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Broadband velocity modulation spectroscopy of HfF⁺: towards a measurement of the electron electric dipole moment

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provides valuable checks to reduce systematic errors [11]. The use of molecular ions makes trapping straightforward, thus enabling measurements with long coherence times.

The general approach for the eEDM experiment with HfF⁺ or ThF⁺ will be to perform a Ramsey-type spectroscopy mea-

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> Prior to this work, very little spectroscopic information was available for either HfF⁺ or ThF⁺. The energies of the lowlying ${}^{3}\Delta_{1}$, ${}^{3}\Delta_{2}$, and ${}^{3}\Delta_{3}$ states were recently measured using pulsed-field ionization zero kinetic energy (PFI-ZEKE) spectroscopy [12, 13]. Additionally, several isoelectronic species including TiO [14, 15, 16], TiF⁺ and TiCl⁺ [17, 18] have been

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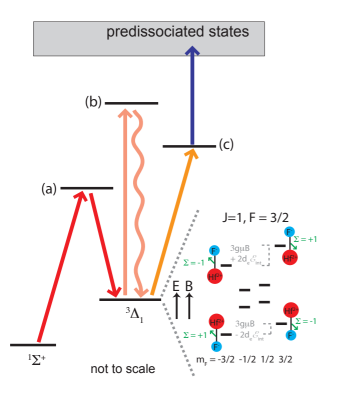


Figure 1: Energy levels of HfF⁺ relevant to the eEDM experiment. The eEDM experiment will utilize the J = 1 level of the ${}^{3}\Delta_{1}$ metastable state. Mixedspin excited states that couple well to both the ground ${}^{1}\Sigma^{+}$ and the ${}^{3}\Delta_{1}$ could be used for population transfer (state (a)). Readout from the ${}^{3}\Delta_{1}$ could be accomplished directly via laser-induced fluorescence (b) or resonantly-enhanced multiphoton photodissociation (c). The inset shows the structure of the J = 1, F = 3/2 manifold of the ${}^{3}\Delta_{1}$ level in applied electric and magnetic fields. The eEDM experiment will measure the difference in energy between the upper (and lower) pairs of $|m_F| = 3/2$ states. The applied magnetic field splits these states by $3g\mu B$, where *B* is the magnitude of the magnetic field, μ is the Bohr magneton, and *g* is the magnetic g-factor. In addition, the electron experiences an effective internal electric field (\mathcal{E}_{int}), which results in an additional splitting with magnitude $2d_e \mathcal{E}_{int}$, where d_e is the eEDM (aligned with the electron spin).

studied. Importantly though, there are no fully characterized optical transitions in either HfF⁺ or ThF⁺ at wavelengths below 1 μ m (above 10⁴ cm⁻¹) where laser-induced fluorescence detection is possible. Due to the large number of electrons involved, quantum calculations are challenging: current high-level calculations have errors of perhaps 1000 cm⁻¹ [9, 19].

The combined lack of experimental data and large theoretical uncertainties necessitated precision spectroscopy over a very broad spectral range. To achieve this, we recently developed frequency-comb velocity-modulation spectroscopy (comb-vms), which provides simultaneously broad bandwidth, high resolution, high sensitivity, and ion specificity [20]. We were able to measure 1000 cm⁻¹ of continuous spectra around 12000 cm⁻¹ using comb-vms and observed four bands of HfF⁺. This information, combined with PFI-ZEKE data and theoretical predictions, was then used to guide specifically targeted scans based on single-frequency (cw-laser) velocitymodulation spectroscopy, which resulted in 15 additional bands with origins ranging from 9950 to 146000 cm⁻¹. We obtained precise molecular constants for the ${}^{1}\Sigma^{+}$, ${}^{3}\Delta_{1}$, ${}^{3}\Pi_{0^{-}}$, ${}^{3}\Pi_{0^{+}}$, ${}^{1}\Pi_{1}$, ${}^{3}\Pi_{1}$, ${}^{3}\Sigma_{0^{+}}$, and ${}^{3}\Phi_{2}$ states, many of which will be of use for the eEDM experiment. We were also able to measure the electronic contributions to isotope splittings. In addition, we discuss the observed Λ -doublings, including the first measurement of the size of the small doubling in the ${}^{3}\Delta_{1}$. This value is important for determining the necessary applied electric field to fully polarize HfF⁺ as well as for estimating other systematic effects in the final eEDM experiment. Finally, the large amount of data enabled refinements in the *ab initio* calculations; we briefly discuss the modifications and compare the new theory with the measurements. This may help to improve the accuracy of other calculations involving high-Z atoms.

Outside of the eEDM experiment, precision spectroscopy of molecular ions is useful in a wide range of fields including fundamental physics, chemistry, and astrophysics. In astrochemistry for example, at least 22 ions (both positive and negative) ranging in complexity from H_3^+ to H_2COH^+ and C_8^- have been identified in interstellar and circumstellar gases [21]. Many of these ions are highly reactive and are believed to be intermediates in a variety of reactions; nonetheless, some rate constants and branching ratios are still not well known [22]. For example, the pathway for formation of a very simple molecular ion, CH⁺, remains elusive [23]. Searches for new species, including efforts to identify the origin of the diffuse interstellar bands [24, 25], benefit from laboratory measurements of optical transitions as well as measurements of rotational constants to aid microwave spectroscopy [26, 27, 28]. On the physical chemistry side, precision spectroscopy of H₃⁺ above the barrier to linearity (near 9913 cm⁻¹) provides rigorous tests for ab initio theory [29, 30, 31]. Many other carbocations are interesting for both their presence as intermediates in reactions such as combustion as well as the challenges they present to theory; for example, the spectrum of CH₅⁺, a highly nonclassical carbocation, still remains unassigned due to both its complexity and spectral interference from other contaminant species [32, 33, 34]. A combination of comb-vms with a recently developed ion-beam spectrometer capable of sensitive, sub-Doppler vms of rotationally cold species ([35]) could enable mass-selective spectroscopy of ions over a broad bandwidth with many potential applications.

2. Experiments

2.1. Comb-vms

Comb-vms combines cavity-enhanced direct frequency comb spectroscopy [37, 38, 39] with velocity-modulation spectroscopy [40, 41, 42, 43] for discrimination between ions and neutral species. This technique could also be readily extended to the detection of radicals by concentration modulation. By resolving every comb mode of a femtosecond Ti:sapphire laser (3 GHz mode spacing) simultaneously over a wide spectral bandwidth, we have high resolution and absolute frequency accuracy with over 1500 channels measured at once. Figure 2 provides an overview of the experimental setup, more details can be found in [20]. Briefly, we couple light from the comb into a ring cavity containing an ac discharge cell, which forms the HfF⁺ and modulates the ions' drift velocity. The resulting

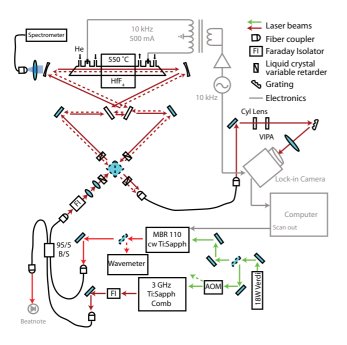


Figure 2: **Setup for comb-vms.** We use a Ti:sapphire comb with a repetition rate of 3 GHz, which is transferred via fiber to the optical cavity. The comb is coupled into one direction of a ring bowtie cavity, which contains an acdischarge cell with Brewster-angled end windows. The cavity finesse is about 100 and the length is matched to be an integer multiple (25) of the comb cavity. We use liquid crystal variable retarders and a polarizing beamsplitter to alternate the direction of propagation through the cavity. The ac-discharge is driven at 10 kHz with about 500 mA. We flow helium gas through the tube at a pressure of about 6 Torr. HfF⁺ ions are created by heating about 0.5 g of HfF₄ powder to 550 °C inside the discharge tube. The cavity transmitted light, which contains the modulated absorption signal, is analyzed using a two-dimensional imaging system and a lock-in camera. A cw Ti:sapphire laser is used to stabilize the comb and serves as a frequency reference in the spectra. More details can be found in [20]. A fiber-coupled grating spectrometer is used to monitor the discharge fluorescence.

modulated Doppler shift produces an absorption signal that is modulated at the discharge frequency. The comb light transmitted through the cavity is then spectrally dispersed using a highresolution (1 GHz), two-dimensional cross-dispersion system [39, 44], which resolves every comb mode. We then image this onto a lock-in camera [45, 46] to demodulate the absorption signal simultaneously on many comb teeth.

