



The spectrum of radioactive water vapor: the H₂¹⁹O radio-isotopologue

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Received: 26 April 2024 / Accepted: 24 July 2024
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Abstract

The absorption spectrum of H₂¹⁹O, a radioactive isotopologue of the water molecule, is predicted using variational nuclear motion calculated based on a high precision potential energy function and ab initio dipole moment surface. Vibrational - rotational energy levels and wave functions, line centers and Einstein coefficients for dipole transitions are calculated. Predicted transition wavenumbers are improved by extrapolating known empirical energy levels of the stable H₂¹⁶O, H₂¹⁷O and H₂¹⁸O isotopologues to H₂¹⁹O. A line list for possible atmospheric application is presented which includes air line broadening coefficients. The calculations span a wide spectral range covering infrared and visible wavelengths, and are appropriate for temperatures up to 1000 K. Windows suitable for observing absorption by H₂¹⁹O are identified and comparisons made with the infrared spectra of water vapor in natural abundance, H₂¹⁵O and H₂¹⁴O.

Keywords Infrared spectrum · Energy levels · Water vapor · Absorption lines

Introduction

Water is one of the most important substances for humanity and its environment. Therefore, there are a large number of scientific works studying water and, in particular, the water molecule, its properties, reactivity, spectra, and isotopic composition.

Measurements and calculations of the absorption, emission, or scattering spectra of radioactive water isotopologues

(radio-isotopologues) are of interest for many reasons. Radio-isotopologues of atmospheric gases, including H₂O, are formed in the atmosphere when the Earth is irradiated by cosmic rays. The interaction of atmospheric molecules with cosmic rays leads, for example, to the accumulation in the atmosphere of triterated water, HT¹⁶O, and the carbon dioxide radio-isotopologue ¹⁴CO₂ [11]. It has been shown that various radio-isotopes of beryllium, carbon, nitrogen, and oxygen can be produced by lightning discharges [3, 29], which also leads to the appearance of radio-isotopologues such as ¹⁶O¹⁵O and H₂¹⁵O in the atmosphere. We [51] previously showed that radio-isotopologue H₂¹⁵O is apparently present in trace amounts in the atmosphere during thunderstorms. Radioactivity occurring during a thunderstorm front is discussed by Bordonskiy [4].

Oxygen-19 is also found in the cooling circuits of nuclear reactors [8, 55], Nemoto et al. [28] discovered that radio-isotopologues of water are formed when a target containing deuterium is irradiated with femtosecond laser pulses, and the mechanism of production of these isotopologues is similar to their production in lightning discharges. Other radio-isotopologues of water, H₂¹⁹O and H₂²⁰O, are produced in the primary cooling circuits of nuclear reactors [43]. The contribution of these isotopic modifications to radioactivity fields at nuclear power plants should be taken into account. Inside water-cooled fusion reactors, water is

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irradiated by neutrons when it passes the reactor core. This causes higher doses around the primary coolant: the main observed nuclides are ^{16}N and ^{19}O [19]. Similarly, these radioisotopes are also found in pressurized water reactors (PWRs) [43, 55] which has raised safety concerns [12, 17]. So far, only nuclear spectra (decay of the ^{19}O atom) have been recorded [8]. It is clear that ^{19}O lives long enough to form water molecules; however, the spectrum of H_2^{19}O has yet to be investigated.

In the 1970 abstract review [1] there is mention of work on ^{19}O and H_2^{19}O (Investigation of the recoil chemistry of ^{19}O and of radiolysis of water by reactor radiation. Blaser W. 1970. 79p. Dep. SFSTI (in German)). This abstract discussed the energies of ^{19}O and some molecules such as H_2^{19}O , $\text{H}^{16}\text{O}^{19}\text{OH}$, $^{16}\text{O}^{19}\text{OH}$ and $^{16}\text{O}^{19}\text{O}$. A number of monographs discussing ^{19}O species, including water, are available [2, 15, 20, 21, 23, 25, 30, 34, 36–39, 44]. In addition, there are a large number of monographs that consider the 6 main isotopes of oxygen from H_2^{14}O to H_2^{19}O .

To address the various scientific and technical problems involved with the presence of radioactive water, detailed information is needed on the spectra of water radio-isotopologues. Since measuring the spectra of short-lived radioactive substances is costly and difficult due to both safety issues and the short lifetime of radio-isotope, the most convenient way to determine their spectrum is to calculate it using a high-precision method benchmarked for the stable isotopologues against the extensive available experimental data. We have previously computed predicted spectra for H_2^{15}O [51, 53] and H_2^{14}O [52]. The present work is aimed at studying the vibrational and rotational states of the short-lived radio-isotopologue of the water molecule H_2^{19}O , and the associated spectrum at infrared (IR) and visible wavelengths. The calculations are carried out by combining two different techniques. The first of them uses a high-precision potential energy surface [6], which includes non-adiabatic and

relativistic corrections, and the variational nuclear-motion code DVR3D [40]. The second step employs a simple quadratic approximation based on using the known empirical energy levels of three stable isotopologues H_2^{16}O , H_2^{17}O and H_2^{18}O to improve the predicted transition frequencies. Calculations of vibrational energy levels of H_2^{19}O as well as other isotopologues of water vapor (H_2^XO , where $X=11, \dots, 26$) were early presented in Ref. [47].

