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Cryogenic buffer gas beams of AlF, CaF, MgF, YbF, Al, Ca, Yb and NO – a comparison

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ABSTRACT

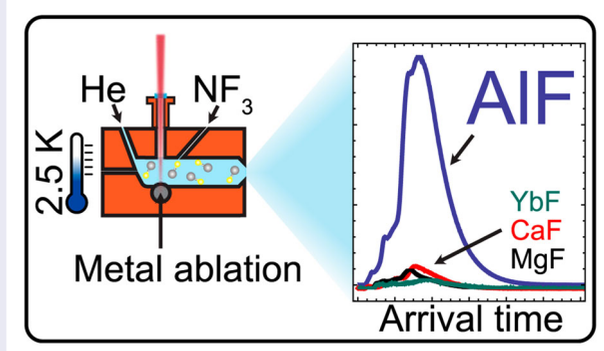
Cryogenic buffer gas beams are central to many cold molecule experiments. Here, we use absorption and fluorescence spectroscopy to directly compare molecular beams of AlF, CaF, MgF and YbF molecules, produced by chemical reaction of laser ablated atoms with fluorine rich reagents. The beam brightness for AlF is measured as 2×10^{12} molecules per steradian per pulse in a single rotational state, comparable to an Al atomic beam produced in the same setup. The CaF, MgF and YbF beams show an order of magnitude lower brightness than AlF and far below the brightness of Ca and Yb beams. The addition of either NF_3 or SF_6 to the cell extinguishes the Al atomic beam, but has a minimal effect on the Ca and Yb beams. NF_3 reacts more efficiently than SF_6 , as a significantly lower flow rate is required to maximise the molecule production, which is particularly beneficial for long-term stability of the AlF beam. We use NO as a proxy for the reactant gas as it can be optically detected. We demonstrate that a cold, rotationally pure NO beam can be generated by laser desorption, thereby gaining insight into the dynamics of the reactant gas inside the buffer gas cell.

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1. Introduction

Cryogenic buffer gas cooling is a versatile technique to produce intense atomic and molecular beams with a low forward velocity [1–4] and allows efficient cooling of the rotational and vibrational degrees of freedom in molecules. Buffer gas cooling is now routinely used for precision spectroscopy and measurements [5–8], to trap molecules using magnetic fields [9–11] and electric fields [12], to study collisions at low temperatures

[13–16], and to provide slow atoms and molecules for magneto-optical [17–24] and electro-optical traps [25], in which the particles can be cooled to μK temperatures.

The species that have been cooled using a cryogenic buffer gas range from light atoms such as Li [26] and K [23] to heavy atoms such as Yb [27], exotic atoms such as Tm, Er and Ho [17], diatomic molecules [28, 29], including radicals [2, 27, 30, 31], small polyatomic molecules [32–35] and large, complex molecules such as

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functionalised arenes [36], Nile Red [37], and C_{60} [38]. Diatomic metal fluoride molecules with a $^2\Sigma^+$ electronic ground state, such as MgF, CaF, SrF, BaF and YbF, have attracted attention due to their suitability for direct laser cooling [39–42] and precision measurements [7, 43]. Buffer gas molecular beams of these molecules have been reported and characterised [4, 13, 43, 44]. Recently, the molecules AlF and AlCl, which have $^1\Sigma^+$ electronic ground states, have been produced for the first time in buffer gas sources and are now the subject of laser cooling efforts [45–49]. These molecules are fundamentally different to the $^2\Sigma^+$ ground state molecules that have been laser cooled thus far, and we recently reported a bright molecular beam of AlF and fast optical cycling on its $A^1\Pi \leftarrow X^1\Sigma^+$ transition [46].

It is often challenging to determine whether the molecular yield in one experiment differs from another because of the choice of molecule, the detection method or conditions of the molecular beam source which may deteriorate over time. Here, we investigate beams of AlF, CaF, MgF, YbF, NO, Al, Ca and Yb in the same apparatus and use absorption and laser induced fluorescence spectroscopy to characterise them. By comparing the brightness of the molecular and atomic beams, we gain new insight in the production efficiency for the various species. We also describe observations which we consistently find produce desirable molecular beam properties and better long-term performance.

2. Experimental setup

2.1. Molecular beam source

Figure 1(a) shows a sketch of the molecular beam source. We use a two-stage closed-cycle He cryocooler (Sumitomo RDK-415DP (with helium pot) with F-50H compressor) to cool the source to about 2.5 K. The first stage cools the aluminium radiation shields to about 40 K. The buffer gas cell, shown in Figures 1(c) and 1(d), is attached to the second stage and surrounded by a copper box, whose internal walls are coated with activated charcoal. The charcoal acts as an efficient sorption pump for helium gas at temperatures below 10 K. The operating pressure of the source chamber under cryogenic conditions is about 1×10^{-8} mbar, measured outside the radiation shields, and typically increases to 3×10^{-8} mbar when 2 standard cubic centimetres per minute (sccm) of helium is flowed through the cell.

To increase the effective pumping area, we attach charcoal-coated copper fins to the top internal face of the copper box. An additional single fin is mounted along the molecular beam direction (shown in Figure 1(b)). While the top-fins have only a minor influence on the

beam properties, the fin along the molecular beam helps maintaining a high molecular beam flux. Saturation of the activated charcoal with reaction products and ‘ablation dust’ – indicated by a discoloration of the charcoal around the aperture – can reduce the downstream flux by up to a factor of 10, without noticeably changing absorption outside the cell. In this case, the fin can be easily replaced. Increasing the distance between the cell and the first charcoal-coated surface mitigates this problem without affecting the downstream flux considerably. Cartridge heaters can be used to heat the set-up to room temperature within about 5 hours.

A sketch of the buffer gas cell is shown in Figures 1(c) and 1(d). It is based on the design presented by Truppe *et al.* [4], with minor changes to facilitate machining and cleaning. The cell is machined from oxygen-free copper, has a circular bore with a 10-mm diameter, and its length is variable between 30 and 60 mm, using extension pieces. Windows on the extensions allow in-cell absorption spectroscopy. For the experiments presented in this study, we used a 40-mm-long cell. The exit aperture of the cell has a diameter of 4 mm. The multi-species metal ablation target is attached to a copper adaptor with a fine thread, which enables rotation of the target via a mechanical vacuum feed-through. We find that translating the target is superior to translating the ablation-laser beam and results in more consistent molecular beam properties. The relatively small bore size of the cell results in a short extraction time for molecules and thus short molecular pulses [4, 50], ideal for Stark deceleration and chirped laser slowing. The helium gas is pre-cooled in two copper pipe bobbins that are attached to each of the two cooling stages (see Figure 1(a)). With a buffer gas flow rate of 1 sccm the in-cell helium density is $7 \times 10^{14} \text{ cm}^{-3}$, where we assumed an ideal vacuum conductance of the exit aperture.

The output of a pulsed Nd:YAG laser (Continuum Minilite II) with a pulse energy of up to 40 mJ and a pulse length of 5–7 ns is gently focused to produce a 0.7-mm-diameter spot size on the target. The Nd:YAG fires with a repetition rate of 1 Hz for all measurements in this study, and firing the ablation light defines $t = 0$ for each molecular pulse. A higher repetition rate leads to an increased heat load, resulting in an increase of the rotational temperature of the molecular beams. We ablate a metal target and introduce a fluorine donor gas (SF_6 , NF_3 , CF_4 , XeF_2) into the cell through a copper capillary that is thermally insulated from the cell. We also use this capillary to inject NO molecules. The temperature of the capillary is kept at about 120 K.

We found over the course of our experiments that the beam properties of AlF are sensitive to the cell having clean internal surfaces. In particular, cleaning the

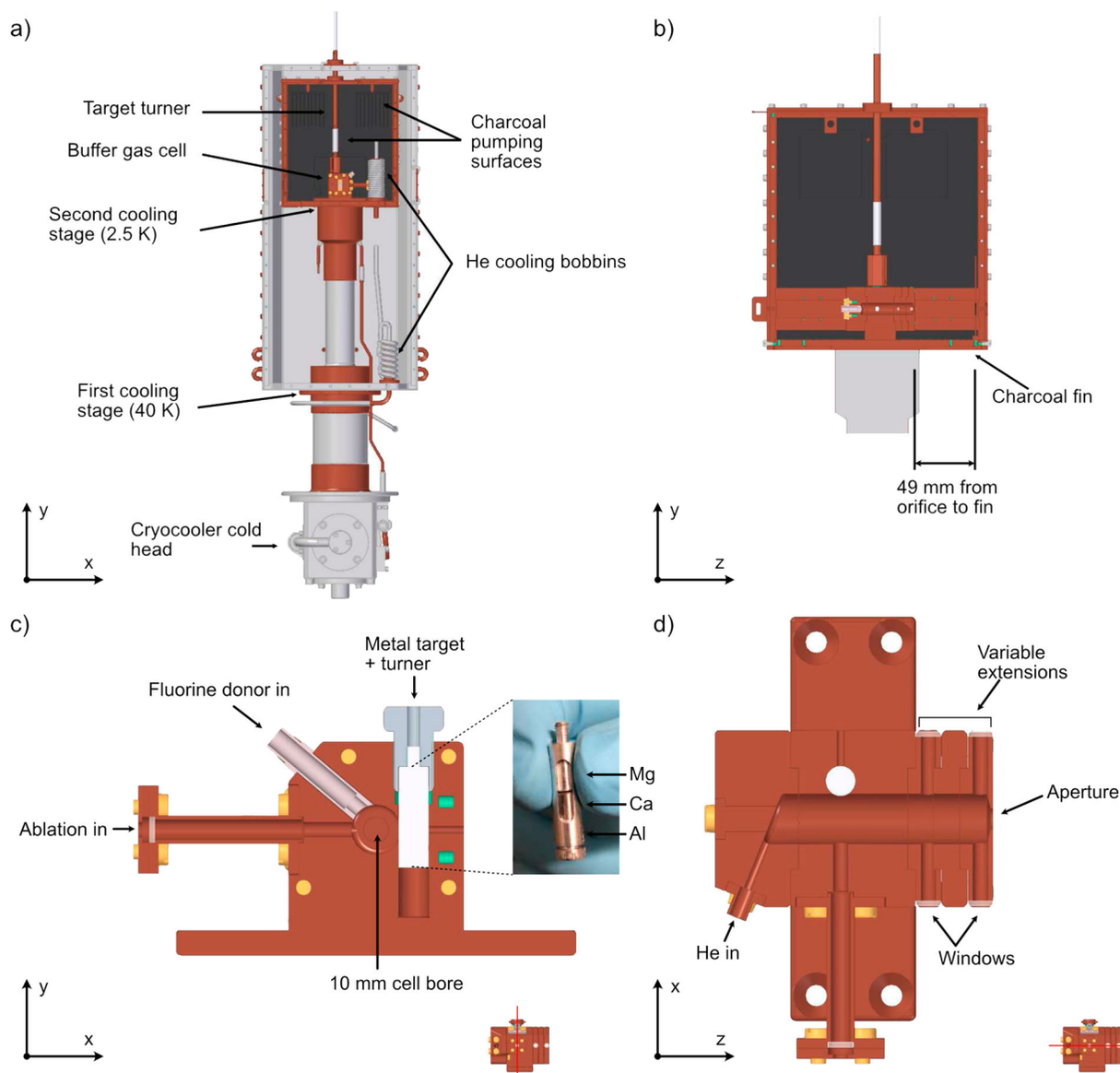


Figure 1. (a) A diagram of the molecular beam source. The molecular beam travels into the page. The front plates have been omitted to give a complete view of the inside of the source chamber. (b) Cut through the charcoal-coated shield along the direction of the molecular beam. (c) Cut through the buffer gas cell, perpendicular to the molecular beam direction, showing the region where the metal is ablated and reacts with a fluorine donor gas. The inset is a photograph of the multi-species metal target. (d) Cut along the molecular beam direction showing the path of the He buffer gas.

cell in an ultrasonic bath with Citranox acidic detergent (Alconox, Inc.) results consistently in a slow beam, with a small spread of arrival times seen in downstream fluorescence. With increasing number of ablation laser pulses, the time-of-flight broadens, and secondary higher velocity peaks appear. Removal of ablation products by wiping the inside of the cell leads to some recovery, but the efficacy of this method reduces gradually. Recovery of the original behaviour is reliably achieved by ultrasonic cleaning to expose the metal surface. We show examples of this behaviour for AlF in Figure 2. Coating the surfaces of the cell with a thin layer of gold to ensure a chemically

inert surface was unsuccessful at preventing the observed degradation.

A schematic of the full experimental setup used for this study is shown in Figure 3(a). We use a multi-species target to quickly change the metal at the ablation spot, and therefore the species observed in the buffer gas beam. We can probe the atoms and molecules by absorption spectroscopy with a weak laser beam, both inside the cell and at a variable distance 0–2 cm from the exit aperture, detecting the transmitted light with a photodiode. The extraction efficiency as measured from the peak absorption of CaF, AlF and MgF inside and directly outside

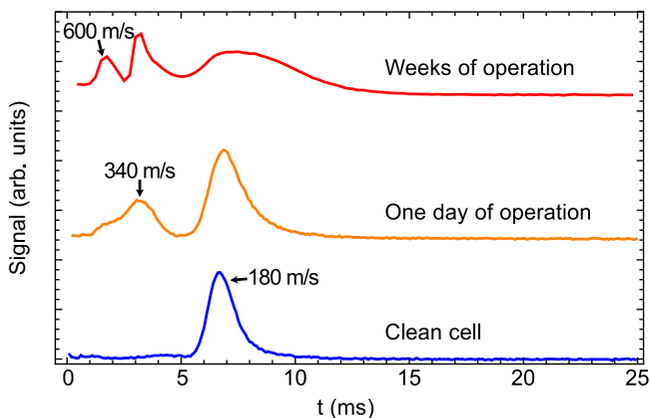


Figure 2. Fluorescence time of flight (TOF) traces of AIF with a buffer gas cell after different periods of use. From bottom to top: Cell freshly cleaned with Citranox detergent; same cell after 1 day of operation using the same ablation spot; same cell geometry after weeks of operation.

the cell is about 10 – 15%. The Doppler width measured inside and outside the cell is the same, but downstream of the cell, an increase of about a factor 2 in the transverse velocity spread is observed. We discuss the implications of this broadening for beam brightness measurements later.