We produce HfF^+ by heating about 0.5 g of HfF_4 powder to about 550 °C in a 1 m long home-built discharge cell. Helium buffer gas is flowed through the discharge such that the total pressure is about 6 torr. By recording the emission of the discharge using a low-resolution grating spectrometer, we were able to reliably monitor molecule production. With each loading of the oven we were able to run for about 3 hours. A 2.5 m long bow-tie optical enhancement cavity consisting of two flat, 98% mirrors (input and output couplers) and two 100 cm radius-of-curvature, low-dispersion, 99.9% high reflectors surrounds the discharge cell. The reflectivity of the input and output couplers was chosen to match the losses from the Brewsterangled windows and thus provide efficient input coupling. The cavity length is actively stabilized to an integer multiple (25) of the frequency-comb laser cavity length, which ensures that each comb component is coupled to a cavity mode. Our feed-

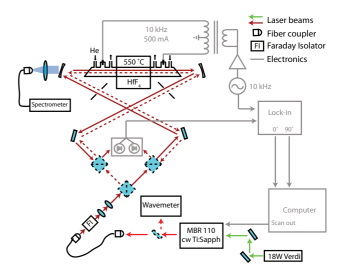


Figure 3: **Setup for cw-vms.** We use a cw Ti:sapphire laser that is tunable from about $9500 - 14800 \text{ cm}^{-1}$ (1050 - 675 nm). After fiber coupling between tables, the laser is split with a 50/50 non-polarising beamsplitter. Each beam goes through an additional 50/50 non-polarizing beamsplitter (to separate the return light) and is then sent in opposite direction is reflected by the beamsplitters and is differenced using an auto-balanced photodetector [36]. This difference signal is then sent to lock-in detectors to record signal in-phase and 90° out-of-phase with the HfF⁺ absorption. We placed irises in the beam paths to reduce the amount of discharge fluorescence observed on the auto-balanced detector; this helped both to reduce noise and primarily to reduce baseline drifts in the spectra.

back loop provides about 20 kHz of bandwidth by using a highspeed, low-range piezo-electric transducer (PzT) [47] and a second long-range PzT. We use liquid-crystal variable retarders and a polarizing beamsplittler to rapidly (50 ms) switch the direction of propagation through the discharge tube while maintaining the comb-cavity lock. This allows us to subtract out slowly varying noise due to drifts in camera pixel offsets and also to improve the rejection of neutral background absorption.

We record about 150 cm⁻¹ of spectrum spread over 1500 channels simultaneously. One comb tooth is locked to a stable cw Ti:sapphire laser, which provides an absolute frequency reference in our spectrum. For each measurement, we average and subtract images for each direction of propagation and also record the power per comb tooth by applying a calibrated amplitude modulation to the laser. In order to fully sample the spectrum, we interleave 30 measurements with the cw laser stepped over 3 GHz. This results in a spectrum that covers 150 cm⁻¹ sampled every 100 MHz with an absolute frequency accuracy of 30 MHz (set by a rubidium referenced wavemeter). When the wavemeter was not calibrated for a particular measurement, we set the uncertainty at 100 MHz. For strong bands with fully resolved isotope structure, absolute accuracy of the determination of the band origin is limited predominantly not by statistical errors but by the absolute knowledge of the cw Ti:sapphire laser frequency we use as a reference. For our purposes we were satisfied with the 30-100 MHz (0.001 - 0.003 cm^{-1}) absolute accuracy of the wavemeter. This could be readily improved if needed. For analysis purposes we then interpolate the spectrum onto a fixed 0.001 cm⁻¹ grid. This al-

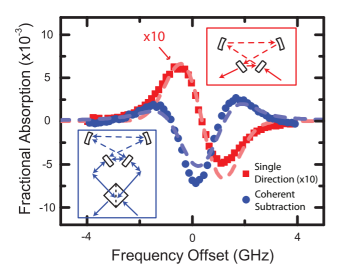


Figure 4: **Comparison of coherent subtraction and single direction measurements.** The signal to noise for coherent subtraction (using a 52/48 beam splitter) surpasses that of single direction measurements by a factor of 10 when the noise is primarily technical light intensity noise. Both measurements of a single N₂⁺ line were made with the cw laser. The dashed lines are a calculation of the expected lineshape (see text). The modification in the lineshape for the coherent subtraction is due to a stray differential phase shift of approximately $\pi/20$ between the two counter-propagating beams, which results in a sensitivity to both absorption and dispersion.

lows us to easily average or combine a collection of different scans. One full scan takes about 30 minutes and results in a single-pass fractional absorption sensitivity of 3×10^{-7} . Since one scan contains 45000 channels, this equates to a sensitivity of 4×10^{-8} Hz^{-1/2} (spectral element)^{-1/2}, which is the sensitivity that a single-frequency laser system would need, in addition to being able to scan 150 cm⁻¹ continuously, to match the performance of the comb-vms system. We recorded spectra over 1000 cm⁻¹ with both the oven on and off to check for contamination.

2.2. Single-frequency vms

The information obtained from the comb scans, combined with theory predictions, allowed us to scan other bands using single-frequency velocity-modulation spectroscopy as sketched in Figure 3. For these measurements we removed the enhancement cavity and counter-propogated beams from the cw Ti:sapphire laser through the discharge tube. We then subtract these two beams using an auto-balanced photodetector [36]. Due to noise from the discharge, we reached a sensitivity of about 5×10^{-8} Hz^{-1/2} with 1 mW on the detector; this noise seemed to be related to optical pickup from the discharge emission, which was reduced using irises, and to non-commonmode amplitude noise from acoustical pickup. Using this technique we were able to find 15 more bands ranging from 9950 to 14600 cm⁻¹ without having to scan the full spectral range. We can continuously scan about 0.5 cm^{-1} in 4 minutes, which is at least 30 times slower than the comb-vms system.

2.3. Coherent Subtraction

We have also investigated a novel technique for differential detection that relies on coherent interference between the two

counter-propogating laser beams instead of subtraction of photocurrents as discussed previously. This is accomplished by coherently splitting and recombining the two beams using one non-polarizing beamsplitter to form a Sagnac-type interferometer containing the discharge cell and cavity if desired (blue inset to Figure 4). The signal is then detected at the destructive interference, or "dark port", of the interferometer. In the regime where laser intensity noise dominates, the signal-to-noise using coherent subtraction increases as the splitting ratio approaches 50:50 between the two beams until detector readout noise or shot noise dominates. Physically, the fractional signal is increasing while the fractional noise level remains constant and dc power are both decreasing. We have tested coherent subtraction versus a single direction of propagation using a photodiode and single-channel lock-in detector with our cw laser and demonstrate a factor of 10 gain in the signal to noise (blue data in Figure 4), which is the predicted improvement with the 52/48 beamsplitter that was used.

