Molecular Fountain

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The resolution of any spectroscopic or interferometric experiment is ultimately limited by the total time a particle is interrogated. Here we demonstrate the first molecular fountain, a development which permits hitherto unattainably long interrogation times with molecules. In our experiments, ammonia molecules are decelerated and cooled using electric fields, launched upwards with a velocity between 1.4 and 1.9 m/s and observed as they fall back under gravity. A combination of quadrupole lenses and bunching elements is used to shape the beam such that it has a large position spread and a small velocity spread (corresponding to a transverse temperature of < 10 μK and a longitudinal temperature of < 1 μK) when the molecules are in free fall, while being strongly focused at the detection region. The molecules are in free fall for up to 266 ms, making it possible, in principle, to perform sub-Hz measurements in molecular systems and paving the way for stringent tests of fundamental physics theories.

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The ability to control atoms using lasers [1–3] has resulted, amongst many other spectacular achievements, in the demonstration of an atomic fountain [4]. In such a fountain, laser cooled atoms are gently pushed upwards on a vertical trajectory and left to fall back under gravity. The long free-fall times permitted by a fountain allow very precise spectroscopy to be performed. In an atomic fountain clock [5], the atoms pass a microwave cavity twice—as they fly up and as they fall back down. The effective interrogation time in such a Ramsey-type measurement scheme includes the entire flight time between the two traversals through the driving field. This interrogation time is typically up to one second. Nowadays, about a dozen fountain clocks are operated at various metrological institutes around the world to realize the SI unit of time, the second, with an accuracy better than 10−15 [6–8]. Atomic fountains are also used in matter-wave interferometry. In such an interferometer, atoms are placed into a superposition of spatially separated atomic states, each of which has an associated quantum-mechanical phase term. When these states are brought back together at a later time, they will interfere with one another in a manner determined by the phase difference accrued between the states. A fountain geometry permits the atoms to remain in the superposition for a long time, allowing a large phase difference to evolve [9]. Such interferometers have been used to determine the Newtonian gravitational constant [10,11], to measure inertial motion [12], and to test quantum superposition over large distances [13]; they have been proposed to be used for testing general relativistic effects [14] and for detecting gravitational waves [15].

A molecular fountain will allow stringent tests of fundamental physics theories [16,17]. The sensitivity of any experiment looking for a frequency shift due to a certain physical phenomenon depends both on the size of the shift, i.e., the inherent sensitivity of the atom or molecule to a certain effect, and on the ability to measure this shift. The complex internal structure of molecules can make them very sensitive to aspects of fundamental physics [18,19]. For instance, in certain heavy molecules such as YbF and ThO, the energy shift due to a permanent electric dipole moment of the electron is thousands of times larger than in heavy atoms [20,21]. Molecules are also used in the study of weak interactions, leading to an anapole moment of the nucleus [22], in the search for a difference in transition frequency between chiral molecules that are each other’s mirror image [23,24], and for constraining a possible variation of the proton-to-electron mass ratio [25,26]. The long interrogation times permitted by a molecular fountain will greatly enhance the sensitivity of these tests. A molecular fountain may also be used for interferometric measurements with molecules in free fall, for instance, to test Einstein’s equivalence principle for rotating objects [27].

Unfortunately, the enhanced sensitivity of molecules comes at a price: The rotational and vibrational degrees of freedom make molecules more difficult to cool [28,29]. Laser cooling, while spectacularly successful at cooling atoms, is much less efficient for molecules [30].

Here, we control the motion of molecules by exploiting the property that polar molecules experience a force in an inhomogeneous electric field [29,31]. A supersonic beam of ammonia molecules is decelerated to rest using a combination of a conventional Stark decelerator, consisting of a series of 100 electrode pairs to which voltages of ±10 kV are applied, and a traveling wave decelerator,
consisting of a series of 336 rings to which oscillating voltages of ±5 kV are applied. For details of the molecular beam decelerator, see Refs. [32,33]. Panel (a) of Fig. 1 depicts the top part of the vertical molecular beam machine, showing the end of the traveling wave decelerator and the quadrupole lens system. In the experiments presented here, ammonia ($^{14}$NH$_3$) molecules in the low-field seeking component of the $J = 1, K = 1$ state are brought to a standstill and trapped inside the traveling wave decelerator, about 10 mm before the end. Once trapped, the molecules are adiabatically cooled to below 1 mK by slowly (∼2 ms) lowering the voltages from 5 to 1 kV. Subsequently, the molecules are launched upwards with a variable acceleration to create a beam with a velocity in the range from 1.4 to 1.9 m/s. Molecules with speed in this range will fly up to 100–180 mm before falling back under gravity.