Table 1 presents data on different oxygen isotopes. It can be seen that after ^{15}O and ^{14}O , ^{19}O has the longest half-life. This also indicates the importance of predicting the spectrum H_2^{19}O .

An key focus of our study is to show that at the current level of development of the theory, expensive and complex experiments can be replaced by calculations without loss of accuracy and reliability. Moreover, calculations within the framework of our semi-empirical approach allow not only the calculation of absorption spectra with high accuracy, but also thermodynamic functions such as the temperature-dependent entropy and heat capacity.

Calculated spectrum and of line list of H_2^{19}O

The theoretical absorption spectrum of H_2^{19}O is obtained via direct variational calculation of line centers and intensities for states with rotational levels with $J \leq 20$ using a high accuracy potential energy surface (PES) [6], and an *ab initio* dipole moment surface (DMS) [22]. The calculations were carried out as follows. First, the vibrational-rotational (VR) energy levels and associated wavefunctions for H_2^{19}O were computed using the DVR3D variational nuclear-motion package [40]. These calculations followed the same procedure used for other water radio-isotopologues [52, 53], but with mass of the oxygen atom set to 19.003577969 Da. The calculations were carried out on the “amun” cluster (UCL,

Table 1 Atomic mass and half-life for known isotopes of oxygen ^XO , $X=13, \dots, 23$ [27]

Nuclide	Atomic mass and Uncertainty [Da]	Half-life and Uncertainty	Discovery Year
^{13}O	$13.024815435 \pm 0.000010226$	$8.58 \text{ ms} \pm 0.05$	1963
^{14}O	$14.008596706 \pm 0.000000027$	$70.621 \text{ s} \pm 0.011$	1949
^{15}O	$15.003065636 \pm 0.0000000526$	$122.266 \text{ s} \pm 0.043$	1934
^{16}O	$15.99491461926 \pm 0.00000000032$	Stable	1919
^{17}O	$16.99913175595 \pm 0.00000000069$	Stable	1925
^{18}O	$17.99915961214 \pm 0.00000000069$	Stable	1929
^{19}O	$19.003577969 \pm 0.00000283$	$26.470 \text{ s} \pm 0.006$	1936
^{20}O	$20.004075357 \pm 0.000000095$	$13.51 \text{ s} \pm 0.05$	1959
^{21}O	$21.008654948 \pm 0.000012882$	$3.42 \text{ s} \pm 0.10$	1968
^{22}O	$22.009965744 \pm 0.000061107$	$2.25 \text{ s} \pm 0.09$	1969
^{23}O	$23.015696686 \pm 0.000130663$	$97 \text{ ms} \pm 8$	1970

London, UK). This procedure has been used to produce a number of line list for H₂O and its isotopologues including BT2, VTT [49], POKAZATEL [33], Conway [10], Hot-Wat78 [32] and VoTe [50].

The line list for H₂¹⁹O, which we call VoTe-19, is designed for use at room temperature over a wide range of wavenumbers. It contains 72 920 states and 106 680 822 (more than one hundred million) transitions in total. The H₂¹⁹O (VoTe-19) line list is available from the ExoMol database www.exomol.com using ExoMol format [41]. Extracts from the States `.states` and Transition `.trans` files are shown in Tables 2 and 3, respectively. The H₂¹⁹O `.trans` files contain Einstein A coefficients (in s⁻¹) together with the upper and lower state ID numbers. The transitions are divided into twelve Transition files according to the following spectroscopic ranges: 0–500, 500–1000, 1000–1500, 1500–2000, 2000–2500, 2500–3500, 3500–4500, 4500–5500, 5500–7000, 7000–9000, 9000–14,000, 14,000–25,000 cm⁻¹.

The State file contains a list of ro-vibrational states of H₂¹⁹O with the state ID numbers, energy term values (in cm⁻¹), uncertainties (in cm⁻¹) and quantum numbers: the provision of which are discussed below. The exact quantum numbers are the total angular momentum J and the total symmetry $\Gamma = A_1, A_2, B_1, B_2$ in the Molecular Symmetry group C_{2v}(M). The total state degeneracy, g_i is given by $(2J+1)$ times the nuclear spin factor, g_{ns} . ¹⁹O has a nuclear

Table 3 Extract from a `.trans` file of the H₂¹⁹O line list

f	i	A_{fi}
53397	54633	0.32305E-23
25152	24212	0.80174E-20
4853	7956	0.24636E-22
64698	61306	0.62528E-22
38048	36839	0.38686E-19
56019	57261	0.80929E-19
19875	20924	0.13715E-18
54551	48696	0.40687E-24
24031	24993	0.62368E-18
29261	28107	0.12623E-17
26198	27331	0.24941E-17

f : Upper state counting number;
 i : Lower state counting number;
 A_{fi} : Einstein-A coefficient (in s⁻¹)

spin of $I = \frac{5}{2}$ which for H₂¹⁹O leads to $g_{ns} = 6, 6, 18,$ and 18 for $\Gamma = A_1, A_2, B_1, B_2,$ respectively.