Coherent subtraction, unlike subtraction of photocurrents, is sensitive to phase as well as amplitude of the light field; thus, the resulting lineshape is dependent on both absorption (approximately first-derivative shaped) and dispersion (approximately second-derivative) and can vary if there are differential phase shifts between the two directions. We performed a simple simulation, shown as dashed lines in Fig. 4, that reproduces the lineshape modification. These simulations were done by applying a cosine modulation in time to the center frequency for both the absorption (assumed Gaussian lineshape) and dispersion terms (obtained via Kramers-Kronig) and then selecting the appropriate frequency term from the Fourier cosine transformation. Repeating this at a variety of simulated laser frequencies generates the lineshape. The amplitude and linewidth of the single-direction simulation was scaled to the measured value and these values were used for the coherent-subtraction simulation. The constant phase offset (most likely due to the beamsplitter) added to the coherent subtraction simulation was chosen to fit the measured lineshape (about $\pi/20$ radians). In addition to the phase sensitivity, which could be useful for some applications, coherent subtraction has several other advantages. First, the dark port can be used to reduce the optical power seen by the detector thus avoiding saturation while still maintaining a shot-noise limited signal-to-noise ratio. Second, the subtraction is performed prior to the detector and is wavelength independent. This means that coherent subtraction can be used with comb-vms or other dispersive detection systems for increased signal-to-noise. We did not implement coherent subtraction with the comb-vms system here because we were close to camera-noise limited, thus the gains would not have been very significant.

3. Results and Discussion

Figure 5(a) shows the spectrum of HfF^+ acquired using the comb-vms system in blue (offset for clarity) as well as all data obtained with single-frequency vms in red. As illustrated in the zoomed region shown in the lower panel of Figure 5(b),

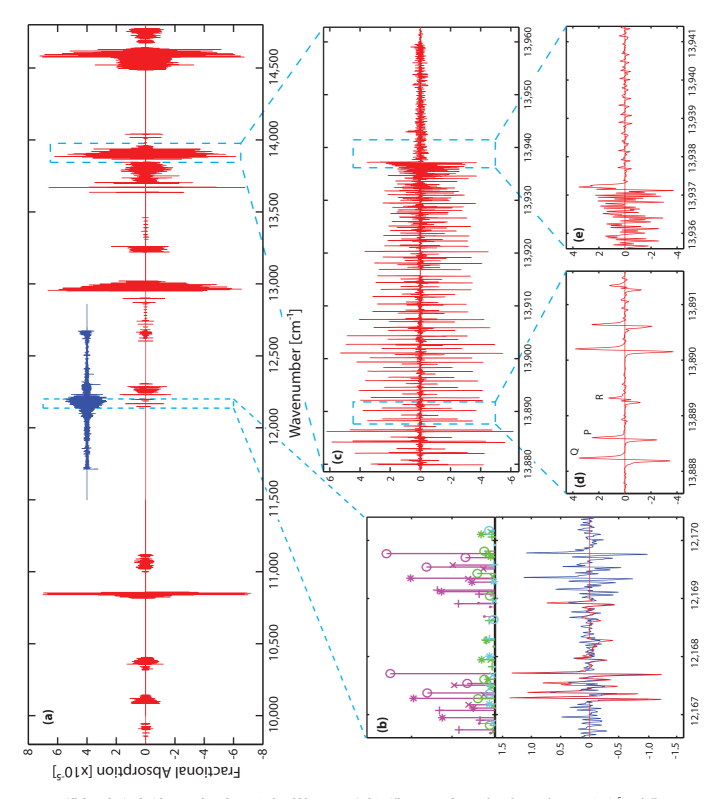


Figure 5: All data obtained with cw- and comb-vms (red and blue, respectively). All y-axes are fractional single-pass absorption (×10⁻⁵) and all x-axes are in cm⁻¹. In (a) we show all of the HfF⁺ spectra acquired to date. The comb-vms data is vertically offset for clarity. The lower panel of (b) shows a zoomed region of the comb data where some lines were also measured with the cw system. The upper panel of (b) plots predicted line positions from the fits; different colors correspond to different bands (magenta, ${}^{1}\Pi_{1} \leftarrow {}^{1}\Sigma^{+}$ (0,1); green, ${}^{3}\Pi_{1} \leftarrow {}^{1}\Sigma^{+}$ (3, 1); cyan, ${}^{3}\Pi_{1} \leftarrow {}^{1}\Sigma^{+}$ (2, 0)) while different symbols correspond to Hf isotopes: $-\circ$, 180 Hf; $-\times$, 178 Hf; -+, 177 Hf; -+, 176 Hf. Inset (c) shows the ${}^{3}\Phi_{2} \leftarrow {}^{3}\Delta_{1}$ (0, 0) band (on the left) and part of the ${}^{3}\Sigma_{0^{+}}^{-} \leftarrow {}^{1}\Sigma^{+}$ (1, 0) on the right. The region in (d) illustrates the cancellation of isotope shifts in $\Delta v = 0$ transitions; the sharpness of P- and Q-branch lines is due to a cancellation between the rotational, vibrational, and electronic contributions to the isotope shift. The splitting in the (higher-J'') R-branch lines that is not observed in the (lower-J'') P- and Q-branch lines is due to Λ -doubling in the ${}^{3}\Delta_{1}$ state. Inset (e) shows a prominent band-head from the ${}^{3}\Phi_{2} \leftarrow {}^{3}\Delta_{1}$ (0, 0) band as well as weaker lines from the ${}^{3}\Sigma_{0^{+}}^{-} \leftarrow {}^{1}\Sigma^{+}$ (1, 0) band.

the spectrum from about 12100-12300 cm⁻¹ is extremely congested due to the presence of many bands, each with five isotopes, and the high temperature of our oven, which results in observed J'' values up to about 70. The dynamic range of frequency-comb velocity-modulation spectroscopy is demonstrated by our ability to identify the overlapping ${}^{1}\Pi_{1} \leftarrow {}^{1}\Sigma^{+}$ (v' = 0, v'' = 1) and ${}^{3}\Pi_{1} \leftarrow {}^{1}\Sigma^{+}$ (3,1) bands despite the difference in linestrengths and an offset in band origin of only ~1 cm⁻¹. Since 150 cm⁻¹ sections are acquired simultaneously when using the comb, relative linestrengths within the region are not influenced by variability in oven and discharge conditions, which significantly helps to disentangle bands. The upper panel of Figure 5(b) shows predicted line positions from fits to three different bands (the fitting is discussed below), which illustrates our ability to resolve each isotope ($-\circ$, ¹⁸⁰Hf; $-\times$, 179 Hf; -*, 178 Hf; -+, 177 Hf; -·, 176 Hf) for multiple bands. Figure 5(b) also demonstrates excellent overlap between the comb and cw spectra.