The slow beam that exits the decelerator is focused using a combination of a linear quadrupole lens and two bunchers. The quadrupole lens consists of four 270-mm-long cylindrical rods with a 20-mm diameter, which are spaced equidistantly on the outside of a 20-mm-diameter circle. The lens is switched on by applying voltages of ±5 kV to adjacent rods, resulting in an electric field that increases linearly away from the molecular beam axis. In electric fields below 10 kV/cm, the Stark shift of $^{14}$NH$_3$ is almost perfectly quadratic and the molecules experience a harmonic potential that results in a transverse oscillation frequency of about 83 Hz. In the longitudinal direction, the molecules are bunched by applying a voltage of 4 kV to two ring electrodes (6 mm aperture) when they pass through each one, resulting in a longitudinal oscillation frequency of about 85 Hz. The time sequence is generated by a computer code that simulates the trajectories through the lens system and whose parameters are optimized using the molecular signal. The largest signal is obtained by focusing the beam at the buncher rings and the detection zone, and collimating the beam (i.e., minimizing the velocity spread) above the second buncher ring using a series of nine voltage pulses of varying duration (six to the quadrupole and three to the buncher). The experiment runs at a repetition rate of 10 Hz when the upgoing beam is recorded, while it runs at 3 Hz for launch velocities between 1.4 and 1.6 m/s and at 2 Hz for launch velocities between 1.7 and 1.9 m/s when the falling beam is recorded.

The red curves in panel (a) of Fig. 1 show a simulation of trajectories through the lens system for a beam launched with a velocity of 1.8 m/s. (b)–(g) Phase-space plots showing the acceptance of the setup in both the longitudinal [(b)–(d)] and transverse directions [(e)–(g)], at three different heights. Note that the axes of panel (g) are scaled by a factor of 10 compared to panels (e) and (f). The grey ellipses show the distribution of the packet of molecules at the exit of the decelerator.

FIG. 1. Experimental setup with simulated trajectories. (a) Schematic view of the top part of the vertical beam machine showing the end of the traveling wave decelerator and the quadrupole lens system. The quadrupole lens consists of four cylindrical rods suspended by two ceramic discs. Two ring electrodes bunch molecules in the $z$ direction. For a view of the inside, part of the quadrupole and the buncher have been cut in the figure. Molecules are ionized by a UV laser and imaged on a phosphor screen located behind a multichannel plate (MCP). The image is recorded using a charge coupled device (CCD) camera and a photo-multiplier tube (not shown). The red curves show a simulation of trajectories through the lens system for a beam launched with a velocity of 1.8 m/s. (b)–(g) Phase-space plots showing the acceptance of the setup in both the longitudinal [(b)–(d)] and transverse directions [(e)–(g)], at three different heights. Note that the axes of panel (g) are scaled by a factor of 10 compared to panels (e) and (f). The grey ellipses show the distribution of the packet of molecules at the exit of the decelerator.
FIG. 2.  Time-of-flight profiles of the rising and falling beams. Number of detected ammonia ions per shot as a function of time after opening the pulsed molecular beam valve for the rising (top panel) and falling (lower panel) beams. Each data point (open circles) is averaged for 400 (1000) shots for the rising (falling) beam. The red lines show fits to Lorentz profiles. Background signals of typically 0.7 ions per shot (rising beam) and 0.15 ions per shot (falling beam) have been subtracted.

