## Second-Scale Coherence Measured at the Quantum Projection Noise Limit with Hundreds of Molecular Ions

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Cold molecules provide an excellent platform for quantum information, cold chemistry, and precision measurement. Certain molecules have enhanced sensitivity to beyond standard model physics, such as the electron's electric dipole moment (eEDM). Molecular ions are easily trappable and are therefore particularly attractive for precision measurements where sensitivity scales with interrogation time. Here, we demonstrate a spin precession measurement with second-scale coherence at the quantum projection noise (QPN) limit with hundreds of trapped molecular ions, chosen for their sensitivity to the eEDM rather than their amenability to state control and readout. Orientation-resolved resonant photodissociation allows us to simultaneously measure two quantum states with opposite eEDM sensitivity, reaching the QPN limit and fully exploiting the high count rate and long coherence.

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Molecular ions are being used in the search for the electron's electric dipole moment (eEDM) [1], with the potential to break the current sensitivity limit [2]. The flexibility of ion traps enables the probing of a coherent superposition at long times, directly improving sensitivity and reducing the susceptibility to systematic uncertainty. This intrinsic advantage provides the freedom to choose molecules such as HfF<sup>+</sup> and ThF<sup>+</sup>, which have enhanced sensitivity to the eEDM, leveraging the long interrogation time to search for physics beyond the standard model. Molecular ions can also benefit precision measurements in related experimental investigations, such as electroweak interaction [3], fundamental symmetry violation [1,2,4–7], variation of fundamental constants [8–10], and dark matter searches [9,11,12]. While directly laser coolable molecules [13–17] and assembled molecules [18,19] are typically chosen for their efficient state preparation and readout rather than their measurement utility, for our molecular ion systems, we must separately devise efficient state preparation and detection methods to fully exploit their amenability to trapping and make the most precise measurement possible. Specifically, we desire to measure coherent quantum-state superpositions at the fundamental limits set by the state lifetime and the quantum projection noise (QPN) limit. A challenge in precision metrology is that harvesting the QPN-limited sensitivity becomes ever harder with increasing count rate, as technical noise becomes proportionally more significant. We present here a noise-immune scheme for extracting the internal quantum phase in large samples of molecular ions.

We employ efficient internal state cooling to significantly enhance the population in the desired science state for the EDM search with both ThF<sup>+</sup> and HfF<sup>+</sup>. To fully utilize the large number of signal ions, we use angularly resolved photodissociation to measure two quantum states of opposite molecular orientation in the same experiment cycle to differentially isolate the coherent superposition signal from technical noise, extracting it near the QPN limit (Fig. 1). For an *e*EDM-like experiment using trapped molecular ions, we demonstrate that these improvements decrease our statistical uncertainty by more than an order of magnitude. Our second-scale coherence is now measured

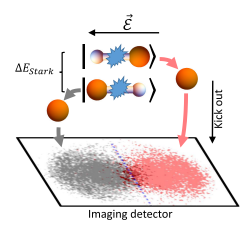


FIG. 1. Molecules are prepared in two oriented states. Upon dissociation, the photofragments are ejected in the direction of the molecular orientation and kicked out toward an imaging detector. The 2D image of Hf $^+$  ions after HfF $^+$  dissociation is pictured. The quantization axis is defined by the applied electric field  $\vec{\epsilon}.$   $\Delta E_{\text{Stark}}$  is the energy difference between the molecular orientations.

near the QPN limit, leading to a statistical sensitivity of 0.3 mHz $\sqrt{h}$  with HfF<sup>+</sup> (corresponding to an *e*EDM uncertainty of  $2.7 \times 10^{-29} e \text{ cm}\sqrt{h}$ ) as compared to Ref. [1] where the best sensitivity was 14.5 mHz $\sqrt{h}$ .

State preparation.—The high multiplicity of closely spaced levels in molecules such as HfF<sup>+</sup> and ThF<sup>+</sup> means that any desired coherent signal is highly diluted at the beginning of an experiment cycle. Careful consideration of the electronic transition branching ratios and leveraging of the ideas developed for optical pumping and laser cooling of molecules [20-24] allows us to compensate for significantly off-diagonal Franck-Condon elements such that we can concentrate the population into the lowest rovibrational quantum state of the eEDM-sensitive target  ${}^{3}\Delta_{1}$ . This increases our signal while reducing the number of harmful spectator ions, which serve as a possible source of systematic uncertainty and limit the achievable coherence. We prepare the desired spin-polarized stretched states,  $|m_F = 3/2, \Omega = \pm 1\rangle$ , which are the initial states in our spin precession measurements, by applying circularly polarized light along the  $\Omega = 0^-$  transition [Fig. 2(a)]. This excited state forms an electronically closed system with  ${}^3\Delta_1$  due to parity symmetry. (The quantum number  $\Omega$ denotes the projection of the total angular momentum onto the internuclear axis.) Overall, including the rotational and vibrational cooling, we accumulate 60% of the total population in the spin stretched states for ThF<sup>+</sup>