Two laser-induced fluorescence (LIF) detectors enable us to measure the brightness of the collimated molecular beam, its velocity distribution and the rotational state distribution. The first detector, LIF-1, is located at $L_1 = 23$ cm downstream of the cell exit. It is primarily used as a region for optical pumping, for instance to determine the velocity of the molecules using a pump-probe method [46]. The second zone, LIF-2, is located $L_2 = 44$ cm downstream of the cell exit and is used for spectroscopy and beam brightness measurements. The entrance to LIF-2 is restricted by a square aperture of size 2×2 mm² to reduce the transverse velocity spread of the molecular beam to about 1.5 ms⁻¹. This reduces the Doppler broadening in our fluorescence excitation spectra and defines the interaction volume with the probe laser beam.

Figure 3(b) shows the collection optics of LIF-2. The fluorescence of the molecular beam is collimated by a 50-mm plano-convex lens (focal length 66 mm at $\lambda = 227$ nm) and concave mirror mounted in vacuum, and focussed onto a Hamamatsu R928 photomultiplier tube (PMT) by a second externally mounted lens. We collimate the excitation laser light to a e^{-2} intensity diameter of 2–3 mm, to ensure that all molecules that enter the detection zone are addressed by the excitation laser light. The aperture of the detector subtends a solid angle $\delta\Omega = 2.1 \times 10^{-5}$ sr at the buffer gas cell exit and the peak optical power reaching the PMT when probing the

brightest atomic beam is about 1 nW. The PMT photocurrent is sent to a transimpedance amplifier with a current-to-voltage conversion factor of 10^5 VA⁻¹.

We use the same optical setup in the fluorescence detector for all species. The wavelengths of the fluorescence light span from 227 to 606 nm and the collection efficiency of the LIF detector varies across this range due to differences in the PMT quantum efficiency and chromatic aberration of the detection optics. To compare the detection efficiency of the PMT across the range of emission wavelengths, we illuminated the PMT directly with about 1 nW of laser light at 227, 360, 452 and 606 nm. This was done by calibrating a $\sim 10^{-3}$ neutral density filter at incident powers of order 1 mW, and then using the filter to attenuate to 1 μ W, measured using a calibrated optical power meter. To account for differences in collection efficiency, we use ray tracing simulations combined with the measured transmission and reflection losses of the optical components.

2.2. Molecular beam brightness measurement

The figure of merit for this study is the number of molecules produced per unit solid angle per pulse, in a given rovibrational level. To estimate this for the different species, we use both absorption and fluorescence measurements.

Absorption measurements have the advantage of directly measuring the column density of the species of interest, but sufficient density is only available inside or near the exit of the buffer gas cell. The Lambert–Beer law relates the density, $n(t)$, to the measured absorption,

$$-\ln(I_t/I_0) = n(t)\sigma(\nu)z. \quad (1)$$

Here, $\sigma(\nu)$ is the absorption cross-section, z is the interaction length, and I_t and I_0 are the intensity transmitted with and without the absorbing species present, respectively. In the absence of Doppler or hyperfine broadening, the maximum value of $\sigma(\nu)$ is the resonant absorption cross section σ_0 , which for a transition $J' \leftarrow J$ is given by [51]

$$\sigma_0 = \frac{\lambda^2}{2\pi} \frac{2J' + 1}{2J + 1} \frac{A_{ji}}{\Gamma}. \quad (2)$$

Here, J and J' are the ground and excited state total electronic angular momentum quantum numbers for atoms and the total electronic angular momentum including the end-over-end rotation for molecules, A_{ji} is the Einstein A coefficient of the resonant transition, and Γ is the total spontaneous decay rate. In the presence of broadening mechanisms, the peak absorption cross-section σ_D can be calculated using the relation $\int \sigma(\nu)d\nu = \sigma_0\Gamma\frac{\pi}{2}$, where

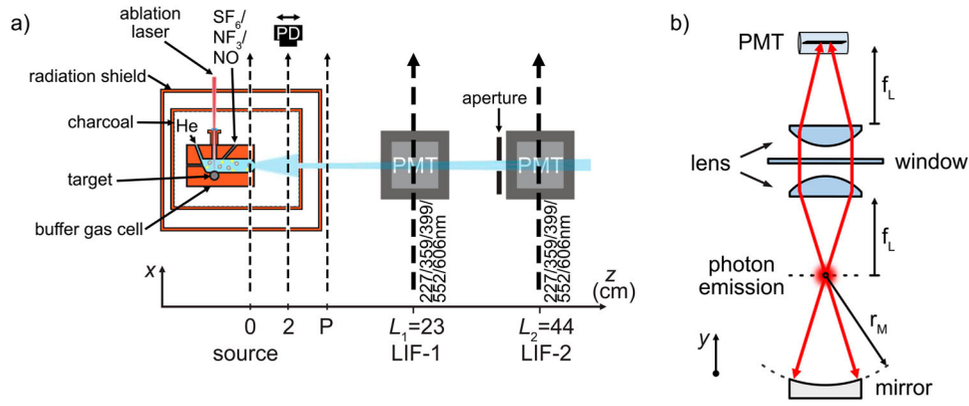


Figure 3. (a) Schematic view of the experimental setup used in this study to characterise the atomic and molecular beams. A photodiode (PD) is used for absorption and photomultiplier tubes (PMTs) for fluorescence spectroscopy. P marks the first downstream position where optical pumping can occur. Closer to the buffer gas cell, collisions with helium redistribute the rotational state populations. (b) Optical layout of the fluorescence detector.

the integral over frequency ν covers the full absorption spectrum.

We estimate the total number, N , by integrating the flux outside the cell over time,

$$N = \nu_f a_{\text{aperture}} \int n(t) dt = \frac{\nu_f a_{\text{aperture}}}{z \sigma_D} \int -\ln(I_t/I_0) dt, \quad (3)$$

where a_{aperture} is the area of the exit aperture of the cell, z is assumed to be the aperture diameter and ν_f is the forward velocity which can be estimated from measurements downstream of the source. The brightness of the beam in absorption, \mathcal{B}_{abs} , is then calculated as

$$\mathcal{B}_{\text{abs}} = \frac{N}{\Delta\theta^2} \quad (4)$$

with $\Delta\theta = 2 \tan^{-1}(\nu_t/2\nu_f)$ the angular divergence of the beam, and ν_t the full-width at half maximum (FWHM) of the transverse velocity distribution, which is obtained from the absorption spectrum.

Fluorescence measurements, on the other hand, are much more sensitive and can be carried out far from the cell, but are typically difficult to calibrate on an absolute scale. For an optically closed transition, the number of photons scattered depends on the effective intensity and interaction time of the laser, and is thus sensitive to the laser beam profile and velocity of the species of interest. If the fluorescence can be saturated by optically pumping population to another state, a known number of photons \mathcal{N} is absorbed and only the overall detection efficiency $\epsilon(\lambda)$ remains uncertain. When exciting from an initial state i to an excited state j , \mathcal{N} is given by

$$\mathcal{N} = \frac{\Gamma}{\Gamma - A_{ji}}, \quad (5)$$

where A_{ji} is the Einstein A coefficient for decay back to the initial state and $\Gamma = \sum_n A_{jn}$ for all states n is the total

decay rate of the excited state j . If N molecules enter the fluorescence detector, the number of photons detected is then simply $N\mathcal{N}\epsilon(\lambda)$. The on-axis brightness is then calculated as

$$\mathcal{B}_f = \frac{N}{\delta\Omega}. \quad (6)$$

This method can be easily applied to diatomic molecules with diagonal Franck–Condon factors, where it is straightforward to optically pump molecules using rotationally open electronic transitions, and \mathcal{N} can be predicted with knowledge of the relevant Hönl–London factors. For atoms, optically open electronic transitions are commonly available, but often the coefficients A_{ji} are not known accurately. It is important to consider the influence of the fluorescence emission pattern on $\epsilon(\lambda)$. Unless stated otherwise, we use linearly polarised light and state fluorescence with the polarisation axis forming an angle $\theta_m = 54.7^\circ$ with the detector axis, the so-called ‘magic angle’.

2.3. Laser systems

We use two different continuous laser systems for this study. To detect Al, Ca, Yb, AlF, MgF and NO, we use a Ti:Sapphire laser (MSquared Solstis), whose output is frequency doubled in successive enhancement resonators containing a nonlinear optical crystal. The linewidth of the fundamental light is less than 400 kHz. A single stage of frequency doubling is sufficient to generate light near 360 nm (MgF) and 399 nm (Yb), and we use two successive doubling stages to generate UV light near 227 nm (AlF, NO, Al, Ca). The second laser system is a Coherent 899 ring dye laser (RDL) which generates light near 606 nm for the detection of CaF and near 552 nm for the detection of YbF. The laser linewidth is around 1 MHz.

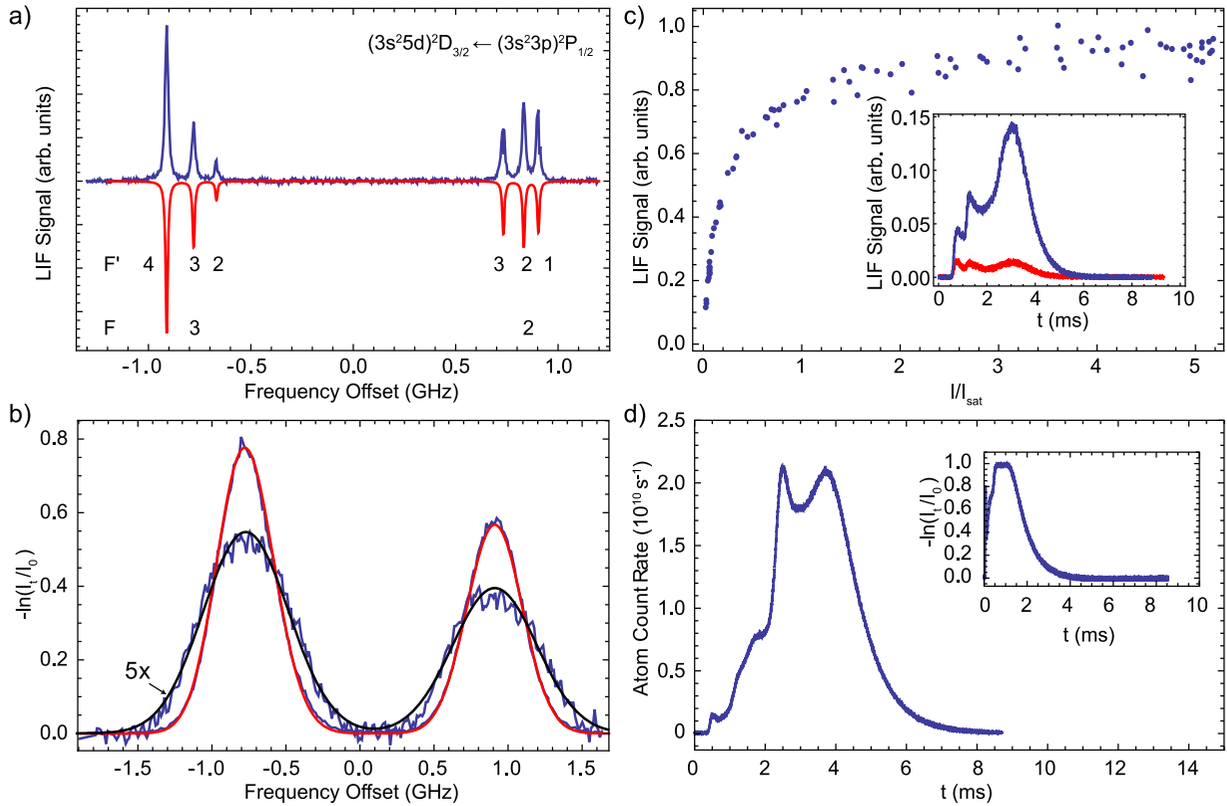


Figure 4. (a) Hyperfine-resolved LIF spectrum of the $(3s^2 5d)^2 D_{3/2} \leftarrow (3s^2 3p)^2 P_{1/2}$ transition of Al near 227.6 nm. The experimental spectrum is shown in blue. A simulated spectrum using Equations (7) and (8) and the known linewidth of 12.8 MHz is shown in red. The ground-state (F) and excited-state (F') hyperfine quantum numbers of the transitions are given below the simulated spectrum. (b) Aluminium absorption spectra recorded directly outside the cell exit and 10 mm downstream (magnified by a factor 5). The solid curves in red and black are fits to the data (blue) using the known spectroscopic constants and a Gaussian lineshape to determine the Doppler broadening. The Doppler broadening increases for the downstream position. (c) Fluorescence saturation curve of the $(3s^2 5d)^2 D_{3/2}, F' = 4 \leftarrow (3s^2 3p)^2 P_{1/2}, F = 3$ line of Al, demonstrating that we can saturate the fluorescence by optical pumping. The inset shows the LIF TOF without (blue) and with (red) a pumping laser on the same transition in LIF-1. (d) Time of flight traces of the Al beam under saturated fluorescence of the $(3s^2 5d)^2 D_{3/2}, F' = 4 \leftarrow (3s^2 3p)^2 P_{1/2}, F = 3$ transition to determine the total flux through a $2 \times 2 \text{ mm}^2$ aperture 440 mm from the source aperture. The inset shows the absorption trace directly outside the cell.

The laser frequencies are monitored with a HighFinesse WS8-10 wavemeter calibrated using a temperature-stabilised HeNe laser. Details on the absolute accuracy of the wavemeter can be found in reference [52].¹

3. Atomic beam measurements

We first discuss measurements on the atomic buffer gas beams of Ca, Al and Yb. These provide an important reference when comparing with the molecular beams.

3.1. Aluminium

To characterise the Al atomic beam, we use the $(3s^2 5d)^2 D_{3/2} \leftarrow (3s^2 3p)^2 P_{1/2}$ transition near 226.4 nm, conveniently close to the $A^1\Pi \leftarrow X^1\Sigma^+$ transition in AlF. Due to the fine structure splitting of the electronic ground state, there exists a $(3s^2 3p)^2 P_{3/2}$ state approximately 112 cm^{-1} ($\sim 160 \text{ K}$) above the ground state. By

probing the nearby $(3s^2 5d)^2 D_{5/2} \leftarrow (3s^2 3p)^2 P_{3/2}$ transition, we find that about 1% of the Al atoms exiting the source initially occupy the $^2 P_{3/2}$ state, and they appear only with high velocities greater than 500 ms^{-1} . We also exclude aluminium dimer formation in the source, as probing the $E^3\Sigma_g^- \leftarrow X^3\Pi_{0u}$ transition in Al_2 near 366.6 nm [53] gave no observable fluorescence in LIF-2. We therefore assume the population in the $(3s^2 3p)^2 P_{1/2}$ state represents the full atomic population.

