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On equilibrium structures of the water molecule

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Equilibrium structures are fundamental entities in molecular sciences. They can be inferred from experimental data by complicated inverse procedures which often rely on several assumptions, including the Born–Oppenheimer approximation. Theory provides a direct route to equilibrium geometries. A recent high-quality *ab initio* semiglobal adiabatic potential-energy surface (PES) of the electronic ground state of water, reported by Polyansky *et al.* [Science **299**, 539 (2003)] and called CVRQD here, is analyzed in this respect. The equilibrium geometries resulting from this direct route are deemed to be of higher accuracy than those that can be determined by analyzing experimental data. Detailed investigation of the effect of the breakdown of the Born–Oppenheimer approximation suggests that the concept of an isotope-independent equilibrium structure holds to about 3×10^{-5} Å and 0.02° for water. The mass-independent [Born–Oppenheimer (BO)] equilibrium bond length and bond angle on the ground electronic state PES of water is $r_e^{\text{BO}} = 0.957\,82$ Å and $\theta_e^{\text{BO}} = 104.48_5^\circ$, respectively. The related mass-dependent (adiabatic) equilibrium bond length and bond angle of H_2^{16}O is $r_e^{\text{ad}} = 0.957\,85$ Å and $\theta_e^{\text{ad}} = 104.50_0^\circ$, respectively, while those of D_2^{16}O are $r_e^{\text{ad}} = 0.957\,83$ Å and $\theta_e^{\text{ad}} = 104.49_0^\circ$. Pure *ab initio* prediction of $J=1$ and 2 rotational levels on the vibrational ground state by the CVRQD PESs is accurate to better than 0.002 cm^{-1} for all isotopologs of water considered. Elaborate adjustment of the CVRQD PESs to reproduce all observed rovibrational transitions to better than 0.05 cm^{-1} (or the lower ones to better than 0.0035 cm^{-1}) does not result in noticeable changes in the adiabatic equilibrium structure parameters. The expectation values of the ground vibrational state rotational constants of the water isotopologs, computed in the Eckart frame using the CVRQD PESs and atomic masses, deviate from the experimentally measured ones only marginally, especially for A_0 and B_0 . The small residual deviations in the effective rotational constants are due to centrifugal distortion, electronic, and non-Born–Oppenheimer effects. The spectroscopic (nonadiabatic) equilibrium structural parameters of H_2^{16}O , obtained from experimentally determined A'_0 and B'_0 rotational constants corrected empirically to obtain equilibrium rotational constants, are $r_e^{\text{sp}} = 0.957\,77$ Å and $\theta_e^{\text{sp}} = 104.48^\circ$.

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I. INTRODUCTION

The Born–Oppenheimer (BO) approximation^{1–3} introduces the separation of nuclear and electronic motions and is the single most important concept in our understanding of almost all of molecular sciences, especially that of chemistry. This is due to the fact that the Born–Oppenheimer approxi-

mation defines the concept of an electronic potential-energy (hyper)surface (PES).^{4–6} Potential-energy (hyper)surfaces, which in the standard approximation are mass-independent, are mathematically defined as total energies of species with respect to their geometric variables responsible for the internal motions. Adiabatic corrections^{7–11} to the BO-PES relax the strict separation of electronic and nuclear degrees of freedom, defining an adiabatic (mass-dependent) PES. Many molecular processes and properties, discussed routinely, only have meaning within the Born–Oppenheimer approximation

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TABLE I. Brief history of the mass-independent spectroscopic equilibrium structure of water. Bond length, r_e , in angstrom; bond angle, θ_e , in degrees.

Year	r_e	θ_e	Comment
1932	...	115	Ref. 23 ^a
1945	0.958 ₄	104.45	Ref. 24 ^b
1956	0.9572(3)	104.52(5)	Ref. 25 ^c
1961	0.956 ₁	104.57	Ref. 26 ^d
1976	0.9575	104.51	Landolt–Börnstein, Ref. 27 ^e
1979		104.48	Ref. 28
1994	0.9578	104.48	Ref. 29
1994	0.957 85(3)	104.542(9)	Ref. 30
1997	0.957 67	104.482	Ref. 31, <i>ab initio</i> PES
1997	0.957 83	104.509	Ref. 31, fitted empirical PES
2003	0.957 82	104.499	Ref. 32, fitted PES of H ₂ ¹⁶ O

^aObtained from the fundamental frequencies of water assumed to be 5309, 1597, and 3742 cm⁻¹ and through the use of an equation derived by Dennison [D. M. Dennison, *Philos. Mag.* **1**, 195 (1926)].

^bBased upon careful analysis of results due to Mecke and co-workers [R. Mecke, *Z. Phys.* **81**, 313 (1933); W. Baumann and R. Mecke, *Z. Phys.* **81**, 445 (1933); K. Freudenberg and R. Mecke, *Z. Phys.* **81**, 465 (1933)], Darling and Dennison [B. T. Darling and D. M. Dennison, *Phys. Rev.* **57**, 128 (1940)], and Nielsen [H. H. Nielsen, *Phys. Rev.* **59**, 565 (1941); H. H. Nielsen, *Phys. Rev.* **62**, 422 (1942)].

^cBenedict *et al.* (Ref. 25) reported equilibrium structures not only for H₂ ¹⁶O but also for HD ¹⁶O and D₂ ¹⁶O, after inclusion of electronic motion effects into the moments of inertia. The D₂ ¹⁶O structural parameters differ from the equilibrium structural parameters of H₂ ¹⁶O by +0.0003 Å and -0.049°. These differences are about an order of magnitude larger than those obtained in the present study (cf. Table IV).

^dKuchitsu and Bartell reported structural estimates for D₂ ¹⁶O, as well, with $r_e=0.957_0$ Å and $\theta_e=104.43^\circ$. The results reported were obtained from the rotational constants of Benedict *et al.* (Ref. 25) and the vibration-rotation interaction constants determined in Ref. 26.

^eThese values are based on ground-state rotational constants from Cook *et al.* [R. L. Cook, F. C. De Lucia, and P. Helminger, *J. Mol. Spectrosc.* **53**, 62 (1974)] and on vibration-rotation interaction constants from Ref. 25.

and for the resulting (adiabatic) PESs. The most important of these is arguably the equilibrium structure, corresponding to the molecular configuration at the very bottom of a potential well.^{12–22} The equilibrium structure is the subject of the present study, employing water, a triatomic molecule of highest relevance, as an example. For a brief historical review of some of the most relevant determinations^{23–32} of the mass-independent, and, in cases, mass-dependent equilibrium structure of the water molecule see Table I. It is clear that by the 1940s, when Herzberg published his influential epic on rovibrational spectra,²⁴ the structure of water was known with remarkable precision. Nevertheless, the arduous task of moving beyond this precision has proved to be non-trivial even for such a small and simple triatomic molecule.