(see Supplemental Material [25]) and a comparable fraction for HfF<sup>+</sup>. We can further prepare a single stretched state with a specific molecular orientation (given by  $m_F\Omega$ ), such as  $|m_F=+3/2,\Omega=+1\rangle$  for diagnostic purposes by depletion of the unwanted state. We label these four relevant fully oriented and fully stretched states A, B, C, and D [Fig. 2(b)]. Additionally, we protect our target science state from contamination by the remaining spectator ions in various internal molecular quantum states via depletion or repumping of the possible decay channels that may occur over the course of the long interrogation time (see Supplemental Material [25]).

With the initial population prepared in states A and C, we initiate the spin precession by applying a  $\pi/2$  pulse, lowering the electric bias field as demonstrated in Ref. [1], to create a coherent superposition in both the upper (A, B) and lower (C, D) Stark doublets. The same pulse works for both orientations simultaneously. The eEDM-like signal contributes to the differential spin precession phase between these doublets. We can compute the phase for the spin polarizations in the upper and lower Stark doublets by reading out the populations  $N_A$ ,  $N_B$ ,  $N_C$ , and  $N_D$  depicted in Fig. 2(b) according to

$$\wp_u = \frac{N_A - N_B}{N_A + N_B} \sim \mathcal{C} \sin \phi^u, \tag{1}$$

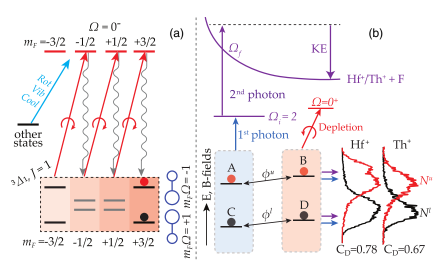


FIG. 2. Preparation of the initial quantum state and orientation-selective photofragmentation of HfF<sup>+</sup> and ThF<sup>+</sup>. (a) We use circularly polarized light via an  $\Omega=0^-$  electronic state to pump to the fully stretched  $m_F$  states of the lowest rovibrational state of  $^3\Delta_1$ . (b) After the spin precession sequence, we project the fully stretched  $m_F$  states by depletion of the other stretched states. This is done with a circularly polarized excitation to the  $\Omega=0^+$  from which the possibility of decay back to  $^3\Delta_1$  is small. Additional molecular species specific optical and microwave fields provide rotational and vibrational cooling (see Supplemental Material [25]). This is followed by two-color photodissociation of the *e*EDM-sensitive  $^3\Delta_1$  states, where the product of  $m_F$  and  $\Omega$  determines the molecular orientation. The first REMPD photon excites the molecules to the fully stretched bound intermediate state  $|\Omega_i=2\rangle$ , which maintains their orientation. The second REMPD photon couples the intermediate state to the continuum states, resulting in oriented photofragments, Hf<sup>+</sup> or Th<sup>+</sup> and F atoms, and determines the kinetic energy (KE). The metallic ions from each molecular orientation are individually mapped onto an imaging detector with the resulting 1D distributions. Their orientation contrasts  $C_D$  (defined in the Supplemental Material [25]) are 78% and 67%, respectively.