In addition, for some of the levels, approximate quantum numbers are given using standard harmonic oscillator - rigid rotor designations: $\nu_1, \nu_2, \nu_3, K_a, K_c$; the provision of these is discussed below. Here ν_1, ν_2, ν_3 are the normal mode vibrational quantum numbers, K_a and K_c are the oblate and

Table 2 Extract from the `.states` file of the H₂¹⁹O line list

i	\tilde{E}/cm^{-1}	g	J	δ/cm^{-1}	ν_1	ν_2	ν_3	K_a	K_c	e/o	Γ_{tot}	$\tilde{E}_D/\text{cm}^{-1}$	Code
1	0.000000	6	0	0.000001	0	0	0	0	0	e	A1	0.000000	IE
2	1585.510477	6	0	0.045301	0	1	0	0	0	e	A1	1585.527013	IE
3	3133.674946	6	0	0.089534	0	2	0	0	0	e	A1	3133.680018	IE
4	3646.583546	6	0	0.104187	1	0	0	0	0	e	A1	3646.545140	IE
5	4640.650684	6	0	0.132590	0	3	0	0	0	e	A1	4640.650684	Ca
6	5215.434906	6	0	0.149012	1	1	0	0	0	e	A1	5215.425140	IE
7	6100.343362	6	0	0.174296	0	4	0	0	0	e	A1	6100.349223	IE
8	6747.186711	6	0	0.192776	1	2	0	0	0	e	A1	6747.174726	IE
9	7179.236738	6	0	0.205121	2	0	0	0	0	e	A1	7179.234201	IE
10	7407.596097	6	0	0.211645	0	0	2	0	0	e	A1	7407.572789	IE

i : state identifier;

\tilde{E} : state term value, DVR3D or pseudo-experimental;

g : state degeneracy;

J : state rotational quantum number;

δ : energy uncertainty;

$\nu_1 - \nu_3$: normal mode vibrational quantum numbers;

K_a and K_c : state oblate and prolate quantum numbers;

e/o even or odd - K_c : state oblate and prolate quantum numbers;

Γ_{tot} : total symmetry in C_{2v}(M)

\tilde{E}_D : state term value, DVR3D;

Code - Ca (Calculated by DVR3D) or IE - Isotopologue Extrapolation (Pseudo-experimental)

prolate rotational quantum numbers (projection of the angular momentum on the a and c molecular axes, respectively).

DVR3D only supplies rigorous quantum numbers which for H_2O correspond to J , parity and whether the state is ortho or para. To provide the approximate rotation and vibration quantum labels, namely ν_1 , ν_2 , ν_3 , K_a and K_c , we matched the H_2^{14}O energies to the assigned states of the parent isotopologue H_2^{16}O as provided in the VoTe line list [50], which was based on the calculations with the same PES [50]. Following VoTe, here we also provide the parity of the K_c quantum number, which can be reconstructed from the (rigorous) values of J and the total symmetry Γ as shown in [7]. ExoMol [41, 42] standard notation is adopted and the value “NaN” is to denote undetermined quantum numbers.

As an attempt to improve the energy levels, and hence transition wavenumbers, predicted for H_2^{19}O , we apply pseudo-experimental corrections given by Eq. (1) below to the set of 3426 empirical energy levels identified as common to all three stable isotopologues of interest. These states are indicated with the label “IE” (isotopologue extrapolation [24]) in contrast to all other, calculated values, labelled with “Ca”. Following the pseudo-experimental extrapolation method proposed in [32, 33], we use the obs.-calc. residuals obtained for the parent isotopologue to obtain empirical corrections to the ro-vibrational energy values of the minor isotopologues of water as follows:

$$E_N^{\text{corr}} = E_N^{\text{calc}} + E_{16}^{\text{obs}} - E_{16}^{\text{calc}}, \quad (1)$$

where E_N^{calc} is a DVR3D energy calculated for a isotopologue, $N = 17, 18, 19$, and $E_{16}^{\text{obs}} - E_{16}^{\text{calc}}$ is an empirical correction estimated as the difference between the calculated and experimental energies of H_2^{16}O . The approach is based on the assumption that the main source of the error is from the inaccuracy of the Born-Oppenheimer PES of water, which should affect all four isotopologues similarly and was shown to work well for H_2^{18}O and H_2^{17}O [32, 33].

The provision of uncertainties in the energy levels is now a formal part of the ExoMol data structure [41, 42]. For the pseudo-experimental values, the uncertainties are estimated as the the obs.-calc. residuals of the H_2^{17}O isotopologue obtained as the difference between the W2020 and DVR3D energy values. For the other levels, formula (2) was used:

$$\text{unc} = \Delta\xi\tilde{E} + \Delta BJ(J + 1), \quad (2)$$

with $\Delta\xi=1/35000 \text{ cm}^{-1}$ and $\Delta B=0.0005 \text{ cm}^{-1}$. Formula (2) was obtained via correlation and auto-correlation methods [5] applied to the W2020 [14] data of the three isotopologues in conjunction with the method of Voronin et al. [52]. For H_2^{16}O , we can use residuee for the 19225 empirical energy levels provided W2020, and calculate the errors using formula (2). For rotational-vibrational states, where the energies of all three H_2^{16}O isotopologues were present

in W2020, we used the double uncertainty of the main isotopologues, since it is very small.

A total 3426 levels were replaced by the pseudo-experimental values. The maximum deviations of these changes do not exceed 0.1 cm^{-1} .

Spectrum of H_2^{19}O for applications

The line list produced was used to model spectra of H_2^{19}O .

