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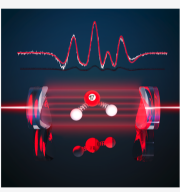


Hyperfine-Resolved Near-Infrared Spectra of H₂¹⁷O

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ABSTRACT: Huge efforts have recently been taken in the derivation of accurate compilations of rovibrational energies of water, one of the most important reference systems in spectroscopy. Such precision is desirable for all water isotopologues, although their investigation is challenged by hyperfine effects in their spectra. Frequency-comb locked noise-immune cavity-enhanced optical-heterodyne molecular spectroscopy (NICE-OHMS) allows for achieving high sensitivity, resolution, and accuracy. This technique has been employed to resolve the subtle hyperfine splittings of rovibrational transitions of H₂¹⁷O in the near-infrared region. Simulation and interpretation of the H₂¹⁷O saturation spectra have been supported by coupled-cluster calculations performed with large basis sets and accounting for high-level corrections. Experimental ¹⁷O hyperfine parameters are found in excellent agreement with the corresponding computed values. The need of including small hyperfine effects in the analysis of H₂¹⁷O spectra has been demonstrated together with the ability of the computational strategy employed for providing quantitative predictions of the corresponding parameters.



■ INTRODUCTION

Hyperfine parameters constitute an important source of information on physicochemical molecular properties related to electron densities and electronic structure. The nuclear quadrupole coupling is the strongest hyperfine interaction in closed-shell molecules. From the associated constants, information on intramolecular interactions and on the ionic or π character of the bonds involving the quadrupolar nucleus can be retrieved. Nuclear spin-rotation interaction and their corresponding constants provide, instead, insight into the paramagnetic part of the nuclear magnetic shielding constants, and this was the motivation for investigating the hyperfine structure (HFS) of the rotational spectrum of water isotopologues containing an ¹⁷O.^{1,2} Nuclear quadrupole coupling and spin-rotation interaction give rise to splittings in the rotational and rovibrational transitions (the HFS mentioned above), thus providing an experimental approach to the abovementioned molecular properties.

However, the hyperfine effects in water tend to be small in view of its closed-shell nature in the electronic ground state, thus requiring a very high resolving power to reveal subtle splittings in its spectrum. Indeed, the HFS of the $H_2^{17}O$ rotational spectrum was resolved only in a few investigations. Besides the work reported in ref 1, there is only the study by De Lucia and Helminger from the 1970s.³ Both works concerned the vibrational ground state and were performed via pure rotational spectroscopy in the microwave domain. However, to the best of our knowledge, the HFS of the ¹⁷O-containing species has never been resolved in vibrational excited states. Water is a crucial reference spectroscopic

system, and a comprehensive characterization is also essential for this isotopic species. As in the main isotopologue, the rovibrational states are separated into two subsets: the ortho and para nuclear-spin isomers. Ortho-para conversion represents an open issue, with the energy separation of the ortho and para states requiring to be known precisely. In the case of $H_2^{17}O$, a precise determination of the rovibrational level structure cannot avoid the full characterization of its HFS because the hyperfine splittings are of the same order of magnitude as the desired accuracy.

In the past decade, the development of intracavity-based optical techniques allowed high resolution and high sensitivity to be achieved also in vibrational spectroscopy. Different groups have applied such approaches to the investigation of vibrational excitation of $H_2^{17}O.^{4-7}$ A large number of spectral lines were accurately measured in a wide spectral region, thereby leading to the revised and updated W2020 database of rovibrational transitions and energy levels of ¹⁷O-containing water.⁸ However, all of these studies were Doppler-limited and no hyperfine effects could be resolved. Recently, the intracavity techniques were further developed and combined with high accuracy calibration, thus allowing the exploitation of the Lamb-dip effect in rovibrational spectra.^{9–12} In only one study,

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such Lamb-dip vibrational spectroscopy was applied to $H_2^{17}O$, although the linewidth, due to collisional and time-of-flight broadening, was still too large to resolve the HFS.¹³

In the present study, we extend our previous work on the saturation spectroscopy of water¹⁴ to the ¹⁷O-containing species, thereby using the advanced capabilities of a frequency-comb locked noise-immune cavity-enhanced optical-heterodyne molecular spectroscopy (NICE-OHMS) setup to resolve, for the first time, the HFS of rovibrational transitions of H₂¹⁷O. NICE-OHMS is a precision-spectroscopy technique that brings a new perspective in the field; having the sensitivity to observe saturation spectra at very low pressures and powers, it allows for the resolution of hyperfine structures in vibrational overtones, while at the same time providing an absolute frequency scale at kHz accuracy. The experimental determination of the hyperfine parameters for the vibrational state involved is supported and guided by state-of-the-art *ab initio* computations.