In an excellent review entitled “Determination of reliable structures from rotational constants,”¹³ Demaison and co-workers set the standards as “reliable structures [are] structures with an uncertainty less than 0.2° for the angles and 0.002 Å for the distances.” This standard of accuracy seems to be at least one order of magnitude less than the expected precision of adiabatic equilibrium structures deduced from high-quality *ab initio* calculations. It is also well known that there are intrinsic and sizable differences between the equilibrium (r_e) structures and the experimentally derived $r_a, r_\alpha, r_c, r_g, r_m, r_s, r_z$, etc., structures.^{13,16–18,33} Several excellent summaries of the approximate relationships

between the large number of observable structures and the equilibrium structure exist.^{13,18,33,34} For example, one of the best compromise when gas-phase electron diffraction (GED) and millimeterwave (MW) spectroscopy are used jointly for structure refinement is a temperature-dependent r_g structure. Due to the fact that the approximate (“diatomic”) relation $r_e - r_g = (3/2) au^2$ holds reasonably well, where a is the usual Morse parameter^{16,35} and u is the vibrational amplitude of the bond concerned, the difference between the r_e and r_g distance types, as well as between r_e and any vibrationally averaged distance type, is substantial, on the order of 0.01 Å.

To arrive at spectroscopic equilibrium structures an inverse, perturbative approach is widely employed.³⁶ As part of this procedure the accurately measured ground vibrational state effective rotational constants are corrected, principally by vibration-rotation interaction constants of different origins and orders, and equilibrium geometries are deduced from the resulting equilibrium rotational constants. This inverse procedure can be employed to fairly large molecules, see, e.g., a recent study on the 17-atom amino acid proline.³⁷ Nevertheless, the ultimate accuracy of this route to equilibrium geometries, which starts from experimental observables (effective rotational constants) and corrects them with appropriate constants, has proved to be, in some sense, insufficient, as diatomic paradigms have shown.^{38–40}

A direct, fully *ab initio* approach comes to the rescue. It starts with first-principles computation of equilibrium geometries. The ultimate accuracy of this direct approach has so far been realized (see, e.g., Refs. 19 and 38–40) only for diatomic and few-electron triatomic⁴¹ cases due to the incompleteness of most presently applicable computational approaches. Nevertheless, with recent significant advances in electronic structure theory and computer hardware, the goal of achieving high precision for mass-independent (Born–Oppenheimer) as well as mass-dependent (adiabatic) equilibrium structures has become possible even for many-electron systems, as shown here.

In the direct approach comparison with experiment is difficult, in fact, impossible if only the equilibrium structure has been determined *ab initio*. If the same level of theory was used to compute a PES, comparison between computed and experimental transition frequencies, most importantly rotational transitions on the ground vibrational state, becomes feasible. Another possibility for comparison between theory and experiment is provided at the level of vibrationally averaged spectroscopic quantities, determined here as the appropriate expectation values using variational vibrational wave functions. The direct approach is helped in the latter case by the fact that close-to-exact solution of the vibrational motion problem, especially for triatomic species, is straightforward to achieve.^{42–44} Both approaches for comparison are pursued in this paper. One must remember, nevertheless, that theory going beyond the diagonal Born–Oppenheimer correction, i.e., consideration of non-BO effects, is still in its infancy, and that the so-called experimental spectroscopic constants are effective parameters, obtained through phenomenological approaches which parametrize the Hamiltonian as flexibly as needed. The subtle relation between the spectroscopic and the Born–Oppenheimer and adiabatic

TABLE II. Born–Oppenheimer and adiabatic equilibrium structures of water from *ab initio* electronic structure computations. Bond length (r_e) in angstrom, bond angle (θ_e) in degrees. The ICMRCI calculations are valence only (freezing the 1s core orbital of O). CBS=complete basis set.

Level of theory/correction	r_e	θ_e	Comment
aug-cc-pV5Z ICMRCI	0.958 75	104.387	Ref. 42
aug-cc-pV6Z ICMRCI	0.958 70	104.411	Ref. 42
CBS ICMRCI	0.958 62	104.422	Ref. 42
Core correlation	−0.000 96	+0.134	Refs. 31 and 42
Relativistic Breit	+0.000 16	−0.074	Refs. 42, 55, and 56
Quantum electrodynamics	<10 ^{−5}	+0.003	Refs. 42 and 64
Best mass-independent <i>ab initio</i>	0.957 82	104.485	This work, Ref. 42
Adiabatic correction (H ₂ ¹⁶ O)	+0.000 03	+0.015	Ref. 42
Best mass-dependent <i>ab initio</i> (H ₂ ¹⁶ O)	0.957 85	104.500	This work, Ref. 42

equilibrium structures is not completely clear, as also exemplified in this study.

In a recent publication⁴² some of us presented *ab initio* adiabatic PESs of the ground electronic state of water which yielded, for the first time for a polyatomic and polyelectronic molecule, so-called spectroscopic accuracy, 1 cm^{−1}, on average for *all* measured rovibrational levels of all isotopologs considered. This remarkable precision could not have been

obtained without the intrinsic *ab initio* determination of highly accurate, mass-dependent equilibrium structures of water. In this paper we provide a detailed analysis of our *ab initio* results for the equilibrium structures of water (Table II). Water is probably the only polyatomic molecule besides H₃⁺ for which adiabatic PESs of almost ultimate accuracy are available.

II. COMPUTATIONAL DETAILS

In this study the DOPI,^{43,45} where DOPI stands for discrete variable representation (D), Hamiltonian in orthogonal (O) coordinates, direct product (P) basis, iterative (I) diagonalization, and the DVR3D⁴⁶ program suites, both employing a discrete variable representation (DVR) for all three vibrational degrees of freedom, have been employed for the determination of the $J=0, 1$, and 2 rovibrational energy levels. Highly converged wave functions were obtained during the nuclear motion calculations, the rovibrational eigenvalues from the two programs, employing somewhat different formalisms, agreed with each other to better than 10^{−3} cm^{−1}. The relevant input files employed for running the more widely utilized DVR3D program are provided in the Supplementary Material (EPAPS, Ref. 113).

