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New techniques for a measurement of the electron's electric dipole moment

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Abstract

The electric dipole moment of the electron (eEDM) can be measured with high precision using heavy polar molecules. In this paper, we report on a series of new techniques that have improved the statistical sensitivity of the YbF eEDM experiment. We increase the number of molecules participating in the experiment by an order of magnitude using a carefully designed optical pumping scheme. We also increase the detection efficiency of these molecules by another order of magnitude using an optical cycling scheme. In addition, we show how to destabilise dark states and reduce backgrounds that otherwise limit the efficiency of these techniques. Together, these improvements allow us to demonstrate a statistical sensitivity of 1.8×10^{-28} e cm after one day of measurement, which is 1.2 times the shot-noise limit. The techniques presented here are applicable to other high-precision measurements using molecules.

1. Introduction

Any local, energy-positive, Lorentz-invariant field theory must conserve CPT, the combined symmetry of parity (P) , time-reversal symmetry (T) and charge conjugation (C) [1, 2]. A permanent electric dipole moment of the electron (eEDM) violates both P and T, and so requires some amount of CP violation to exist. The Standard Model already includes a small amount of CP violation, leading to a very small eEDM: *d*^e ∼ 10[−]⁴⁴ e cm [3]. However, more CP violation is required to explain the observed asymmetry between matter and antimatter in the Universe $[4, 5]$. Theoretical models such as supersymmetry attempt to explain beyond-Standard-Model physics by introducing new sources of CP violation. These lead to larger values for the eEDM: $d_e \sim 10^{-27}$ e cm to 10^{-30} e cm [6], which are accessible to modern precision measurement techniques. Therefore, searching for the eEDM can restrict these new theories, and can also help reveal new physics beyond the Standard Model.

Experiments that search for the eEDM measure the energy shift due to the linear Stark interaction of the eEDM with an electric field. Such experiments typically use paramagnetic atoms or molecules which greatly enhance the strength of this interaction. These systems are also sensitive to any electron–nucleon interaction that violates P and T, but here we adopt the single-source approach of assuming that any P, T-violating signal is due to the eEDM. The energy shift for a paramagnetic system can be expressed as $\Delta E = -d_e E_{\text{eff}}$, where E_{eff} is an effective electric field calculated from atomic and molecular theory which can be much larger than applied fields in the laboratory. For atoms, it turns out that $|E_{\text{eff}}| \approx 8Z^3\alpha^2 E_{\text{ext}}$ where *Z* is the nuclear charge, α is the fine structure constant and E_{ext} is the externally applied electric field [7]. The dependence of the interaction energy on $Z³$ implies that heavy atoms are much better at measuring *d_e*, and indeed early measurements used heavy atoms such as Cs and Tl [8, 9], which have enhancement

factors of $E_{\text{eff}}/E_{\text{ext}} = 120$ and -585 respectively⁴. The linear dependence of E_{eff} on E_{ext} indicates that the atoms are only weakly polarised in the external electric field. In polar molecules, the interaction energy is larger because it is easier to polarise these molecules in an electric field. It is more appropriate to write $E_{\text{eff}} = \eta E_{\text{eff,max}}$ for molecules, where η is the degree of polarisation of the molecule, and $E_{\text{eff,max}}$ is the maximum effective field seen by the electron when the molecule is fully polarised, $\eta = 1$. The latter is typically in the range 10 GV cm^{-1} to 100 GV cm^{-1} , which is much larger than electric fields that can be applied in the laboratory.

In 2011, the precision of atomic measurements was surpassed in an experiment using YbF, setting a new upper limit⁵ of $|d_e| < 1.06 \times 10^{-27}$ e cm [10]. The enhancement of YbF was $E_{\text{eff}} \approx -14.5$ GV cm⁻¹. Crucially, a systematic effect which is large for atoms—the Zeeman interaction with the motional magnetic field mimicking the EDM interaction—is highly suppressed in molecules due to their strong tensor polarisability [11]. In 2014, the ACME collaboration pushed the limit down to $|d_e|$ < 8.7 × 10⁻²⁹ e cm using a beam of ThO molecules in an Ω -doublet state [12]. Molecules in this state are fully polarised in a small applied electric field, giving $E_{\text{eff}} = E_{\text{eff,max}} \approx 84 \text{ GV cm}^{-1}$. The Ω-doublet can also be used conveniently for internal co-magnetometry. In 2017, a measurement using trapped HfF^+ molecular ions $(E_{\text{eff}} \approx 23 \text{ GV cm}^{-1})$ reported the limit $|d_e| < 1.3 \times 10^{-28}$ e cm [13]. This experiment benefited from the long coherence times available in a molecular ion trap, but was limited by the relatively low number of ions trapped. In 2018, the ACME collaboration improved on their limit, reaching $|d_e| < 1.1 \times 10^{-29}$ e cm [14]. This last result constrains any new physics arising from T-violating effects to energy scales above 3 TeV [14].