THEORY AND COMPUTATIONS

Each rovibrational energy level of $H_2^{17}O$, including the HFS caused by the ¹⁷O nuclear spin, is uniquely labeled by seven quantum numbers. They can be classified as vibrational quantum numbers, $(v_1 \ v_2 \ v_3)$, representing the excitation of the symmetric stretching, bending, and antisymmetric stretching modes, respectively, and the rotational quantum

Table 1. Equilibrium Values, Vibrational Corrections, and Vibrationally Corrected Values of the Oxygen Quadrupole-Coupling Tensor (NQCT) (MHz)

	Xaa	$\chi_{ m bb}$	Xcc
Equilibrium valu	es ^a		
$CCSD(T)^{b}$	-8.809	-1.066	+9.876
ΔT^{c}	-0.012	-0.001	+0.012
ΔQ^d	+0.011	-0.001	-0.010
ΔREL^{e}	-0.009	-0.014	+0.023
sum ^f	-8.819	-1.082	+9.901
Vibrational corre	ctions (DVR-QAK) ^g	
(000)	+0.007	-0.223	+0.216
(040)	-0.067	+0.177	-0.111
(120)	+0.023	-0.353	+0.329
(021)	+0.067	-0.415	+0.348
(200)	-0.005	-0.698	+0.704
(101)	+0.023	-0.735	+0.712
(002)	+0.040	-0.760	+0.719
Vibrationally cor	rected values ^h		
(000)	-8.812	-1.305	+10.117
(040)	-8.886	-0.905	+9.790
(120)	-8.796	-1.435	+10.230
(021)	-8.752	-1.497	+10.249
(200)	-8.824	-1.780	+10.605
(101)	-8.796	-1.817	+10.613
(002)	-8.779	-1.842	+10.620

^{*a*}Computed at the semiexperimental geometry. ^{*b*}Computed employing the aug-cc-pCV6Z basis set. ^CDifference of the NQCT at CCSDT/aug-cc-pCVQZ and CCSD(T)/aug-cc-pCVQZ levels. ^{*d*}Difference of the NQCT at CCSDTQ/aug-cc-pCVTZ and CCSDT/aug-cc-pCVTZ levels. ^{*e*}Computed via DPT2 at the CCSD(T)/aug-cc-pCV6Z level. ^{*f*}Sum of CCSD(T), ΔT , ΔQ , and ΔREL . ^{*g*}Difference of vibrationally averaged and equilibrium value at the CCSD(T)/aug-cc-pCVQZ level. ^{*h*}Equilibrium values augmented by the corresponding vibrational corrections.

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Table 2. Equilibrium Values, Vibrational Corrections, and Vibrationally Corrected Values of the Oxygen Spin-Rotation Tensor (SRT) (kHz)

	C_{aa}	C_{bb}	C_{cc}
Equilibrium values ^a			
$CCSD(T)^{b}$	-22.25	-25.20	-17.48
ΔT^{c}	-0.10	-0.11	-0.06
ΔQ^d	+0.05	+0.08	+0.03
sum ^e	-22.31	-25.22	-17.50
Vibrational correction	s (DVR-QAK) ^f		
(000)	-6.35	-2.78	-1.02
(040)	-25.86	-2.38	+3.13
(120)	-22.93	-5.53	-0.10
(021)	-20.05	-6.82	-1.24
(200)	-18.58	-7.79	-3.59
(101)	-16.80	-8.73	-4.40
(002)	-14.45	-9.67	-4.96
Vibrationally corrected	d values ^g		
(000)	-28.66	-28.01	-18.52
(040)	-48.16	-27.61	-14.37
(120)	-45.23	-30.75	-17.60
(021)	-42.35	-32.04	-18.74
(200)	-40.89	-33.01	-21.09
(101)	-39.10	-33.96	-21.91
(002)	-36.75	-34.89	-22.46