The nuclear masses employed in the variational nuclear motion calculations are given in caption to Table III. For the

TABLE III. $J=1$ and 2 rotational term values (J_{K,K_c}), in cm^{−1}, for the ground vibrational states of water isotopologs. 1 cm^{−1}=2.997 924 58×10⁴ MHz. The *ab initio* CVRQD PESs are taken from Ref. 42. The fitted PESs for H₂¹⁶O, H₂¹⁷O, and H₂¹⁸O are due to Shirin *et al.* (Ref. 79). The fitted PES used for D₂O is from S. V. Shirin, N. F. Zobov, O. L. Polyansky, J. Tennyson, T. Parekunnel, and P. F. Bernath, J. Chem. Phys. **120**, 206 (2004). Nuclear masses, in u, used in the calculations and in the determination of the adiabatic correction surfaces: $m(\text{H})=1.007\,276$, $m(\text{D})=2.013\,553$, $m(^{16}\text{O})=15.990\,526$, $m(^{17}\text{O})=16.994\,742$, and $m(^{18}\text{O})=17.994\,771$.

	H ₂ ¹⁶ O			H ₂ ¹⁷ O		
	CVRQD	Fitted	Expt. ^a	CVRQD	Fitted	Expt. ^b
1 ₀₁	23.795	23.794	23.7944	23.774	23.774	23.7735
1 ₁₁	37.138	37.137	37.1371	36.931	36.931	36.9311
1 ₁₀	42.372	42.372	42.3717	42.187	42.187	42.1869
2 ₀₂	70.094	70.091	70.0908	70.007	70.005	70.0047
2 ₁₂	79.499	79.496	79.4964	79.229	79.227	79.2273
2 ₁₁	95.178	95.176	95.1759	94.973	94.971	94.9705
2 ₂₁	134.903	134.901	134.9016	134.146	134.145	134.1453
2 ₂₀	136.166	136.164	136.1639	135.432	135.431	135.4312
	H ₂ ¹⁸ O			D ₂ ¹⁶ O		
	CVRQD	Fitted	Expt. ^c	CVRQD	Fitted	Expt. ^d
1 ₀₁	23.756	23.755	23.7549	12.117	12.117	12.117
1 ₁₁	36.749	36.749	36.7486	20.257	20.258	20.258
1 ₁₀	42.024	42.023	42.0234	22.682	22.683	22.683
2 ₀₂	69.930	69.927	69.9274	35.877	35.878	35.878
2 ₁₂	78.991	78.989	78.9886	42.067	42.068	42.068
2 ₁₁	94.791	94.789	94.7886	49.337	49.338	49.338
2 ₂₁	133.478	133.476	133.4758	73.669	73.672	73.672
2 ₂₀	134.785	134.783	134.7830	74.135	74.137	74.137

^aReference 75.

^bReference 76.

^cReference 77.

^dReference 78.

calculation of vibrationally averaged rotational constants atomic masses have been used.

III. THE *AB INITIO* CVRQD POTENTIAL-ENERGY SURFACE OF WATER

The high-accuracy semiglobal adiabatic *ab initio* ground electronic state PESs of the water isotopologs, utilized heavily in this study, were first described in Ref. 42. Since few details are provided in that publication, the potentials are discussed here briefly, giving reference also to the codes employed in their computation. Components of this *ab initio* potential, denoted as CVRQD throughout this paper, were obtained as follows: (a) the final PES is built upon augmented correlation consistent polarized valence X zeta (aug-cc-pVXZ),^{47,48} $X=4, 5$, and 6 , valence-only internally contracted multireference configuration interaction⁴⁹ (ICMRCI) calculations, extrapolated to the complete basis set (CBS) limit, over a grid containing more than 350 points, employing the program package MOLPRO;⁵⁰ (b) added to this is the core correlation correction surface of Partridge and Schwenke³¹ determined at a different set of points at the averaged coupled pair functional⁵¹ (ACPF) level employing the program MOLPRO; (c) added to this is a relativistic correction surface obtained by first-order perturbation theory as applied to the one-electron mass-velocity (MV) and one- and two-electron Darwin terms (MVD2),^{52–56} calculated using the program packages ACESII (Refs. 57 and 58) and DALTON,⁵⁹ supplemented by a correction obtained from the inclusion of the Breit term in the electronic Hamiltonian and using four-component Dirac–Hartree–Fock wave functions, utilizing the program package BERTHA;^{60,61} (d) added to this is a correction surface due to effects from quantum electrodynamics (QED),^{62,63} in particular, the one-electron Lamb shift, determined by a simple scaling procedure;⁶⁴ and (e) finally, adding to this the adiabatic correction surface obtained at the MRCI level.⁴² It should also be noted that a nonadiabatic correction was also employed in Ref. 42 for the calculation of (ro)vibrational band origins, based on a two-term adjustment of the vibrational kinetic-energy operator^{42,65} employing parameters taken from a ground-breaking study by Schwenke.⁶⁶

Because of an often favorable error compensation between incompleteness of the basis set and deficiencies in the treatment of electron correlation, as well as the use of approximate Hamiltonians,^{6,54,67} lower-level electronic structure computations often result in surprisingly small errors for a particular property. This useful practical approach is not pursued here as, instead, this study focuses on the convergence of electronic structure theory to obtain the best technically possible equilibrium geometries of the isotopologs of water. Therefore, the large number of results from intermediate levels of theory, a few of which are given in Table II, is not utilized directly in this study. Note finally that *ab initio* computation of equilibrium geometries is a somewhat unbalanced procedure⁶⁸ due to the almost complete cancellation of two large terms of opposite sign, the approximate electronic and the possibly exact nuclear first derivatives, contributing to the vanishing forces at equilibrium.