Many new ideas are now emerging on how to improve measurement sensitivity even further. In present experiments, the total uncertainty is typically dominated by the statistical uncertainty, so methods to improve statistical sensitivity are needed. Laser cooling is a technique that can significantly extend coherence times and brightness of atomic or molecular beams, thereby lowering the statistical sensitivity. The use of laser cooling is evident in proposed eEDM experiments using atoms such as Fr [15], diatomic molecules such as YbF [16] or BaF [17], or polyatomics such as YbOH [18]. Transverse laser cooling of molecular beams of YbF and YbOH has been demonstrated [19, 20] and experiments with ultracold molecules in beams, fountains, or traps now seem feasible. Other ideas include the use of polar molecules embedded in a rare-gas matrix [21]. In this paper, we demonstrate new state preparation and state detection techniques using 174YbF molecules, which increase the statistical sensitivity to the eEDM by a factor of 20 over our previous methods described in reference [22]. The new techniques implemented here can also be applied to other experiments using molecules and, since they rely strongly on scattering many photons from a particular molecular state, are also closely related to the techniques used to apply laser cooling to molecules [23].

2. Measurement method

The relevant energy levels of 174 YbF for the new state preparation and detection schemes are shown in figure 1. We use the molecular population in the lowest three rotational states of the ground electronic and vibrational state, $X^2\Sigma^+(\nu=0, N=0, 1, 2)$, where ν and *N* are vibrational and rotational quantum numbers, respectively. The parity of the rotational states is given by (−1)*^N*, and is indicated in parentheses in figure 1. The interaction between the electron spin, the spin-1/2 fluorine nucleus, and the molecule rotation splits the rotational states into hyperfine states with total angular momenta $F = N + 1$, *N*, *N*, *N* − 1. Because there are two hyperfine states with $F = N$, we distinguish them using the notation N_h and N_ℓ to refer to the states that lie higher and lower in energy, respectively. For $N = 0$, the molecule rotation is absent and so the hyperfine states are simply $F = 0$, 1. Each hyperfine level has $(2F + 1)$ Zeeman sublevels. The *N* = 0 state is used for the eEDM measurement, as will be discussed later. In the experiment, we excite molecules to the electronically excited state $A^2\Pi_{1/2}(\nu'=0, J'=1/2)$, where J' is the total electronic angular momentum of the excited state. This state is split into an Ω−doublet, labelled by *e* and *f*, which have parities +1 and −1 [24]. These are further split by the hyperfine interaction into states $F = 0, 1$, but these are typically unresolved. To ease notation, we use $O(0)$ and $P(1)$ to refer to optical transitions connecting $N = 0$ to f and $N = 1$ to e respectively, as shown in figure 1.

An overview of the experiment is shown in figure 2. A pulsed supersonic beam of YbF molecules, which has a forward velocity of 600 m s⁻¹ and a temperature of 2 K, is produced by ablating a Yb target with a

 4 The sign indicates whether the atomic EDM is parallel (+) or antiparallel (−) to the eEDM.

⁵ All eEDM limits quoted here are at the 90% confidence level.

pulsed Nd:YAG laser into a supersonically expanding gas jet of Ar and SF₆ [25]. The number of molecules participating in the experiment is increased by transferring population from the $N = 0, 1, 2$ states into the $(N, F) = (0, 0)$ state by the new optical pumping scheme, which will be described in detail in section 3. The molecules enter a region of uniform electric and magnetic fields, $E = E\hat{z}$ and $B = B\hat{z}$. In the electric field, the $m_F = 0$ sublevel of $F = 1$ is shifted downwards relative to the $m_F = \pm 1$ states by the Stark interaction. The magnetic field breaks the degeneracy of the $m_F = \pm 1$ states via the linear Zeeman interaction, shifting the energies of the states by $g\mu_B m_F B$, where $g \simeq 1$. A non-zero eEDM, d_e , will further shift the energies of the states by $-d_e m_F E_{\text{eff}}$, where $E_{\text{eff}} = \eta E_{\text{eff,max}}$ and for YbF, $E_{\text{eff,max}} \approx -26$ GV cm⁻¹ [26]. The polarisation factor η depends on the strength of the electric field applied. In our experiment, with $E = 10 \text{ kV cm}^{-1}$, $\eta = 0.558$, so $E_{\text{eff}} \approx -14.5 \text{ GV cm}^{-1}$. A π -pulse of rf magnetic field, polarised along the *x*-axis, then transfers the population into an equal superposition of $m_F = +1$ and $m_F = -1$ states in $F = 1$. The molecules evolve in the *E* and *B* fields for a time $\tau = 800 \mu s$, and another rf π -pulse is applied which projects any population remaining in the original superposition back into the $F = 0$ state. When the rf pulses are perfect π -pulses and exactly on resonance, the probability of finding a molecule in the $F = 0$ or *F* = 1 state is $p_0 = \cos^2(\phi_B + \phi_E)$ or $p_1 = \sin^2(\phi_B + \phi_E)$, where the phases $\phi_B = g\mu_B B\tau/\hbar$ and $\phi_E = -d_e E_{\text{eff}} \tau / \hbar$ are due to the Zeeman and eEDM interactions. The populations in $F = 0$ and $F = 1$ states are then sequentially measured by laser-induced fluorescence detection in two separate detectors labelled A and B. The detection scheme will be discussed in greater detail in section 4.