The lifetime of the $(3s^2 5d)^2 D_{3/2}$ excited state, τ_{ex} is 12.3 ns ($\Gamma/2\pi = 12.8 \text{ MHz}$) [54]. To our knowledge, the hyperfine structure of the $(3s^2 5d)^2 D_{3/2} \leftarrow (3s^2 3p)^2 P_{1/2}$ transition has not been measured. The nuclear spin $I(I_{\text{Al}} = 5/2)$ couples with the electronic total angular momentum J to give eigenstates labelled by the total angular momentum quantum number F . Figure 4(a) shows a hyperfine-resolved experimental spectrum of the $(3s^2 5d)^2 D_{3/2} \leftarrow (3s^2 3p)^2 P_{1/2}$ transition, obtained using about $100 \mu\text{W}$ of linearly polarised laser light. The larger

splitting ~ 1.5 GHz arises from the known ground state hyperfine interaction, and the interaction in the excited state leads to a further splitting of a few hundred MHz. The hyperfine levels are shown in Figure 5. We fit the hyperfine energies, $E_i(J, I, F)$, to the equation [55],

$$E_i = \frac{A_i C}{2} + B_i \frac{\frac{3}{4}C(C+1) - I(I+1)J(J+1)}{2I(2I-1)J(2J-1)}, \quad (7)$$

where A_i is the interaction strength between the nuclear spin I and the electronic angular momentum J , B_i is the quadrupole interaction coefficient, necessarily zero for $J < 1$, and $C = F(F+1) - I(I+1) - J(J+1)$. The index i labels the electronic state. For the ground state, we find $A = 501.6(1.2)$ MHz, consistent with the literature value [56]. For the excited state, we determined $A = -33.9(4)$ MHz and $B = 1.9(2.5)$ MHz. Shown pointing downwards in red in Figure 4 is a simulated spectrum using the known natural linewidth, where we assumed equal population of the ground state Zeeman sub-levels and isotropic emission of the fluorescence light. Under this assumption, the relative intensities of the hyperfine lines, $I_{F',F}$, are given by

$$I_{F',F} = \left| \sqrt{(2F'+1)(2F+1)} \times \begin{Bmatrix} J' & F' & I \\ F & J & 1 \end{Bmatrix} \right|^2. \quad (8)$$

This formula describes the data well. The fitted FWHM of the lines is found as 19 MHz, slightly larger than the measured linewidth, which we attribute to Doppler broadening, residual Zeeman shifts due to a small ambient magnetic field (~ 1 G) in the detection region, and the effect of optical pumping.

The measured hyperfine structure can then be used to determine the Doppler broadening present in absorption measurements. Figure 4(b) shows two such absorption spectra, directly outside the exit of the buffer gas cell and 10 mm further downstream. There and in the following absorption plots, we show the optical depth $-\ln(I_t/I_0)$ on the y -axis. We fit the data using the measured excited state hyperfine structure convolved with a Doppler broadening term with FWHM Γ_D . The transverse velocity width, $v_t = \Gamma_D \lambda / (2\pi)$, is approximately 90 ms^{-1} at $z = 0$ mm and 150 ms^{-1} at $z = 10$ mm. This increase of about a factor 1.7 indicates collisions of the Al atoms with (somewhat hotter) He buffer gas outside the cell, increasing the divergence of the beam. Using the downstream value for the transverse velocity spread, we estimate the brightness of the atomic beam as $4.1 \times 10^{12} \text{ sr}^{-1}$ per pulse.

To measure the Al beam brightness in fluorescence, we use the scheme shown in Figure 5. We excite the $F' = 4 \leftarrow F = 3$ hyperfine line, which optically pumps 95% of the population to the $(3s^2 3p)^2 P_{3/2}$ state. The remaining

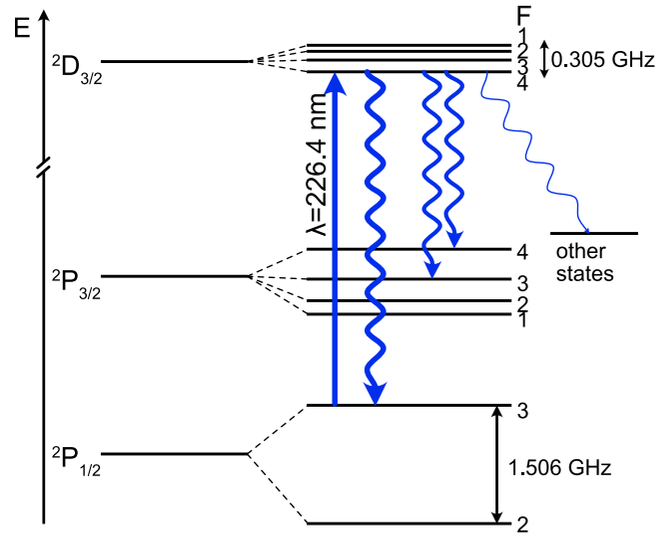


Figure 5. Scheme of the relevant electronic states and hyperfine levels of Al. Electronic energies are not to scale, the relative hyperfine splittings are shown to scale. We use the ${}^2D_{3/2}, F' = 4 \leftarrow {}^2P_{1/2}, F = 3$ transition to detect Al atoms.

5% of the atomic population is pumped to higher lying 2P and 2F electronic states, which cannot directly decay back to the electronic ground state by dipole allowed transitions. We neglect the effect of these states other than the fact that they slightly increase the optical pumping probability. From the ratio of the Einstein A coefficients of these transitions, we calculate that we optically pump population in the $F = 3$ hyperfine level after absorbing on average 6.1(1.1) photons. In Figure 4(c), the fluorescence signal on this transition in LIF-2 is plotted against the laser intensity expressed in terms of the two-level saturation intensity $I_{\text{sat}} = \pi \hbar c \Gamma / (3\lambda^3)$, where Γ is the spontaneous decay rate and λ is the transition wavelength. The fluorescence saturates, demonstrating that we can use this transition to accurately determine the average number of photons emitted by the atoms. The inset shows the change in the fluorescence signal when the same laser light is used in LIF-1 to optically pump the atoms before detecting the remaining atoms in LIF-2. The pumping efficiency is approximately 90%, increasing with arrival time at the detector. After accounting for the $F = 2$ ground state population which is not detected, we arrive at a total brightness of Al atoms of $3.0 \times 10^{12} \text{ sr}^{-1}$ per pulse. This is within 30% of the number derived from absorption measurements in the source when we assume the transverse velocity as measured 10 mm downstream from the source. If, instead, we used the transverse velocity spread as measured right outside the cell, we would overestimate the beam brightness by a factor of 3.5. This clearly shows that absorption measurements close to the

cell can be misleading and typically will provide an upper bound to the beam brightness observed downstream.

3.2. Calcium

To detect calcium, we use the $(3p^64snp)^1P_1 \leftarrow (3p^64s^2)^1S_0$ transition at 227.6 nm. In Figure 6(a), the relevant energy levels for our study of the Ca atomic beam are shown. Exciting the atoms to the 1P_1 state primarily pumps the population to the metastable $(3p^63d4s)^1D_2$ state, with emission of a 452-nm photon. There are additional decay channels to higher lying 1S and 2D states, but these emit at wavelengths longer than 940 nm, outside the detection range of the PMT. The direct decay channel back to the 1P_0 state emits a photon at 227.6 nm. The Einstein A coefficient for this decay channel has been measured as $A(^1S_0) = 2.84(39) \times 10^7 \text{ s}^{-1}$ [57]. We collect both 227 nm and 452 nm photons and account for the detector sensitivity at the different wavelengths. The two decay paths have different fluorescence emission patterns, which are demonstrated by monitoring the LIF as a function of the angle of linearly polarised excitation light, θ . Figure 6(b) shows the detected fluorescence against θ with and without a filter blocking the visible light covering the PMT. The emission pattern of the 227 nm decay is proportional to $\sin^2 \theta$, whereas the 452 nm emission is proportional to $1 + \frac{1}{6} \sin^2 \theta$, and almost isotropic. This leads to the reduced contrast in the fluorescence as a function of θ when detecting both the visible and UV fluorescence compared to the UV alone. The solid lines in Figure 6(b) fit to the function $a + b \sin^2 \theta$, and from these measurements and the detector efficiencies, we estimated that the ratio of Einstein A coefficients of the two transitions, $A(^1D_2)/A(^1S_0) = 2 \pm 0.3$. Using the measured value for $A(^1S_0)$, we find a total decay rate of $A(^1S_0) + A(^1D_2) = 8.5(1.4) \times 10^7 \text{ s}^{-1}$, larger than the measured spontaneous emission rate of $\Gamma = 1/\tau = 6.37 \times 10^7 \text{ s}^{-1}$ [58]. We assume that atoms in the Ca beam emit on average $\mathcal{N}_{227} = 0.5$ and $\mathcal{N}_{452} = 1$ photons at the two detection wavelengths when the fluorescence is saturated, but note that these values are approximate.

The fluorescence spectrum is shown in Figure 7(a), where we clearly resolve four naturally abundant Ca isotopes. The transitions of the two low-abundance bosonic isotopes, ^{42}Ca and ^{44}Ca , are shifted relative to ^{40}Ca by +1.43(1) GHz and +2.755(8) GHz respectively. We hereon focus on ^{40}Ca , which is relevant for comparison with ^{40}CaF .

In Figure 7(b), absorption spectra of the most abundant isotope peak of the $(3p^64snp)^1P_1 \leftarrow (3p^64s^2)^1S_0$ transition are shown directly at the cell orifice and 20 mm downstream. We again observe an increase in

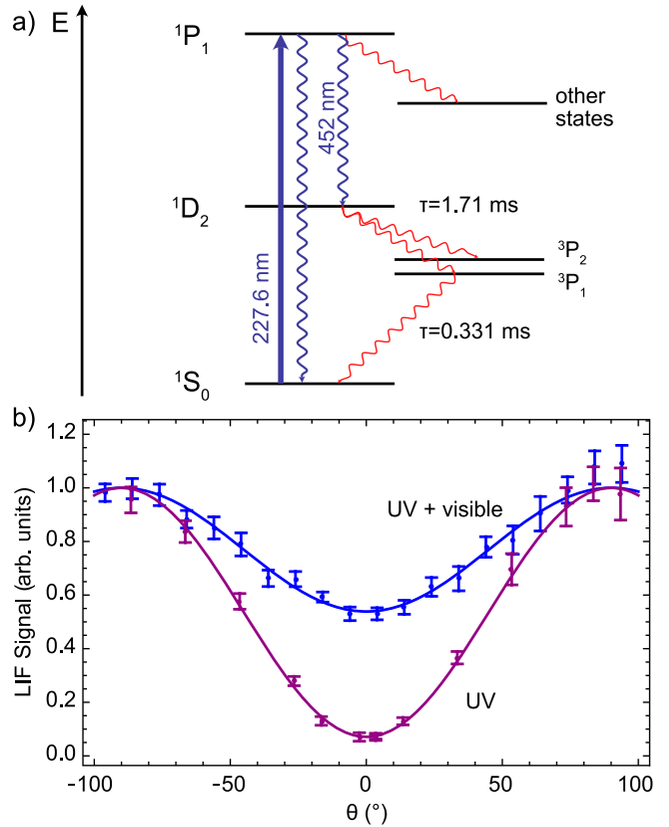


Figure 6. (a) Scheme of the relevant energy levels of calcium. (b) Polarisation dependence of fluorescence of the Ca atomic beam. The UV fluorescence, isolated with a 227-nm bandpass filter, shows strong anisotropy due to the dipole emission pattern. The combined fluorescence in the UV and visible shows reduced emission anisotropy as the visible light emission is nearly isotropic.

the transverse velocity width at the downstream position, from 75 ms^{-1} to 120 ms^{-1} .

In Figure 7(c), the fluorescence signal in LIF-2 is plotted against peak laser intensity, demonstrating saturation of the fluorescence. The inset to the figure shows a pump-probe measurement, from which it appears that we are only able to pump 70% of the atoms. However, the finite lifetime of the 1D_2 state of 1.71 ms [59] is comparable to the flight time between the detectors. Partial repopulation of the ground state then occurs via the 3P_1 state, which is populated with a probability of 83% and decays to the 1S_0 state with a lifetime of 0.331 ms. Consistent with this, the pumping efficiency increases for earlier arrival times at the detector, where the flight time between pump and probe is shorter. We therefore conclude that the population can be fully optically pumped for the purposes of measuring the beam brightness.