IV. THE DIRECT COMPUTATIONAL ROUTE TO BORN–OPPENHEIMER AND ADIABATIC EQUILIBRIUM STRUCTURES

The direct route to the computation of accurate, converged Born–Oppenheimer and adiabatic equilibrium structures employs systematically improved levels of electronic structure theory. This is due to the fact that for polyelectronic systems none of the limits of *ab initio* electronic structure theory⁶ can be reached without some sort of extrapolation and approximation. As detailed in Sec. III, the ultimate mass-independent (Born–Oppenheimer) equilibrium structure of water, with an OH bond length of $r_e^{\text{BO}}=0.957\,82\text{ \AA}$ and a HOH bond angle of $\theta_e^{\text{BO}}=104.48_5^\circ$, has been determined⁴² from a large set of often expensive *ab initio* electronic structure computations, relying in part on well-established extrapolation and approximation techniques. These computations result in a point-wise representation of the ground electronic state PES of water. Consequently, the Born–Oppenheimer and adiabatic equilibrium structural parameters of the water isotopologs, reported throughout this paper, have been obtained by interpolations employing fitted PESs.⁴² The use of this procedure means that six-digit accuracy in the *ab initio* determination of the equilibrium structural parameters of the isotopologs of water cannot be expected.

The convergence of the valence-only ICMRCI treatment with respect to the one-particle basis set expansion is relatively fast (cf. Table II). Even more important for the purposes of the present study are the incremental contributions to the equilibrium structure of water, also listed in Table II. The significant contribution of core correlation to the equilibrium structure of water is not at all surprising.^{69,70} The related results of Martin,⁷⁰ obtained at a less complete level of theory, seem to provide a slight underestimation of the core correlation effect for the bond length. Accuracy on the order of 10^{-4} \AA in r_e can only be achieved if relativistic effects are taken into account. For θ_e , due to the more substantial change in the electronic structure of water upon bending, relativistic effects have more than half of the contribution of core correlation. This is in line with what has been observed during computational investigations of the barrier to linearity of water.^{67,71,72} Note also the opposite sign of the core correlation and relativistic contributions, resulting in an improved apparent accuracy of the valence-only treatment. The adiabatic contribution, while almost negligible for the equilibrium bond length, is substantial for the bond angle. Even consideration of quantum electrodynamic effects, namely, the one-electron Lamb shift,⁶⁴ has a noticeable contribution to the bond angle, in the sixth significant digit, at the border of the claimed precision of the present study. In summary, the direct approach, though computationally expensive, provides a way to obtain highly accurate Born–Oppenheimer and adiabatic equilibrium structures in a controlled way.

Table IV summarizes the effects of isotopic substitution on the equilibrium structures of the water isotopologs. It is clear from this table, and from Table II as well, that the concept of mass-independent (Born–Oppenheimer) equilibrium structures seems to be valid for water to about

TABLE IV. Isotopic effects, based on the CVRQD potential, on the equilibrium structural parameters of water isotopologs. Bond lengths in angstrom, bond angles in degrees. The underlying mass-dependent equilibrium structure of H_2^{16}O is $r_e^{\text{ad}}(^{16}\text{OH})=0.957\,854\,\text{\AA}$ and $\theta_e^{\text{ad}}(\text{H}^{16}\text{OH})=104.500^\circ$, as given in Table II. The geometry parameters obtained from the empirical potential (see Ref. 79) are the same as those given above within the claimed precision. It is basically meaningless to give further digits in the reported geometry parameters as they would not represent the underlying *ab initio* values accurately due to loss of precision caused by the fitting of the DBOC surfaces.

	Δr_e	$\Delta \theta_e$
H_2^{17}O	$<10^{-6}$	$<10^{-3}$
H_2^{18}O	$<10^{-6}$	$<10^{-3}$
D_2^{16}O	-0.000 019	-0.010

$3 \times 10^{-5}\,\text{\AA}$ and 0.02° . It is also clear, and it reflects simple intuition, that the effect of H to D substitution is much larger than that accompanying the ^{16}O to ^{18}O change. Note also that the intrinsic adiabatic contributions to the $r(^{16}\text{OH})$ and $r(^{16}\text{OD})$ equilibrium bond lengths are positive (Table II) and negative (Table IV), respectively, and on the order of $10^{-5}\,\text{\AA}$. Therefore, this intrinsic adiabatic effect is about two orders of magnitude smaller than the easily explicable substantial negative $r_0(\text{OH}) \rightarrow r_0(\text{OD})$ change in the vibrationally averaged distance.

Note, finally, that at the all-electron augmented correlation consistent polarized core-valence quadruple zeta (aug-cc-pCVQZ) coupled cluster singles, doubles (triples) [CCSD(T)] level, which gains its usual very high accuracy^{73,74} from favorable error compensation, the optimized structural parameters, $r_e^{\text{BO}}=0.958\,10\,\text{\AA}$ and $\theta_e^{\text{BO}}=104.481^\circ$, deviate very little, $3 \times 10^{-4}\,\text{\AA}$ and 0.004° , from the ultimate r_e^{BO} and θ_e^{BO} of this study.

Overall, we feel confident that the *ab initio* structural parameters of water, based on the CVRQD PESs of Ref. 42, should be nearly exact in the Born–Oppenheimer and adiabatic limits.

V. ROTATIONAL EIGENVALUES

As mentioned above, ultimately the accuracy of the *ab initio* equilibrium structure(s) of water can be probed most directly and meaningfully by computing the observable rotational levels with the aid of the related PESs computed at the same level of theory.

The CVRQD adiabatic PESs reproduce excellently the lowest rotational levels for the isotopologs of water considered, H_2^{16}O , H_2^{17}O , H_2^{18}O , and D_2^{16}O . The levels corresponding to the rotational quantum numbers $J=1$ and 2 are collected in Table III and exemplify this statement. The $J=1$ term values deviate from the experimentally determined ones^{75–78} by less than $0.001\,\text{cm}^{-1}$. The maximum deviation for the $J=2$ CVRQD term values is $0.003\,\text{cm}^{-1}$. The related mean and maximum deviations for the fitted potential of Ref. 79, obtained through elaborate adjustment of the CVRQD PESs to all observed rovibrational levels of several isotopologs of water to better than $0.05\,\text{cm}^{-1}$, are very similar. The minuscule differences between the CVRQD *ab initio* and the fitted rotational term values should not be regarded

