

# A Direct Evaluation of the Partition Function and Thermodynamic Data for Water at High Temperatures

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The rovibrational partition function of the water molecule is calculated using a classical statistical mechanics approach and a hybrid method recently proposed by Prudente et al. [*J. Phys. Chem. A* 2001, 105, 5272], which corrects the classical results. The phase-space integrals are solved using a Monte Carlo technique. For temperatures between 500 and 6000 K, the results are compared with previous approximate and exact quantum calculations. Estimates of some thermodynamic quantities for gas-phase water as a function of temperature are also reported and compared with previous results. The calculated partition function, Gibbs enthalpy, Helmholtz function, entropy, and specific heat at constant pressure indicate that the hybrid scheme can provide accurate thermodynamic data for polyatomic molecules at high temperatures.

## 1. Introduction

Accurate thermodynamic data of gas-phase polyatomic systems are of great importance in chemistry and physics. In principle, the partition function, and hence other thermodynamic properties, can be evaluated exactly in quantum statistical mechanics by summing directly over all of the energy levels of the system (so-called sum-over-states). Although an enormous advance has been made in recent years along this line of approach, the calculation of rovibrational states is currently feasible only for systems with a few degrees of freedom,<sup>1–5</sup> which limits the applicability of the sum-over-states approach to small molecules.<sup>6–9</sup> To overcome this problem, the partition function and related thermodynamic data have been traditionally calculated by fitting effective Hamiltonians to experimental data.<sup>10–13</sup> However, the accuracy of the results obtained by using the traditional methods is expected to be poor at high temperatures<sup>8,14</sup> and for floppy (anharmonic) systems.<sup>15</sup>

For the above reasons, several procedures have been proposed as routes to the direct sum-over-states approach and fitting of experimental data. These include the hybrid analytic/direct summation method of ab initio calculations,<sup>14</sup> Fourier path–integral Monte Carlo methods,<sup>16–18</sup> and classical statistical mechanics (CSM) methods both with consideration of quantum,<sup>19</sup> semiclassical,<sup>20</sup> and semiempirical<sup>21</sup> corrections and without consideration of such corrections.<sup>15,22–25</sup> In a previous paper,<sup>26</sup> we surveyed briefly the most popular classical methods, which employ corrections of various types, and proposed a novel scheme (hybrid LCP/QFH), which consists of adding an effective potential to the classical Hamiltonian to mimic quantum effects. Such a method blends the advantages of the linear classical path<sup>27</sup> (LCP) and the quadratic Feynman–Hibbs<sup>28</sup> (QFH) methods while avoiding their undesirable features. In fact, preliminary calculations for diatomic molecules<sup>26</sup> have shown that the hybrid LCP/QFH method performs generally better than previous approaches for moderate and high temperatures.

A major goal of this work is to extend the hybrid LCP/QFH calculations of the rovibrational partition function and thermodynamic properties [e.g. the Gibbs enthalpy function (*gef*), Helmholtz function (*hcf*), entropy (*S*), and specific heat capacity at constant pressure (*C<sub>p</sub>*)] of gas-phase triatomic systems described by realistic potential energy surfaces. Thus, we envisage a simple and relatively inexpensive computational scheme amenable to generalization to multidimensional systems and that can provide accurate internal partition functions (and other thermodynamic data) for such polyatomic systems. Conversely to previous work,<sup>15,23,26,29</sup> the multidimensional phase-space integrals that appear in the classical formalism will be evaluated by using a crude Monte Carlo method to sample the coordinates, while the Barker Monte Carlo algorithm<sup>30</sup> will be used to sample the conjugate momenta (see later).

As a case study, we consider the H<sub>2</sub>O molecule in its ground electronic state. Indeed, water is the most common polyatomic molecule in the universe, being fundamental to life and an essential constituent of the Earth's atmosphere. Its thermodynamic data at high temperatures is therefore of great importance for modeling combustion, exhaust gases, and the atmosphere of cool stars, just to mention a few examples. Moreover, the H<sub>2</sub>O molecule is representative of molecular systems with a deep potential well and commonly plays the role of a benchmark system both for bound-state and for reactive scattering calculations [O(<sup>1</sup>D) + H<sub>2</sub> reaction]. It is also known for the poor results of the partition functions that are obtained even at moderate and high temperatures when using CSM.<sup>20,23</sup> Because of relatively simple and fundamental characteristics of water, there are many predictions of its partition function and thermodynamic properties,<sup>7,8,10–14,18,20,23,31</sup> including the more recent and accurate estimation obtained by Vidler and Tennyson.<sup>32</sup> This will allow a detailed test of our method.