Figure 7(d) shows a time of flight trace of the Ca atoms in LIF-2. We calculate that 1.9×10^8 atoms pass

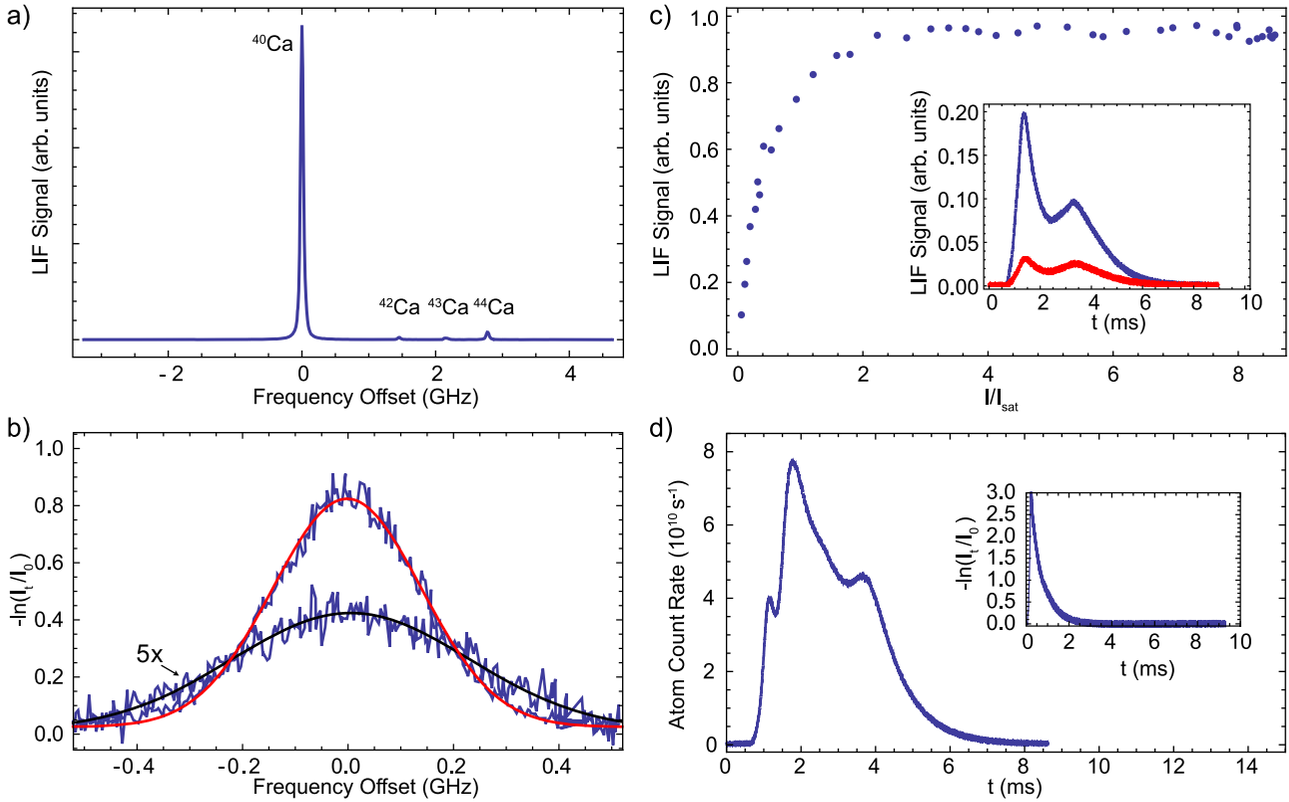


Figure 7. (a) LIF spectrum of the $(3p^64snp)^1P_1 \leftarrow (3p^64s^2)^1S_0$ line of Ca. The observed isotope peaks are labelled. (b) Absorption spectra of the $(3p^64snp)^1P_1 \leftarrow (3p^64s^2)^1S_0$ transition in ^{40}Ca , recorded directly outside the cell exit (Gaussian fit shown in red) and 20 mm downstream (black). (c) Fluorescence saturation curve of the $(3p^64snp)^1P_1 \leftarrow (3p^64s^2)^1S_0$ transition of Ca. The inset shows the LIF TOF without (blue) and with (red) a pumping laser tuned the same transition frequency. (d) Fluorescence and absorption TOF traces of the Ca beam, using light resonant with the $(3p^64snp)^1P_1 \leftarrow (3p^64s^2)^1S_0$ transition of ^{40}Ca .

through the detector, corresponding to an on-axis brightness of $9.2 \times 10^{12} \text{ sr}^{-1}$ per pulse. Using the absorption measurement (shown in the inset of Figure 7 (d)), we find 7.4×10^{12} per pulse, in reasonable agreement. We conclude that the Ca atomic beam is brighter than the Al atomic beam, by about a factor 2.5.

3.3. Ytterbium

We use the $(6s6p)^1P_1 \leftarrow (6s^2)^1S_0$ transition near 399 nm to study the Yb atomic beam. The excited state lifetime $\tau_{\text{Yb}} = 5.464 \text{ ns}$ ($\Gamma/2\pi = 29.13 \text{ MHz}$) [60]. An absorption spectrum taken outside the cell is shown in Figure 8(a). Ablation of Yb metal is highly efficient and the absorption saturates for the most abundant isotopes. We are able to measure the absorption of the ^{168}Yb isotope (Figure 8(a), inset), and use this to estimate the number of atoms and to extract the Doppler width of the beam. The transverse velocity width FWHM at the cell orifice is 42 ms^{-1} and 60 ms^{-1} when measured 18 mm downstream. After taking into account the relative natural abundances of $^{168}\text{Yb} : ^{174}\text{Yb}$ (1:245), we estimate the

brightness of the ^{174}Yb isotope in absorption is $1.2 \times 10^{13} \text{ sr}^{-1}$ per pulse.

Figure 8(b) shows a time-of-flight profile obtained when probing the ^{174}Yb isotope with $40 \mu\text{W}$ of laser power, to be compared later with the ^{174}YbF molecule. The $(6s6p)^1P_1 \leftarrow (6s^2)^1S_0$ transition is optically closed, and therefore not suitable for precise fluorescence measurements of the beam brightness. We simply estimate the brightness observed downstream by modelling the interaction of atoms with the mean forward velocity with the intensity profile of the probe light. Assuming a velocity of 150 ms^{-1} , we estimate the beam brightness as $5.0 \times 10^{12} \text{ sr}^{-1}$ per pulse, in LIF-2, in reasonable agreement with the absorption measurements. We note that the total brightness considering all isotopes is at least 1.6×10^{13} per steradian per pulse, making the Yb beam the brightest of the three atomic species tested.

3.4. Addition of fluorine donor gases

Figure 9 shows the fluorescence TOF profiles of the atomic beams before and directly after introducing a flow of 0.002 sccm NF_3 to the cell. For the Al atoms, there is

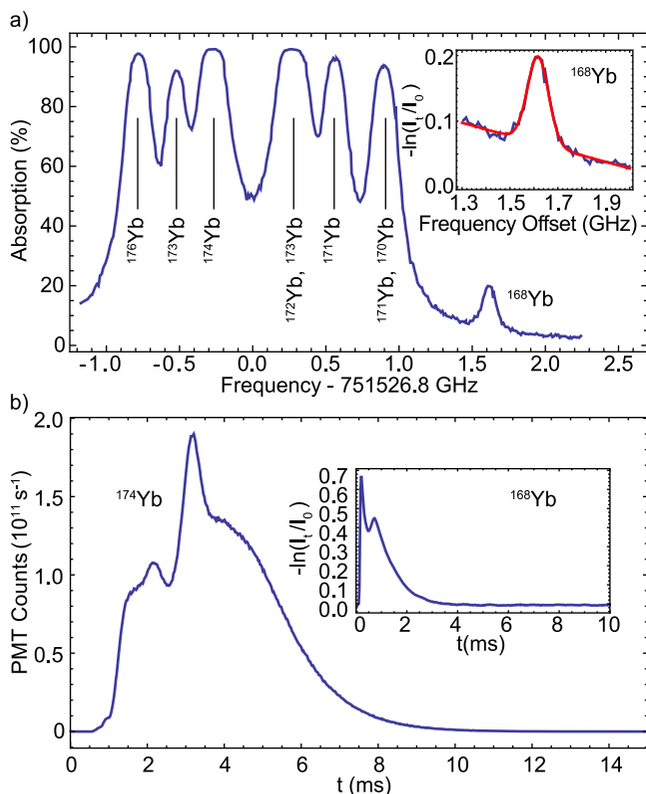


Figure 8. (a) Absorption spectrum of the $(6s6p)^1P_1 \leftarrow (6s^2)^1S_0$ transition, for all stable isotopes of ytterbium. The inset shows a zoom-in on the ^{168}Yb peak. The red line is a fit to determine the Doppler width against the background absorption of more abundant isotopes. (b) Fluorescence TOF traces of the ytterbium beam, using light resonant with the $(6s6p)^1P_1 \leftarrow (6s^2)^1S_0$ transition for the ^{174}Yb beam. Because the transition is optically closed, we show the PMT count rate which is proportional to the atomic density inside the detection volume. The inset shows an absorption trace for the ^{168}Yb isotope taken directly outside the cell exit.

an evident loss in signal, especially for the atoms with late arrival times in the detector. We typically observe a reduction of around 80% in the total number of atoms reaching the detector in LIF-2. For Ca and Yb, the effect of the reactant gas is much less pronounced. We observe a 20% and 10% reduction in the atomic signal, respectively and obtain qualitatively similar results when using SF_6 as the reactant gas. However, the atomic beam speeds up upon excess addition of SF_6 ; we discuss observable effects on the molecular beam thermalisation later. Excited state chemistry can increase the yield of the species of interest in the source. This has been demonstrated for the reaction of Yb and Ca with water and alcohols to form YbOH and CaOH [24, 61, 62] and for the reaction of carbon atoms with hydrogen molecules to form CH [63]. We found no influence on CaF production when exciting the Ca atoms with 40 mWcm^{-2} of $(3p^64snp)^1P_1 \leftarrow (3p^64s^2)^1S_0$ light longitudinally through the cell.

The exact process through which the reaction between the atoms and the reactant gas happens is largely unknown, but available experimental data provides some useful information. The activation energies of the reactions $\text{Al} + \text{NF}_3 \rightarrow \text{AlF} + \text{NF}_2$ and $\text{Al} + \text{SF}_6 \rightarrow \text{AlF} + \text{SF}_5$ have been experimentally measured as 2990 K and 4800 K respectively [64]. This can be compared to the kinetic energy of the ablated Al atoms, which decreases exponentially with the number of Al-He collisions [3]. Measurements of the Doppler width of the Al atoms in the absence of the buffer gas suggested an initial temperature of $3400 \pm 1000 \text{ K}$. Even assuming an initial Al temperature of 10^4 K , the typical energy for the reaction falls below the activation energy after 4 (SF_6) and 7 (NF_3) Al-He collisions. This suggests that for a two-body gas phase reaction, the reaction must be highly efficient to generate an appreciable number of molecules.

A number of observations suggest that the reactant gas in fact remains frozen in the cell and is brought into the gas phase in the ablation process. First, removing the flow of reactant gas into the cell with a tap results in a slow drop of the molecular beam signal, over 200 shots, but never fully to zero. However, this requires active ablation of the target, if both the ablation light and the reactant gas flow are removed for the duration of several hundred shots, and the ablation light is then introduced again, the signal decay begins from its original value. We have often observed a sizeable fraction of signal for ~ 30 minutes of continued operation with the reactant gas tap completely shut.

To investigate this further, we replaced the flow of reactant gas with 0.01 sccm of nitric oxide (NO) and fired the ablation laser at the metal target to reproduce the usual conditions in the cell.² NO can be optically detected via the $A^2\Sigma^+, v' = 0 \leftarrow X^2\Pi_{1/2}, v = 0$ transition near 226.2 nm, and its freezing point (109 K) is intermediate between SF_6 (223 K) and NF_3 (66 K). The hyperfine structure of this transition has been resolved in a number of studies [65–69], and absorption spectroscopy in a room-temperature cell was recently used to measure hyperfine structure of high- J lines of the $A^2\Sigma^+, v' = 0 \leftarrow X^2\Pi_{3/2}, v = 0$ transition [70]. Here, we probe the two lowest energy levels of the $X^2\Pi_{1/2}, v = 0$, clearly resolving the hyperfine structure. A level scheme for NO is shown in Figure 10(a). The Franck–Condon matrix for this transition is highly non-diagonal [65], and to a good approximation we can assume a single photon absorption will optically pump an NO molecule to other rovibrational levels of the $X^2\Pi$ state.

With the buffer gas cell cooled to 2.5 K, we observe a beam of buffer gas cooled, ground-state NO molecules, which we assume are desorbed upon firing the ablation laser (Figure 10(b)). The mean forward velocity of this

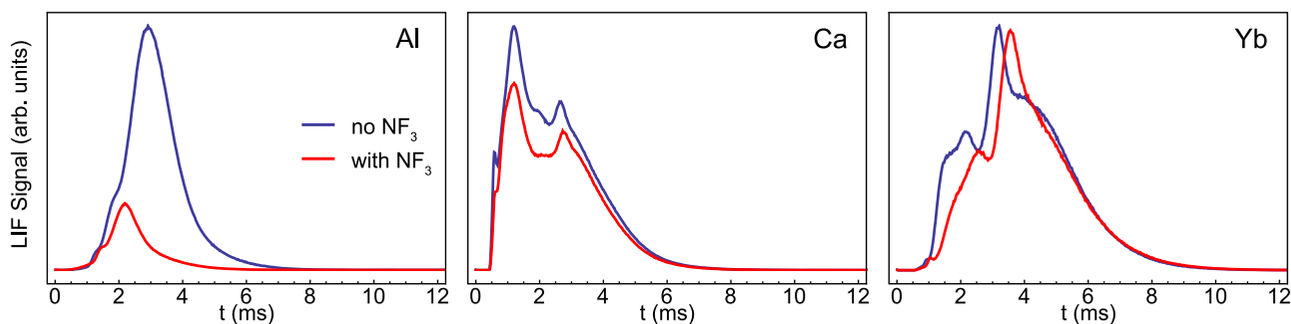


Figure 9. LIF signal traces of the Al, Ca and Yb atom beams without (blue) and with (red) NF_3 flow into the buffer gas cell, demonstrating the strong effect of a fluorine donor gas on the number of aluminium atoms in the beam.