as estimates of the uncertainty in the *ab initio* prediction of the structural parameters of the water isotopologs as (a) adjustment of the CVRQD PESs was performed with equal weights for low- and high-energy regions and thus the fit was affected very little by the lowest part of the PES; and (b) as part of this work, direct inclusion of linear terms in the PES, representing changes in the equilibrium structure of water, was tested with using a variety of spectroscopic data; the fits all proved to be divergent. The latter statement holds not only for the adjustment of the CVRQD PESs but also for the best previous spectroscopic PES of H_2^{16}O .³² Furthermore, note that many of the $J=2$ CVRQD and fitted variational results bracket the experimental term values. This means that it is highly unlikely that any fitting attempt to reproduce the rotational term values even more accurately by a fitted PES will result in any significant change in the adiabatic equilibrium structures of water. To wit, 201 experimental levels for the (000), (100), (010), and (001) states of H_2^{16}O , H_2^{17}O , and H_2^{18}O were included in a separate fit that resulted in a standard deviation of only $0.0035\,\text{cm}^{-1}$. Since no linear terms were included in the fit, the related adiabatic equilibrium structural parameters are the same as in the CVRQD PESs, namely, $r_e^{\text{ad}}(^{16}\text{OH})=r_e^{\text{ad}}(^{17}\text{OH})=r_e^{\text{ad}}(^{18}\text{OH})=0.957\,85\,\text{\AA}$ and $\theta_e^{\text{ad}}(\text{H}^{16}\text{OH})=\theta_e^{\text{ad}}(\text{H}^{17}\text{OH})=\theta_e^{\text{ad}}(\text{H}^{18}\text{OH})=104.50_0^\circ$.

Precision of the *ab initio* prediction of higher J transitions by the CVRQD PESs is exceptionally good, as well.

VI. SPECTROSCOPIC EQUILIBRIUM STRUCTURES

Vibrationally averaged effective rotational constants are the principal structural results obtained from fitting of appropriate effective rovibrational Hamiltonians^{80–86} to spectroscopic data. One can recover these effective constants from theoretical rovibrational computations basically in two ways. The traditional route goes through second-order vibrational perturbation theory (VPT2) formulas and the lowest-order vibration-rotation interaction constants α_i , obtained from a cubic force field expansion of the PES.^{35,87–91} The difference between the effective and the equilibrium rotational constants is approximated in this indirect route as a sum of vibration-rotation interaction constants. The resulting spectroscopic equilibrium structural parameters are denoted here collectively as r_e^{sp} and θ_e^{sp} , though, as detailed below, several meaningful variants can be defined. Note that centrifugal distortion terms need to be considered before the experimental rotational constants are converted into ground-state moments of inertia for structure analysis.^{81,85} These corrected rotational constants are usually referred to as A' , B' , and C' . In the alternative, direct route one computes the effective rotational constants as expectation values employing vibrational wave functions from converged variational nuclear motion calculations. It is important to emphasize that pure vibrational wave functions for triatomic molecules can be determined with little numerical effort.^{43,45} Following the second route provides the *vibrational* difference between effective and equilibrium rotational constants. This second, direct route is recommended for computation of vibrationally averaged rotational constants whenever its use is feasible. To

obtain effective rotational constants following the second route one should use coordinates given in the Eckart system of axes since effective rovibrational Hamiltonians employed during fitting of experimental spectra are derived this way.^{92,93} In this case the inertia tensor \mathbf{I} is not diagonal, and the rotational constants are calculated by vibrational averaging of the diagonal elements of \mathbf{I}^{-1} . Consideration of the Coriolis contribution results in change in only one of the rotational constants, C .⁹⁴ The significant difference between the principal axes system (PAS) and Eckart coordinate system vibrationally averaged rotational constants for water, employing up to sextic force field representations of the PES, has recently been demonstrated.⁴³ First-principles vibrationally averaged rotational constant results of this study, obtained in the coordinates in the Eckart system and by inclusion of the Coriolis contribution, are given in Table V.

It is instructive to compare the vibrational corrections obtained from the direct variational approach to those computed from the indirect, traditional approach, based on VPT2. To facilitate this comparison we computed the VPT2 vibrational corrections to the ground-state rotational constants of H_2^{16}O from the best, empirically adjusted *ab initio* vibration-rotation interaction constants of Ref. 95, corresponding to a refined anharmonic force field and thus representing an excellent set of physically meaningful constants obtained through VPT2. These values as well as the spectroscopic results from Ref. 25 are also given in Table V. By comparing the spectroscopic and the *ab initio* ΔA , ΔB , and ΔC values, where, e.g., ΔB stands for the difference between the vibrationally averaged and the equilibrium B values, it becomes clear that the spectroscopic values²⁵ are much closer to the high-quality variational predictions of this study than the *ab initio* values derived from the vibration-rotation interaction constants of Ref. 95. In fact, the largest discrepancy between the spectroscopic²⁵ and the present *ab initio* CVRQD values is 560 MHz for ΔA , while it is 3270 MHz between the spectroscopic and the perturbational (VPT2) *ab initio* results. Note that for prototypical semirigid molecules, for which there is at least two orders of magnitude difference between vibration-rotation interaction constants of consecutive order, the vibration-rotation interaction constants can be computed straightforwardly from the vibrationally averaged rotational constants of the lowest vibrational states. However, for water, for which the corresponding difference is less than tenfold, this procedure does not result in accurate lowest-order vibration-rotation interaction constants.

The computed and measured ground-state inertia defects (Δ_{pl}^0) of the planar water molecule exemplify further the difficulties in comparing measured and computed effective constants (Table V). The inertia defect of $\Delta_{\text{pl}}^0 = 0.050\,64\, \text{u}\text{\AA}^2$ corresponds to rotation-vibration constants obtained in Ref. 25. The inertia defect corresponding to one of the most recent set of effective ground-state rotational constants⁹⁶ is $0.050\,92\, \text{u}\text{\AA}^2$ ($0.0515\, \text{u}\text{\AA}^2$ with the uncorrected rotational constants). If the rotation constants are not corrected for distortion terms, the corresponding inertia defect is $0.051\,49\, \text{u}\text{\AA}^2$, in good agreement with the above number. However, if the rotation constants are corrected for distortion terms, the inertia defect becomes substantially smaller,

TABLE V. Effective ground-state rotational constants (A_0 , A'_0 , B_0 , B'_0 , C_0 , and C'_0 , in MHz), ground-state inertia defects (Δ_{pl}^0 , in $\text{u}\text{\AA}^2$), and differences between effective and equilibrium rotational constants (ΔA , ΔB , and ΔC , in MHz), in the ground vibrational states of symmetric isotopologs of water. $1\, \text{cm}^{-1} = 2.997\,924\,58 \times 10^4\, \text{MHz}$ and $h/8\pi^2c = 505\,379\, \text{MHz}\, \text{u}\text{\AA}^2$. $\Delta_{\text{pl}}^0 = I_c - I_a - I_b$, where I_α , $\alpha = a, b, c$, is the appropriate moment of inertia. Atomic masses have been employed in the calculations. Note that during the refinement of the H_2^{16}O PES $m(^{16}\text{O}) = 15.990\,726\, \text{u}$ has been employed.