Since we were fitting a variety of different transitions with various combinations of Hund's cases, we chose to fit each band with the general expression given in Eq. 1

$$\nu(\nu',\nu'',J',J'',s',s'') = \nu_0 + F_{\nu'}(J',s') - F_{\nu''}(J'',s'')$$
(1)

where the rotational energies, F_v are

$$F_{\nu}(J,s) = (B_{\nu} - s\frac{k_{\nu}}{2})J(J+1) - (D_{\nu} - s\frac{k_{D,\nu}}{2})J^{2}(J+1)^{2}.$$
 (2)

We include rotation (B), centrifugal distortion (D), Λ -doubling (k), and distortion of the Λ -doubling (k_D) for each vibrational level, v; s is an e/f-symmetry term where s = +1 for e-symmetry and -1 for f-symmetry. k and k_D are generic constants for the Λ -doubling since we have fit states with several different values of Λ and Ω . In Σ_0 states, only one symmetry term is chosen and k and k_D are both set to 0. v_0 includes both the electronic and vibrational energies:

$$\nu_0 = T'_e - T''_e + E_{vib},$$
(3)

$$E_{vib} \approx (\omega'_e(v'+1/2) - \omega_e x'_e(v'+1/2)^2) - (\omega''_e(v''+1/2) - \omega_e x''_e(v''+1/2)^2).$$
(4)

Here T_e is defined as the energy difference between the minimum of the potential curve from the minimum of the X¹ Σ^+ curve and ω_e and $\omega_e x_e$ are the usual vibrational constant and anharmonic correction, respectively.

Due to the congested nature of the spectra, fits were performed to a manually identified list of isolated lines (position only) for each band and for isotope in bands with well resolved isotope structure. Figure 6(b) shows the residuals from a representative fit; root-mean-square values for the residuals were typically < 0.005 cm⁻¹ with no apparent structure. For most $\Delta v = 0$ transitions the fits were performed to estimated isotope averaged line centers and then the rotational constants were shifted by reduced mass to the ¹⁸⁰Hf values. The ³ $\Phi_2 \leftarrow {}^{3}\Delta_1$ bands exhibited line doubling at high-J due to Λ -doubling (discussed in detail below), for these bands the center of the doublet was used for the fits. Similarly, the ${}^{1}\Pi_{1} \leftarrow {}^{3}\Delta_{2}$ (0, 0) band exhibited doubling of every transition (i.e., 6 branches) due to large Λ -doubling in the ${}^{1}\Pi_{1}$; for this band all six branches were fit simultaneously by assuming the splitting to each transition was $k' \times J'(J' + 1)$. A summary of the fitted constants for each band is given in Table 1.

Assignments of the observed bands were not particularly straightforward, partially because the presence or absence of low-J lines usually could not be determined due to the high temperature and complicating isotope structure. In fact, only for the ${}^{1}\Pi_{1} \leftarrow {}^{1}\Sigma^{+}$ (0, 0) band near 13000 cm⁻¹ were we able to directly establish that $\Omega' = 1$ and $\Omega'' = 0$ since we observed an R(0) line but no P(1) line. We also measured several bands with no Q-branch, which were assigned as ${}^{3}\Pi_{0^{+}} \leftarrow {}^{1}\Sigma^{+}$ and ${}^{3}\Sigma_{0^{+}}^{-} \leftarrow {}^{1}\Sigma^{+}$. This information, combined with data from [12], isotope shifts, and Λ -doubling structure provided the assignments in Table 1. Due to possible ambiguities, several bands are still unassigned as indicated.

From the fitted constants for the 16 assigned bands, we can determine constants for the $X^1\Sigma^+$, ${}^3\Delta_1$, ${}^3\Pi_{0^-}$, ${}^3\Pi_{0^+}$, ${}^1\Pi_1$, ${}^3\Pi_1$, ${}^{3}\Sigma_{0+}^{-}$, and ${}^{3}\Phi_{2}$ states as given in Table 2. T_{0} and $\Delta G_{1/2}$ are directly obtained from the measurements: T_0 is defined as the energy of the v = 0 level of an excited state relative to the $X^{1}\Sigma^{+}$ v = 0 level, and $\Delta G_{1/2}$ is the energy difference between the v = 0 and v = 1 levels. All other (equilibrium) constants are extracted from the data. The rotational constant for a given vibrational level is given to first order in v as $B_v = B_e - \alpha_e (v + \omega_e)$ 1/2), where B_e is the equilibrium rotational constant. Since we have measured at least two vibrational levels for each state, we can determine $\alpha_e \approx B_v - B_{v+1}$, and then use this to obtain B_e for each state. T_e , ω_e , and $\omega_e x_e$ are determined using Eqs. 3 and 4. For the ${}^{3}\Pi_{1}$ state, we were able to calculate ω_{e} and $\omega_e x_e$ directly; for all other states we assumed a Morse potential so that $\omega_e x_e = \alpha_e^2 \omega_e^2 / 36B_e^3 + \alpha_e \omega_e / 3B_e + B_e$ (i.e., the Pekeris relationship).

In addition, we have observed isotope shifts of the state origins, called δT_e in Table 2, that we attribute to Hf electronic isotope shifts (due to the finite nuclear charge radius) [48, 49, 50]. We define the isotope shift between ¹⁸⁰HfF⁺ and ¹⁷⁸HfF⁺ for a given spectral line as

$$\delta E = E^{180} - E^{178}$$

= $(1 - \frac{\mu_{180}}{\mu_{178}})E_{rot} + (1 - \sqrt{\frac{\mu_{180}}{\mu_{178}}})E_{vib} + \delta T'_e - \delta T''_e.$ (5)

Here μ_{180} and μ_{178} are the reduced masses, E_{rot} is the rotational energy in ¹⁸⁰HfF⁺ (i.e., offset from the band origin), and E_{vib} is the vibrational energy in ¹⁸⁰HfF⁺. For bands with well resolved isotope structure (i.e., $\Delta v \neq 0$), $\delta T'_e - \delta T''_e$ was determined either by fitting both v_0^{180} and v_0^{178} or by measuring δE near the origin and then subtracting the vibrational contribution. For $\Delta v = 0$ bands, we found the frequency where the lines were the sharpest, indicating that $\delta E \approx 0$ (as illustrated by the Q- and P-branch lines in Figure 5(d)); this location relative to the band origin is used for E_{rot} in Eq. 5, which then gave $\delta T'_e - \delta T''_e$. Values of $|\delta T'_e - \delta T''_e|$ were $\lesssim 0.1 \text{ cm}^{-1}$ for all transitions. To obtain the δT_e values given in Table 2, we

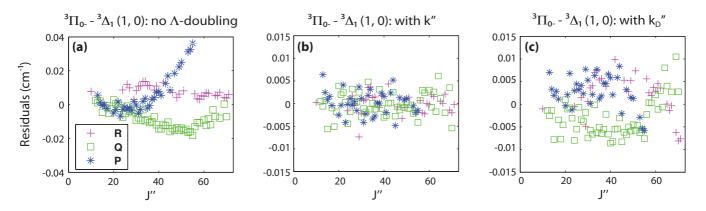


Figure 6: **Residuals for three fits to the** ${}^{3}\Pi_{0} \leftarrow {}^{3}\Delta_{1}$ (1, 0) **band.** The residuals are only plotted for assigned lines in each branch: R-branch (+), Q-branch(square), and P-branch (*). (a) Fit with no A-doubling terms included. Notice the large structure to the residuals. (b) Only k'' included, i.e., only A-doubling proportional to J^{2} in the lower state. No structure in the residuals; the scatter is primarily due to error in identification of the line center. (c) Only k''_{D} included, i.e., only A-doubling proportional to J^{4} in the lower state. There is some slight structure to the residuals, indicating that the primary contribution to the A-doubling goes as J^{2} .

set $\delta T_e''(X^1\Sigma^+) = 0$ and calculated relative shifts of the other states. The simplistic model of Eq. 5 neglects vibrationalband-specific perturbations or other more complicated leveldependent effects; however, results were consistent across different transitions, which supports the assignment of the effect to electronic states, not individual bands. Except for the $X^1\Sigma^+$ state the isotope shifts are all fairly similar with an average value of about -0.06 cm⁻¹(-1.7 GHz), which agrees well with the value for the Hf⁺ 5d²6s² \rightarrow 5d²6s6p of -1.8 GHz [50].