^{*a*}Computed at the semiexperimental geometry. ^{*b*}Computed employing the aug-cc-pCV6Z basis set. ^{*c*}Difference of the SRT at CCSDT/ aug-cc-pCVTZ and CCSD(T)/aug-cc-pCVTZ levels. ^{*d*}Difference of the SRT at CCSDTQ/aug-cc-pCVDZ and CCSDT/aug-cc-pCVDZ levels. ^{*e*}Sum of CCSD(T), ΔT , and ΔQ . ^{*f*}Difference of vibrationally averaged and equilibrium value at the CCSD(T)/aug-cc-pCVQZ level. ^{*g*}Equilibrium values augmented by the corresponding vibrational corrections.

numbers. These are the total angular momentum for the endover-end rotation *I*, the pseudoquantum numbers K_{1} and K_{c} used in the designation of levels in asymmetric-top rotors, and *F* the total angular momentum accounting for the coupling of *I* with the ¹⁷O nuclear spin (I = 5/2). The latter coupling, which is caused by the interactions mentioned above, splits each rotational level into different but closely spaced sublevels. For ortho-H₂¹⁷O, additional splittings are due to the hydrogen spins (only spin-rotation interaction), with the involved interaction being too small to produce measurable effects in the present experiment. The splittings of the energy levels generate the already mentioned HFS of the spectrum. While the HFS due to ¹⁷O exhibits several components typically separated by a few hundred kHz, that due to the hydrogen nuclei is expected only to give rise to a small broadening of the Lamb-dips, as already observed in a previous study on $H_2^{16}O.^1$

To simulate the HFS of rovibrational transitions, the hyperfine parameters of each vibrational state involved in the transition are required. In particular, relying on the additivity approximation, vibrationally averaged parameters are evaluated as the sum of their equilibrium value and the corresponding vibrational correction, with the former term computed by means of a composite scheme. The hyperfine parameters under consideration are the oxygen (¹⁷O) quadrupole-coupling constants, χ_{ii} , and the spin-rotation constants of both oxygen and hydrogen, C_{ii} (*i* and *j* denote the principal inertia axes).

The nuclear quadrupole coupling and nuclear spin-rotation constants have been computed adopting a similar protocol as

Table 3. Equilibrium Values, Vibrational Corrections, and Vibrationally Corrected Values of the Hydrogen Spin-Rotation Tensor (SRT) (kHz)

	C_{aa}	C_{bb}	C_{cc}	C_{ab}^{a}	$C_{ba}{}^{a}$
Equilibrium values ^b					
$CCSD(T)^{c}$	-35.45	-31.78	-33.98	-47.81	-21.01
ΔT^d	+0.00	+0.00	-0.01	-0.03	-0.02
ΔQ^{e}	-0.01	+0.00	-0.01	-0.03	+0.00
sum ^f	-35.47	-31.78	-33.99	-47.87	-21.03
Vibrational corrections (DVR-QAK) ^g					
(000)	+1.24	+0.62	+1.51	-1.29	-0.09
(040)	+6.22	+0.43	+3.83	-26.56	+3.13
(120)	+4.24	+0.94	+3.08	-9.84	+1.30
(021)	+4.87	+1.65	+4.54	-9.07	+0.61
(200)	+2.69	+1.53	+2.65	-0.24	-0.32
(101)	+3.01	+2.08	+3.85	-0.05	-0.93
(002)	+3.10	+2.60	+4.88	+0.22	-1.51
Vibrationally corrected values ^h					
(000)	-34.23	-31.16	-32.48	-46.16	-21.13
(040)	-29.24	-31.36	-30.16	-74.44	-17.91
(120)	-31.23	-30.85	-30.91	-57.71	-19.73
(021)	-30.60	-30.13	-29.45	-56.94	-20.43
(200)	-32.78	-30.25	-31.34	-48.11	-21.35
(101)	-32.46	-29.70	-30.14	-47.93	-21.97
(002)	-32.37	-29.19	-29.11	-47.66	-22.55