The paper is organized as follows. In section 2, we discuss the calculation of the internal partition function and related thermodynamic properties by using the standard classical statistical mechanics and hybrid LCP/QFH methods and summarize the Monte Carlo procedure utilized to evaluate the involved multidimensional phase-space integrals. The details of the calculations and results are presented in section 3, while some conclusions are in section 4.

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## 2. Methodology

**2.1. General.** The molecular partition function is usually expressed as

$$Q = Q_{\text{tr}} Q_{\text{elec}} Q_{\text{rovib}} \quad (1)$$

where  $Q_{\text{tr}}$ ,  $Q_{\text{elec}}$ , and  $Q_{\text{rovib}}$  are the translational, electronic, and rovibrational contributions, respectively. Although  $Q_{\text{tr}}$  can be calculated using the ideal gas formalism<sup>33</sup> and  $Q_{\text{elec}}$  can be assumed to be unity (because no electronic excited states are involved;<sup>8,14</sup> for a discussion on this issue, see ref 34),  $Q_{\text{rovib}}$  has to be evaluated from the potential energy surface by using quantum statistical mechanics or an approximate procedure. In this work, we use two formulations based on classical statistical mechanics. In the first approach,  $Q_{\text{rovib}}$  assumes the standard classical form<sup>33,35</sup>

$$Q_{\text{rovib}}^{\text{CM}}(T) = \frac{\exp(\beta\epsilon_0)}{h^n} \int \int_{\mathcal{B}} \exp[-\beta H^{\text{CM}}(\mathbf{q}, \mathbf{p})] \, d\mathbf{q} \, d\mathbf{p} \quad (2)$$

where  $H^{\text{CM}}(\mathbf{q}, \mathbf{p})$  is the classical Hamiltonian,  $\epsilon_0$  is the zero-point energy of the system,  $\beta = 1/(k_{\text{B}}T)$ ,  $k_{\text{B}}$  is the Boltzmann constant,  $T$  is the temperature,  $h$  is the Planck constant,  $n$  is the number of degrees of freedom,  $\mathbf{q}$  is the generalized coordinate vector, and  $\mathbf{p}$  is the corresponding conjugate momenta. In the second approach, we employ the hybrid LCP/QFH method, which corrects the classical rovibrational partition function by adding an effective potential to the classical Hamiltonian. One has<sup>26</sup>

$$Q_{\text{rovib}}^{\text{LCP/QFH}}(T) = \frac{\exp(\beta\epsilon_0)}{h^n} \int \int_{\mathcal{B}} \exp\{-\beta[H^{\text{CM}}(\mathbf{q}, \mathbf{p}) + V^{\text{eff}}(\mathbf{q})]\} \, d\mathbf{q} \, d\mathbf{p} \quad (3)$$

where the effective potential is given by

$$V^{\text{eff}}(\mathbf{q}) = \beta A \nabla^2 V(\mathbf{q}) + \beta^2 A (\nabla \cdot V(\mathbf{q}))^2 \quad (4)$$

with  $A = \hbar^2/(48\mu)$ . The first term ( $\times 2$ ) of eq 4 is the quadratic Feynman–Hibbs (QFH) approximation<sup>28</sup> of the Feynman path integral formulation, while the second one ( $\times 2$ ) is the linear classical approximation (LCP) due to Miller.<sup>27</sup> We have shown in a previous paper<sup>26</sup> that the QFH method generally underestimates the values of the quantum partition function, while the LCP approximation overestimates the values at low and moderate temperatures. Note that the subscript  $\mathcal{B}$  in eqs 2 and 3 implies that the hypervolume of integration is restricted to phase-space regions corresponding to a bound-state situation:<sup>36</sup>  $0 \leq H^{\text{CM}}(\mathbf{q}, \mathbf{p}) \leq D_e$ , where  $D_e$  is the classical dissociation energy of the molecule with the minimum of the potential energy surface assumed as the reference energy. In turn, the factor  $\exp(-\beta\epsilon_0)$  in eqs 2 and 3 is required to compare with previous results for water, which have been calculated by assuming the zero-point energy ( $\epsilon_0$ ) as the reference energy.