Due to our ability to see high-J lines, we were able to measure precise values for the Λ -doubling, parametrized by k and k_D in Eq. 2, in several different electronic levels. In Hund's case (a) k can be related to the more standard Λ -doubling parameters defined in [51] for Π states (*o*, *p*, and *q*) and to \tilde{o}_{Δ} as defined in [52] for ³ Δ states; k was not measurable in Φ states. The values of k and k_D were useful in assigning many transitions and also provide some insight into inter-state interactions. For the eEDM measurement, the most important Λ -doubling parameter is the splitting of the J = 1 levels in the ${}^{3}\Delta_{1}$, as this goes into determining several different potential systematic effects. In a case (a) ${}^{3}\Delta_{1}$ state, the Λ -doubling is expected to be given by $\pm \tilde{o}_{\Delta} \times J(J+1)$, where the upper and lower signs refer to the two e/f-symmetry levels [52]. We can make a rough estimate the value of \tilde{o}_{Δ} by a simple scaling from that measured in TiO or WC [53]:

$$\tilde{o}_{\Delta} \approx \frac{\zeta^2 (\text{Hf}^{2+}) B^2 (\text{Hf}F^+)}{\zeta^2 (\text{WC/TiO}) B^2 (\text{WC/TiO})} \times \tilde{o}_{\Delta} (\text{WC/TiO}).$$
(6)

This scaling relationship arises because the \tilde{o}_{Δ} term results from the application of two spin-orbit operators (which scale as the atomic spin-orbit, ζ) and two L-uncoupling operators, which scale as the rotational constant B [52, 53]; however, it neglects changes in state order and spacing. Using this, with ζ obtained from [54], we predict $\tilde{o}_{\Delta} \approx 50$ kHz. The Λ -doubling in the ${}^{3}\Delta_{1}$ can be observed at high-J in the ${}^{3}\Phi_{2} \leftarrow {}^{3}\Delta_{1}$ (0, 0) band by comparing neighboring P-, Q-, and R-branch lines, as shown in Figure 5(d). Since the R-branch lines have the highest J, there is noticeable doubling in these lines compared to the P- or Qbranch lines, which indicates that this doubling is not due to isotope splitting. In this band, the lines are not sufficiently well resolved to permit an accurate determination of the Λ -doubling constant, but by fitting the ${}^{3}\Pi_{0^{-}} \leftarrow {}^{3}\Delta_{1}$ (1, 0) transition, we were able to precisely determine the sign and size of the Λ -doubling. As shown in Figure 6, the best fit was obtained with k'' instead of k''_{D} , implying that the Λ -doubling goes mainly as J(J + 1) as expected (adding k''_{D} to the fit with k'' resulted in no improvement). This fit gives a value of $k'' = -1.23(6) \times 10^{-5} \text{ cm}^{-1}$, which gives $\tilde{o}_{\Delta} = k''/2 = 6.2(3) \times 10^{-6} \text{ cm}^{-1} = 185(9) \text{ kHz}$. The precision on \tilde{o}_{Δ} is only about an order of magnitude worse than that obtained for TiO using pure rotational spectroscopy [15], illustrating the ability to obtain high-resolution spectroscopic information from broad-bandwidth ro-vibronic spectra.

A-doubling in the ³Π₀ can arise from interactions with both ¹Σ⁺ and ³Σ⁺ states, which result in a splitting of the Ω = 0⁻ and 0⁺ levels that is independent of *J*. We measured this splitting to be 189.099(4) cm⁻¹ with the 0⁺ (e-symmetry) higher in energy than 0⁻ (f). To the extent that the X¹Σ⁺ state is a pure (*sσ*)² configuration, we can assume that this splitting is from a higherlying ³Σ⁺ state with an (*sσdσ*) configuration. The spin-orbit interaction between the ³Σ⁺₀ (f) and the ³Π₀- can be estimated by

$$\frac{(\frac{1}{\sqrt{2}}\langle\sigma_1(\beta)\pi(\beta)|\frac{1}{2}\hat{a}l^+s^-|\sigma_1(\beta)\sigma_2(\alpha)\rangle)^2}{\Delta E} \approx \frac{\frac{3}{4}\zeta^2(5d)}{\Delta E}.$$
 (7)

We have used $\langle \pi(\beta)|l^+s^-|\sigma_2(\alpha)\rangle \approx \sqrt{6}$ in the pure-precession model [51, 55]. Using the measured splitting and the atomic spin-orbit coefficient, we estimate $\Delta E \approx 11000 \text{ cm}^{-1}$, which gives an approximate location of the ${}^{3}\Sigma_{0^{-}}^{+}$ state to be 21200 cm⁻¹. This is close to the predicted position of 21694 cm⁻¹ from the new calculations (see Section 4). In a Hund's case (a) basis, this splitting corresponds to the more familiar 2(o + p + q) [51].

The cause of the Λ -doubling in the ${}^{3}\Pi_{1}$ state is primarily interaction with the ${}^{3}\Sigma_{0^{-}}^{+}$ state through the ${}^{3}\Pi_{0}$. This type of interaction is characterized by the parameter *p* in [51]. With esymmetry above f in ${}^{3}\Pi_{0}$ and ${}^{3}\Pi_{0}$ located below ${}^{3}\Pi_{1}$, we would expect e above f in ${}^{3}\Pi_{1}$, which corresponds to a negative value of k as observed. The magnitude of k is expected to be about $4Bp/A_{\Pi}$, where A_{Π} is the spin-orbit parameter for the ³ Π manifold and

$$p \approx C \langle {}^{3}\Pi_{1} | BL^{+} | {}^{3}\Sigma_{0^{-}}^{+} \rangle \langle {}^{3}\Sigma_{0^{-}}^{+} | \hat{a}L^{-}S^{+} | {}^{3}\Pi_{0} \rangle / (E_{\Pi} - E_{\Sigma})$$
(8)

$$\approx CB\zeta(5d)\langle \pi^+|l^+|\sigma_2\rangle^2/(E_{\Pi}-E_{\Sigma}).$$
(9)

Here, C is a numerical factor dependent on the spin [51]. Again using the pure-precession model, we estimate $k \approx 8 \times 10^{-4}$ cm⁻¹, in reasonable agreement with experiment.