A 296 K, room temperature H_2^{19}O line list was generated using the intensity threshold of $>10^{-30} \text{ cm/molecule}$ for atmospheric applications containing 250 224 lines. The line list is provided as part of the supplementary material in a HITRAN-like format [16], see Table 4.

The HITRAN database provides data for seven isotopologues of water using the following isotopologue codes: 11(H_2^{16}O), 12(H_2^{18}O), 13(H_2^{17}O), 14(HD^{16}O), 15(HD^{18}O), 16(HD^{17}O), 17(D_2^{16}O). We adopted 18 as a code for H_2^{15}O [53] and 10 for H_2^{14}O [52]; here we use 19 for H_2^{19}O . None of this number have yet to be adopted by HITRAN. The line list consists of the isotopologue code (19) (Molecule number 1 + Isotopologue number 9), line positions, 296 K intensities (cm/molecule), Einstein coefficients (s^{-1}), air-broadened widths (γ_{air} , $\text{cm}^{-1}/\text{atm}$), self-broadened widths (γ_{self} , $\text{cm}^{-1}/\text{atm}$), temperature dependence component of air n_{air} , line shifts (set to 0.0 cm^{-1}) and ro-vibrational quantum number. The line broadening parameters were evaluated using the J and ‘ JJ –dependency’ methods [46, 48]. For the temperature-dependence exponent n_{air} (unitless) we assumed the water vapor air-broadened half-widths from Table 7 of HITRAN2004 [35].

Partition function, $Q(T)$, for H_2^{19}O as a function of temperature, $T(\text{K})$.

Using vibrational-rotational energy levels, the partition function of H_2^{19}O was calculated for different temperatures up to 1000 K using

$$Q(T) = \sum_i g_i \exp\left(-\frac{E_i}{kT}\right). \quad (3)$$

Fig. 1 shows a comparison of partition functions $Q(T)$ of the H_2^{19}O , H_2^{16}O and H_2^{15}O species for the temperature range from 1 to 1000 K. The main difference is the nuclear spin factor, which is 6 times larger for H_2^{19}O than for H_2^{16}O . Furthermore, the partition function of H_2^{16}O is significantly more complete, with the levels from [50] covering the rotational excitations up to $J = 50$.

In principle, with a set of levels up to $25,000 \text{ cm}^{-1}$ and up to $J = 20$, our partition function of H_2^{19}O should be

Table 4 Examples spectrum of H₂¹⁹O for applications

C	$\tilde{\omega}$	Intensity	A.coeff	γ_{air}	γ_{self}	E''	k_T	shift	$\nu'_1 \nu'_2 \nu'_3$	$J' K'_a K'_c$	$\nu''_1 \nu''_2 \nu''_3$	$J'' K''_a K''_c$
19	0.049770	0.179E-26	0.643E-12	0.0808	0.495	444.534074	0.64	0.0	0 0 0	5 2 3	0 0 0	6 1 6
19	0.100849	0.393E-29	0.107E-10	0.0808	0.495	1911.474732	0.73	0.0	0 1 0	5 1 5	0 1 0	4 2 2
19	0.109513	0.193E-29	0.151E-10	0.0808	0.495	2111.656650	0.73	0.0	0 1 0	5 3 3	0 1 0	4 4 0
19	1.535666	0.138E-28	0.197E-07	0.0284	0.207	3063.921281	0.36	0.0	NaN NaN NaN	14 NaN NaN	0 0 0	15 3 12
19	1.612058	0.103E-29	0.187E-07	0.0537	0.309	3490.471234	0.53	0.0	0 1 0	9 7 2	0 1 0	8 8 1
19	2.167510	0.142E-29	0.272E-06	0.0808	0.495	3574.837000	0.73	0.0	0 2 0	5 2 4	0 2 0	4 3 1
19	2.662134	0.110E-25	0.296E-06	0.0870	0.523	1893.370213	0.77	0.0	0 1 0	4 2 3	0 1 0	3 3 0
19	2.695240	0.275E-27	0.197E-06	0.0808	0.495	2380.912733	0.64	0.0	0 1 0	5 5 1	0 1 0	6 4 2
19	2.819662	0.725E-26	0.504E-06	0.0989	0.478	1729.193474	0.77	0.0	0 1 0	2 2 0	0 1 0	3 1 3
19	3.023634	0.221E-25	0.447E-06	0.0870	0.523	1807.870770	0.77	0.0	0 1 0	4 1 4	0 1 0	3 2 1
19	3.075450	0.109E-27	0.202E-06	0.0537	0.309	2887.401244	0.45	0.0	0 1 0	9 3 6	0 1 0	10 2 9
19	3.321044	0.118E-28	0.221E-06	0.0677	0.372	3072.715103	0.53	0.0	0 1 0	7 7 1	0 1 0	8 6 2
19	3.343575	0.359E-28	0.226E-06	0.0677	0.372	3072.692605	0.53	0.0	0 1 0	7 7 0	0 1 0	8 6 3
19	3.581622	0.146E-26	0.462E-06	0.0808	0.495	2380.028898	0.64	0.0	0 1 0	5 5 0	0 1 0	6 4 3
19	3.707662	0.120E-27	0.259E-06	0.0324	0.224	2725.588226	0.36	0.0	NaN NaN NaN	13 NaN NaN	0 0 0	14 3 11
19	4.391607	0.707E-29	0.457E-06	0.0232	0.198	3418.617109	0.38	0.0	0 0 0	15 6 10	0 0 0	16 3 13