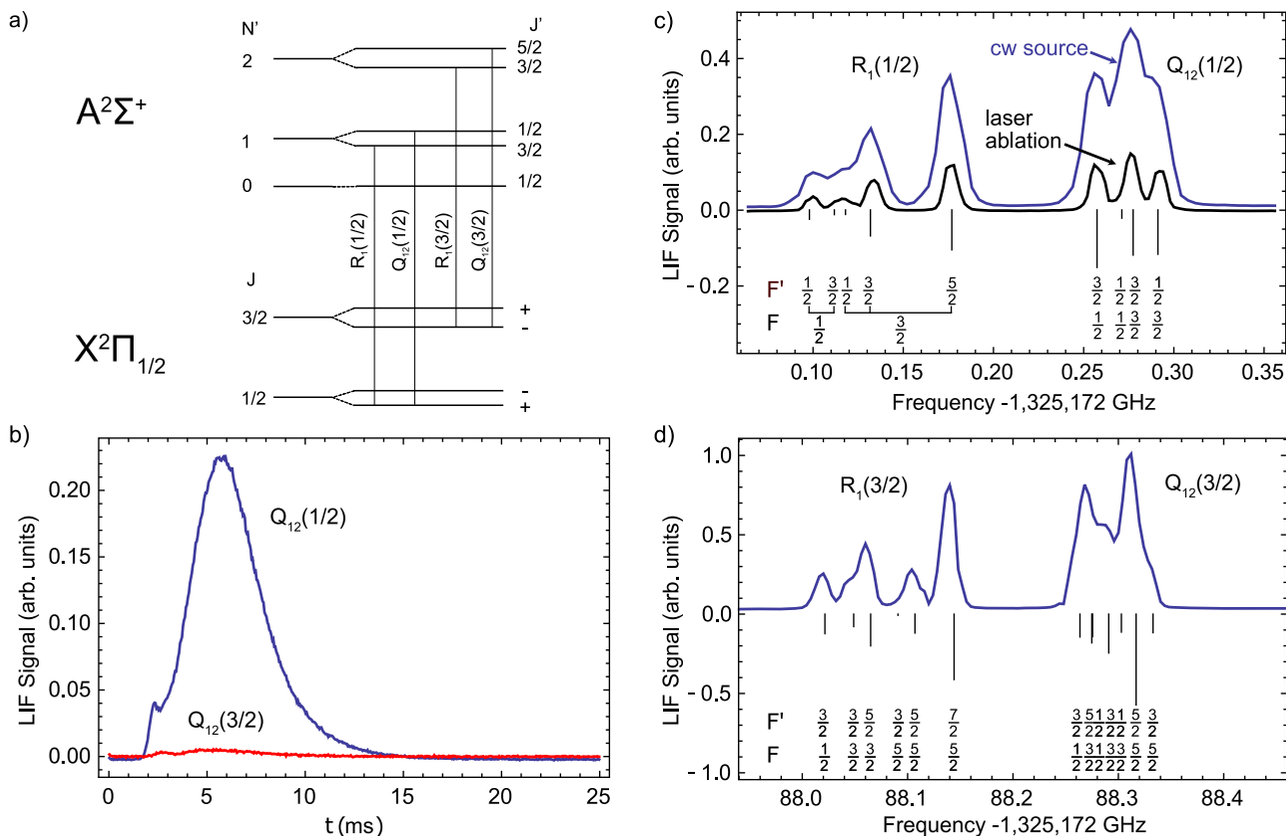


Figure 10. Experiments with nitric oxide (NO) flowed into the buffer gas cell through the fluorine donor gas inlet. (a) Scheme of the relevant energy levels (hyperfine structure omitted). (b) Firing the ablation laser generates a pulse of rotationally cold, slow NO molecules. (c) A spectrum of the $R_1(1/2)$ and $Q_{12}(1/2)$ lines taken using the pulsed desorption method with the buffer gas cell at 2.5 K, and a spectrum using a continuous beam with the buffer gas cell at 70 K. (d) A spectrum of the $R_1(3/2)$ and $Q_{12}(3/2)$ lines taken using the continuous source shown on the same vertical scale as (c).

beam is about 120 ms^{-1} and we estimate the brightness as 10^{12} molecules per steradian per pulse, with a stability of better than 10% (standard deviation). The signal in the $J = 3/2$ first rotationally excited state is a factor 50 lower. We note that this method of generating an intense beam of slow, rotationally pure NO molecules may be of use in cold collision experiments [71, 72].

We also observe a bright, continuous beam of cold NO molecules without ablation light, provided the second

stage of the cryocooler is heated to above about 60 K; below this temperature the signal completely disappeared. Spectra of the $R_1/Q_{12}(1/2)$ lines taken with the desorbed and continuous beams are shown in Figure 10(c). Stick spectra shown pointing downwards are the individual hyperfine lines as predicted using the ground state parameters of [66] and the excited state parameters of [70]. The simulated line positions and the experimental spectra are in good agreement. The fitted FWHM of

the lines is larger by about a factor 1.7 for the continuous beam, from which we deduce a forward velocity of 200 ms^{-1} . By comparison of the relative fluorescence signals, we find that the continuous beam produces the same number of molecules in the $J = 1/2$ level in a 5-ms time interval as a single desorption pulse. A spectrum for the $R_1/Q_{12}(3/2)$ line using the continuous source is also shown in Figure 10(d). The probe laser power is equal for both spectra, showing that the $J = 3/2$ level has greater population under these conditions. The experiments with NO suggest that a similar effect occurs for the fluorine donor gas during the production of metal fluoride molecules. The density of the reactant gas increases around the time of ablation by vaporisation of ice inside the cell. This may explain how reactant gas sources can be efficient despite the high activation energy of the reaction.

4. Results: molecular beams

In this section, we determine the brightness of molecular beams of AlF, ^{40}CaF , ^{24}MgF and ^{174}YbF and discuss the influence of experimental parameters on the molecular beam. For CaF, MgF and YbF, we probe molecules in the ground rotational state using the $Q_1(0)$ line of the $A^2\Pi_{1/2}, v' = 0 \leftarrow X^2\Sigma^+, v = 0$ band. For these molecules, the vibrational branching is small compared with rotational branching, and we expect that at high laser intensities each molecule scatters on average $\mathcal{N} = 3$ photons before being optically pumped to the second rotationally excited state in the vibronic ground state. The total emission pattern for the $Q_1(0)$ lines is isotropic. To probe the AlF beam, we use the $A^1\Pi, v' = 0 \leftarrow X^1\Sigma^+, v = 0 R(J)$ lines, which optically pumps molecules from rotational state J to $J + 2$ in the vibronic ground state. For the $R(0)$ line, we also expect $\mathcal{N} = 3$ photons per molecule in saturated fluorescence. The emission pattern has been measured to be slightly anisotropic by $\sim 20\%$, so we use light which is linearly polarised parallel to the detector direction. In this configuration, the relative number of photons scattered on $P(J)$ and $R(J)$ lines has been experimentally found to be in agreement with the Hönl–London factors [46].

4.1. Comparison of molecular beams

The left panels of Figure 11 show the absorption spectra recorded directly after the buffer gas cell orifice for all four molecular species. The peak absorption of the AlF beam, presented in Figure 11(a), is an order of magnitude higher than for the other monofluoride species. The absorption measured 20 mm downstream is shown magnified in Figure 11(a). This measurement reveals

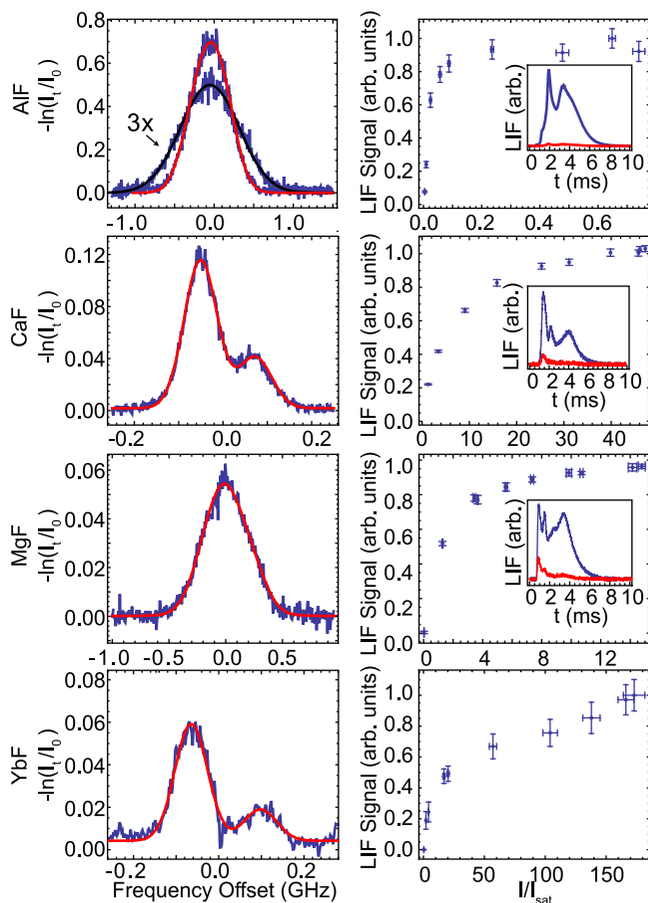


Figure 11. Left column: absorption spectra recorded directly outside the cell orifice. The solid red curves are fits based on the known energy structure of the molecules and using a Gaussian lineshape. The enlarged spectrum shown for AlF is measured 20 mm downstream. Right column: saturation curves on the rotationally open transitions. For AlF, CaF and MgF, the insets show the fluorescence signal without (blue) and with (red) optical pumping applied in LIF-1, demonstrating that the molecules are indeed pumped away from the addressed transition. In the case of YbF, high rotational lines from other isotopologues prevent saturation.

a broadening of the transverse velocity spread and is comparable to the atomic beams. We extract the transverse velocity widths by fitting the absorption spectra to Doppler-broadened line shapes and use these to predict the brightness of each molecular beam.

The right panels in Figure 11 show saturation of the fluorescence. The insets show the results of pump-probe measurements for AlF, CaF and MgF demonstrating that the molecules are optically pumped. For AlF, the large transition linewidth and small hyperfine structure in the $J = 0$ ground state permit efficient saturation of the fluorescence at $I/I_{\text{sat}} < 1$ and we observe a pumping efficiency of 97%. For CaF, MgF and YbF, a relatively high laser intensity of $I/I_{\text{sat}} > 40$ is required to power-broaden the transition enough to cover the ground state spin-rotation and hyperfine structure. We verify this

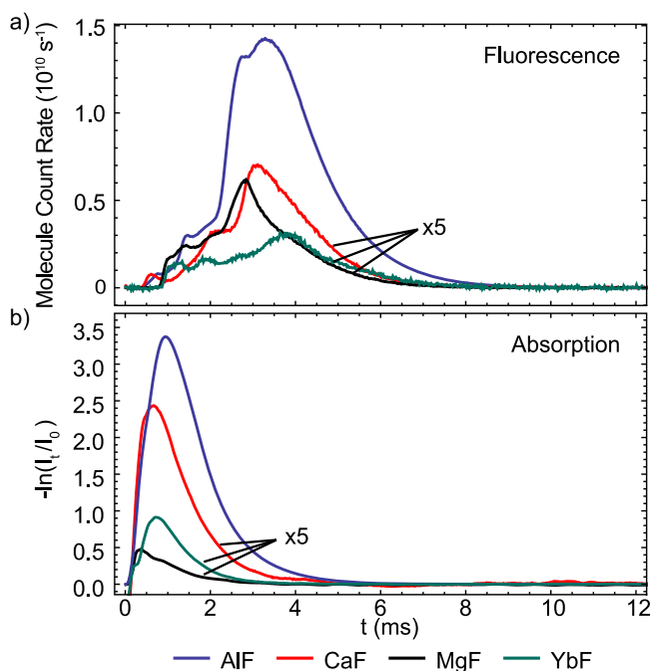


Figure 12. (a) Comparison of the molecule count rates in the ground rotational state for all species, obtained in saturated fluorescence, as a function of arrival time in the detector. We plot the count rate as the number of molecules entering the detector, limited by the 2×2 mm² square aperture. (b) Comparison of the optical depth of the molecular beams, recorded at the buffer gas cell orifice. In both (a) and (b), the MgF, CaF and YbF signals are magnified by a factor 5.

by fluorescence spectra at high power and pump-probe measurements. In the case of YbF, the background due to rotational lines of other molecular isotopologues leads to an increase in the fluorescence above $I/I_{\text{sat}} = 100$ and we assume our conversion of fluorescence signal to molecule number is uncertain to within a factor 2.

Figure 12(a) shows a comparison of the maximum fluorescence signals obtained – under identical experimental conditions – for all four molecules. The MgF, CaF and YbF signals are magnified by a factor 5. From these signals and the absorption measurements shown in Figure 12(b), we calculated the respective brightness in absorption and fluorescence. In Table 1, we summarise our estimates of the brightness of each atomic and molecular beam source. For the molecules, the given values correspond to the number of molecules in the ground rotational state. In both absorption and fluorescence, AlF is the brightest molecular species by about a factor 10, whereas aluminium produces the lowest brightness atomic beam by about a factor 2.

For AlF, we also recorded the relative rotational state populations by saturated fluorescence of successive R(*J*) lines up to $J = 3$. The resulting state distribution is presented in Figure 13(a). It is poorly described by a thermal

Table 1. Properties of the atomic and molecular beams, as determined from fluorescence and absorption experiments. For the molecular species, we quote the brightness as observed in the ground rotational state. The transverse velocities v_t are derived from the absorption spectra taken directly outside the cell exit aperture; where possible we state the transverse spread downstream of the cell in brackets. The combined uncertainties of the detection efficiencies lead to error bars for the LIF brightnesses of around 30%. In the case of Yb and YbF, the given values are expected to be accurate within a factor 2 because we cannot use open transitions saturated by optical pumping.

Species	v_t (ms ⁻¹)	\mathcal{B}_f (sr ⁻¹ per pulse)	\mathcal{B}_{abs} (sr ⁻¹ per pulse)
Al	90 (150)	3.0×10^{12}	4.1×10^{12}
⁴⁰ Ca	75 (120)	9.2×10^{12}	7.4×10^{12}
¹⁷⁴ Yb	40 (60)	5.0×10^{12}	1.2×10^{13}
AlF	74 (160)	1.6×10^{12}	2.1×10^{12}
⁴⁰ CaF	53 (125)	1.4×10^{11}	1.1×10^{11}
²⁴ MgF	112	1.1×10^{11}	1.1×10^{11}
¹⁷⁴ YbF	50	0.7×10^{11}	2.1×10^{11}

Boltzmann distribution; we show expected distributions for $T = 1.15$ K and 2.5 K as solid lines in the figure. The combined beam brightness for $0 \leq J \leq 3$ is 3×10^{12} sr⁻¹ per pulse, and consistent with the atomic aluminium beam measurement. This supports the idea that NF₃ reacts highly efficiently with aluminium atoms to produce AlF. Assuming a similar rotational distribution for CaF, we estimate a beam brightness which is 3% of the atomic Ca beam. The helium flow has a significant effect on the rotational thermalisation in the AlF beam, as illustrated in Figures 13(b) and 13(c). The $J = 0$ population is maximised at a He flow rate of about 1 sccm, but it appears population is redistributed to higher rotational states as the flow rate increases. Whilst we did not ourselves attempt to model the in-cell dynamics, we note that a number of approaches have been explored [73–77].