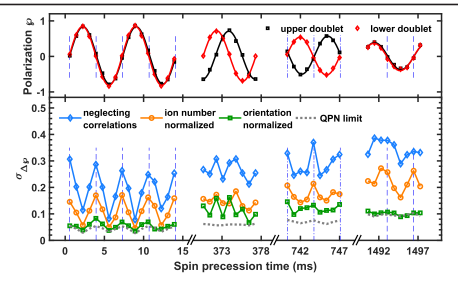


FIG. 3. Demonstration of a differential measurement at the QPN limit. (Upper) Spin precessions of the upper and lower doublets [Eqs. (1) and (2)], which have  $\sim$ 0.6 Hz frequency difference caused by different magnetic moments. (Lower) The scatter in  $\Delta_{\wp}$ . From the scatter of  $\wp_u$  and  $\wp_l$  determined independently, we would anticipate  $\sigma_{\Delta_{\wp}}$  (blue diamonds) to be 5 times above the QPN limit (dotted line). With normalization by each cycle's ion production noise (orange circles),  $\sigma_{\Delta_{\wp}}$  is still more than 2 times the QPN limit. By extracting  $\Delta_{\wp}$  with simultaneous orientation-selective detection (green squares), we completely eliminate all common-mode noise to bring  $\sigma_{\Delta_{\wp}}$  close to the QPN limit when the fringes of the upper and lower doublets are in phase. Spin precession phase noise dominates the out-of-phase measurements ( $\sim$ 745 ms). At early times ( $\sim$ 5 ms), dips in  $\sigma_{\Delta_{\wp}}$  are observed at maximum or minimum polarizations [Eq. (3)]. Vertical dashed lines mark the zero crossings when the spin precessions are in phase, while dash-dotted lines mark them for the out-of-phase case.

$$\wp_l = \frac{N_C - N_D}{N_C + N_D} \sim \mathcal{C} \sin \phi^l, \tag{2}$$

where C is the spin precession contrast, and  $\phi^u$  and  $\phi^l$  are the spin precession phases of the upper and lower doublets, respectively (Fig. 3). The coherence time measured in this dataset is 1.9(1) s for both molecular orientations and the state lifetime is limited by collisions to 1.14 s.

*Noise-immune state detection.*—The most broadly applicable high-yield methods for state-sensitive molecular detection such as resonant photoionization and photodissociation rely on pulsed lasers and are frequently contaminated with noise well in excess of QPN. Moreover, they generally allow for the detection of only a single state in a closely spaced multiplet, precluding the possibility of differential measurements. We use resonanceenhanced multiphoton dissociation (REMPD) followed by mass spectrometry to distinguish the Hf<sup>+</sup> (Th<sup>+</sup>) photodissociation products from background HfF<sup>+</sup> (ThF<sup>+</sup>) [31,32]. Laser fluctuations give rise to excess noise on the detected ion number N with standard deviation  $\alpha N$ , while QPN scales as  $N^{1/2}$  [33]. For instance, our experimental cycle of molecule creation, preparation, and detection involves five pulsed lasers, four of which are frequency doubled. Even with careful monitoring of laser frequencies and intensities, we observe shot-to-shot fractional noise in excess of  $\alpha \sim 0.2$ . For  $N \le 25 \sim 1/\alpha^2$ , the QPN limit is reached. For increased N, such as our typical sample size  $N \gtrsim 500$ , excess noise from our lasers dominates and the signal-to-noise ratio stops improving [34].

To circumvent these limitations, we developed orientation-selective photofragment imaging, which allows us to count the ion populations in two distinct molecular orientations in a single cycle (Fig. 1). The inherent correlation between these two populations that are prepared and read out by the same laser pulses allows us to make differential measurements at the QPN limit while detecting hundreds of ions in each cycle. Not only are all common-mode fluctuations arising from ionization, state preparation, and detection canceled by the simultaneous detection of the two molecular orientations, but spin precession phase noise, which may arise from fluctuations in our magnetic bias field, is suppressed as well. Differential measurement is often employed for shot-noise-limited detection in precision metrology [35,36]. For high detection efficiency of molecules, action spectroscopy and ion detection provide high yields but were not amenable to simultaneous multiple state detection. Here we implement our new angle-resolved approach to achieve both high detection efficiency and differential noise suppression.

We can extract information about the molecular orientation if the intermediate and final states inherit the spatial orientation of the ground state, which maps the molecular orientation to momentum-space anisotropy of the photofragments [37–39]. Choosing the first REMPD photon such that it excites the molecules to a bound intermediate state

with  $\Omega_i=2$  maintains the molecular orientation for  $M_i\neq 0$ , where M is the projection of total angular momentum on the electric bias field. For the stretched states,  $m_F$  and M are mapped one to one. We use states with  $|M|=|\Omega|$  for the highest orientation contrast between states with opposite orientations (see Supplemental Material [25]). The second REMPD photon couples the intermediate state to dissociating states, breaking HfF<sup>+</sup> (ThF<sup>+</sup>) molecules into Hf<sup>+</sup> (Th<sup>+</sup>) ions and neutral F atoms. The spatial distributions of the charged photofragments are shown in Figs. 1 and 2(b). Experimentally, we obtain similar orientation contrasts for both HfF<sup>+</sup> and ThF<sup>+</sup>.