C. molecule num.=1 +isotopologue number=9(19)

$\tilde{\omega}$ Transition wavenumber, cm⁻¹;

Intensity at 100% abundance, cm/molecule;

Einstein A-coefficient, s⁻¹;

γ_{air} Air-broadened width, cm⁻¹/atm;

γ_{self} Self- broadened width, cm⁻¹/atm;

E'' lower-state Energy, cm⁻¹;

k_T Temperature dependence (of air width), dimensionless;

Pressure shift, always zero in our case;

Upper vibrational quanta, ν'_1, ν'_2, ν'_3 ;

Upper local quanta, J', K'_a, K'_c ;

Lower vibrational quanta, $\nu''_1, \nu''_2, \nu''_3$;

Lower local quanta, J'', K''_a, K''_c

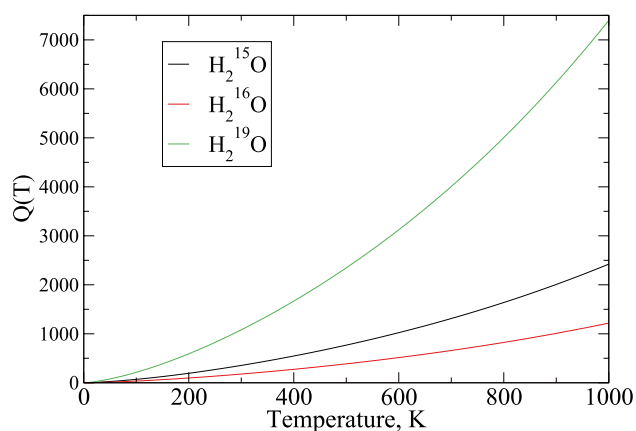


Fig. 1 Partition function $Q(T)$ for H₂¹⁹O, H₂¹⁶O and H₂¹⁵O water for temperature range from 1 to 1000 K

appropriate for up to 800 K. Using our results for temperatures higher than this will lead to resulting spectra becoming increasingly incomplete, for more details about partition function see [18, 45]. Our partition function for H₂¹⁹O as in presented as a table in the Supplementary materials (Part-Fun-H219O-J20.dat).

Estimated possibility of detecting H₂¹⁹O in laboratory conditions

The isotopologues of H₂O have characteristic absorption fingerprints in the IR spectral region and can be analyzed quantitatively and qualitatively using spectroscopic methods. The advantage of high resolution spectroscopy is that it is a non-destructive and non-invasive method which allows simultaneous monitoring of several species in real time.

To reveal spectral windows in which H₂¹⁹O could be detected under laboratory conditions, we simulated the

transmission of main isotopologues of H₂O (H₂¹⁶O, H₂¹⁸O, H₂¹⁷O, HD¹⁶O, HD¹⁸O, HD¹⁷O) in natural abundance with the addition H₂¹⁹O at high spectral resolution for $T = 294$ K, atmospheric pressure of 1013 mbar, partial pressure of H₂O 18,800 ppm and path length of 100 m. The H₂¹⁹O/H₂O ratio was varied from 0.1 to 1%. Transmission was calculated using a line-by-line method [26] with a boxcar apparatus function.

The absorption line parameters, needed for the line-by-line transmission calculation, are the line intensity, S (cm⁻¹/molecule cm⁻²), position of the line centre, ν^* (cm⁻¹), lower-state energy, E (cm⁻¹), air- and self-broadened half-widths γ_{air} and γ_S (cm/atm), and temperature-dependence exponent for $\gamma_{\text{air}} - n_{\text{air}}$. The H₂O absorption lines parameters of main isotopologues were taken from the HITRAN2020 spectroscopic database [16]; our calculated absorption line parameters were used to simulate the transmission by H₂¹⁹O. Laboratory spectrometers measure the intensity (radiance) of source I , attenuated by gaseous medium at the path length L with the transmission function T and apparatus function f with the spectral resolution $\Delta\nu$. The intensity at wavenumber ν is determined by

$$I(\nu) = \int_{\Delta\nu} I_0(\nu') T(\nu') f(\nu' - \nu) d\nu' \\ = \int_{\Delta\nu} I_0(\nu') \exp(-K(\nu')L) f(\nu' - \nu) d\nu'. \quad (4)$$

The absorption coefficient, $K(\nu)$, includes the contributions of all spectral lines in the spectral interval considered and depends on pressure, p , and temperature, t :

$$K(\nu, p, t) = \sum_{j=1}^M \sum_{i=1}^N S_{ij}(t) g(\nu_{ij}^*, \nu, p, t) n_j, \quad (5)$$

where n_j is the concentration of the j th gas (isotopologue); M is the number of gases (isotopologues); N is the number of spectral lines belonging to the j th gas; and $g(\nu_{ij}^*, \nu, p, t)$ is the absorption line profile function. A Voigt line profile [13, 31] is usually employed for atmospheric conditions. Line profiles were calculated using broadening line parameters γ and the air temperature-dependence exponent, n_{air} . Transmission spectra of H₂O and H₂¹⁹O with a H₂¹⁹O/H₂O ratio of 0.5%, calculated with spectral resolution of 0.02 cm⁻¹ and 5 cm⁻¹ in the 0–15,000 cm⁻¹ spectral region, are shown in Fig. 2. It was found that use high spectral resolution of about 0.02 cm⁻¹ allows the H₂¹⁹O lines to be revealed against of the background H₂O lines at this H₂¹⁹O/H₂O concentration ratio.