The order of magnitude difference in molecule yield between AlF and the other monofluorides can be explained by the electronic structures of the atoms and molecules. Calcium and magnesium are group II metals with a fully occupied *s* orbital, ytterbium is a lanthanoid with fully occupied *f* and *s* orbitals, but aluminium is a group III metal with an unpaired *p* electron, making it a reactive radical. The gas-phase aluminium atoms react efficiently with the fluorine donor gas, whereas for the other metal species, only a small percentage of the atoms react, as demonstrated in Figure 9. While all studied monofluorides are stable molecules in the gas phase, only AlF possesses no unpaired electrons, making it much less reactive than the radicals CaF, MgF and YbF. The latter three are prone to the formation of difluorides. Theoretical calculations for CaF predict that a considerable amount of CaF₂, comparable to or more than the amount of CaF, is formed in the reaction of ablated calcium and

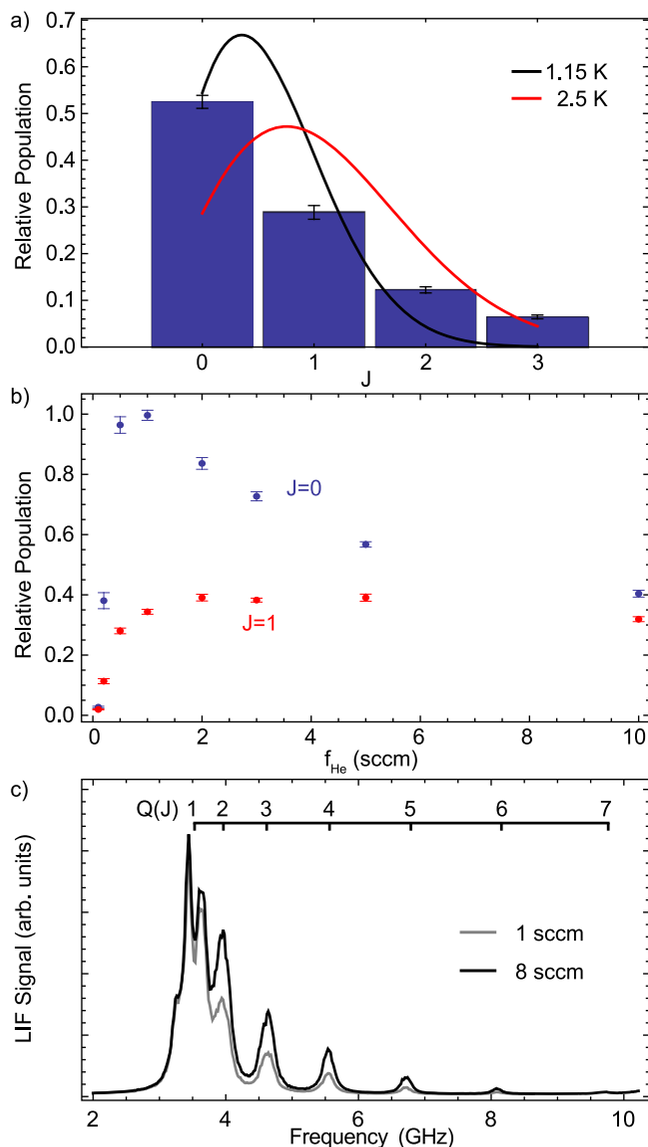


Figure 13. (a) Distribution of the AlF population over the four lowest rotational levels in the vibrational ground state of the $X^1\Sigma^+$ state. The solid lines show Boltzmann distributions at 1.15 K and 2.5 K, respectively. (b) Influence of the helium buffer gas flow rate f_{He} on the fluorescence signal in LIF-2, detected on the R(0) line (blue) and the R(1) line (red). (c) Q-branch scans at He buffer gas flows of 1 sccm and 8 sccm.

a fluorine donor gas, while the difluoride production is suppressed for Al [78].

4.2. Velocity distribution of the molecular beam

4.2.1. Pump-probe method

We determined the forward velocity distributions of the Al and AlF beams, using the pump-probe method described in [46]. Briefly, applying optical pumping light in LIF-1 removes the signal in LIF-2; a short ($\sim 10 \mu\text{s}$) removal of the pump light results in an appearance of

signal in LIF-2, in such a way that time of arrival in LIF-2 correlates with velocity. Repeatedly switching off the pump light at appropriate intervals as the molecules transit through LIF-1 then allows estimating the forward velocity distribution, $f(v)$, using a single molecular pulse with a high signal to noise ratio. We show the results of these experiments in Figure 14 for Al and AlF, where the velocity distributions for AlF are shown for $J = 0-3$, and normalised such that the integral $\int f(v)dv = 1$ for each J . The finite transit time through the laser beams corresponds to a velocity uncertainty of below 5%. Overall the velocity distribution of the molecular and atomic beams is similar, and in the case of AlF, the ground rotational state contains the slowest molecules.

4.2.2. Stark decelerator

To gain additional insight into the molecular buffer gas source, we replaced the LIF detection zones with a Stark decelerator to map out the longitudinal phase space distribution (z, v_z) for AlF [79]. The decelerator [80] consists of 132 electrode pairs spaced at 5.5 mm along the z -axis, whose orientation alternates between $\pm 45^\circ$ to the x - and y -axes in the x - y plane. We operate the decelerator in the $S = 1$ guiding mode (i.e. at a phase angle of 0), where the electrode configuration is switched synchronously with a molecule travelling at velocity v_z , which reaches the position of the decelerator entrance z_0 at a variable in-coupling time t_i . In this way, molecules in a small region of phase space ($z_0 \pm \delta_z, v_z \pm \delta_v$) at time t_i are guided through the chain of electrode pairs and arrive at a defined time in the LIF detection zone, located at a short distance from the decelerator output. The acceptance region (δ_z, δ_v) is determined by the longitudinal electrode spacing, the peak electric field between the electrodes and the number of used electrode stages. Along the z -axis, the peak electric field is about 100 kVcm^{-1} , corresponding to a potential depth of 0.5 K for AlF molecules in low field seeking states of the $J = 1$ level. From the trap depth we can estimate an upper bound for δ_v of 13 ms^{-1} , whereas trajectory simulations of the molecules give $\delta_v = 10 \text{ ms}^{-1}$.

Figure 14(c) shows a false colour plot of the LIF signal as a function of the scanned parameters t_i and v_z , for AlF molecules in the $J = 1$ state. The white line in the figure marks the relation $v_z = z_0/t_i$ with $z_0 = 30 \text{ cm}$, representing ballistic motion from the target at the time of the ablation to the decelerator entrance. The buffer gas cooled pulse of molecules reaches the decelerator with an average velocity of about 170 ms^{-1} , 0.7 ms later than the ballistic curve due to collisions and thus thermalisation with the buffer gas in the cell. A faster part of the beam arrives close to the ballistic trajectory curve near 320 ms^{-1} , which presumably results from molecules which leave the

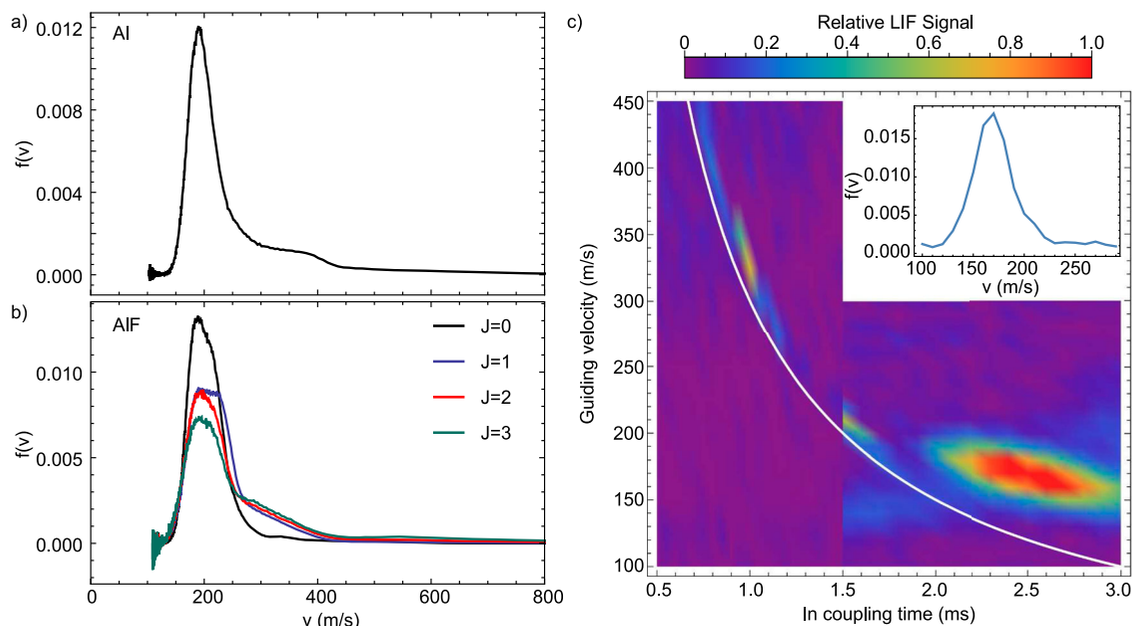


Figure 14. Velocity distributions, determined using a pump-probe method, of (a) the Al atomic beam and (b) the AIF molecular beams, for the four lowest rotational states in the $X^1\Sigma^+$, $v = 0$ ground electronic state. (c) Phase space distribution of the AIF molecular beam, measured by velocity guiding $J = 1$ molecules in a Stark Decelerator. The white curve shows the relation $v_z = z_0/t_i$ with $z_0 = 300$ mm. The molecules which are well thermalised arrive with low velocities, and delayed by about 0.7 ms relative to this ballistic trajectory. The inset shows the velocity distribution calculated by summing over all t_i for $100 < v_z < 300$.

cell shortly after the ablation without thermalising with the helium in the cell. This tends to occur over the course of several thousand shots necessary for such a measurement. In the inset of the figure, the velocity distribution is computed by integrating over all in-coupling times for $100 < v < 300 \text{ ms}^{-1}$. The distribution is narrower than measured by the pump-probe method in panel (b), with its peak velocity about 40 ms^{-1} slower. The two different measurements were taken months apart, with the decelerator measurement using a freshly cleaned cell, and this can account for such a difference.

4.3. Influence of the ablation laser fluence

The influence of the energy of the Nd:YAG infrared ablation laser on the number of atoms and molecules is shown in Figure 15. The signal measured in LIF-2 is shown as a function of the ablation pulse energy and laser fluence using the measured spot size of 0.7 mm. We observe that the threshold energy for observing Al and AIF is between 10 and 15 mJ, and the signal from both species increases up to the maximum available pulse energy of 40 mJ. In particular, the AIF yield is proportional to the ablation energy, and both would appear to benefit from increased pulse energy. For Ca and Yb, the threshold is 5–10 mJ, and we observe an optimum in respective molecular production at about 25 mJ. Above this pulse energy, the atomic signal either plateaus or weakly

increases, but the number of molecules decreases. This may be due to incomplete thermalisation with the buffer gas, or changing reaction dynamics at higher ablation energies.

The differences in the phase explosion threshold energy for the metals used in our study likely arise from the difference in the latent heat of vaporisation of the atomic species, where the value for Al is roughly a factor 2 larger than for Ca, Mg and Yb. The reflectivity of the bulk metal surfaces at room temperature – values for cryogenic temperatures are not reported – is around 93% for Al, Ca, and Mg, but 70% for Yb [81]. The latter is a possible explanation for the large number of vaporised Yb atoms.

4.4. Influence of the fluorine donor gas

While the inert gas SF_6 is commonly used in experiments with monofluorides, theory suggests that the use of the more reactive and corrosive molecule NF_3 has advantages [78]. We find that both produce a similar number of molecules when observed downstream of the source, within the range of day-to-day fluctuations of source operation with one fluorine donor gas. Overall, the response of molecule yield to the gas flow rates was not reliably reproducible from day to day, and strongly dependent on the flow rate history during a measurement sequence. We attribute this to probable freezing effect of

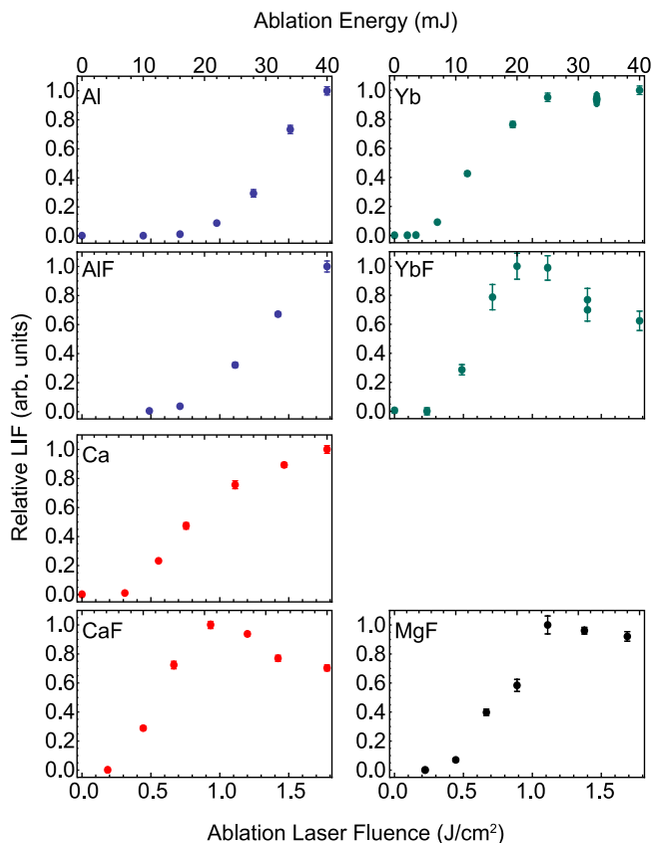


Figure 15. Fluorescence signals of all investigated species as a function of the ablation laser pulse energy (upper x -axes) and laser fluence (lower x -axes). We plot the relative LIF for the atomic beams and for the rotational ground state of the molecules.

the reactant gas discussed previously, and to the ‘poisoning’ effect discussed below. However, we typically find NF_3 performs well with flow rates 0.001 sccm or below, but that SF_6 requires about 10 times more flow to give comparable signal. This is illustrated in Figure 16 for the case of MgF . Operating the source with NF_3 is more reliable and results in a slower beam for a longer period of operation. SF_6 can produce a similar beam brightness with a freshly cleaned cell, but after 1 day of operation, the TOF profile shifts to higher velocities. This effect is particularly striking for AlF , where the forward velocity increases steadily with the number of ablation shots when SF_6 is used. We do not observe this effect when only flowing SF_6 or only ablating Al for a day. We assume that this speed-up is caused by the buildup of ablation products and sulphur-containing inorganic compounds on the internal buffer gas cell wall, thus hindering efficient (re-)thermalisation of the helium buffer gas. This effect is less pronounced for the other molecules. A larger internal volume of the cell is likely to improve this, but typically results in long temporal pulses. We find that extracting the molecular beam through the aperture into an extension that has the same bore diameter but no aperture

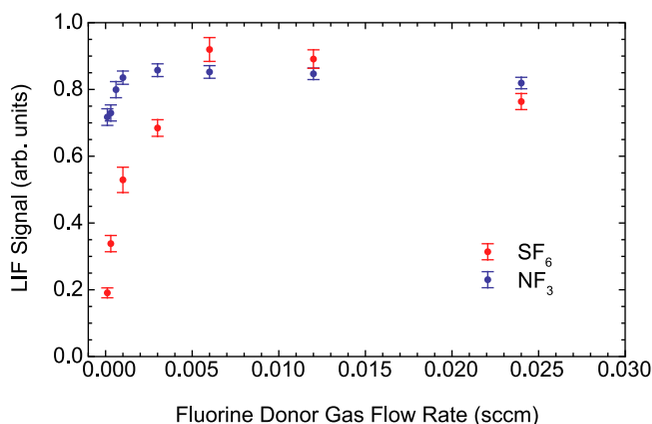


Figure 16. Influence of the fluorine donor gas flow rate on the $\text{MgF } Q_1(0)$ fluorescence signal in LIF-2.

reduces the forward velocity and significantly improves the long-term stability of the beam without affecting the beam brightness. This indeed indicates that the helium thermalisation is affected by ablation products in the main cell. An additional mesh on the extension reduces the mean forward velocity of AlF by up to 70 ms^{-1} , but reduces the beam brightness [30, 46].