Isotopomer	H_2^{16}O	H_2^{18}O	D_2^{16}O
Measured [spectroscopy (Refs. 82–86)] ^{a,b}			
A_0	835 839.9	825 367.32	462 292.4
A'_0	835 783.3		
B_0	435 354.5	435 353.81	217 979.9
B'_0	435 044.5		
C_0	278 133.3	276 950.50	145 303.3
C'_0	278 446.9		
Δ_{pl}^0	0.048 64	0.051 64	0.066 43
ΔA	14 960		4887
ΔB	−2248		−809
ΔC	−7165		−2698
Computed [CVRQD PES (Ref. 42)]			
A_0	835 390.0	824 928.7	462 097.8
ΔA	14 400.4	14 162.2	5 463.5
B_0	434 825.1	434 835.4	217 894.7
ΔB	−2286.1	−2272.8	−866.3
C_0	278 699.4	277 502.6	145 436.5
ΔC	−6543.1	−6494.4	−2467.7
Δ_{pl}^0	0.046 13	0.046 31	0.061 88
Computed [Fitted PES (Ref. 79)]			
A_0	835 433.6	824 971.1	462 006.6
ΔA	14 444.0	14 204.5	5390.5
B_0	434 825.2	434 835.5	217 917.4
ΔB	−2286.0	−2272.7	−835.0
C_0	278 703.3	277 506.5	145 438.0
ΔC	−6539.1	−6490.5	−2460.3
Δ_{pl}^0	0.046 13	0.046 31	0.061 87
Computed [VPT2 (Ref. 95)] ^c			
ΔA	11 690		
ΔB	−2700		
ΔC	−7050		
Δ_{pl}^0	0.046 31		

^a A'_0 , B'_0 , and C'_0 values are only available for H_2^{16}O . For the other isotopologs contributions from the centrifugal distortion terms have not been removed, hindering direct comparison with the computed values.

^bThe ΔA , ΔB , and ΔC values reported correspond to vibration-rotation interaction constants taken from Ref. 25. The calc. II ΔA , ΔB , and ΔC values of Kuchitsu and Bartell (Ref. 26) are 10 500, −3300, and −7800 MHz for H_2^{16}O and 4500, −1200, and −2700 MHz for D_2^{16}O .

^cVPT2=second-order vibrational perturbation theory. The underlying vibration-rotation interaction constants correspond to Set III of Table IV of Ref. 95.

$0.048\,64\, \text{u}\text{\AA}^2$.⁸² The computed CVRQD and VPT2 values are $0.046\,13$ and $0.046\,31\, \text{u}\text{\AA}^2$, respectively. At the same time the underlying computed CVRQD and VPT2 ΔA and ΔC constants deviate from each other substantially. The CVRQD ΔA and the VPT2 ΔC constants are close to their respective spectroscopic counterparts while the remaining two constants disagree by as much as 10%–20%. Note that for water the largest contribution to Δ_{pl}^0 , almost 90%, comes from ΔC . The sizable differences between the VPT2 and the variational CVRQD ΔA , ΔB , and ΔC values call attention to the fact that the accuracy of semispectroscopic equilibrium structures, i.e., those obtained through corrections of experi-

TABLE VI. Spectroscopic equilibrium structural parameters for two isotopologs of water obtained using different ground-state rotational constants (A'_0 and B'_0), rotational constant corrections (ΔA and ΔB), and atomic masses.

A'_0, B'_0	$\Delta A, \Delta B$	H_2^{16}O		D_2^{16}O	
		r_e	θ_e	r_e	θ_e
CVRQD	CVRQD	0.957 854	104.500	0.957 834	104.490
Expt. ^a	CVRQD	0.957 618	104.499	0.957 567	104.483
Expt. ^a	Expt. ^b	0.957 766	104.483		

^aReference 82 for H_2^{16}O and Ref. 85 for D_2^{16}O , see Table V for the numerical values. No correction for centrifugal distortion effects is made in the case of the D_2^{16}O rotational constants, so they do not correspond to A'_0 and B'_0 . Only the A_0 and B_0 combination is used here as these rotational constants are unaffected by the Coriolis effect (for further discussion see Tables 13.10 and 13.18 of Ref. 36).

^bReference 25.

mental rotational constants by *ab initio* VPT2 vibration-rotation interaction constants, an approach^{37,89,97} gaining popularity, might be compromised when high accuracy is sought, on the order of 10^{-4} Å for bond lengths, for small and light species.

There are two plausible explanations for the small but significant remaining discrepancies observed for the measured, the variationally computed, and the VPT2 A_0 , B_0 , and C_0 constants of Table V. The first one recalls that the spectroscopic constants, though apparently of very high precision, are phenomenological in nature and incorporate several “small” effects. It is clearly demonstrated by the data of Table V that ΔA , ΔB , and ΔC corrections computed from an excellent cubic force field representation of the PES of water⁹⁵ are rather inaccurate. It is also clear that centrifugal distortion effects are extremely large in water.^{36,82} The other explanation suggests that the remaining differences between the computed CVRQD and the spectroscopic ground-state rotational constants are due to shortcomings of the theory, related principally to three factors. First, though the CVRQD PESs are of very high accuracy, they are still not the ultimate adiabatic PESs of water isotopologs and minor improvements of the adiabatic potentials are feasible. Second, no account of the presence of electrons in the molecule is given, coupling of the electronic angular momentum with the overall rotation of the nuclear framework⁹⁸ has not been considered and, consequently, no electronic contributions to the effective rotational constants have been computed. Third, it must be investigated whether nonadiabatic effects,^{65,66} not considered so far explicitly and the hardest to ascertain, have a noticeable contribution.

To investigate the first possible theoretical shortcoming the latest semispectroscopic, fitted PES of water⁷⁹ has been employed in the expectation value calculations with results presented in Table V. The smallness of the adjustments to the lower part of the fully *ab initio* CVRQD PES is demonstrated repeatedly in this study by the excellent agreement between the CVRQD and the fitted effective rotational constant results. The maximum unsigned deviation is only 90 MHz (0.003 cm^{-1}), while the mean deviation is only on the order of 20 MHz. The small changes in the ground-state rotational constants are due to the slightly different form of the wave function (see Sec. V). Therefore, further improvement of the *ab initio* PES of Ref. 42 through fitting to ob-

served transitions does not seem to hold promise in obtaining significant changes in the values of the rotational constants.