Transmission spectra of H₂O and H₂¹⁹O with H₂¹⁹O/H₂O ratio of 0.5% are shown in Fig. 2.

By varying the H₂¹⁹O/H₂O ratio, spectral windows suitable for H₂¹⁹O detection under laboratory conditions were identified in regions from far to near IR. Examples of the

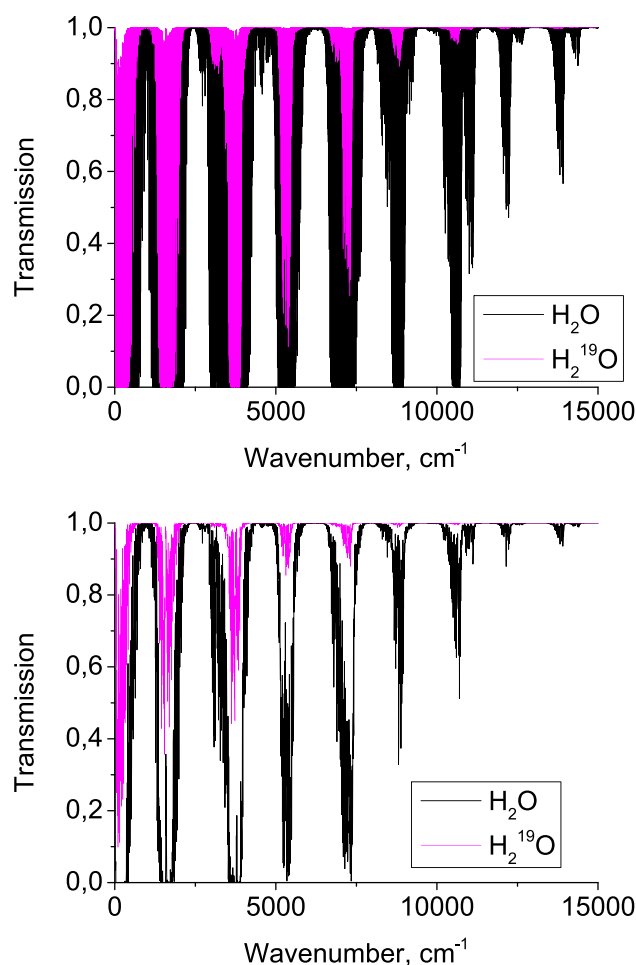


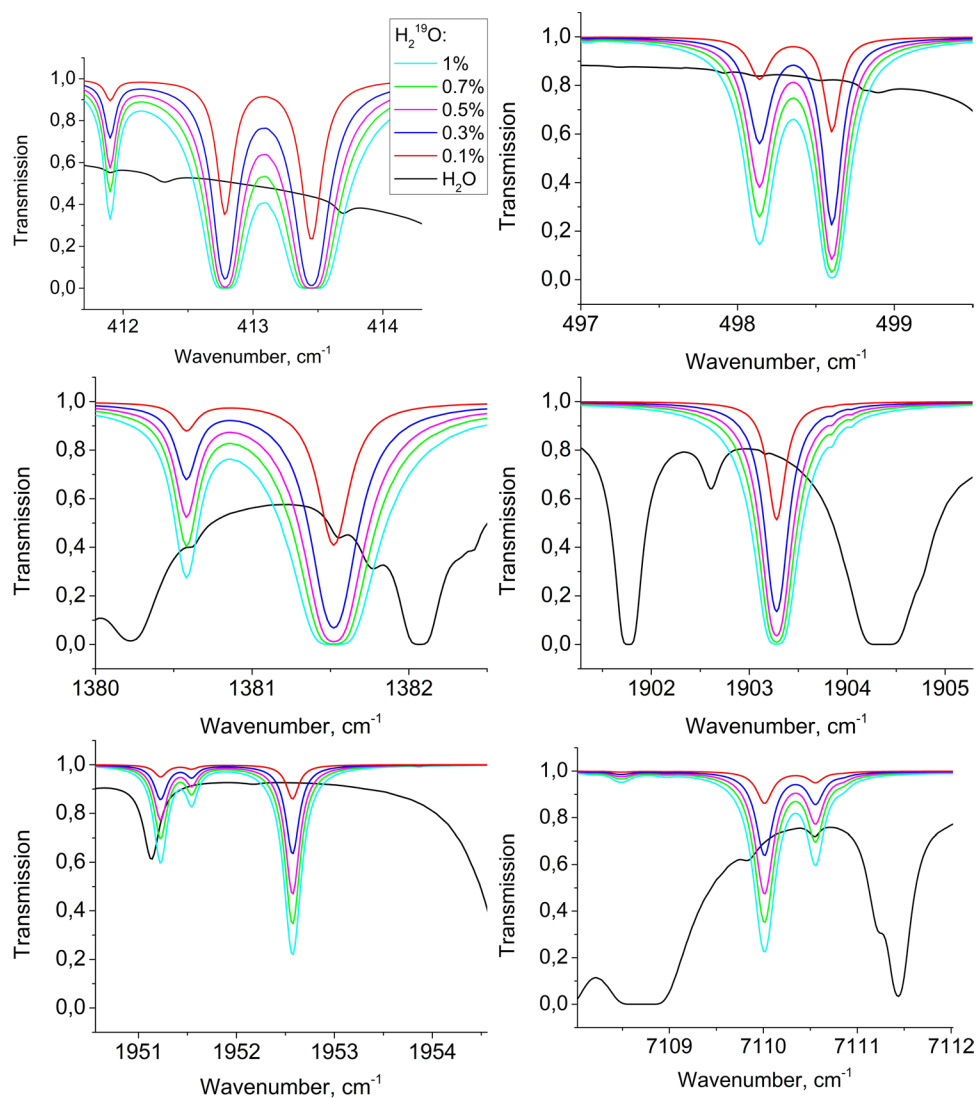
Fig. 2 Transmission spectra of H₂O and H₂¹⁹O, simulated with spectral resolution of (upper panel) 0.02 cm⁻¹ and (lower panel) 5 cm⁻¹; the H₂¹⁹O/H₂O ratio is 0.5%