^aPlease note that for the other hydrogen, the C_{ab} and C_{ba} terms are inverted in sign. ^bComputed at the semiexperimental geometry. ^cComputed employing the aug-cc-pCV6Z basis set. ^dDifference of the SRT at CCSDT/aug-cc-pCVTZ and CCSD(T)/aug-cc-pCVTZ levels. ^eDifference of the SRT at CCSDTQ/aug-cc-pCVDZ and CCSDT/aug-cc-pCVDZ levels. ^fSum of CCSD(T), ΔT , and ΔQ . ^gDifference of vibrationally averaged and equilibrium value at the CCSD(T)/aug-cc-pCVQZ level. ^hEquilibrium values augmented by the corresponding vibrational corrections.

in refs 1, 2. The so-called equilibrium values were evaluated at the semiexperimental equilibrium geometry^{*a*} (r(OH) = 0.9575)Å, \angle (HOH) = 104.51°)¹⁵ using the coupled-cluster singles and doubles approach augmented by a perturbative treatment of triple excitations $(CCSD(T))^{16-19}$ in conjunction with the aug-cc-pCV6 $Z^{20,21}$ basis set. The CCSD(T) values have been augmented by higher-level corrections. These latter were evaluated as the difference between coupled-cluster singles, doubles, and triples $(CCSDT)^{22-24}$ and CCSD(T) employing the aug-cc-pCVXZ (X = T, Q)²⁵⁻²⁷ basis sets (ΔT term), as well as the difference between coupled-cluster singles, doubles, triples, and quadruples $(CCSDTQ)^{22,23}$ and CCSDT employing the aug-cc-pCVXZ (X = D, T)^{25-27} basis sets (ΔQ term). In the calculations of spin-rotation tensors, perturbationdependent basis functions were used to ensure fast basis-set convergence, as described in refs 28, 29. Scalar relativistic corrections to the nuclear quadrupole-coupling tensors (Δ REL term) have been evaluated using second-order direct perturbation theory $(DPT2)^{30}$ at the CCSD(T)/aug-ccpCV6Z level.

As in previous studies focusing on the (000) vibrational ground state,^{1,2} the DVR-QAK³¹ scheme has been employed to compute the vibrational corrections to equilibrium values. In this approach, the treatment of vibrational effects is based on the variation principle and on the use of the so-called Watson Hamiltonian,³² given in terms of rectilinear dimensionless normal coordinates, and fully accounts for Coriolis interactions and anharmonic effects in the potential. Converged vibration-

ally averaged property values were determined by evaluating the corresponding expectation values over the vibrational wave function of the corresponding vibrational state, using a multidimensional Gauss—Hermite quadrature. For the actual computations, 11–15 quadrature points per each mode have been used to evaluate the matrix elements over the anharmonic part of the potential and 11–15 harmonic-oscillator basis functions per each mode have been used as a product basis. The vibrational corrections were calculated as the difference between the vibrationally averaged and the equilibrium values, both at the CCSD(T) level in conjunction with the aug-ccpCVQZ^{25–27} basis set. A value of -25.58(22) mb was used for the ¹⁷O nuclear quadrupole moment.³³

All computations were carried out with all electrons included in the correlation treatment and using the CFOUR program package;^{34,35} for some calculations, the parallel version of CFOUR³⁶ has been employed. All CCSDT and CCSDTQ results were obtained with the string-based many-body code MRCC^{37,38} interfaced to CFOUR.

The equilibrium values and their vibrational correction terms are presented in Table 1 for the $\chi_{ii}(^{17}\text{O})$'s and in Tables 2 and 3 for the C_{ij} 's of oxygen and hydrogen, respectively. These parameters have been computed for the (000) vibrational ground state as well as for the (040), (120), (021), (021), (200), (101), and (002) vibrational states. All these levels lie around 7000 cm⁻¹, which is the frequency region explored in the present experiment.