Next, let us investigate the effects due to the presence of electrons in the molecule. They can be approximated most easily through changes in the masses of the nuclei, most importantly that of H, in the nuclear motion calculations. Two related comments. First, using atomic masses $M^\beta = m^\beta + Z^\beta m_e$ or scaled atomic masses⁹⁹ in computations, where m^β and Z^β are the nuclear mass and atomic number of atom β , respectively, and m_e is the electronic mass, corresponds to absorbing some part of the nonadiabatic effect into the adiabatic approximation. Second, atomic rather than nuclear masses are employed in the analysis of molecular spectra. Table VI contains equilibrium distances and angles of water isotopologs obtained from using either experimental A'_0 and B'_0 or CVRQD ground-state effective rotational constants corrected with the seemingly dependable CVRQD or experimental ΔA and ΔB constants. (ΔA and ΔB constants do not depend noticeably on whether nuclear or atomic masses were used during their evaluation.) The results obtained show that the use of experimental versus CVRQD constants changes the equilibrium structural parameters substantially, $r_e(^{16}\text{OH})$ by up to $0.000\,24$ Å and $\theta_e(\text{H}^{16}\text{OH})$ by up to 0.017° . The best spectroscopic equilibrium structure of water is $r_e^{\text{sp}} = 0.957\,77$ Å and $\theta_e^{\text{sp}} = 104.48^\circ$.

As to electronic contributions related to the rotational g factor,^{36,98} they are usually regarded to be small. This general wisdom can be checked for water as it is one of the few polyatomic systems for which the electronic effect has been known for quite some time^{100,101} and the underlying⁹⁸ rotational g -tensor elements have been verified repeatedly.^{102,103} Based on the data in Refs. 100 and 101, the electronic corrections to $\text{H}_2^{16}\text{O}(\text{D}_2^{16}\text{O})$ are as follows: $\Delta_A^{\text{el}} = 294(91)$ MHz, $\Delta_B^{\text{el}} = 168(42)$ MHz, and $\Delta_C^{\text{el}} = 99(26)$ MHz. The corrections are considerably smaller for D_2^{16}O than for H_2^{16}O since they basically scale linearly both with the g factors and with the effective values of the rotational constants. This fact is partially responsible for the apparently higher accuracy of the effective rotational constant predictions of D_2^{16}O as compared to H_2^{16}O . These corrections, when added to the computed CVRQD A_0 and B_0 constants, not affected by Coriolis effects, largely bridge the experimental–computational deviations. To wit, for $\text{H}_2^{16}\text{O}(\text{D}_2^{16}\text{O})$ the A_0 deviations decrease from

393(195) to 99(104) MHz, while the B_0 deviations decrease from 219(85) to 51(43) MHz. On the other hand, the electronic correction increases the deviation between the measured and computed results for C_0 . Note that correcting the rotational constants for electronic effects changes $r_e(^{16}\text{OH}/^{16}\text{OD})$ by $-0.000\,24/-0.000\,09$ Å and $\theta_e(\text{H}^{16}\text{OH}/\text{D}^{16}\text{OD})$ by $-0.017/-0.000^\circ$.

In summary, none of the factors considered so far seem to be able to eliminate completely the gap between the variationally computed theoretical and the spectroscopic ground-state rotational constants. Iterative refinement of the CVRQD *ab initio* PES through fitting to observed rovibrational levels does not seem to be able to reduce the discrepancies between computed and experimental rotational constants. Consideration of the electronic effects does not make the discrepancies between the *ab initio* and the spectroscopic ground-state rotational constants vanish though serves to reduce them substantially. Nevertheless, combination of these effects, especially centrifugal distortion correction of the experimental effective rotational constants and electronic effects on the computed rotational constants, makes the apparent discrepancies between the exceedingly high-quality *ab initio* rotational constant predictions and experiment comfortably small.

Finally, one must address the effect of nonadiabaticity^{38–40,104–107} on the equilibrium structure of water, though at present time, apart from H_3^+ and perhaps water, this can be done only for diatomics with substantial confidence.

It has been shown for light diatomic hydrides, namely, for LiH, BeH, BH, and CH^+ (Refs. 38, 40, and 105) that the nonadiabatic effect, defined in a somewhat strange sense as the difference between the adiabatic equilibrium bond length and that obtained from spectroscopy, i.e., from the experimental fitted effective molecular rotational constant, is substantial for the equilibrium bond lengths. In fact, for BH and CH^+ the nonadiabatic correction is estimated^{38,40} this way to be large and similar in magnitude, $+0.002\,34$ and $+0.002\,75$ Å, respectively, while the well-defined adiabatic corrections appear to be considerably smaller, on the order of $+0.000\,65$ Å. For the heavier hydrides the nonadiabatic distance correction seems to decrease in magnitude, e.g., for NH it is estimated to be less than $0.000\,70$ Å.⁴⁰ Nevertheless, as widely appreciated, the adiabatic correction decreases just as fast or faster, for NH the adiabatic correction on r_e^{BO} is estimated to be $+0.000\,27$ Å.⁴⁰ Thus, based on the results on these diatomic paradigms one can assume that the adiabatic [diagonal Born–Oppenheimer corrections (DBOC)] effect is small for r_e of the diatomic radical OH and similarly in H_2O . This is supported by the results in Table II, where the adiabatic correction for r_e of $\text{H}_2^{16}\text{O}(\text{D}_2^{16}\text{O})$ is only $+0.000\,03$ ($+0.000\,01$) Å.

One somewhat striking result for $r_e(\text{OH})$ of water must be mentioned. For the pairs XH/XD, with $X=\text{Be}, \text{B}, \text{C}$, and N , it has been clearly established that $r_e^{\text{ad}}(\text{XH}) > r_e^{\text{ad}}(\text{XD})$.^{38–40} For example, $r_e^{\text{ad}}(^{12}\text{CH}) = 1.118\,06$ Å and $r_e^{\text{ad}}(^{12}\text{CD}) = 1.117\,92$ Å,¹⁰⁸ whereas $r_e^{\text{BO}}(^{12}\text{CH})$ appears to be³⁹ $1.117\,77$ Å. The present calculations support this trend, in water $r_e^{\text{ad}}(^{16}\text{OH}) = 0.957\,85$ Å versus $r_e^{\text{ad}}(^{16}\text{OD}) = 0.957\,83$ Å.