In a single experimental cycle, we can now simultaneously detect the populations of two states with the same  $m_F$ . In two adjacent cycles, we measure  $N_A$  and  $N_C$  followed by  $N_B$  and  $N_D$ , which we use to compute the spin polarizations according to Eqs. (1) and (2). Figure 3 shows experimental measurements of the phase evolution with up to 1.5 s interrogation time. The different magnetic moments of the upper and lower doublet states give rise to a beating between the spin precession fringes.

Near zero crossings when the spin precessions of the upper and lower doublets are in phase,  $\phi^u - \phi^l \approx 2n\pi$  (vertical dashed lines in Fig. 3), the phase difference is proportional to the spin polarization difference,  $\Delta \wp = \wp_u - \wp_l \sim \mathcal{C}(\phi^u - \phi^l)$ . The *e*EDM sensitivity is ultimately determined by the measurement uncertainty of this spin polarization difference ( $\Delta \wp$ ). If the spin precessions of the upper and lower doublets are evaluated independently, ignoring any correlations, we anticipate the total scatter to be  $(\sigma_{\wp_u}^2 + \sigma_{\wp_l}^2)^{1/2}$ , depicted by the blue diamonds in Fig. 3, which is 5 times higher than the QPN limit (dotted gray line) due to excess noise in ion production and detection. Expressed in terms of the spin polarizations [Eqs. (1) and (2)], the QPN limit is

$$\sigma_{\Delta\wp}^2 = \left(\frac{2}{N_t}\right) [(1+\wp_u)(1-\wp_u) + (1+\wp_l)(1-\wp_l)], \quad (3)$$

where  $N_t$  is the total number of detected dissociated ions in two adjacent cycles [33].  $N_t$  is 1400, 1000, 800, and 400, respectively, in the four time segments of Fig. 3. One standard method to reduce excess noise from initial molecular ion production is to normalize for the varying total HfF<sup>+</sup> number, which is measured simultaneously with the Hf<sup>+</sup> number [40]. With ion number normalization (orange circles in Fig. 3),  $(\sigma_{\wp_u}^2 + \sigma_{\wp_l}^2)^{1/2}$  still remains more than 2 times higher than the QPN limit due to excess noise from photodissociation.

Instead of monitoring and correcting for each source of technical noise, we use simultaneous detection of both molecular orientations to normalize all common-mode noise in each experimental cycle. One might expect that a complete normalization of the differential spin polarization  $\Delta \wp$  requires simultaneous detection of all four states involved. However, simultaneously detecting two states (A and C or B and D) is adequate to remove most of the excess noise. Fluctuations in ion production, state preparation, and detection all give rise to excess noise that is positively correlated between  $N_A$  and  $N_C$  or between  $N_B$ and  $N_D$ . Furthermore, when the two spin precession fringes are in phase, common-mode phase fluctuations contribute to positive correlations between the same pairs of states. When the spin precession fringes are simultaneously near a zero crossing, we obtain the highest phase sensitivity (vertical dashed lines in Fig. 3). Positive correlations between pairs of these populations cancel excess noise, resulting in a measurement of  $\Delta \wp$  at the QPN limit, represented by green squares in Fig. 3, both at early and late times ( $\sim 1.5$  s).

When the spin precessions of the upper and lower doublets are out of phase, the positive correlations from ion production, state preparation, and detection should still cancel to first order at the zero crossings (vertical dashdotted lines in Fig. 3). However, common-mode phase noise correlates negatively and cannot be eliminated simultaneously. This phase noise brings the scatter of  $\Delta \wp$  significantly higher than the QPN limit, as shown in Fig. 3 at  $t \approx 742$  ms. In our experiment, the early time and late time in-phase measurements are optimal for achieving a QPN-limited eEDM sensitivity, canceling both laser noise and magnetic field noise. The out-of-phase measurements allow us to characterize shot-to-shot phase noise such as arising from magnetic field fluctuations.

In summary, we demonstrate a spin precession measurement at the quantum projection noise limit with hundreds of ions and an interrogation time of 1.5 s. We leverage quantum-state-resolved photochemistry with molecular orientation doublets, a unique demonstration of controlled photofragmentation, to differentially isolate the coherence signal, taking full advantage of the increased count rate of the efficient state preparation scheme. Particularly, for the two quantum states detected by the angle-resolved photodissociation, the electron is polarized with opposite sign, such that the simultaneous differential measurement is sensitive only to the parity odd components such as the eEDM. Overall, this scheme has significantly increased the statistical sensitivity of our eEDM measurement and may also assist other studies of fundamental symmetries [9] or stereochemistry [41].

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