EXPERIMENT

The measurements were performed in the near-infrared region employing the NICE-OHMS apparatus developed for

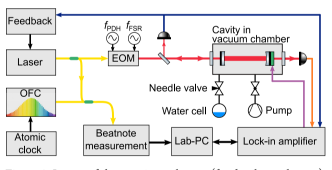


Figure 1. Layout of the experimental setup (for details see the text).

saturation spectroscopy and already employed for HD³⁹ and the main water isotopologue.¹⁴ In the present setup, shown schematically in Figure 1, a high-finesse (~150 000) cavity has been employed together with an infrared diode laser operating at 1.4 μ m. This laser is modulated at 305 MHz, equivalent to the free spectral range (FSR) of the cavity, for generating the side-band signals, and at 20 MHz for the cavity-lock via a Pound-Drever-Hall (PDH) stabilization scheme. In addition to the first layer of modulation at 305 MHz, the second layer of modulation is applied by dithering one of the cavity mirrors at a low frequency of 415 Hz. The doubly modulated spectroscopic signal is demodulated by a powerful lock-in system (Zurich Instruments; HF2LI), where the 1st and 3rd harmonics (1f and 3f signals, respectively) of the dither modulation are extracted. A stabilized optical frequency-comb (OFC), referenced to a Cs atomic clock, is employed to stabilize the infrared laser and the optical cavity, and to provide a frequency scale accurate to the 1 kHz level.

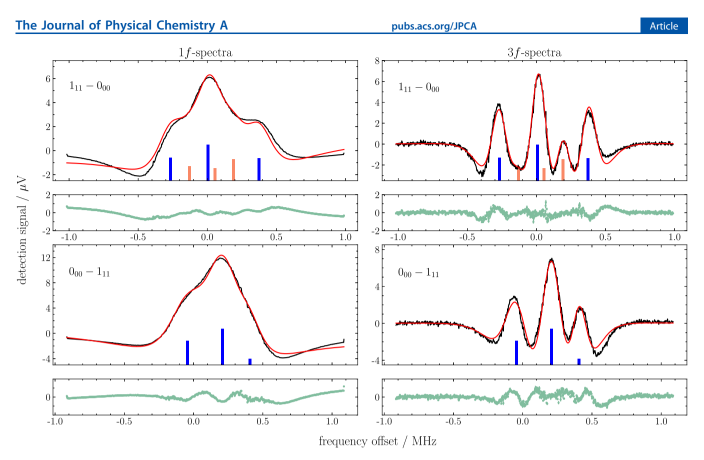


Figure 2. Modeling results for the *If* and *3f* spectra recorded for the *JKaKc* = $1_{11} \leftarrow 0_{00}$ (top panels) and the *JKaKc* = $0_{00} \leftarrow 1_{11}$ (bottom panels) transitions of the (200) \leftarrow (000) overtone. The black and red lines plot the experimental data and the "best-fit" model, respectively. The stick spectra denote the positions and relative intensities of the hyperfine components (in blue) and crossover resonances (orange). The fit residuals (light green) are shown in the boxes below each panel.

The intracavity high-power laser enabled the saturation of several rovibrational transitions of $H_2^{17}O$ and thus the exploitation of the Lamb-dip effect with extremely narrow dip profiles. Indeed, thanks to the highly reflective mirrors, the intracavity power can be increased up to 150 W. However, the intracavity power is matched to the oscillator strength to operate in the weakly saturating regime, thus avoiding significant power broadening. This results in intracavity powers below 1 W. The resulting linewidth of the resolved hyperfine components is mainly limited by the transit time of the molecules across the laser beam, overall yielding a width of ~400 kHz (full width at half-maximum).

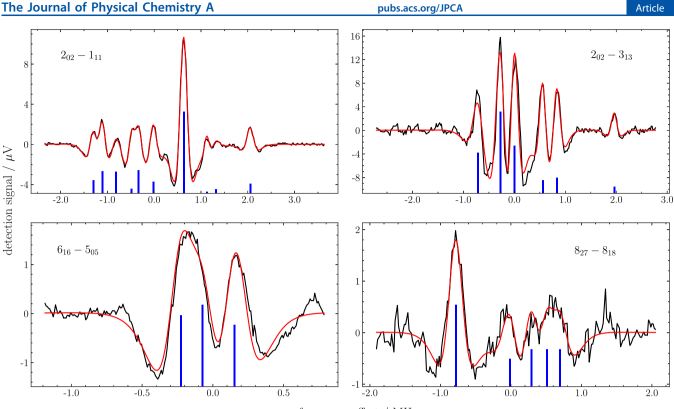
For the spectral recordings, an enriched $H_2^{17}O$ sample (Cambridge Isotopes; 20% $H_2^{17}O$ isotopic purity) has been used. The measurements have been carried out under steady gas flow conditions, at pressures in the range of 0.1–0.3 Pa, measured by a capacitance pressure gauge.