We call the number of photons counted in each detector s_A and s_B , and combine them to form a quantity called the asymmetry, defined as

$$
\mathcal{A} = \frac{s_A - s_B}{s_A + s_B}.\tag{1}
$$

This quantity is immune to shot-to-shot fluctuations in the number of molecules from the source, N_{mol} . In the ideal detection case, we have

$$
s_{A} = N_{\text{mol}} \epsilon p_{0} = N_{\text{mol}} \epsilon \cos^{2} \phi,
$$

\n
$$
s_{B} = N_{\text{mol}} \epsilon p_{1} = N_{\text{mol}} \epsilon \sin^{2} \phi,
$$
\n(2)

where $\phi = \phi_B + \phi_E$ and ϵ is the number of photons detected per molecule, which we define to be the detection efficiency. The asymmetry is $A = \cos 2\phi$ in this case. We define the contrast, C, as the amplitude of the cos 2ϕ term in the asymmetry,

$$
\mathcal{A} = \mathcal{C} \cos 2\phi. \tag{3}
$$

In the perfect experiment, we have $C = 1$. We discuss imperfections which reduce C in section 5.

Figure 2. Overview of the experiment and interferometer states. (a) Pulsed beams of YbF molecules are produced with equal populations in the four $N = 0$ sublevels. (b) Population is optically pumped using microwave, rf and optical fields into $F = 0$. (c) An equal superposition of $m_F = +1$ and $m_F = -1$ is created by an rf pulse. (d) The two populated levels accumulate a relative phase due to interaction with *E* and *B* fields. (e) Population in the original superposition is projected back into *F* = 0 by an rf pulse. (f) Population in $F = 0$ is detected. (g) Population in $F = 1$ is detected.

The experiment measures ϕ_E by reversing the direction of the external electric field *E* and measuring the resulting change in A , from which we can extract d_e . The uncertainty of the measurement is therefore given by

$$
\sigma_{d_{e}} = \frac{\hbar}{E_{eff}} \sigma_{\phi}
$$
\n
$$
= \frac{\hbar}{E_{eff}} \left| \frac{\partial \phi}{\partial A} \right| \sigma_{A}
$$
\n
$$
= \frac{\hbar}{E_{eff}} \frac{1}{2C |\sin 2\phi|} \sigma_{A}.
$$
\n(4)

To minimise the uncertainty, we use the magnetic field *B* to set $\phi_B = \pi/4$. At this phase, provided that $\epsilon \ll 1$, the uncertainty in the asymmetry can be shown to be limited by Poissonian statistics of photon-counting, $\sigma_A = 1/\sqrt{s_T}$, where the total count $s_T = s_A + s_B$. We will elaborate on this in section 4, as optical-cycling detection can lead to noise in excess of this simple shot-noise limit [27]. In our experiment, we estimate the detection efficiency to be $\epsilon \approx 0.06$, so we can use the shot noise of the detected photons as that of the experiment. The shot-noise limited sensitivity of the experiment can then be written as

$$
\xi_{d_e} = \frac{\hbar}{2E_{\text{eff}}\tau C \sqrt{s_{\text{T}}}}.\tag{5}
$$

This shows that to maximize the sensitivity of the experiment, we should maximize the coherence time, contrast and total photon count. The rest of the paper will focus on new techniques we have implemented to measure both s_A and s_B , and to increase the count rate and contrast.

3. State preparation

Figure 3 shows the relevant transitions in YbF for the state preparation scheme. We want to increase the number of molecules in the $X^2\Sigma^+(v=0, N=0, F=0)$ ground state, which is the starting point for the interferometer. Doing so increases N_{mol} and therefore the total signal s_T .

We optically pump population from the even-parity states $(N, F) = (0, 1), (2, 1), (2, 2_{\ell})$ and $(2, 2_h)$ to (0, 0) via the odd-parity $A^2\Pi_{1/2}(\nu'=0, J'=1/2, f)$ state. The (2, 2_h) and (2, 1) states are closely spaced in energy, and are addressed by a single laser frequency. Due to parity and angular momentum selection rules, the only loss channel is spontaneous decay into higher vibrational states, $\nu > 0$, which has a branching ratio⁶ of 0.072. Population in $(2, 3)$ does not participate in the optical pumping scheme because the state has too much angular momentum. The *N* = 1 states have odd parity, and therefore cannot be optically pumped into the even-parity state $(0, 0)$. Instead, we couple all the hyperfine states in $N = 1$ to the $(0, 1)$ state using one microwave field and two rf fields, as shown in figure 3, so that the population in $N = 1$ can then be optically pumped into $(0, 0)$ as well.

Figure 4 shows the state preparation region. The molecular beam travels vertically along the *y*-axis. The pump lasers are a set of counter-propagating elliptical beams which travel along the *x*-axis. These beams have 1/*e* ² diameters of 23.5× 2.2 mm along the *y* and *z* axes, providing an interaction time of 40 μs with the molecules. The peak laser intensities are approximately 120 mW cm⁻², 340 mW cm⁻² and 140 mW cm⁻² for the frequency components addressing the $(N, F) = (0, 1)$ state, the $(2, 2_h)$ and $(2, 1)$ states, and the

⁶ The Franck–Condon factor for the A($\nu = 0$) $\rightarrow X(\nu = 0)$ transition was measured to be 0.928 [28].

 $(2, 2_{\ell})$ state, respectively. There is also a clean-up beam, which co-propagates with one of the pump lasers in order to remove background signal in the detectors, as explained in section 5.

A microwave horn delivers a maximum of 19 dBm of power at the frequency resonant with the $(0, 1) \rightarrow (1, 2)$ transition. A set of resonant rf coils surround the molecular beam path, with resonances tuned to 30.9 MHz and 161.2 MHz. The two rf fields are first generated by voltage-controlled oscillators, combined and amplified, then sent into the machine to a drive coil which couples the rf to the resonant coils. The excited resonant coils then generate rf radiation which drive M1 transitions between the hyperfine levels in $N = 1$. Both the microwave and rf powers are optimised empirically to maximise the population in $(0, 0)$.

