simple exponential decay over a decade in ion number indicates that the loss involves a single ion, rather than collisions between ions. We also measure the trap fluorescence for identical ion and MOT isotopes. In the latter case, we observe a small initial decay, likely due to heating by collisions interrupting the ions' micromotion, but no exponential loss (Fig. 2, inset). This confirms that the measured trap loss for different ion and MOT isotopes is indeed due to charge-exchange collisions, and not caused by heating from ion-atom collisions that interrupt the ion's micromotion [28].

To accurately determine the charge-exchange rate coefficient, we vary the atomic density at the ion trap location by moving the MOT with a magnetic bias field. We determine the local atomic density at the ions' location by



FIG. 1 (color online). Setup for trapping neutral (MOT) and singly charged (surface planar Paul trap) Yb in the same spatial volume. Stray dc field compensation and trapping along \hat{z} for the Paul trap is provided by dc electrodes. The traps are imaged along both the (1,0,1) and (0,1,1) axes.



FIG. 2 (color online). Typical ¹⁷²Yb⁺ ion-crystal fluorescence decay and exponential fits for no (circles), moderate (squares), and high (triangles) ¹⁷⁴Yb atomic density at the ion trap site. Inset: ¹⁷²Yb⁺ (dashed line) and ¹⁷⁴Yb⁺ (solid line) population evolution in the presence of ¹⁷⁴Yb. Both traces are normalized by the same peak value.





FIG. 3 (color online). (a) (1,0,1) camera image of the ion crystal (blue or dark gray) and cross-section showing highly non-Gaussian shape of crystal (red or gray). (b) (0,1,1) camera image of the ion crystal. (c) Typical low-overlap setting between MOT (colored or shaded contours) and $1/e^2$ contour of ions (white). (d) Same as (c) but for a higher overlap setting.

taking images of both the MOT cloud and the ions on charge-coupled device (CCD) cameras along two different directions (see Fig. 3), with the total atom number calibrated by means of an absorption measurement. We then calculate the average atomic density experienced by the ions $\langle n \rangle = \int p(\mathbf{r})n(\mathbf{r})d\mathbf{r}$, where $p(\mathbf{r})$ is the normalized ion distribution and $n(\mathbf{r})$ is the local atomic density. The inelastic rate coefficient *K* is obtained by a linear fit of the observed ion decay rate constant $\Gamma = 1/\tau$ vs $\langle n \rangle$.

We note that the ions' kinetic energy is determined by their micromotion throughout the rf cycle [29]. Any ion displacement from the zero of the oscillating electric field, be it due to ion-crystal size or a dc electric field, results in micromotion whose energy easily exceeds that of the thermal motion in the secular potential of the Paul trap. To investigate the dependence of the rate coefficient K on average center-of-mass collision energy E, we vary the latter by loading different numbers of ions into the trap or by intentionally offsetting the ion crystal from the zero of the rf quadrupole field with a dc electric field.

The rate coefficient K(E) is shown in Fig. 4. In the region $E \ge 30 \ \mu \text{eV}$ we calculate *E* from the observed Doppler broadening of the ion fluorescence. For well-compensated traps containing small crystals or single ions, where the Doppler broadening is smaller than the natural linewidth, we determine *E* from the observed correlation between *rf* drive signal and ion fluorescence [29]. For all energies investigated here, the atoms' contribution to the collision energy is negligible.

The data point at the lowest energy $E = 3.1 \ \mu eV$, being an average of 22 single-ion measurements, has large uncertainty in K. It also has large energy uncertainty because the micromotion is sufficiently well compensated for the fluorescence correlation signal to be dominated by noise.

FIG. 4 (color online). Charge-exchange rate coefficient *K* with statistical uncertainty as a function of average collision energy *E* in the center-of-mass frame. Circles, green (or light gray) diamonds, and blue (or dark gray) diamonds represent 172 Yb⁺ + 174 Yb, 172 Yb⁺ + 171 Yb, and 174 Yb⁺ + 172 Yb, respectively. The solid line indicates the theoretical Langevin rate coefficient *K*_{ce} [22], the dashed line assumes a contribution from the *P*-state polarizability [31] (see text). The black arrow indicates the ionization isotope shift between 174 Yb and 172 Yb.