Among the vibrational states investigated computationally, the (200) state has been chosen in view of the intensity of its transitions. A number of saturation spectra were recorded for the (200) \leftarrow (000) rovibrational band at 7200 cm⁻¹. In Figure 2, two spectra are shown: the *JKaKc* = 1₁₁ \leftarrow 0₀₀ *R*-transition from the ground state para-level and the *JKaKc* = 0₀₀ \leftarrow 1₁₁ Ptransition probing the lowest para-level in the (200) vibrationally excited state. Both *If* and *3f* recordings are depicted, thereby demonstrating the superior resolution obtainable with the *3f* demodulation scheme. Figure 3 shows four additional Lamb-dip spectra recorded in the high-resolution *3f* mode and involving transitions for both *para-* and *ortho*-H₂¹⁷O.

RESULTS AND DISCUSSION

Initially, we simulated the spectra using the experimental hyperfine constants previously determined for the ground state,¹ and the *ab initio* values computed in this work for the (200) upper state. Simulations of the Lamb-dip spectra based on these "first guess" values were already able to well reproduce the experimental recordings. Nevertheless, as found in the Lamb-dip investigation of the ground-state rotational spectrum,¹ additional features due to crossover resonances were occasionally observed. These are also referred to as "ghost transitions" and are a well-known effect in Lambdip saturation spectra.⁴⁰ They appear in the case of transitions with a common state (either upper or lower) and partially overlapping Doppler profiles. Here, we adopt a phenomenological treatment of such features, whose transition frequencies are given by the arithmetic mean of the frequencies of the two "interacting" transitions.

Once the simulations had been refined by including the observed crossovers, a custom Python3 routine was used to accurately model the spectral line profiles and to retrieve the experimental transition frequencies. This code is an adaptation of the tool used in ref 41 to model the astrophysical spectra of the amidogen radical isotopologues. To reproduce the recorded spectra, the *1f* and *3f* profile functions are computed for each hyperfine component (also including crossovers), they are summed up, and the resulting profile is optimized versus the experimental spectra in a least-squares fashion. The *1f* and *3f* profile functions are defined as derivatives of the typical dispersive NICE-OHMS signals⁴²



frequency offset / MHz

Figure 3. Experimental recordings and modeling results for the 3*f* spectra of the $JKaKc = 2_{02} \leftarrow 1_{11}$ (top-left), $JKaKc = 2_{02} \leftarrow 3_{13}$ (top-right), $JKaKc = 6_{16} \leftarrow 5_{05}$ (bottom-left), and $JKaKc = 8_{27} \leftarrow 8_{18}$ (bottom-right) transitions of the (200) \leftarrow (000) overtone. The two top spectra pertain to the para-species, while the two bottom spectra to the ortho-species. The black and red lines plot the experimental data and the "best-fit" model, respectively. The stick spectra (blue bars) show the positions and relative intensities of the hyperfine components.

Table 4. ¹⁷O Hyperfine Parameters for H₂¹⁷O in the (200) Excited Vibrational Level

parameter ^a	unit	exp. ^b	theory ^c
Xaa	MHz	-8.86(9)	-8.824
$\chi_{ m bb}$	MHz	-1.86(14)	-1.780
C_{aa}	kHz	-48(16)	-40.89
$C_{\rm bb}$	kHz	-38(9)	-33.01
C_{cc}	kHz	-19(4)	-21.09
no. data		30	
rms	kHz	10.8	

^{*a*}Being the nuclear quadrupole-coupling tensor traceless, only two χ 's are given. ^{*b*}Values in parentheses denote 3σ uncertainties in the unit of the last quoted digit. ^{*c*}Results retrieved from Tables 1 and 2.