We attribute most of the Λ -doubling in the ${}^{1}\Pi_{1}$ to interaction with the nearly-degenerate nominal $^{3}\Sigma_{0^{+}}^{-}$ state. It is reasonable that the ${}^{1}\Pi$ and ${}^{3}\Sigma^{-}$ are not pure $(s\sigma d\pi)$ and $(d\delta)^{2}$ configurations, which would lead to increased spin-orbit interaction between ${}^{1}\Pi_{1}$ and ${}^{3}\Sigma_{0^{+}}^{-}$. Estimating the magnitude of the Λ -doubling in this case is difficult due to uncertainty in the purity of the configurations. Qualitatively though, we would expect the effect of a ${}^{3}\Sigma^{-}$ state above the ${}^{1}\Pi_{1}$ state would be to push e-symmetry below f as observed. In addition we observed A-doubling that was strongly vibrationallevel dependent $(3.69 \times 10^{-4} \text{ cm}^{-1} \text{ for the } v = 0 \text{ compared}$ with 2.68×10^{-4} cm⁻¹ for the v = 1), which is indicative of nearly-degenerate interacting states. We also measured a Λ doubling term proportional to J^4 (denoted k_D). This term can be explained by substituting $\Delta E \rightarrow \Delta E + (B_{\Pi} - B_{\Sigma})J(J+1)$ in the Λ -doubling denominator; the first two terms in the Taylor expansion give the J(J + 1) and the $J^2(J + 1)^2$ components. From this we can estimate that $k_D \approx 3k(B_{\Pi} - B_{\Sigma})/(E_{\Pi} - E_{\Sigma})$, which gives $k_D \approx 2 \times 10^{-8} \text{ cm}^{-1}$.

4. Theory

In earlier theoretical study of HfF⁺ [19], the correlation calculations of the spectroscopic constants were performed in two different ways: in the first series 10 electrons from 5d, 6s shells of Hf and 2s, 2p of F were correlated while in the second series 5s, 5p outer core electrons of Hf and 1s of F were also correlated. The rest of the 1s - 4d inner-core electrons of Hf were excluded from the explicit treatment using the generalized relativistic effective core potential (GRECP) method³ [56]. While it was shown that the inclusion of the additional electrons in the 20-electrons case contributed significantly to the excitation energies, such 20-electron calculations were performed only for one internuclear Hf-F distance in a small space of manyelectron basis functions due to computer limitations. This calculation was then applied for other distances as a "core correction". In the present work a new 20-electron relativistic correlation calculation was carried out in order to consider electronic states that were not investigated in [19] as well as to overcome the bottlenecks of the previous studies of HfF⁺. Eventually 23 electronic states (with excitation energies up to 22000 cm^{-1}) were considered in the present study instead of ten states in [19].

Some modifications were made in the computational procedure to achieve better accuracy. We used a direct multireference configuration-interaction approach accounting for spin-orbit effects (SODCI) [57, 58] as a method to treat both correlation and relativistic effects simultaneously. Onecomponent basis functions (orbitals) are required in this method to construct many-electron spin-adapted functions (SAF's). Instead of the orbitals obtained within the complete active space self-consistent field method (used in [19]), the natural orbitals of a one-electron density matrix averaged over the density matrices of the states of interest were used. The latter density matrices were calculated within the scalar-relativistic coupled-clusters method with single and double cluster amplitudes (CCSD) using the CFOUR code [59]. From numerical investigation [60] it is known that the use of natural orbitals can provide faster convergence in terms of the number of SAF's required to account for a given part of correlation energy.

In the SODCI calculation the same atomic basis sets for Hf and F were used as in our previous study [19], i.e., a generallycontracted basis set for Hf consisting of 6 s-, 5 p-, 5 d-, 3 fand 1 g- type contracted functions (denoted as $\{6,5,5,3,1\}$) and an ANO-I $\{4,3,2,1\}$ basis set for F [61]. However, a correction on the extension of the basis set was applied additionally in the present work. To evaluate the correction, two scalar-relativistic calculations were performed using the coupled clusters method with single, double and perturbative triple cluster amplitudes: (i) in the same basis set that was used at the SODCI stage, and (ii) with an extended basis set for Hf (produced by uncontracting d- and adding g-, h-, i- type basis functions) and an ANO-L $\{7,7,4,3\}$ [61] basis for F.

To compute potential curves the described calculations were performed at 14 points in the range of 3.0-4.2 a.u. As shown in Table 3, the new calculations are in remarkable agreement with all of the experimental results. These improvements should enable more accurate calculations in other species with heavy atoms, where relativistic effects are extremely important.

5. Outlook and Conclusions

As shown in Figure 1 and discussed in the introduction, several different optical transitions may be necessary for the eEDM experiment. We have measured suitable candidates for all of these: (a) can be accomplished using the ${}^{3}\Pi_{0^{+}}$ intermediate, (b) could use the ${}^{3}\Phi_{2}$ level, and (c) could use one of the ${}^{3}\Pi_{0}$ levels or the ${}^{3}\Phi_{2}$ level. Another important value for the eEDM measurements is the size of the Λ -doubling in the ${}^{3}\Delta_{1} J = 1$ level; it is the opposite parity levels in this state that are mixed in an electric field to polarize the molecule. The necessary electric field for full polarization, \mathcal{E}_{pol} , is approximately where the Stark energy is larger than the energy difference between the two parity states: thus $\mathcal{E}_{pol} \approx \omega_{ef}/2\pi d_{mf}$ [7], where ω_{ef} is the Λ -doubling splitting (in angular frequency) and d_{mf} is the molecular-frame electric dipole moment of the molecule (4.3 Debye for HfF⁺ [10]). We can use our measurement of \tilde{o}_{Δ} to estimate that $\omega_{ef} = 2\pi \times 4\tilde{o}_{\Delta} \approx 2\pi \times 740$ kHz, which is at least an order magnitude larger than previously predicted [10]. Nonetheless, this means that \mathcal{E}_{pol} is still under 1 V/cm, which is

³60-electron core GRECP for Hf is available at www.qchem.pnpi.spb.ru

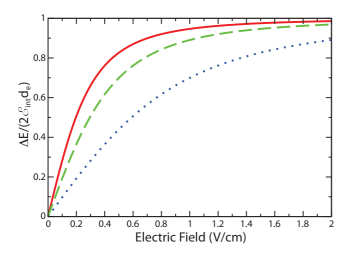


Figure 7: The eEDM induced Stark splitting (ΔE) for J = 1 hyperfine levels of the ${}^{3}\Delta_{1}$ state of HfF⁺: between $F = 3/2 m_{F} = \pm 3/2$ levels (solid line), between $F = 3/2 m_{F} = \pm 1/2$ levels (doted line), and between $F = 1/2 m_{F} = \pm 1/2$ (dashed line).

important for the eEDM experiment as larger fields tend to lead to more issues with systematic errors. As can be seen in Figure 7, numerical results for ${}^{3}\Delta_{1} J = 1$ hyperfine levels confirm that at 1 V/cm the splitting in the $|m_{F}| = 3/2$ levels due to eEDM related Stark shift is almost saturated at $2d_{e}\mathcal{E}_{int}$. This saturation occurs as the molecule becomes fully polarized. These results were obtained as described in [62].

We can also use the improved theory calculations to better estimate the radiative lifetime, another important consideration for the eEDM experiment. Using the measured energy separation and calculated dipole moments, we estimate the lifetime of ${}^{3}\Delta_{1} v = 0$ to be about 2 s [19]. This is encouraging, as it sets the ultimate limit on the coherence time achievable in the experiment. The high precision that we obtain for rotational constants should enable microwave spectroscopy with minimal searching, which will then be able to resolve the fine and hyperfine structure in the ${}^{3}\Delta_{1}$ level. These measurements are important for estimating systematics and for checking the accuracy of the calculated effective electric field that the electron experiences (\mathcal{E}_{int}) . We predict that the ${}^{1}\Sigma^{+}(v=0) J' = 1 \leftarrow J'' = 0$ transition in ¹⁸⁰HfF⁺ will be at 18.290(2) GHz and the ${}^{3}\Delta_{1}(v = 0)$ $J' = 2 \leftarrow J'' = 1$ transition doublet will be at centered at 35.869(2) GHz. To get these frequencies, we used the best measurements of B_e and ω_e from Table 3 to calculate $D_e = 4B_e^3/\omega_e^2$, then used the measured B_0 and calculated D_e to determine the microwave frequencies.