windows are presented in Fig. 3, where one can observe a distinct contribution of the H₂¹⁹O lines against a background of H₂O absorption. The spectral windows for perspective H₂¹⁹O detection are located in the 411–414, 497–499, 1380–1383, 1902–1905, 1950–1954 and 7109–7112 cm⁻¹ wavenumber regions.

In our previous studies [52, 53] the spectra of H₂¹⁵O and H₂¹⁴O radio-isotopologues were investigated, and the spectral windows, suitable for H₂¹⁴O and H₂¹⁵O detection, were identified. It is interesting to compare these results with ones obtained for H₂¹⁹O. This comparison reveals overlapping absorption lines of the radio-isotopologues and allows one to estimate how their interference might impact detections. The transmission spectra of H₂¹⁵O and H₂¹⁴O with the concentrations of 0.5% relative to H₂O content are shown in Fig. 4 in the spectral windows where the H₂¹⁹O can be detected.

As can see in Fig. 4, the H₂¹⁹O absorption lines overlap with H₂¹⁵O lines in the window of 1380–1383 cm⁻¹, that interfere with the H₂¹⁹O detection in this spectral range.

Fig. 3 Spectral windows for the detection of H_2^{19}O at different relative concentration of H_2^{19}O . The $\text{H}_2^{19}\text{O}/\text{H}_2\text{O}$ ratio varies from 0.1 to 1%



Moreover, the H_2^{14}O lines can influence on the H_2^{19}O measurement in the $7109\text{--}7112\text{ cm}^{-1}$ window. According to the transmissions comparisons, the most suitable spectral ranges for detecting H_2^{19}O in laboratory conditions are $412\text{--}414\text{ cm}^{-1}$ and $497\text{--}499\text{ cm}^{-1}$.