Figure 5 shows the results of the state preparation scheme. Using one of the two detectors, we measure the population in $(N, F) = (0, 0)$ by scanning the frequency of a weak probe laser around the $F = 0$ component of the Q(0) transition, and detecting the laser-induced fluorescence. When we add only the $N = 0$ pump laser, which was the method used in the previous eEDM measurement [10, 22], the signal increases by a factor 1.8 \pm 0.2 over the signal from the thermal population in $F = 0$. The origin of the error bar is shot-to-shot fluctuations in the number of molecules. Numerical simulations using optical Bloch equations (OBEs) [29] predict a factor of 1.9, in good agreement with the measurement. When we add the $N = 2$ pump laser, the enhancement factor grows to 5.9 ± 0.6 . Adding the microwave field increases this to

8.7 \pm 0.8, and adding the rf fields increases this further to 11.8 \pm 1.2. This last factor is again in agreement with OBE simulations which predict a signal increase of 10.5.

4. State detection

A simple way to measure the population in one of the two hyperfine states is to excite molecules in that state and detect the resulting laser-induced fluorescence using the $Q(0)$ transition (see figure 1). This method, which we call Q(0) detection, is the one used in the previous YbF eEDM measurement. It has two disadvantages. First, there is no closed transition from *N* = 0, so each molecule scatters just a small number of photons before it is optically pumped to a different state. From OBE simulations, we calculate that an $(N, F) = (0, 0)$ molecule scatters only 1.2 photons on average. Second, the population in one of the two hyperfine sublevels is partially pumped into the other, so the measurement of one population changes the other.

Here we implement a better detection method using the $P(1)$ transition which scatters many more photons and also allows us to measure s_A and s_B independently. Our $P(1)$ detection scheme is shown in detail in figure 6. We detect the molecules in $F = 0$ and $F = 1$ sequentially, in detectors A and B. In detector A, a resonant microwave field couples the states $(N, F) = (0, 0)$ and $(1, 1_f)$. A probe laser together with sidebands, tuned to the the $P(1)$ transition, is used to detect the molecules. In detector B, the same is carried out except with resonant microwaves coupling the states $(0, 1)$ and $(1, 2)$ instead, which detects population in $F = 1$. The $P(1)$ transition is rotationally closed: molecules can only either decay back to the $N = 1$ states in the $\nu = 0$ manifold, or to higher vibrational states. The branching ratio for the decay to $\nu > 0$ states is $\Gamma_{\nu>0} = 0.072$, so each molecule scatters, on average, $1/(1 - \Gamma_{\nu>0}) = 13.8$ photons before it becomes dark to the probe laser.

The two detectors are illustrated in figure 7. Each includes two microwave horns, although only one is used when detecting the molecules. The other horn is included so that the standing wave pattern of the microwave field can be changed, allowing us to test for systematic effects in the experiment. In each detector, we can apply either microwave frequency, so the order of detecting the $F = 0$ and $F = 1$ populations can be reversed to check for systematic effects. An optical fibre delivers light of all three probe frequencies to the upper detector and this same light is routed to pass through the lower detector to ensure some common-mode rejection of fluctuations in the light. A second fibre delivers another probe beam to the lower detector, which counter propagates along the same path. Each beam has a circular Gaussian profile and a 1/*e*2-radius of 6 mm. The combined peak intensities of the two beams are 80 mW cm−2, 160 mW cm[−]² and 80 mW cm[−]² for the three different frequencies which address the *F* = 1, *F* = 0/*F* = 2, and $F = 1_h$ hyperfine levels respectively, as shown in figure 6. The middle frequency addresses two hyperfine levels as they are very closely spaced in energy (see figure 1).

The resulting laser-induced fluorescence from the molecules is collected by a large spherical mirror and imaged onto a PMT by a pair of aspheric lenses. With such a high laser power in the detection region, it is imperative that we minimise the background signal due to laser scatter. Although this background can be measured⁷ and subtracted away, there is still the increase in uncertainty due to the noise in the laser scatter. We minimise this background by blackening the interior surfaces of the chamber δ , using optical baffles,

 $\frac{7}{7}$ This is done by measuring the PMT signal when the molecules are not in the detection region.

⁸ Some surfaces were blackened using soot from an acetylene flame, and others were blackened using a black paint from Alion Science (MH2200).

extending the input and output arms of the laser ports, and angling the port windows. With these measures, the scatter comes down to about 50 (100) photons in 1 μs in the lower (upper) detection region, equivalent to 5% (10%) of the molecular signal averaged over the time window used in the data analysis. The increase in uncertainty in the measurement due to noise in the laser scatter is then negligible.

To ensure that the optical cycling due to the rotationally-closed nature of the $P(1)$ transition is maximised, we use three probe laser frequencies (see figure 6) so that we excite molecules from all of the $N = 1$ hyperfine levels. Furthermore, some of the P(1) transitions have $F \geq F'$, where F, F' are the total angular momenta of the hyperfine states in the ground and excited electronic states. It is well known that if the driving laser has a static polarisation, such a configuration has dark states, which will significantly reduce the number of photons scattered per molecule [30]. We destabilise these dark states by modulating the polarisation of the probe beams between *y* and *z* (with circular in between) using an electro-optic modulator (EOM). We use a rate of 0.9 MHz because our OBE simulations give a very broad optimum centered on this frequency. We also counter-propagate the probe beams in order to create polarisation gradients in the *x*-direction. The molecules then experience a further varying laser polarisation as they fly through the detection region since they have non-zero transverse velocity.