A temperature-dependent estimate of the thermodynamic quantities considered in this work can be obtained from the partition function and its first and second moments, the rovibrational contributions of which are defined by<sup>14</sup>

$$Q'_{\text{rovib}} = T \frac{dQ_{\text{rovib}}}{dT} \quad (5)$$

$$Q''_{\text{rovib}} = T^2 \frac{d^2 Q_{\text{rovib}}}{dT^2} + 2Q'_{\text{rovib}} \quad (6)$$

where the rovibrational partition function is expressed as in eqs 2 or 3. In principle, the moments could be obtained through numerical differentiation of  $Q_{\text{rovib}}^{\text{CM}}$  or  $Q_{\text{rovib}}^{\text{LCP/QFH}}$ , although it is more advantageous to differentiate them analytically. For the standard classical approach, the resulting expressions are

$$Q'_{\text{rovib}}^{\text{CM}} = \frac{1}{h^n} \int \int_{\mathcal{B}} \beta \tilde{H}^{\text{CM}}(\mathbf{q}, \mathbf{p}) \exp[-\beta \tilde{H}^{\text{CM}}(\mathbf{q}, \mathbf{p})] \, d\mathbf{q} \, d\mathbf{p} \quad (7)$$

$$Q''_{\text{rovib}}^{\text{CM}} = \frac{1}{h^n} \int \int_{\mathcal{B}} [\beta \tilde{H}^{\text{CM}}(\mathbf{q}, \mathbf{p})]^2 \exp[-\beta \tilde{H}^{\text{CM}}(\mathbf{q}, \mathbf{p})] \, d\mathbf{q} \, d\mathbf{p} \quad (8)$$

where  $\tilde{H}^{\text{CM}}(\mathbf{q}, \mathbf{p}) = H^{\text{CM}}(\mathbf{q}, \mathbf{p}) - \epsilon_0$ , while for the hybrid LCP/QFH approach, one has

$$Q'_{\text{rovib}}^{\text{LCP/QFH}} = \frac{1}{h^n} \int \int_{\mathcal{B}} [\beta \tilde{H}^{\text{CM}}(\mathbf{q}, \mathbf{p}) + 2\beta^2 A \nabla^2 V(\mathbf{q}) + 3\beta^3 A (\nabla \cdot V(\mathbf{q}))^2] \exp\{-\beta[\tilde{H}^{\text{CM}}(\mathbf{q}, \mathbf{p}) + V^{\text{eff}}(\mathbf{q})]\} \, d\mathbf{q} \, d\mathbf{p} \quad (9)$$

$$Q''_{\text{rovib}}^{\text{LCP/QFH}} = \frac{1}{h^n} \int \int_{\mathcal{B}} \{[\beta \tilde{H}^{\text{CM}}(\mathbf{q}, \mathbf{p}) + 2\beta^2 A \nabla^2 V(\mathbf{q}) + 3\beta^3 A (\nabla \cdot V(\mathbf{q}))^2]^2 - 2\beta^2 A \nabla^2 V(\mathbf{q}) - 6\beta^3 A (\nabla \cdot V(\mathbf{q}))^2\} \exp\{-\beta[\tilde{H}^{\text{CM}}(\mathbf{q}, \mathbf{p}) + V^{\text{eff}}(\mathbf{q})]\} \, d\mathbf{q} \, d\mathbf{p} \quad (10)$$

The ideal gas thermodynamic functions as a function of temperature can be obtained in terms of  $Q_{\text{rovib}}$ ,  $Q'_{\text{rovib}}$  and  $Q''_{\text{rovib}}$  as follows:

### The Gibbs enthalpy function

$$\text{gef}(T) = -\frac{[G(T) - H_0]}{T} = R \ln Q_{\text{rovib}} + \text{gef}^{\text{tr}}(T) + \frac{H_0}{T} \quad (11)$$

### The Helmholtz function

$$\text{hcf}(T) = H(T) - H_0 = RT \frac{Q'_{\text{rovib}}}{Q_{\text{rovib}}} + \text{hcf}^{\text{tr}}(T) - H_0 \quad (12)$$

### The entropy

$$S(T) = R \frac{Q'_{\text{rovib}}}{Q_{\text{rovib}}} + R \ln Q_{\text{rovib}} + S^{\text{tr}}(T) \quad (13)$$