In Figure 17, we use the compact Q-branch of AlF as a convenient means to probe the rotational distribution reaching LIF-2, and thereby gain insight into the thermalisation dynamics in the cell. We plot a series of Q-branch spectra, separated by the arrival time of the molecules in the detection region, incremented in 0.5 ms steps. Thus we can see the time-dependent rotational distribution of the molecular beam. The upper blue set of spectra show typical behaviour when using NF_3 gas. The middle (black) set of spectra are measurements taken with a clean cell and an SF_6 flow rate of 0.0006 sccm, and are similar to the NF_3 data. The red spectra are measurements with a flow rate of 0.005 sccm after 1 day of source operation without cleaning afterwards. These spectra show that at a higher flow rate and in the used cell, the molecules arrive earlier at the detector, and are distributed over many rotational states; we indeed observe molecules in $J = 13$ arriving 1 ms after the ablation pulse. The inset of the figure shows TOF fluorescence traces when saturating the $R(0)$ line, demonstrating how the molecular pulse shifts towards higher velocities. We find that the onset of this reduced thermalisation occurs more rapidly as the flow rate increases, and that good thermalisation can only be recovered by cleaning ablation products from the buffer gas cell. The rotational spectra for different molecule arrival times also serve to show that translational and rotational energy of the AlF molecules are correlated – fast molecules are rotationally hotter than slow ones.

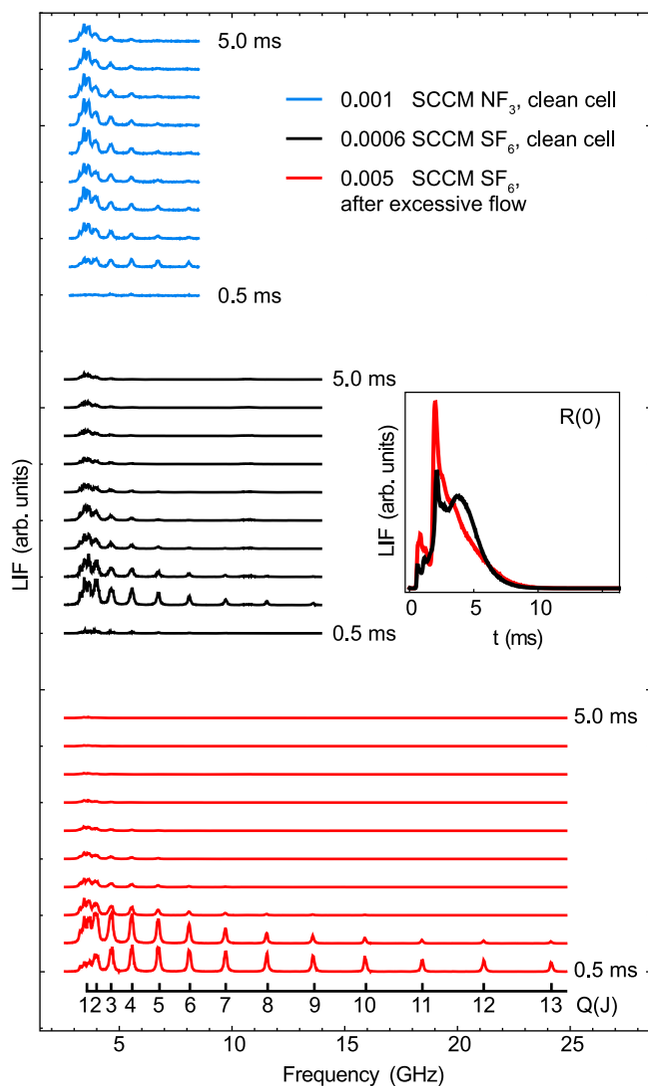


Figure 17. Q-branch spectra of the $A^1\Pi \leftarrow X^1\Sigma^+$ transition in AlF selected by arrival time at the detector, in 0.5 ms increments. The upper blue spectrum is recorded with a clean buffer gas cell using NF_3 gas (blue, top), whereas the lower two spectra are recorded with a clean (black, middle) and contaminated (red, bottom) cell using SF_6 . The ruler inside the figure marks the different $Q(J)$ -lines. The inset shows the signal obtained for the $J = 0$ population using the $R(0)$ line, for the two SF_6 examples.

Sulphur hexafluoride and nitrogen trifluoride are not the only possible fluorine donors. We performed preliminary experiments using tetrafluoromethane (CF_4) as fluorine donor gas that lead to similar molecule yields as for NF_3 and SF_6 , and we therefore did not further pursue this avenue. Solid xenon difluoride (XeF_2), theoretically predicted to be a good candidate fluorine donor molecule [78], did not prove as a viable option. At room temperature, the compound evaporates quickly into a hazardous gas. Nevertheless, we connected a reservoir containing XeF_2 crystals to the buffer gas cell and

adjusted the flow with a needle valve. We observed efficient molecule production, but the relatively high flow rate resulted in a very fast molecular beam.

5. Conclusion and outlook

We have presented a series of experiments comparing buffer gas cooled beams of Al, Yb, Ca, and their monofluorides AlF, CaF, MgF and YbF formed by reaction of laser-ablated atoms with a fluorine donor gas inside a buffer gas cell. We find that the molecular beam brightness of AlF is about 1 order of magnitude larger than for the other monofluorides, and when multiple rotational states are considered, we observe a similar beam brightness to the atomic Al beam. This is qualitatively consistent with the near complete loss of atomic signal upon introducing the reactant gas into the cell. In contrast, the Ca and Yb atomic beams are relatively unaffected by the fluorine reagents, and brighter than the Al beam. The molecular yield of CaF, MgF and YbF suggests a reaction efficiency on the few percent level. This difference in reactivity is explained by the radical character of aluminium and the stability of the AlF molecule, while the other molecules are radicals that are formed from less reactive atoms.

We demonstrated high-flux pulsed and continuous molecular beams of buffer gas cooled NO, and performed CW UV spectroscopy of low- J lines of the $A^2\Sigma^+, v' = 0 \leftarrow X^2\Pi_{1/2}, v = 0$ transition. The low forward velocity and rotational state purity of the pulsed beam provide a useful starting point for low-energy collision experiments with NO. Our observations also provide insight about the vaporisation of ice in the cell, and the pulsed desorption method may be applicable to other diatomic molecules (e.g. H_2 , O_2).

We conclude with some guidance for others wishing to develop cryogenic buffer gas beam sources. First, we find it useful to measure the molecular beam brightness with both absorption and fluorescence methods, noting that absorption directly outside the cell leads to an overestimate. A large discrepancy between the two measurements indicates insufficient cryopumping or deteriorating charcoal. Second, monitoring the atomic buffer gas beams, especially in sources using reactant gases, provides a useful reference when studying the molecular beam properties. Finally, thermalisation inside the cell is influenced by both the choice of reactant gas, and the condition of the internal cell surfaces. NF_3 reacts more efficiently to form fluoride molecules as a significantly lower flow rate is required to produce the same molecular beam brightness. The lower freezing point also allows cooling the capillary that feeds the fluorine donor gas, which reduces the thermal heat load on the cell. We also

find that the molecular beam parameters, especially for ALF, are more consistent when using NF₃.

Notes

1. Probing the Mg atoms would require light at 285 nm to excite the $^1P_1 \leftarrow ^1S_0$ transition, which was not available for this study.
2. We note that the NO experiments were done with a slightly different LIF detector to that used during the rest of this study ($L_2 = 61$ cm, with an 8 mm diameter molecular beam aperture), and so we do not make a quantitative comparison with the other molecular beams.

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Data availability statement

The data presented can be accessed from Zenodo (<https://doi.org/10.5281/zenodo.7063976>) and may be used under the Creative Commons Attribution 4.0 International license.

Disclosure statement

No potential conflict of interest was reported by the author(s).

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References

- [1] D.R. Willey, R.L. Crownover, D.N. Bittner and F.C. De Lucia, *J. Chem. Phys.* **89** (4), 1923–1928 (1988). doi:10.1063/1.455089
- [2] S.E. Maxwell, N. Brahm, R. DeCarvalho, D.R. Glenn, J.S. Helton, S.V. Nguyen, D. Patterson, J. Petricka, D. DeMille and J.M. Doyle, *Phys. Rev. Lett.* **95** (17), 173201 (2005). doi:10.1103/PhysRevLett.95.173201
- [3] N.R. Hutzler, H.I. Lu and J.M. Doyle, *Chem. Rev.* **112**, 4803–4827 (2012). doi:10.1021/cr200362u
- [4] S. Truppe, M. Hambach, S.M. Skoff, N.E. Bulleid, J.S. Bumby, R.J. Hendricks, E.A. Hinds, B.E. Sauer and M.R. Tarbutt, *J. Mod. Opt.* **65** (5–6), 648–656 (2018). doi:10.1080/09500340.2017.1384516
- [5] D. Patterson, M. Schnell and J.M. Doyle, *Nature* **497** (7450), 475–477 (2013). doi:10.1038/nature12150
- [6] V. Andreev, D.G. Ang, D. DeMille, J.M. Doyle, G. Gabrielse, J. Haefner, N.R. Hutzler, Z. Lasner, C. Meisenhelder, B.R. O'Leary, C.D. Panda, A.D. West, E.P. West and X. Wu, *Nature* **562** (7727), 355–360 (2018). doi:10.1038/s41586-018-0599-8
- [7] X. Alauze, J. Lim, M.A. Trigatzis, S. Swarbrick, F.J. Collings, N.J. Fitch, B.E. Sauer and M.R. Tarbutt, *Quantum Sci. Technol.* **6** (4), 044005 (2021). doi:10.1088/2058-9565/ac107e
- [8] N.H. Pilgram, A. Jadbabaie, Y. Zeng, N.R. Hutzler and T.C. Steimle, *J. Chem. Phys.* **154** (24), 244309 (2021). doi:10.1063/5.0055293.
- [9] R. DeCarvalho, J.M. Doyle, B. Friedrich, T. Guillet, J. Kim, D. Patterson and J.D. Weinstein, *Eur. Phys. J. D* **7** (3), 289–309 (1999). doi:10.1007/s100530050572
- [10] W.C. Campbell and J.M. Doyle, in *Cold Molecules. Theory, Experiment, Applications*, edited by Roman V. Krems, Bretislav Friedrich and William C. Stwalley, Chap. 13 (CRC Press, Boca Raton, 2009), pp. 473–508.
- [11] H.I. Lu, I. Kozyryev, B. Hemmerling, J. Piskorski and J.M. Doyle, *Phys. Rev. Lett.* **112** (11), 113006 (2014). doi:10.1103/PhysRevLett.112.113006
- [12] P. Aggarwal, Y. Yin, K. Esajas, H.L. Bethlem, A. Boeschoten, A. Borschevsky, S. Hoekstra, K. Jungmann, V.R. Marshall, T.B. Meijknecht, M.C. Mooij, R.G.E. Timmermans, A. Touwen, W. Ubachs and L. Willmann, *Phys. Rev. Lett.* **127** (17), 173201 (2021). doi:10.1103/PhysRevLett.127.173201
- [13] S.M. Skoff, R.J. Hendricks, C.D.J. Sinclair, J.J. Hudson, D.M. Segal, B.E. Sauer, E.A. Hinds and M.R. Tarbutt, *Phys. Rev. A* **83** (2), 023418 (2011). doi:10.1103/PhysRevA.83.023418
- [14] B.C. Sawyer, B.K. Stuhl, M. Yeo, T.V. Tscherebul, M.T. Hummon, Y. Xia, J. Kos, D. Patterson, J.M. Doyle and J. Ye, *Phys. Chem. Chem. Phys.* **13**, 19059–19066 (2011). doi:10.1039/c1cp21203f
- [15] X. Wu, T. Gantner, M. Koller, M. Zeppenfeld, S. Chervenkov and G. Rempe, *Science* **358** (6363), 645–648 (2017). doi:10.1126/science.aan3029
- [16] M. Koller, F. Jung, J. Phrompao, M. Zeppenfeld, I.M. Rabey and G. Rempe, *Phys. Rev. Lett.* **128** (20), 203401 (2022). doi:10.1103/PhysRevLett.128.203401
- [17] B. Hemmerling, G.K. Drayna, E. Chae, A. Ravi and J.M. Doyle, *New J. Phys.* **16** (6), 063070 (2014). doi:10.1088/1367-2630/16/6/063070
- [18] J.F. Barry, D.J. McCarron, E.B. Norrgard, M.H. Steinecker and D. DeMille, *Nature* **512** (7514), 286–289 (2014). doi:10.1038/nature13634
- [19] S. Truppe, H.J. Williams, M. Hambach, L. Caldwell, N.J. Fitch, E.A. Hinds, B.E. Sauer and M.R. Tarbutt,