Figure 8 shows the improvement in the signal from one of the detectors when we use the methods described above to increase the optical cycling of the molecules. In all the cases, the total laser power is 100 mW. For the counter-propagating case, we have used 50 mW per beam. The results are summarized in table 1. When sidebands are added to the probe laser, the signal detected increases by a factor of 2.3 ± 0.3 , in agreement with our simulations. When we also modulate the polarization of the probe lasers, the signal detected is a factor of 3.8 \pm 0.4 higher than the single-frequency case. This is lower than the OBE simulation prediction of 5.2, suggesting that the polarization of the probe light is not being fully modulated after passing through the optical fibres. If we counter-propagate (keeping the total power constant), the increase in signal is 4.9 ± 0.7 times, which agrees with the predicted increase when the dark states are fully destabilized. The OBE simulations additionally show that even with full polarization modulation, we have yet to reach the maximum possible average of 13.8 photons per molecule. This is due to insufficient interaction time of the molecules with the probe lasers in the detection region. A larger probe beam diameter could have been used, but it would have resulted in an unacceptably large amount of laser scatter in our detection chamber, and also made it more challenging to image all the signal photons onto the PMTs.

Lasner and DeMille have carefully considered the signal-to-noise ratio obtained in an experiment such as ours, where each molecule scatters many photons before being pumped to a dark state, and each photon

8

Table 1. Comparison of signal increase due to detection techniques with that predicted by OBE simulations.

has a certain probability of being detected [27]. They find that the expression for the shot noise has to be corrected because the probability distribution of the detected photons per molecule is broader than that of a Poissonian distribution. In our case, this correction is small because the number of photons detected per molecule is much smaller than 1. If the detection efficiency can be improved in the future, it will be important to take proper account of the excess noise factor discussed in reference [27].

To conclude this section, the average of 11.9 photons scattered per molecule using this $P(1)$ detection is an order of magnitude higher than the 1.2 photons scattered per molecule using the previous $Q(0)$ detection method. Moreover, $P(1)$ detection allows us to detect both $F = 0$ and $F = 1$ molecules independently, which doubles the number of useful molecules in the experiment compared to Q(0) detection, further improving our sensitivity. The next section discusses imperfections in our P(1) detection scheme and describes methods to overcome them.

5. Improving the contrast

We have shown in section 2 that in the ideal case, the photon counts in detectors A and B are given by $s_A = N_{\text{mol}} \epsilon p_0 = N_{\text{mol}} \epsilon \cos^2 \phi$ and $s_B = N_{\text{mol}} \epsilon p_1 = N_{\text{mol}} \epsilon \sin^2 \phi$. If there are imperfections in the two rf π -pulses, due to imperfect pulse area or frequency, then the probability of detecting a molecule in $F = 0$ and $F = 1$ after the second pulse has the form [22]

$$
p_0 = a_1 \cos^2 \phi + a_R \cos \phi + a_{C,0},
$$

\n
$$
p_1 = a_1 \sin^2 \phi - a_R \cos \phi + a_{C,1},
$$
\n(6)

where a_I is the amplitude of the main interference signal we want to measure, a_R is the amplitude of the Ramsey fringe that arises from the residual coherence between $F = 0$ and $F = 1$, and $a_{C,i}$ is a constant term that is different for the two states. The amplitudes are constrained such that $p_0 \geqslant 0, p_1 \geqslant 0,$ and $p_0 + p_1 = 1.$

Now let us consider imperfections in the detection scheme and how they affect the signals. Let the lower detector be A, i.e. the one that detects population in $F = 0$. First, the optical cycling process does not completely deplete the population in $N = 1$, and so we write the fraction of population left in $N = 1$ after detector A as *f*left. Next, we include the observation that there is some off-resonant driving of the *F* = 1 state when driving the $F = 0$ microwave transition, and vice versa, and denote the fraction of population that is off-resonantly driven as *f*or. We account for the difference in detection efficiencies of the two regions by a fraction f_ϵ , so that the detection efficiencies are $(1 \pm f_\epsilon)\epsilon$ in the two regions. In addition to the background due to laser scatter from the apparatus, which we have discussed earlier, we have another constant background term due to scattering from molecules in other states, which do not participate in the experiment. We represent the latter as a fraction of the total number of molecules, $f_{\text{bg},A/B}$. The measured photon counts can then be written as

$$
s_{A} = N_{\text{mol}}\epsilon a_{I}(1+f_{\epsilon})(\cos^{2}\phi + f_{R}\cos\phi + f_{\text{or}}\sin^{2}\phi + f_{bg,A}),
$$