We are currently working on improving the comb-vms system and applying it towards characterization of ThF^+ , which has several advantages over HfF^+ for the eEDM experiment. However for this species, the *ab initio* calculations are even more challenging, leading to larger theoretical uncertainties. The first spectroscopy of ThF^+ using PFI-ZEKE plus some LIF (unassigned) has recently been published [13], but, as was the case with HfF^+ , the higher excited states remain uncharacterized. By using highly non-linear fiber to broaden the comb spectrum [63] and a high repetition rate Ti:sapphire amplifier [64], we hope to cover over 2000 cm⁻¹ with comb-vms. This ability to rapidly cover thousands of cm⁻¹ with high sensitivity and high resolution will be a powerful new tool for the study of ions and radicals for many applications. In addition, applications to other many other spectral regions are possible by using different comb sources combined with non-linear optics for covering the near-IR [65, 66] or mid-IR [67, 68, 69, 70, 71]. These sources plus readout systems using different cameras or optical up-conversion [72] create the possibility of ion spectroscopy anywhere from the visible to the mid-IR.

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Table 1: **Fitted constants for observed transitions in** ¹⁸⁰**HfF⁺ in cm**⁻¹**.** Quoted errors are at 95% confidence level, statistical only. For bands recorded with the wavemeter uncalibrated we include an additional 0.006 cm⁻¹uncertainty in v_0 . Internal consistencies in the data suggest that for the $\Delta v = 0$ bands, which typically suffer from partially resolved isotope structure, systematic errors in the fit quantities may exceed the quoted statistical uncertainties in some cases by a factor of two or three. Values without uncertainties were fixed in the fits. All constants are given for the vibrational levels involved in the transition and are not equilibrium values. The A-doubling terms, k'', k', and k'_D are generic terms proportional to J(J + 1) in the ground and excited states and $J^2(J + 1)^2$ in the excited state respectively (see text for more details).

	ν_0	$B^{\prime\prime}$	B'	$D^{\prime\prime}$	D'	$k^{\prime\prime}$	k'	k'_D
	Ū			$[10^{-7}]$	$[10^{-7}]$	$[10^{-4}]$	$[10^{-4}]$	$[10^{-9}]$
$^{1}\Pi_{1} \leftarrow ^{1}\Sigma^{+} (0, 1)$	12217.369(2)	0.30335(2)	0.28115(3)	1.88(8)	1.81(8)	_	3.69(2)	9.7(7)
$^{1}\Pi_{1} \leftarrow {}^{1}\Sigma^{+} (1, 2)$	12136.012(3)	0.30180(5)	0.27973(5)	1.80(14)	1.74(14)	_	2.68(4)	7.4(1.3)
$^{1}\Pi_{1} \leftarrow ^{1}\Sigma^{+} (0, 0)$	13002.189(12)+	0.30474(20)	0.28104(20)	1(2)	1(2)	_	3.55(4)	0
${}^{3}\Pi_{1} \leftarrow {}^{1}\Sigma^{+} (2, 0)$	12304.400(3)	0.30481(5)	0.28096(5)	1.78(12)	1.78(12)	_	-3.82(1)	0
${}^{3}\Pi_{1} \leftarrow {}^{1}\Sigma^{+} (3, 1)^{\circ}$	12216.901(10)	0.30335*	0.27958(20)	1.8(3)	1.8(3)	_	-3.8(2)	0
${}^{3}\Pi_{1} \leftarrow {}^{1}\Sigma^{+} (0, 1)$	10109.877(6)	0.30333(6)	0.28382(6)	1.9(5)	1.9(5)	_	-3.77(5)	0
${}^{3}\Pi_{0^{+}} \leftarrow {}^{1}\Sigma^{+} (0, 0)$	10401.723(13)+	0.30500(10)	0.28437(11)	2.3(1.2)	2.2(1.2)	_	_	_
${}^{3}\Pi_{0^{+}} \leftarrow {}^{1}\Sigma^{+} (1, 0)$	11114.653(7)	0.30482(4)	0.28281(4)	1.71(14)	1.72(13)	_	_	_
${}^{3}\Sigma_{0^{+}}^{-} \leftarrow {}^{1}\Sigma^{+} (0, 0)$	13254.302(7)	0.30478(10)	0.28967(9)	1.7(5)	1.9(5)	_	_	_
${}^{3}\Sigma_{0^{+}}^{-} \leftarrow {}^{1}\Sigma^{+} (1, 0)$	13953.799(6)	0.30483(5)	0.28808(4)	1.78(12)	1.98(13)	_	_	_
${}^{3}\Phi_{2} \leftarrow {}^{3}\Delta_{1} (0, 0)$	13933.340(12)+	0.29900(10)	0.27720(10)	2.2(3)	2.1(3)	†	-0.01(3)	_
${}^{3}\Phi_{2} \leftarrow {}^{3}\Delta_{1} (1, 0)$	14613.782(7)	0.29888(5)	0.27590(5)	1.71(15)	1.59(15)	Ť	0	_
${}^{3}\Phi_{2} \leftarrow {}^{3}\Delta_{1} (2, 1)$	14534.839(7)	0.29745(8)	0.27488(8)	1.7(4)	1.6(4)	†	0	_
$^{3}\Pi_{0^{+}} \leftarrow ^{3}\Delta_{1} (1,0)$	10137.723(7)	0.29887(5)	0.28277(5)	1.70(20)	1.66(20)	-0.14(2)	_	_
${}^{3}\Pi_{0^{-}} \leftarrow {}^{3}\Delta_{1} (1,0)$	9948.624(6)	0.29891(2)	0.28272(2)	1.85(5)	1.81(4)	-0.123(6)	_	_
$^{1}\Pi_{1} \leftarrow ^{3}\Delta_{2} (0, 0)$	10852.757(13)+	0.29915(9)	0.28102(9)	1.7(4)	1.7(4)	0	3.59(3) [‡]	_
? $\Delta v = 0$	13729.918(17)+	0.29860(60)	0.28680(60)	3(3)	3(3)	0	-0.2(3)	_
? $\Delta v = 0$	13822.270(14)+	0.29980(10)	0.28860(10)	1.9(2)	2.2(2)	0	0	_
? $\Delta v = 0$	14636.162(13)+	0.29620(30)	0.27600(30)	1.7(1.5)	1.8(1.5)	0	0	_

 $^{\circ 3}\Pi_1 \leftarrow {}^{1}\Sigma^+(3,1)$ values assigned using a manual fit to multiple isotopes due to challenges of determining the line centers of the weaker lines in the dense spectrum.

* Value fixed to the fitted value from the ${}^{1}\Pi_{1} \leftarrow {}^{1}\Sigma^{+}$ (0, 1) transition.

⁺ Individual isotopes were not fully resolved, thus the fit was done to averaged line positions. We report isotope corrected values with the error due to the uncertainty in the isotope shifts.

 $^{\dagger 3}\Phi_2 \leftarrow {}^{3}\Delta_1$ transitions exhibited line-doubling at high J", but the doublets were not resolved well enough to accurately determine k".

^{*} Each transition was a doublet with a splitting given by $k' \times J'(J' + 1)$ due to Λ -doubling in ${}^{1}\Pi_{1}$.