- Nat. Phys. **13** (12), 1173–1176 (2017). doi:10.1038/nphys4241
- [20] L. Anderegg, B.L. Augenbraun, E. Chae, B. Hemmerling, N.R. Hutzler, A. Ravi, A. Collopy, J. Ye, W. Ketterle and J.M. Doyle, Phys. Rev. Lett. **119** (10), 103201 (2017). doi:10.1103/PhysRevLett.119.103201
- [21] A.L. Collopy, S. Ding, Y. Wu, I.A. Finneran, L. Anderegg, B.L. Augenbraun, J.M. Doyle and J. Ye, Phys. Rev. Lett. **121** (21), 213201 (2018). doi:10.1103/PhysRevLett.121.213201
- [22] J.C. Shaw and D.J. McCarron, Phys. Rev. A **102**, 041302 (2020). doi:10.1103/PhysRevA.102.041302
- [23] Z. Lasner, D. Mitra, M. Hiradfar, B. Augenbraun, L. Cheuk, E. Lee, S. Prabhu and J. Doyle, Phys. Rev. A **104**, 063305 (2021). doi:10.1103/PhysRevA.104.063305
- [24] N.B. Vilas, C. Hallas, L. Anderegg, P. Robichaud, A. Winnicki, D. Mitra and J.M. Doyle, Nature **606** (7912), 70–74 (2022). doi:10.1038/s41586-022-04620-5
- [25] A. Prehn, M. Ibrügger, R. Glöckner, G. Rempe and M. Zeppenfeld, Phys. Rev. Lett. **116** (6), 063005 (2016). doi:10.1103/PhysRevLett.116.063005
- [26] V. Singh, K.S. Hardman, N. Tariq, M.J. Lu, A. Ellis, M.J. Morrison and J.D. Weinstein, Phys. Rev. Lett. **108**, 203201 (2012). doi:10.1103/PhysRevLett.108.203201
- [27] N.E. Bulleid, S.M. Skoff, R.J. Hendricks, B.E. Sauer, E.A. Hinds and M.R. Tarbutt, Phys. Chem. Chem. Phys. **15** (29), 12299 (2013). doi:10.1039/c3cp51553b
- [28] N.R. Hutzler, M.F. Parsons, Y.V. Gurevich, P.W. Hess, E. Petrik, B. Spaun, A.C. Vutha, D. DeMille, G. Gabrielse and J.M. Doyle, Phys. Chem. Chem. Phys. **13** (42), 18976 (2011). doi:10.1039/c1cp20901a
- [29] E.B. Norrgard, E.R. Edwards, D.J. McCarron, M.H. Steinecker, D. DeMille, S.S. Alam, S.K. Peck, N.S. Wadia and L.R. Hunter, Phys. Rev. A **95**, 062506 (2017). doi:10.1103/PhysRevA.95.062506
- [30] H.I. Lu, J. Rasmussen, M.J. Wright, D. Patterson and J.M. Doyle, Phys. Chem. Chem. Phys. **13** (42), 18986–18990 (2011). doi:10.1039/c1cp21206k
- [31] M.T. Hummon, M. Yeo, B.K. Stuhl, A.L. Collopy, Y. Xia and J. Ye, Phys. Rev. Lett. **110** (14), 143001 (2013). doi:10.1103/PhysRevLett.110.143001
- [32] D. Patterson, E. Tsikata and J.M. Doyle, Phys. Chem. Chem. Phys. **12** (33), 9736–9741 (2010). doi:10.1039/c002764b
- [33] P.B. Changala, B. Spaun, D. Patterson, J.M. Doyle and J. Ye, Appl. Phys. B **122** (12), 292 (2016). doi:10.1007/s00340-016-6569-7
- [34] N.R. Hutzler, Quantum Sci. Technol. **5** (4), 044011 (2020). doi:10.1088/2058-9565/abb9c5
- [35] T. Gantner, M. Koller, X. Wu, G. Rempe and M. Zeppenfeld, J. Phys. B At., Mol. Opt. Phys. **53** (14), 145302 (2020). doi:10.1088/1361-6455/ab8b42
- [36] C.E. Dickerson, H. Guo, G.Z. Zhu, E.R. Hudson, J.R. Caram, W.C. Campbell and A.N. Alexandrova, J. Phys. Chem. Lett. **12** (16), 3989–3995 (2021). doi:10.1021/acs.jpcclett.1c00733
- [37] J. Piskorski, D. Patterson, S. Eibenberger and J.M. Doyle, Chem. Phys. Chem. **15** (17), 3800–3804 (2014). doi:10.1002/cphc.v15.17
- [38] P.B. Changala, M.L. Weichman, K.F. Lee, M.E. Fermand and J. Ye, Science **363** (6422), 49–54 (2019). doi:10.1126/science.aav2616
- [39] E.S. Shuman, J.F. Barry, D.R. Glenn and D. DeMille, Phys. Rev. Lett. **103** (22), 223001 (2009). doi:10.1103/PhysRevLett.103.223001
- [40] V. Zhelyazkova, A. Cournol, T.E. Wall, A. Matsushima, J.J. Hudson, E.A. Hinds, M.R. Tarbutt and B.E. Sauer, Phys. Rev. A **89** (5), 053416 (2014). doi:10.1103/PhysRevA.89.053416
- [41] X. Yang, C. Li, Y. Yin, S. Xu, X. Li, Y. Xia and J. Yin, J. Phys. B At. Mol. Opt. Phys. **50** (1), 015001 (2017). doi:10.1088/1361-6455/50/1/015001
- [42] J. Lim, J.R. Almond, M.A. Trigatzis, J.A. Devlin, N.J. Fitch, B.E. Sauer, M.R. Tarbutt and E.A. Hinds, Phys. Rev. Lett. **120** (12), 123201 (2018). doi:10.1103/PhysRevLett.120.123201
- [43] R. Albrecht, M. Scharwaechter, T. Sixt, L. Hofer and T. Langen, Phys. Rev. A **101** (1), 013413 (2020). doi:10.1103/PhysRevA.101.013413
- [44] J.F. Barry, E.S. Shuman, E.B. Norrgard and D. DeMille, Phys. Rev. Lett. **108** (10), 103002 (2012). doi:10.1103/PhysRevLett.108.103002
- [45] S. Truppe, S. Marx, S. Kray, M. Doppelbauer, S. Hof-säss, H.C. Schewe, N. Walter, J. Pérez-Ríos, B.G. Sartakov and G. Meijer, Phys. Rev. A **100** (5), 052513 (2019). doi:10.1103/PhysRevA.100.052513
- [46] S. Hof-säss, M. Doppelbauer, S.C. Wright, S. Kray, B.G. Sartakov, J. Pérez-Ríos, G. Meijer and S. Truppe, New J. Phys. **23** (7), 075001 (2021). doi:10.1088/1367-2630/ac06e5
- [47] J.C. Shaw, S. Hannig and D.J. McCarron, Opt. Express **29** (23), 37140–37149 (2021). doi:10.1364/OE.441741
- [48] J.R. Daniel, C. Wang, K. Rodriguez, B. Hemmerling, T.N. Lewis, C. Bardeen, A. Teplukhin and B.K. Kendrick, Phys. Rev. A **104**, 012801 (2021). doi:10.1103/PhysRevA.104.012801
- [49] T.N. Lewis, C. Wang, J.R. Daniel, M. Dhital, C.J. Bardeen and B. Hemmerling, Phys. Chem. Chem. Phys. **23** (39), 22785–22793 (2021). doi:10.1039/D1CP03515K
- [50] S. Truppe, H.J. Williams, N.J. Fitch, M. Hambach, T.E. Wall, E.A. Hinds, B.E. Sauer and M.R. Tarbutt, New J. Phys. **19** (2), 022001 (2017). doi:10.1088/1367-2630/aa5ca2
- [51] D. Budker, D.F. Kimball and D.P. DeMille, *Atomic Physics. An Exploration Through Problems and Solutions*, 2nd ed. (Oxford University Press, Oxford, 2008).
- [52] M. Doppelbauer, S.C. Wright, S. Hof-säss, B.G. Sartakov, G. Meijer and S. Truppe, J. Chem. Phys. **156** (13), 134301 (2022). doi:10.1063/5.0081902
- [53] Z. Fu, G.W. Lemire, G.A. Bishea and M.D. Morse, J. Chem. Phys. **93** (12), 8420–8441 (1990). doi:10.1063/1.459280
- [54] D.E. Kelleher and L.I. Podobedova, J. Phys. Chem. Ref. Data **37** (2), 709–911 (2008). doi:10.1063/1.2734564
- [55] L. Davis, B.T. Feld, C.W. Zabel and J.R. Zacharias, Phys. Rev. **76** (8), 1076–1085 (1949). doi:10.1103/PhysRev.76.1076
- [56] E.S. Chang, J. Phys. Chem. Ref. Data **19** (1), 119–125 (1990). doi:10.1063/1.555870
- [57] W.H. Parkinson, E.M. Reeves and F.S. Tomkins, J. Phys. B At. Mol. Phys. **9** (2), 157–165 (1976). doi:10.1088/0022-3700/9/2/006
- [58] E.A. Den Hartog, J.E. Lawler, C. Sneden, J.J. Cowan, I.U. Roederer and J. Sobeck, Astrophys. J. Suppl. Ser. **255** (2), 27 (2021). doi:10.3847/1538-4365/ac04b1

- [59] M. Mills, P. Puri, Y. Yu, A. Derevianko, C. Schneider and E.R. Hudson, *Phys. Rev. A* **96** (3), 033402 (2017). doi:10.1103/PhysRevA.96.033402
- [60] Y. Takasu, K. Komori, K. Honda, M. Kumakura, T. Yabuzaki and Y. Takahashi, *Phys. Rev. Lett.* **93**, 123202 (2004). doi:10.1103/PhysRevLett.93.123202
- [61] M.D. Oberlander and J.M. Parson, *J. Chem. Phys.* **105** (14), 5806–5816 (1996). doi:10.1063/1.472457
- [62] A. Jadbabaie, N.H. Pilgram, J. Kłos, S. Kotochigova and N.R. Hutzler, *New J. Phys.* **22** (2), 022002 (2020). doi:10.1088/1367-2630/ab6eae
- [63] K. Ikejiri, H. Ohoyama, Y. Nagamachi and T. Kasai, *Chem. Phys. Lett.* **401** (4), 465–469 (2005). doi:10.1016/j.cplett.2004.11.105
- [64] J.K. Parker, N.L. Garland and H.H. Nelson, *J. Phys. Chem. A* **106** (2), 307–311 (2002). doi:10.1021/jp012895n
- [65] R. Engleman, P. Rouse, H. Peek and V. Baiamonte, Beta and Gamma Band Systems of Nitric Oxide, Division of Technical Information Extension, U.S. Atomic Energy Commission 1969.
- [66] W. Meerts and A. Dymanus, *J. Mol. Spectrosc.* **44** (2), 320–346 (1972). doi:10.1016/0022-2852(72)90109-9
- [67] J.A. Gray, R.L. Farrow, J.L. Durant and L.R. Thorne, *J. Chem. Phys.* **99** (6), 4327–4333 (1993). doi:10.1063/1.466086
- [68] K.L. Reid, S.P. Duxon and M. Towrie, *Chem. Phys. Lett.* **228** (4), 351–356 (1994). doi:10.1016/0009-2614(94)00963-5
- [69] M. Brouard, H. Chadwick, Y.P. Chang, B.J. Howard, S. Marinakis, N. Screen, S.A. Seamons and A.L. Via, *J. Mol. Spectrosc.* **282**, 42–49 (2012). doi:10.1016/j.jms.2012.11.003
- [70] P. Kaspar, F. Munkes, P. Neufeld, L. Ebel, Y. Schellander, R. Löw, T. Pfau and H. Kübler, preprint, arXiv:2206.15243 (2022).
- [71] M. Kirste, X. Wang, H.C. Schewe, G. Meijer, K. Liu, A. van der Avoird, L.M.C. Janssen, K.B. Gubbels, G.C. Groenenboom and S.Y.T. van de Meerakker, *Science* **338** (6110), 1060–1063 (2012). doi:10.1126/science.1229549
- [72] Z. Gao, T. Karman, S.N. Vogels, M. Besemer, A. van der Avoird, G.C. Groenenboom and S.Y.T. van de Meerakker, *Nat. Chem.* **10** (4), 469–473 (2018). doi:10.1038/s41557-018-0004-0
- [73] O. Schullian, J. Loreau, N. Vaeck, A. van der Avoird, B. Heazlewood, C. Rennick and T. Softley, *Mol. Phys.* **113** (24), 3972–3978 (2015). doi:10.1080/00268976.2015.1098740
- [74] M.J. Doppelbauer, O. Schullian, J. Loreau, N. Vaeck, A. van der Avoird, C.J. Rennick, T.P. Softley and B.R. Heazlewood, *J. Chem. Phys.* **146** (4), 044302 (2017). doi:10.1063/1.4974253
- [75] O. Schullian and B.R. Heazlewood, *Mol. Phys.* **117** (21), 3076–3087 (2019). doi:10.1080/00268976.2019.1602740
- [76] Y. Takahashi, D. Shlivko, G. Woolls and N.R. Hutzler, *Phys. Rev. Res.* **3** (2), 023018 (2021). doi:10.1103/PhysRevResearch.3.023018
- [77] O. Schullian, H.S. Antila and B.R. Heazlewood, *J. Chem. Phys.* **156** (12), 124309 (2022). doi:10.1063/5.0083033
- [78] X. Liu, W. Wang, S.C. Wright, M. Doppelbauer, G. Meijer, S. Truppe and J. Pérez-Ríos, *J. Chem. Phys.* **157** (7), 074305 (2022). doi:10.1063/5.0098378
- [79] H. Bethlem, G. Berden and G. Meijer, *Phys. Rev. Lett.* **83** (8), 1558–1561 (1999). doi:10.1103/PhysRevLett.83.1558
- [80] D.P. Engelhart, F. Grätz, R.J.V. Wagner, H. Haak, G. Meijer, A.M. Wodtke and T. Schäfer, *Rev. Sci. Instrum.* **86** (4), 043306 (2015). doi:10.1063/1.4918797
- [81] K.H. Hellwege and J.L. Olsen, Datasheet from Landolt-Börnstein – Group III Condensed Matter · Volume 15B: ‘Metals: Electrical Resistivity, Thermoelectrical Power and Optical Properties’ in SpringerMaterials (doi:10.1007/b19992) Springer-Verlag Berlin Heidelberg, accessed 2022-06-02.