\n
$$
s_{B} = N_{\text{mol}}\epsilon a_{I}(1-f_{\epsilon})(\cos(1-f_{\text{or}})(\sin^{2}\phi - f_{R}\cos\phi) + f_{\text{left}}\cos^{2}\phi + f_{bg,B}),
$$
\n(7)

where we have omitted terms with products of small fractions. Here $f_R = a_R/a_I$ is the Ramsey term amplitude as a fraction of the signal amplitude. The asymmetry, as defined in equation (1), can now be written as

$$
\mathcal{A} = \frac{s_A - s_B}{s_A + s_B}
$$
\n
$$
\approx \left(\frac{f_{\epsilon}}{2} + f_{\text{or}} - \frac{3f_{\text{left}}}{4} + f_{\text{bg},A} - f_{\text{bg},B}\right) + 2f_R \cos \phi + \left(1 - f_{\text{or}} - f_{\text{left}} - f_{\text{bg},A} - f_{\text{bg},B}\right) \cos 2\phi - \frac{1}{4} \left(2f_{\epsilon} + f_{\text{left}}\right) \cos 4\phi\right)
$$
\n
$$
\equiv \mathcal{A}_{\text{c}} + \mathcal{A}_{R} \cos \phi + \mathcal{C} \cos 2\phi + \mathcal{A}_{\text{d}} \cos 4\phi,
$$
\n(8)

where again we have neglected all terms proportional to a product of two or more small quantities. If the order of detecting the molecules is reversed, we obtain a very similar expression to equation (8).

In line with our earlier definition of the contrast in equation (3), we have set C to be the coefficient of the cos 2ϕ term in the expression for A. In addition, there is a constant term \mathcal{A}_c , a Ramsey term with amplitude A_R and a frequency-doubled term with amplitude A_d . The Ramsey component can be removed using the methods described in reference [22], and since all the other imperfections are small, the $\cos 2\phi$ term dominates. The sensitivity of the experiment depends on how well we can measure a change in A from a corresponding change in the phase ϕ , which is given by

$$
\left|\frac{\partial \mathcal{A}}{\partial \phi}\right|_{\phi=\pi/4} = \left|(-2\mathcal{C}\sin 2\phi - 4\mathcal{A}_d\sin 4\phi)\right|_{\phi=\pi/4} = 2\mathcal{C} = 2\left(1 - f_{\text{or}} - f_{\text{left}} - f_{\text{bg,A}} - f_{\text{bg,B}}\right),\right.\tag{9}
$$

Table 2. Improvements in contrast. We show the contrast for the cases where either $F = 0$ or $F = 1$ population is detected first. Unless stated, the total laser power is 100 mW (if using counter-propagating beams, this corresponds to 50 mW per beam).

| | Contrast $(F = 0$ first) | Contrast $(F = 1$ first) | | |
|----------------------------------|-----------------------------|-----------------------------|--|--|
| No improvements | 0.35 | 0.37 | | |
| Clean-up beam (CU) | 0.44 | 0.45 | | |
| Counter-propagating beams (CP) | 0.39 | 0.40 | | |
| Polarization modulation (PM) | 0.45 | 0.43 | | |
| $PM+CU$ | 0.56 | 0.53 | | |
| $PM + CP + CU$ | 0.58 | 0.55 | | |
| $PM + CP + CU$ at 80 mW per beam | 0.62 | 0.61 | | |
| | | | | |

where we have set $A_R = 0$ and ϕ is set to $\pi/4$ to maximise the slope, as in section 2. The sensitivity of the experiment can therefore be maximised by minimising the imperfection terms. We now look at each of these terms.

We minimise the off-resonant driving from the microwaves by optimising the detection microwave powers to achieve the best contrast. The fraction of off-resonant population after optimisation is measured to be $f_{\text{or}} = 0.04 \pm 0.01$. To minimise the leftover population, we destabilise the dark states in the *N* = 1 levels by modulating the probe laser polarisation and counter-propagating the beams, as discussed in section 4. After applying these methods, we measure $f_{\text{left}} = 0.17 \pm 0.02$.

The background scatter from the molecular beam mainly comes from two sources. The first source is molecular population in $N = 1$ that remained after the optical pumping step. We show this in figure $9(a)$, where we operated the usual optical pumping scheme, but scanned a weak probe laser in the detection region around the $174P(1)$ transitions⁹. After optical pumping, there remains a significant amount of population in the $F = 0/F = 2$ hyperfine levels of $N = 1$ in ¹⁷⁴YbF. About 17% of the population is left in $N = 1$ after optical pumping.

The other isotopologues of YbF are the second source of background from molecular scattering. We show this in figure 9(b). In the figure, we note that the $^{174}P(1)$ lines overlap with the $^{172}P(2)$ lines, so the latter can contribute to background scatter since our detection method requires high laser power, leading to very large power broadening. The natural abundances of 174YbF and 172YbF are 32% and 22% respectively, so this is a significant source of background. We demonstrate this by applying a pump laser tuned to the ¹⁷⁴P(1) transitions (the same laser frequencies as the probe lasers). The $N = 1$ levels are depleted, though

⁹ Here, we use the superscript to denote the relevant isotope of Yb that the transition refers to.

Table 3. Fit parameters for the data shown in figure 10.

| Measuring | | | $\mathcal{A}_{\rm R}$ | | $\phi_{\rm b}$ (rad) |
|--------------------------|----------|-------|-----------------------|----------|----------------------|
| $F = 0$ population first | 0.623 | 0.054 | -0.012 | -0.041 | -0.21π |
| $F = 1$ population first | -0.609 | 0.045 | 0.001 | -0.026 | 0.29π |

not completely, and there are other peaks in the vicinity that are also pumped out. These other peaks contribute to the background we observe in the detectors.