Table 5. Absolute Transitions Frequencies of Rovibrational Transitions in the (200) \leftarrow (000) Overtone Band of H₂¹⁷O

line	frequency (MHz)
$1_{11} \leftarrow 0_{00}$	216 698 971.729(20)
$0_{00} \leftarrow 1_{11}$	214 541 043.881(22)
$2_{02} \leftarrow 1_{11}$	216 570 396.811(10)
$2_{02} \leftarrow 3_{13}$	213 423 435.893(10)
$6_{16} \leftarrow 5_{05}$	218 793 162.770(34)
$8_{27} \leftarrow 8_{18}$	218 532 715.457(17)

$$P(\nu)_{1f} = \frac{4 A[\gamma^2 - 4(\nu - \nu_0)^2]}{[\gamma^2 + 4(\nu - \nu_0)^2]^2}$$

$$P(\nu)_{3f} = \frac{3072 \, A \gamma^2 (\nu - \nu_0)^2}{\left[\gamma^2 + 4(\nu - \nu_0)^2\right]^4} - \frac{96 \, A}{\left[\gamma^2 + 4(\nu - \nu_0)^2\right]^2}$$

The adjustable parameters are the line position ν_{0} , the line intensity *A*, and the width of each component γ . If necessary, a cubic polynomial is also fitted to reproduce the spectral background. The final agreement between the modeled and experimental spectra is fairly good, as shown in Figures 2 and 3. In these plots, the optimized positions and intensities of the hyperfine components and crossovers are depicted as stick spectra.

From the line profile analysis of the spectra recorded, the "best-fit" frequencies were retrieved as explained above; then, they were assigned to the corresponding transitions in terms of quantum numbers and analyzed using the SPFIT routine of the CALPGM suite⁴³ and adopting the standard Watson Hamiltonian in the I^r representation.⁴⁴ For blended transitions, the retrieved frequencies were assigned to the intensityweighted average of the involved components. The analysis of 30 hyperfine frequencies led to the very first determination of the nuclear quadrupole-coupling and spin-rotation constants of ¹⁷O for vibrationally excited $H_2^{17}O$. Their values, together with their 3σ uncertainties, are collected in Table 4. From this table, a very good agreement between experiment and theory is noted, with the computed values lying well within the experimental uncertainty. As already pointed out in ref 1, the level of theory employed is able to provide a quantitative prediction of hyperfine parameters. The same accuracy is therefore expected for all computational data reported in Tables 1-3. Since the inclusion of the spin-rotation constants

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of hydrogen has a negligible effect on the simulation of the rovibrational spectra of $ortho-H_2^{17}O$, such interaction was not considered in the final analysis.

As an additional finding, the absolute frequencies of the unperturbed rovibrational transitions recorded in the present study can be retrieved from the analysis. They can be obtained by subtracting the energy contribution due to the hyperfine interactions from the measured hyperfine components. The results of this analysis are presented in Table 5. The absolute accuracy of these transitions is improved by about 2 orders of magnitude with respect to the line frequencies listed in the W2020 database⁸ for $H_2^{17}O$.

CONCLUSIONS

The present study reports on the first observation of hyperfineresolved rovibrational transitions of the (200) \leftarrow (000) overtone band of H₂¹⁷O, recorded in the saturation regime by means of the NICE-OHMS technique. Six transitions, four pertaining to the para-species and two to the ortho-species, were investigated and the nuclear quadrupole-coupling and spin-rotation constants of oxygen derived for the (200) vibrational level. A very good agreement, within 3σ , is found with the corresponding computations. As pointed out in refs 1, 2 and confirmed in the present study, quantitative predictions of the hyperfine parameters can be provided, which, however, cannot be obtained routinely. For the first time, the variational DVR-QAK scheme has been extended to compute the vibrational corrections for excited states. Finally, the unperturbed frequencies of the observed rovibrational transitions of H₂¹⁷O were determined at an unprecedented accuracy of 20 kHz, corresponding to a relative uncertainty of 10^{-11} . These figures supersede by far the quality of the data available in the W2020 database for $H_2^{17}O$.

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Notes

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ADDITIONAL NOTE

"This is obtained by a least-squares procedure involving experimental ground-state rotational constants of various isotopologues computationally corrected for vibrational contributions (from anharmonic force-field calculations).

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