The other data points show observable Doppler broadening and were measured with up to a few thousand ions, resulting in smaller statistical uncertainties. We estimate systematic uncertainties to be a factor of 2 in K due to the difficulty of absolute MOT density calibration, and $\pm 50\%$ in E, due to the nonthermal energy distribution of the micromotion.

The *ab initio* calculated value for the polarizability $\alpha_S = 143 \text{ a.u.} = \epsilon_0 \times 2.66 \times 10^{-28} \text{ m}^3$ [22] for the atomic ${}^{1}S_{0}$ ground state yields $K_{ce} = 5.8 \times 10^{-10}$ cm³ s⁻¹ for ground-state collisions (Fig. 4 solid line). Our experimentally measured value of K = 6×10^{-10} cm³ s⁻¹ in the semiclassical region around $E \sim$ 100 μ eV is thus in good agreement with the Langevin model. The measured K at the highest average collision energy E = 4 meV, close to the transition to the classical scattering region, is somewhat larger than the classical prediction [22,23]. Given the rapid Doppler cooling of the ions, the discrepancy is probably not due to collisioninduced heating and trap loss. Rather, we speculate that it may be caused by averaging our nonthermal (micromotion) energy distribution over an oscillating cross section in the transition region [30]. Doubly excited collisions $Yb^{+*} + Yb^{*}$ cannot occur as the excitation light for ions and atoms is modulated out of phase. For $Yb^{+*} + Yb$ collisions, the Langevin cross section, depending only on the ion's charge and the atom's polarizability, is unchanged, and the charge-exchange process remains resonant, yielding the same value of σ_{ce} . Yb⁺ + Yb^{*} collisions cannot contribute for our collision energies since the atomic transition is tuned out of resonance with the MOT

light at an ion-atom distance $R_0 \approx 30$ nm, larger than $b_c = 2-12$ nm, and modeling of collision trajectories shows that the time required to move between $r = R_0$ and $r = b_c$ exceeds several excited-state lifetimes. For illustration, we have indicated in Fig. 4 the small change in K_{ce} if Yb⁺ + Yb^{*} collisions with $\alpha_P = 500$ a.u. for the 1P_1 state [31] were to contribute at our measured MOT excited-state fraction.

The theoretical value $K_{ce} = \sigma_L v/2$, corresponding to equal binding probability of the electron to the two nuclei, should apply when the collision energy far exceeds the isotope shift of the ionization potential. The latter can be estimated from spectroscopic data of a transition to a state with low electron probability density at the nucleus, such as $4f^{14}6s^2 \rightarrow 4f^{14}6s10d$ [32]. It follows that the ¹⁷²Yb⁺ + $^{174}Yb \rightarrow ^{172}Yb + ^{174}Yb^+$ reaction should be exothermic and release $\Delta E = 2.9 \ \mu eV = h \times 0.7$ GHz. At the substantially larger collision energy $E = 0.21 \text{ meV} \gg \Delta E$ we have investigated various isotope combinations, $^{172}Yb^+$ + 174 Yb, 174 Yb⁺ + 172 Yb, and 174 Yb⁺ + 171 Yb, and find that they all display the same *K* (Fig. 4). We speculate that the data point 172 Yb⁺ + 174 Yb \rightarrow 172 Yb + 174 Yb⁺ at the lowest energy $E = 3.1 \ \mu eV$, corresponding to exothermic collisions with $E \approx \Delta E$, may exhibit an increased rate coefficient compared to K_{ce} .

While it may be impossible to compensate stray fields well enough to reach the *s*-wave scattering limit $E_s =$ 4 peV [6,22], Feshbach and other collision resonances may be observable well above E_s [8]. All the Yb isotopes used here have been cooled to quantum degeneracy in an optical dipole trap [33,34], which would allow the investigation of collision processes between an ion and a Bose-Einstein condensate or Fermi gas. Alternatively, to avoid the large resonant charge-exchange cross section observed here, a different species such as Rb could be used for sympathetic cooling of ions [7,10], or for studying ion impurities in a Bose-Einstein condensate [13].

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