This background due to other molecules can be minimized by introducing some 'clean-up' light into the optical pumping region. This clean-up light is obtained by picking off a small amount of probe laser light, broadening its frequency spectrum with an overdriven EOM such that its bandwidth is about 200 MHz, and directing this light into the optical pumping region. Since the $^{172}P(2)$ transitions are not rotationally-closed, whereas the $^{174}P(1)$ transitions are, we remove the background without adversely affecting the efficiency of the regular optical pumping scheme. With the clean-up beam in place, we measure the contributions of the background signal (from both sources) to be $f_{\text{bg,A}} = 0.17 \pm 0.02$ and $f_{\text{bg,B}} = 0.06 \pm 0.01$.

Based on our measurements of the parameters appearing in equation (9), we expect the contrast to be $|\mathcal{C}| = 0.56 \pm 0.03$. In table 2, we show the improvement in contrast after applying the methods described in this section: (i) polarisation modulation of the probe lasers (PM), (ii) counter-propagating beams in the detector (CP) and (iii) clean-up beam in the optical pumping region (CU). The contrast was measured by fitting interference curves, obtained by scanning the magnetic field, to equation (8). We have achieved a contrast of $|C| = 0.62$ for the detector configuration where we measure $F = 0$ population first, and a contrast of $|C| = 0.61$ for the opposite configuration that measures $F = 1$ population first, which is consistent with our estimation. Figure 10 shows the interference curves and fits with all the improvements and maximum laser power. We have added a phase offset ϕ_b to account for the ambient background magnetic field. The fit parameters are shown in table 3. The other terms in the asymmetry, A_c , A_R and A_d , are small compared to the contrast.

6. Sensitivity of the experiment

We now investigate the statistical sensitivity for an eEDM measurement when we use the new techniques described above. We call a single pulse of the molecular beam a 'shot'. From each shot, we obtain two pulses of photon counts, $s_A(t)$ and $s_B(t)$, at the two detectors. The lower and upper detectors are at distances L_1 and *L*² from the source. The molecules reaching the upper detector at time *t* arrive in the lower detector at the earlier time $t' = (L_1/L_2)t$, so we calculate the asymmetry as $A(t) = (s_A(t') - s_B(t))/(s_A(t') + s_B(t))$. The function $A(t)$ represents the asymmetry for molecules with different arrival times and hence different velocities.

As described in reference [22], we collect 4096 consecutive shots into a 'block'. In each block, we switch a total of nine different parameters between two values, which allow us to calculate the eEDM, protect against systematic errors and noise, and optimise parameter values. To measure the eEDM, three parameters are crucial: E, which sets the direction of the electric field; B, which sets the magnetic field such that the interferometer phase is $\pm \pi/4$; and δ B, which adds or subtracts a small magnetic field ΔB to the one set by *B*. For a switched parameter $X = \{+1, -1\}$, we define the asymmetry correlated with the parameter as its 'channel' $\{X\}$. These channels retain the time-dependence of $\mathcal{A}(t)$. It can be shown that from a block, we obtain the eEDM as follows [22]:

$$
d_{e}(t) = \frac{g\mu_{B}\Delta B}{\eta E_{\text{eff}}} \frac{\{\text{E} \cdot \text{B}\}}{\{\delta \text{B}\}}.
$$
\n(10)

The time-dependence of d_e reminds us that for each block, we obtain measurements of d_e over a range of arrival times of the molecules.

To demonstrate the sensitivity of the experiment, we took $N_{\text{blocks}} = 212$ blocks of data over a total of 22 hours, which we define to be one day of measurement time. We choose to analyze d_e between molecule arrival times of 2675 μ s and 2905 μ s, chosen to maximize the final sensitivity. These correspond to molecules with forward velocities between 590 m s[−]¹ and 640 m s[−]1. The resolution of data acquisition is 10 μ s, so we obtain 23 values of $d_e(t_i)$ per block, where $\{t_i\}$ are the arrival time bin centre values. A fixed, but unknown offset is added to all the values of d_e to blind the analysis procedure. Figure 11(a) shows the distribution of eEDM values with their mean subtracted. The line is a normal distribution with zero mean and standard deviation equal to the average statistical uncertainty for each eEDM value, which is given by

$$
\sigma_{d_{\rm e}, \text{single}} = \sqrt{N_{\rm v}} \,\sigma_{d_{\rm e}},\tag{11}
$$

where $N_v = 212 \times 23 = 4876$ is the total number of values of d_e we include in the analysis, and σ_{d_e} is the estimated standard error on determining the mean of *d*e.

The standard error, σ_{d_e} , is obtained as follows. First, we divide the N_v values of d_e into 23 equal datasets, grouped by their arrival time. We estimate the means of *d*^e for each arrival time bin using the bootstrapped 10% trimmed mean [31]. The trimmed mean is used because it is more robust to outliers, and bootstrapping is a random resampling technique used to easily obtain standard errors for parameter estimates even for non-normal distributions [31].

For each value of t_i , we have a set of N_{blocks} values of d_e . From this set, we randomly draw N_{blocks} values of d_e to form a synthetic set, which we call a bootstrap set. We allow the same value to be drawn more than once. We then calculate the trimmed mean for the bootstrap set, which we call the bootstrap estimate of the trimmed mean. The procedure is repeated 5000 times, from which we obtain 5000 bootstrap estimates. The standard error on the trimmed mean is the standard deviation of these bootstrap estimates. We carry out the same procedure for each value of t , and finally take the weighted mean of the values of d_e for the different *t*'s, where the weights are given by the inverse squares of the standard errors. The final uncertainty for *d*^e is then given by the standard propagation of errors. We find that

$$
\sigma_{d_e} = \frac{1.8 \times 10^{-28}}{\sqrt{N_{\text{days}}}} \text{ e cm},\tag{12}
$$

where N_{days} is the number of days of measurement (where each day corresponds to 22 hours of measurement time). This is a factor of 20 better than the per-day sensitivity from the previous measurement [22].