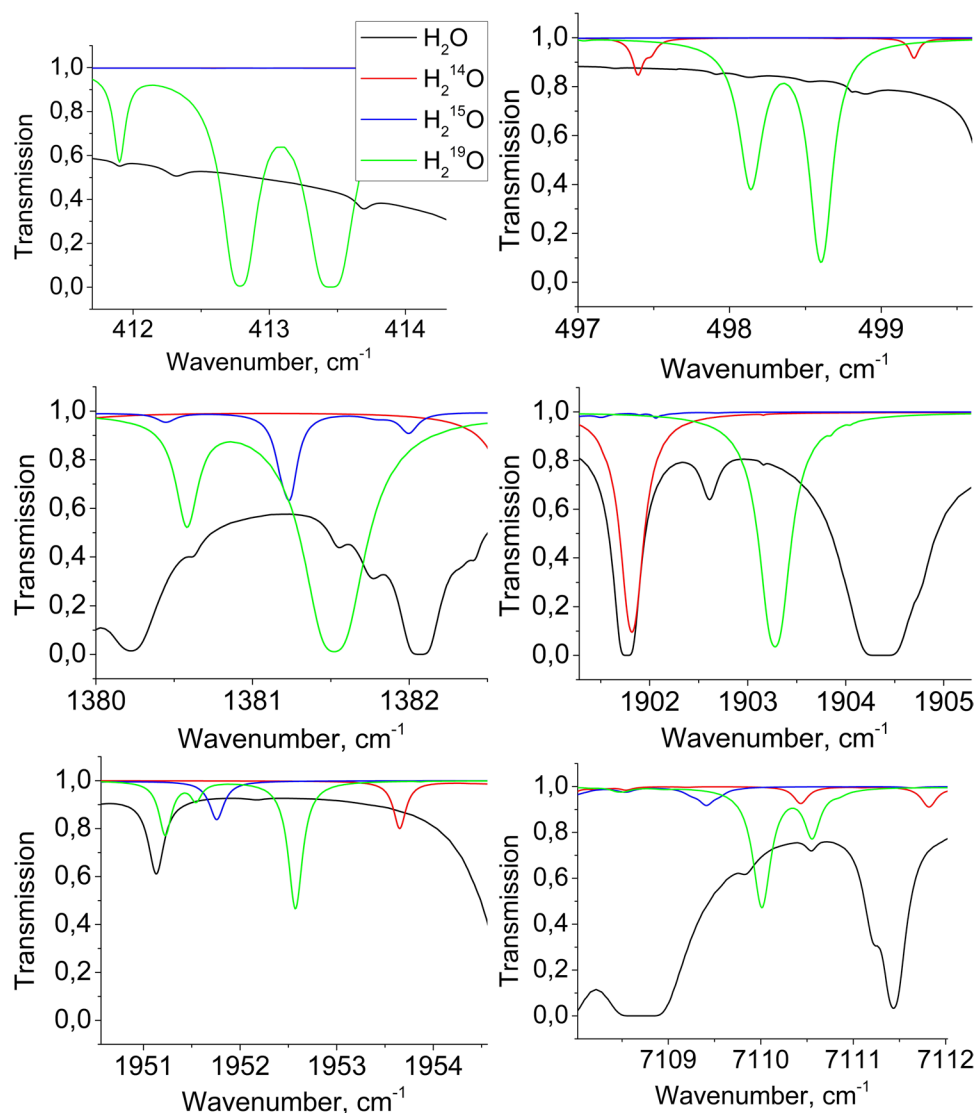
Conclusion

The first comprehensive calculation of vibrational-rotational spectrum of the water isotopologue H_2^{19}O up to $25,000\text{ cm}^{-1}$ is presented. An empirical line list for the H_2^{19}O is created in the ExoMol-style using calculations based on a high-precision intramolecular potential energy surface which includes allowance for relativistic and non-adiabatic corrections, and *ab initio* dipole moment surfaces. The line list is computed using a spectroscopic model optimised for H_2^{16}O and covers

all transitions up to $J=20$, which means it should be valid for temperatures up to 800 K. The quality of the line list was improved through isotopologue extrapolation of the experimentally-determined energies of H_2^{16}O , H_2^{17}O and H_2^{18}O . We hope that these theoretical prediction of H_2^{19}O spectra will lead to experimental measurements.

Transmission spectra of H_2^{19}O were simulated varying their concentration. A comparison of the simulated spectra of H_2^{19}O with the transmission spectra of water vapor in natural abundance (H_2^{16}O , H_2^{18}O , H_2^{17}O , HD^{16}O , HD^{18}O , HD^{17}O), and H_2^{14}O H_2^{15}O radio-isotopologues was carried out to find the spectral ranges are best suited for H_2^{19}O detection in laboratory conditions. It was revealed that the spectral windows relevant for the H_2^{19}O detection are placed in the regions of $411\text{--}414\text{ cm}^{-1}$, $497\text{--}499\text{ cm}^{-1}$, $1380\text{--}1383\text{ cm}^{-1}$, $1902\text{--}1905\text{ cm}^{-1}$, $1950\text{--}1954\text{ cm}^{-1}$, and $7109\text{--}7112\text{ cm}^{-1}$, with the two far infrared windows showing the least interference with the other

Fig. 4 Transmission spectra of H_2O in natural abundance and radio-isotopogues: H_2^{14}O , H_2^{15}O and H_2^{19}O at concentrations of 0.5% of H_2O



radio-isotopologues. As ^{19}O decays, the concentration of the H_2^{19}O isotopologue will diminish, and its absorption will also decrease. The rate of decrease in absorption at a given wavelength, observed in real time, will provide additional evidence that it the absorptions are due to H_2^{19}O .

Our predicted spectrum of H_2^{19}O can be a useful tool for for astrophysics, plasma chemistry, nuclear reactor safety, medical procedures and aid the detection of this radio-isotopologue. It can be expect that H_2^{19}O water plays a greater role in fast-paced complex chemical and nuclear processes than previously assumed. For example, it is possible that short-lived water vapor radio-isotopologues may be responsible for part of the anomalous radiation that is sometimes recorded during thunderstorms [9].

Vibrational-rotational spectroscopy is an effective, non-destructive tool for studying the properties of pollutants and radioactive substances in real time mode. Such studies

can help to understand the physicochemical processes involved and to improve the safety of nuclear reactors.

Acknowledgements The authors thank S.N. Yurchenko (UCL) and Olga Naumenko for useful consultations and prof. Flávio C. Cruz (UNICAMP) for funding acquisition. Boris Voronin is grateful for the financial support of the grant - FAPESP (2022/08772-1). The work was partially supported by the MSHE RF (V.E. Zuev IAO SB RAS).

Author Contributions Boris A. Voronin: Term, Validation, Investigation, Writing - Original Draft; Jonathan Tennyson: Supervision, Conceptualization, Resources, Writing - Review & Editing, Methodology; Tatyana Yu. Chesnokova: Investigation, Data Curation, Writing - Review & Editing; Aleksei V. Chentsov: Formal analysis, Visualization; Aleksandr D. Bykov: Conceptualization, Validation.

Data Availability File partition-function-H2190.dat containing only 2 columns, can be found in the supplementary materials. In the first column, the temperature in K, from 1 to 1000 K, in the second column, the value of the partition function at that temperature. File H2190-spectra-296K-E30.dat, see Table (4), can be found in the supplementary materials. At <https://ftp.iao.ru/pub/VTT/H2190/> there are

15 files: 12 files containing transitions divided into ranges (format is described in the table 3); one file with the energy levels and identification; one file for atmospheric applications in a format like HITRAN-04 database and one file as an archive for easy download. The full dataset is given as part of the ExoMol database at www.exomol.com. Spectra can be generated using program ExoCross [54].

Declarations

Conflict of interest The authors declare no conflict of interest.

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