For the spectroscopic equilibrium bond lengths the trend for the XH species appears to be the same, for example,^{39,108,109} $r_e^{\text{sp}}(^{12}\text{CH}) = 1.119\,79$ Å versus $r_e^{\text{sp}}(^{12}\text{CD}) = 1.118\,88$ Å. Nevertheless, for r_e^{sp} one finds an opposite result for the OH radical,¹¹⁰ $r_e^{\text{sp}}(^{16}\text{OH}) = 0.969\,63$ Å $<$ $r_e^{\text{sp}}(^{16}\text{OD}) = 0.969\,68$ Å, i.e., the ^{16}OD correction is $+0.000\,05$ Å. As Table VI shows, in water $r_e^{\text{sp}}(^{16}\text{OD}) < r_e^{\text{sp}}(^{16}\text{OH})$ in all cases considered. This apparent discrepancy can be resolved by noting that the spectroscopic data for OH and OD (Ref. 110) were determined in different ways. New experimental work^{111,112} and perhaps an accurate computational determination of adiabatic corrections to r_e should help clarifying this issue. In summary, the nonadiabatic effect on $r_e(^{16}\text{OH})$ seems to be significant, as can be judged by comparing $r_e^{\text{ad}}(^{16}\text{OH}) = 0.957\,85$ Å and $r_e^{\text{sp}}(^{16}\text{OH}) = 0.957\,77$ Å. No similar result can be derived for $r_e(^{16}\text{OD})$ as no corresponding experimental A'_0 and B'_0 rotational constants are available to us.

The diatomic paradigms provide no estimate for the adiabatic equilibrium bond angle correction. Fortunately, there seems to be very little uncertainty (cf. Table VI) in the equilibrium bond angle of the water isotopologs. The changes in θ_e occurring upon using different rotational constants and correction factors are minuscule, the data presented in Table VI suggest that we do know θ_e of water for almost five significant digits.

VII. CONCLUSIONS

Equilibrium structures are fundamental entities in molecular sciences yet they are inferred from experimental data by complicated procedures which often rely on several assumptions, including the Born–Oppenheimer approximation. Theory provides a direct route to equilibrium geometries. In this paper, on the example of water, we have explored the direct computational route to the various equilibrium structures of nonlinear polyelectronic and polyatomic systems. In accordance with the well-established diatomic paradigms,¹⁰⁴ three equilibrium structure types have been deduced, namely, Born–Oppenheimer (mass-independent), adiabatic (mass-dependent), and spectroscopic (nonadiabatic) structures. The former two equilibrium structure types have a clear physical meaning, they are based on the concept of potential-energy surfaces. Spectroscopic equilibrium structures are less well defined and they correspond to effective spectroscopic constants, derived by fitting an effective Hamiltonian, preferably that given by Watson,⁸⁰ to appropriate regions of the rovibrational spectrum of water and corrected for centrifugal and perhaps electronic distortion effects.

It is shown that state-of-the-art electronic and nuclear motion calculations, when allowance is made for their coupling, are capable of producing equilibrium structures for polyatomic systems somewhat more accurate than any of the experimental/empirical procedures. In particular, the mass-independent (Born–Oppenheimer) equilibrium structure of the ground electronic state of water is found to be $r_e^{\text{BO}}(\text{OH}) = 0.957\,82$ Å and $\theta_e^{\text{BO}}(\text{HOH}) = 104.485^\circ$. The concept of mass-independent equilibrium structures seems to be valid to about 3×10^{-5} Å and 0.02° for water. The adiabatic, i.e., mass-dependent, equilibrium structural parameters are

$r_e^{\text{ad}}(^{16}\text{OH})=0.957\,85\text{ \AA}$ and $\theta_e^{\text{ad}}(\text{H}^{16}\text{OH})=104.500^\circ$ for H_2^{16}O and $r_e^{\text{ad}}(^{16}\text{OD})=0.957\,83\text{ \AA}$ and $\theta_e^{\text{ad}}(\text{D}^{16}\text{OD})=104.490^\circ$ for D_2^{16}O . It is believed that these equilibrium structural parameters should be nearly exact both in the Born–Oppenheimer and adiabatic limits, conservative error limits are $\pm 0.000\,10\text{ \AA}$ and $\pm 0.010^\circ$.

Our fully *ab initio* adiabatic CVRQD PESs,⁴² incorporating equilibrium geometries some of which are reported above, reproduce the $J=1$ and 2 rotational term values of four isotopologs of water with an average accuracy of 0.002 cm^{-1} . A laborious adjustment of the *ab initio* CVRQD PESs, with equal weights for the low- and high-lying rovibrational states, results in only a minuscule improvement of these rotational frequency predictions. Inclusion of linear terms of the PES in the fitting, representing changes in the equilibrium structure of water, resulted in divergent refinement attempts. These observations indicate that refinements of the presently available high-accuracy *ab initio* adiabatic PESs of water,⁴² even if the refinement would be geared toward the lowest-lying levels, are unlikely to offer improvements on the underlying adiabatic equilibrium structural parameters. Adiabatic effects are small for the structural parameters of water. Nonadiabatic effects seem to be larger than the adiabatic ones for the structures of isotopologs of water. Therefore, the inverse route to Born–Oppenheimer or adiabatic equilibrium geometries is seriously compromised.

Another possibility for judging the computed adiabatic equilibrium geometries and the CVRQD PESs is offered by comparison of computed and experimental vibrationally averaged rotational constants. Spectroscopic vibration-rotation interaction constants, though apparently of very high precision, are somewhat phenomenological in nature and do not correspond very closely to the vibration-rotation interaction constants that can be obtained from a cubic force field representation of the PES via VPT2. The sizable differences between the best *ab initio* and the spectroscopic ΔA , ΔB , and ΔC values call attention to the fact that the accuracy of semi-spectroscopic equilibrium structures obtained through corrections of spectroscopic rotational constants by *ab initio* vibration-rotation interaction constants might be compromised, at least for small and light species.

The quality of our results suggests that more studies of a similar nature are to be performed if really accurate values of Born–Oppenheimer, adiabatic, and spectroscopic equilibrium geometry parameters are needed. This recommendation is based on the fact that while the indirect, (semi)empirical route cannot eliminate all errors, the direct route, when feasible, is successful: theory is now capable of determining physically relevant adiabatic equilibrium structures, in the present case those of water, more accurately than can be meaningfully derived from available experimental data.

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