It is useful to compare the experimental sensitivity to the shot-noise-limited sensitivity given in equation (5). Since we had subtracted away the background due to laser scatter from the apparatus, we now need to add in the noise in this background. Assuming that the latter is just shot noise, we get

$$
\xi_{d_{\rm e}} = \frac{\hbar}{2E_{\rm eff}\tau C} \sqrt{\frac{s_{\rm T} + s_{\rm scatter}}{s_{\rm T}^2}},\tag{13}
$$

where *s*_{scatter} is the total laser scatter measured in both detectors. To calculate the shot noise for a block, we sum over the contributions from all the shots, but retain the dependence on the arrival time *ti*:

$$
\xi_{d_{\rm e},\rm block}(t_i) = \frac{\hbar}{2E_{\rm eff}\tau C(t_i)} \frac{1}{N_{\rm shots}} \sqrt{\sum_{k=1}^{N_{\rm shots}} \frac{s_{\rm T,k}(t_i) + s_{\rm scatter},k}{(s_{\rm T,k}(t_i))^2}},\tag{14}
$$

where $s_{T,k}$ and $s_{scatter,k}$ are the total signal and laser scatter measured in the *k*th shot in the block, $N_{\text{shots}} = 4096$ is the number of shots in a block, and $\mathcal{C}(t_i)$ is the measured contrast as a function of arrival time from the block. For the dataset presented in this paper, this shot-noise-limited sensitivity is shown by the red dashed line in figure 11(b). In the same figure, the blue circles are the experimental uncertainties of d_e for each molecule velocity, $v_i = L_2/t_i$. This was calculated using the same bootstrapping method presented earlier. The increase in uncertainty for the molecules with higher speeds is currently under investigation. For the rest of the molecules, we are very close to the shot-noise limit. Carrying out a

weighted sum of the shot-noise uncertainties, where the weights are the same as those used for the eEDM values, gives

$$
\xi_{d_{\rm e}} = \frac{1.5 \times 10^{-28}}{\sqrt{N_{\rm days}}} \,\text{e cm},\tag{15}
$$

indicating that the statistical sensitivity of the experiment is 1.2 times above the shot-noise limit.

7. Conclusion and outlook

The new state preparation and detection techniques presented in this paper, together with better collection optics in the detectors, have increased the number of photons detected by a factor of 400 since the previous eEDM measurement [22]. The state preparation technique used two separate lasers to address the $N = 0$ and $N = 2$ molecular populations, and a combination of microwave and rf fields to connect the $N = 1$ population to $N = 0$. By leaving out the $F = 0$ component of the Q(0) laser, we are able to optically pump population into the desired ground state. After the interferometer region, the molecules in $F = 0$ and $F = 1$ were separately detected by connecting them to the $N = 1$ levels and the detection efficiency was maximised by using the rotationally-closed P(1) transition. By modulating the polarisation of the detection lasers, counter-propagating the beams, and introducing a clean-up beam in the optical pumping region, we were able to maximise the contrast of the detector. These techniques are also important in the field of laser cooling molecules, where it is also necessary to scatter a large number of photons per molecule. Finally, we have demonstrated that the statistical sensitivity of the experiment is 1.2 times its shot-noise limit. Our current sensitivity is given by equation (12) and is 20 times better than in our previous measurement. With this sensitivity, we would expect a statistical uncertainty at the 10^{-29} e cm level in a measurement consisting of 100 days of eEDM data. This is at a similar level of sensitivity to the current leading eEDM experiments [13, 14], but with a different molecular species and experimental setup.

Further upgrades to the present apparatus are also possible. For example, we plan to introduce light tuned to the R(1) transition just before each detection region, in order to remove population left behind in $N = 1$ and therefore reduce the background. We also plan to use isotopically-pure ¹⁷⁴Yb for our molecular source, which will increase the number of useful molecules by a factor of three, since 174 Yb has a natural abundance of only 32%. This will also remove the background signal due to transitions in other isotopologues of YbF, as discussed in section 5. Furthermore, we have recently developed a buffer gas source of YbF and implemented transverse laser cooling [19]. This source provides a higher flux of molecules which have a lower forward velocity. Using a laser-cooled beam, much longer coherence times become feasible. The slower beam will also have a longer interaction time with the detection lasers, thereby giving a larger signal. Finally, the detection efficiency can be improved further by adding vibrational repump lasers to increase the number of photons scattered per molecule. The large expected increase in signal will also then make it favourable to switch from PMTs to silicon photomultipliers (SiPMs) which have a higher quantum efficiency, therefore increasing our overall detection efficiency. With all these proposed improvements, we expect to be able to probe the eEDM at the 10[−]³¹ e cm level. The data presented can be accessed from Zenodo [\(https://doi.org/10.5281/zenodo.3751773\)](https://doi.org/10.5281/zenodo.3751773) and may be used under the Creative Commons Attribution 4.0 International license.

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