At the detection point, ammonia molecules in the $J = 1$, $K = 1$ state are ionized via a $(2 + 1)$ REMPI scheme using pulsed UV laser radiation with wavelength 322 nm. The resulting ions are focused onto a 2D detector using a series of ion lenses. By applying appropriate voltages to the ion optics, ions with the same initial velocity but a different position are focused at the same position on the detector (a technique known as velocity map imaging [34]). A mask on the back side of the phosphor screen is used to transmit light from only the central part of the image onto a photomultiplier tube, thereby collecting light originating from molecules with a small transverse velocity. In this way we can discriminate signal from the rising and falling beam from signal originating from thermal gas in the chamber [35]. A CCD camera is used to optimize the position of the mask.

Figure 2 shows the ion signal as a function of time (open circles), catching the packets of molecules going up (upper panel) and coming down (lower panel). The molecules are launched upwards with a variable acceleration, set by the applied time sequence, to create beams with a velocity from 1.4 to 1.9 m/s. The velocity of the launched molecules and the time they are in free flight ($\Delta t$) are indicated above the TOF profiles. There is a one-to-one correspondence between the measured time-of-flight (TOF) profiles in the upper and lower panels; i.e., the leftmost profile in the upper panel, showing molecules with a velocity of 1.4 m/s while flying upwards, belongs to the rightmost profile in the lower panel, showing these molecules as they fall down. When the velocity of the beam is increased, the signal from the rising beam also increases as the divergence of the beam decreases. The signal from the falling beam, however, decreases as the molecules are launched with a higher velocity; faster molecules fly up higher before
reaching the apex of their trajectory; hence, they spend more time in free fall and spread out more before being recaptured. The signal from the falling beam is maximally 10% of the signal from the rising beam with a launch velocity of 1.5 m/s and drops to below 2% with a launch velocity of 1.9 m/s. This behavior is well reproduced by trajectory simulations when an exponential decay of the number of detected molecules with a 1/e decay time of 350 ms is included to account for losses due to collisions with background gas, nonlinearity of the focusing forces, and Majorana transitions. The shapes of the measured TOF profiles are determined by the focusing elements and are well described by Lorentzian fits (red curves). As expected from the trajectory simulations, the width of the TOF profiles of the falling beam is smaller than that of the rising beam. For instance, the beam that is launched upwards with a velocity of 1.8 m/s has a TOF profile with a full width at half maximum (FWHM) of 0.39 ms (corresponding to a spatial spread of 0.56 mm), whereas the TOF profile of the falling beam at this launch velocity has a FWHM of only 0.23 ms (corresponding to a spatial spread of 0.33 mm).

In conclusion, we have demonstrated a molecular fountain based on a Stark-decelerated molecular beam which enables the study of molecules in free fall for up to 266 ms. A Ramsey-type measurement during such a time interval would yield a line width of about 3 Hz. With the obtained count rate of 0.1 ions per cycle (one cycle takes 333 ms), a measurement time of 400 seconds would suffice to determine the line center to within 1 Hz. This offers the opportunity to measure inversion, and rotational and vibrational transitions in ammonia with unprecedented accuracy. As these transitions have different dependence on the proton-to-electron mass ratio $\mu$, laboratory measurements can be used to set a stringent limit on a possible time variance of $\mu$ due to cosmological expansion [26] or as a result of dark matter [36]. Our method relies on adiabatic cooling and compression of a slow molecular beam using electrostatic lenses and can be applied to any polar molecule that is available at sufficiently high phase-space density, including heavy molecules like SrF [37], BaF, and YbF [38], which are used in low-energy tests of particle physics models.

The molecular fountain demonstrated here took many years to develop. We thank all those who have contributed, in particular, Rob Kortekaas, Congsen Meng, Steven Hoekstra, Leo Huisman, Imko Smid, Jacques Bouma, Joost Buïjs, Ruud van Putten, Ruth Buning, Masatoshi Kajita, Boris Sartakov, André van Roij, Heinz Junkes, Géorg Heyne, Henrik Haak, and Gerard Meijer. This research has been supported by NWO via a VIDI Grant, by the ERC via a Starting Grant, and by the Netherlands Foundation for Fundamental Research of Matter (FOM) (Project No. 12PR2972 and program “Broken mirrors and drifting constants”).

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