Table 2: **Derived constants for observed states in** ¹⁸⁰**HfF**⁺ **in cm**⁻¹. Values for T₀ are given to the $\nu = 0$ levels, with the X¹ Σ^+ $\nu = 0$ level set to 0. Values for T_e are to the minimum of the potential curves again with X¹ Σ^+ set to 0. $\Delta G_{1/2}$ is only given for states where we had a direct measurement. All equilibrium values assume a Morse potential unless otherwise noted. Quoted uncertainties are 95% and are statistical, i.e., they do not account for any deviation from the Morse potential. $\delta T_e = T_e^{180} - T_e^{-178}$ is an electronic isotope shift relative to ¹ Σ^+ for each state as discussed in the text.

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	T_0	$\Delta G_{1/2}$	T_e	B_e	ω_e	$\omega_e x_e$	$\alpha_e~[10^{-3}]$	δT_e
$^{1}\Sigma^{+}$	0	784.820(12)	0	0.30558(3)	790.76(11)	2.97(5)	1.50(2)	0
$^{3}\Delta_{1}$	976.930(10)	_	991.83(74)	0.29963(5)	760.9(1.5)	2.78(21)	1.45(8)	-0.049(12)
$^{3}\Delta_{2}$	2149.432(18)	_	—	-	_	_	_	-0.055(12)
$^{1}\Pi_{1}$	13002.189(12)	_	13046.04(21)	0.28186(5)	702.9(4)	2.70(15)	1.42(6)	-0.071(6)
${}^{3}\Pi_{1}$	10894.697(14)	_	10933.77(6)+	0.28454(8)	712.382(21)+	$2.51(6)^+$	1.43(4)	-0.054(7)
$^{3}\Pi_{0^{-}}$	—	_	10248.34(36)*	0.28517(17)*	_	_	_	-0.033(16)
$^{3}\Pi_{0^{+}}$	10401.723(13)	712.930(15)	10437.44(36)	0.28518(13)	719.5(7)	3.3(3)	1.63(11)	-0.047(7)
${}^{3}\Sigma_{0^{+}}^{-}$	13254.302(7)	699.497(9)	13296.93(28)	0.29046(14)	705.5(5)	3.0(3)	1.59(10)	-0.106(7)
${}^{3}\Phi_{2}$	14910.270(16)	680.442(14)	14963.15(33)	0.27778(14)	684.5(7)	2.05(31)	1.16(14)	-0.044(16)

⁺ Calculated without assuming a Morse potential.

* Assuming ω_e , $\omega_e x_e$, and α_e are the same as for the ${}^3\Pi_{0+}$.

State	Constant	This Work	[12]	[19]	New theory
$^{1}\Sigma^{+}$	T_e	0	0	0	0
	B_e	0.30558(3)	0.304(10)	0.3082	0.309
	ω_e	790.76(11)	791.2(1.0)	751	792
$^{3}\Delta_{1}$	T_e	991.83(74)	993(2)	1599	1229
	B_e	0.29963(5)	0.301(10)	0.2994	0.301
	ω_e	760.9(1.5)	761.3(2.0)	718	754
$^{3}\Delta_{2}$	T_e	2149.432(16) [†]	2151.7(20) [†]	2807	2394
	B_e	0.29915(9)*	0.300(10)	0.2997	0.302
	ω_e	_	762.3(2.0)	719	766
$^{3}\Delta_{3}$	T_e	_	3951(2)	4324	3995
	B_e	_	0.308(10)	0.3004	0.301
	ω_e	-	761.5(2.0)+	721	757
$^{1}\Delta_{2}$	T_e	_	_	11519	10610
	B_e	_	_	0.2981	0.298
	ω_e	-	-	696	747
³ Π ₀ -	T_e	10248.34(36)	_	11910	10400
	B_e	0.28517(17)	_	0.2848	0.286
	ω_e	-	-	689	716
$^{3}\Pi_{0^{+}}$	T_e	10437.44(36)	_	12196	10658
	B_e	0.28518(13)	_	0.2854	0.285
	ω_e	719.5(7)	-	699	724
${}^{3}\Pi_{1}$	T_e	10933.77(6)	_	12686	11058
	B_e	0.28454(8)	-	0.2835	0.285
	ω_e	712.382(17)	_	687	712
${}^{3}\Pi_{2}$	T_e	_	_	14438	13452
	B_e	-	-	0.2848	0.287
	ω_e	-	-	703	745
$^{1}\Pi_{1}$	T_e	13046.3(3)	_	14784	13493
	B_e	0.28186(5)	_	0.2805	0.283
	ω_e	702.9(4)	-	679	699
${}^{3}\Sigma_{0^{+}}^{-}$	T_e	13296.93(28)	_	_	13773
-	\mathbf{B}_{e}	0.29046(14)	_	_	0.292
	ω_e	705.5(5)	_	_	716

Table 3: Summary of states. Comparison of derived molecular constants from this work and from the experiments of Barker et al. [12] with the old theoretical calculations of Petrov et al. [19] and the improved theory discussed here. The theoretical values of B_e from [19] were computed from the equilibrium bond length. Experimental uncertainties are converted to 95% (2σ).

 $^{+}\Delta G_{1/2}.$

 $^{\dagger} T_0.$ $^{*} B_0.$

	Tabl	e 3: Summary of sta	ates. (con	tinued)	
State	Constant	This Work	[12]	[19]	New theory
$^{3}\Sigma_{1}^{-}$	T_e	_	_	_	14757
1	B_e	_	_	_	0.292
	ω_e	-	-	-	711
${}^{3}\Phi_{2}$	T_e	14963.15(33)	_	_	15284
	B_e	0.27778(14)	_	_	0.278
	ω_e	684.5(7)	_	-	671
${}^{3}\Phi_{3}$	T_e	_	_	_	17457
	B_e	_	_	_	0.277
	ω_e	_	_	-	658
$^{1}\Gamma_{4}$	T_e	_	_	_	18312
·	\mathbf{B}_{e}	_	_	_	0.289
	ω_e	-	_	_	641
${}^{3}\Pi_{0^{-}}$	T_e	_	_	_	19167
0	\mathbf{B}_{e}	_	_	_	0.276
	ω_e	_	_	_	691
$^{3}\Pi_{1}$	T_e	_	_	_	19332
	\mathbf{B}_{e}	_	_	_	0.279
	ω_e	-	_	_	698
${}^{3}\Pi_{0^{+}}$	T_e	_	_	_	20074
110+	\mathbf{B}_{e}	_	_	_	0.280
	ω_e	_	_	_	748
1					
$^{1}\Sigma^{+}$	T_e	_	_	_	20330
	B_e	_	_	_	0.288
	ω_e	-	-	_	610
$^{3}\Pi_{2}$	T_e	-	_	_	20338
	B_e	_	_	_	0.277
	ω_e	_	_	-	665
$^{3}\Phi_{4}$	T_e	_	_	_	20769
	B_e	_	_	_	0.283
	ω_e	-	_	_	740
${}^{3}\Sigma_{1}^{+}$	T_e	_	_	_	21415
	B_e	_	_	_	0.276
	ω_e	-	_	_	665
${}^{3}\Sigma_{0^{-}}^{+}$	T_e	21200°	_	_	21694
0	\mathbf{B}_{e}	_	_	_	0.277
	ω_e	_	_	_	658

Table 3: Summary of states. (continued)

° Estimated from Λ -doubling in the ${}^{3}\Pi_{0}$ (see text).