### The specific heat capacity at constant pressure

$$C_p(T) = R \left[ \frac{Q''_{\text{rovib}}}{Q_{\text{rovib}}} - \left( \frac{Q'_{\text{rovib}}}{Q_{\text{rovib}}} \right)^2 \right] + C_p^{\text{tr}} \quad (14)$$

where  $\text{gef}^{\text{tr}}(T)$ ,  $\text{hcf}^{\text{tr}}(T)$ ,  $S^{\text{tr}}(T)$ , and  $C_p^{\text{tr}}(T)$  are the translational contributions, which, for an ideal gas, assume the form<sup>10</sup>

$$\text{gef}^{\text{tr}}(T) = R \left[ \frac{3}{2} \log M + \frac{5}{2} \log T + \log \left( \frac{k_{\text{B}} (2\pi k_{\text{B}})^{3/2}}{p h^2} \right) \right] \quad (15)$$

$$\text{hcf}^{\text{tr}}(T) = \frac{5}{2} RT \quad (16)$$

$$S^{\text{tr}}(T) = \text{gef}^{\text{tr}}(T) + \frac{5}{2} R \quad (17)$$

$$C_p^{\text{tr}}(T) = \frac{5}{2} R \quad (18)$$

with  $R$  being the gas constant and  $p$  the pressure. The  $H_0$  constant, which appears in  $\text{gef}(T)$  and  $\text{hcf}(T)$ , is the reference enthalpy at the JANAF reference temperature of 298.15 K,<sup>10</sup>

that is,  $H(298.15)$ . Because the results obtained within the classical framework have poor accuracy at low temperatures, we will utilize in our calculations the value of  $H_0 = 9904.1$  J mol<sup>-1</sup> derived by Vidler and Tennyson<sup>32</sup> from both theoretical and experimental energy levels. Such a value is close to the one reported in the JANAF tables ( $H_0 = 9904$  J mol<sup>-1</sup>) and can be compared with the calculated values of Martin et al.<sup>14</sup> and Harris et al.<sup>8</sup> (respectively,  $H_0 = 9902$  J mol<sup>-1</sup> and  $H_0 = 9895.4$  J mol<sup>-1</sup>). Although arbitrary, such a choice leads to errors smaller in magnitude than the statistical errors inherent to our Monte Carlo calculations, even at low temperatures.

**2.2. Classical Hamiltonian and Effective Potential.** The next step consists of obtaining the expressions for the classical Hamiltonian ( $H^{\text{CM}}$ ) and the effective potential ( $V^{\text{eff}}$ ) used to calculate the rovibrational partition function of a system with three structureless particles;  $m_i$  will be the mass of the  $i$ -th particle, and  $\mathbf{X}_i$  will be its position vector with respect to the space-fixed axes. The rovibrational motion of the three particles relative to the center of mass of the system can be described by using mass-weighted Jacobi vectors<sup>37</sup>

$$\mathbf{r} = d^{-1}(\mathbf{X}_3 - \mathbf{X}_2) \quad (19)$$

$$\mathbf{R} = d \left( \mathbf{X}_1 - \frac{m_2 \mathbf{X}_2 + m_3 \mathbf{X}_3}{m_2 + m_3} \right) \quad (20)$$

and their corresponding conjugate momenta  $\mathbf{P}_r$  and  $\mathbf{P}_R$ , which defines a 12-dimensional (12D) phase space. In eqs 19 and 20,  $d$  is the mass scaling or normalizing factor,

$$d = \left[ \left( \frac{m_1}{\mu} \right) \left( 1 - \frac{m_1}{M} \right) \right]^{1/2} \quad (21)$$

and

$$\mu = \left( \frac{m_1 m_2 m_3}{M} \right)^{1/2} \quad (22)$$

is the three-body reduced mass;  $M = m_1 + m_2 + m_3$  is the total mass of the system. With the use of this phase-space coordinates set, the expression of the classical Hamiltonian assumes the form

$$\begin{aligned} H(\mathbf{R}, \mathbf{r}, \mathbf{P}_R, \mathbf{P}_r) &= \frac{\mathbf{P}_R^2}{2\mu} + \frac{\mathbf{P}_r^2}{2\mu} + V(\mathbf{R}, \mathbf{r}) \\ &= \frac{P_R^2}{2\mu} + \frac{P_r^2}{2\mu} + V(R, r, \theta) \end{aligned} \quad (23)$$

where  $R = |\mathbf{R}|$ ,  $r = |\mathbf{r}|$ , and  $\theta = \cos^{-1}(\mathbf{R} \cdot \mathbf{r} / (Rr))$  are the internal mass-weighted Jacobi coordinates,  $P_R = |\mathbf{P}_R|$ , and  $P_r = |\mathbf{P}_r|$ . Note that the classical Hamiltonian depends explicitly only on the variables  $R$ ,  $r$ ,  $\theta$ ,  $P_R$  and  $P_r$ . Note further that the interatomic distances ( $r_{12}$ ,  $r_{13}$ ,  $r_{23}$ ) and internal mass-weighted Jacobi coordinates are related by

$$\begin{aligned} r_{23} &= dr \\ r_{13}^2 &= \left( \frac{m_2 dr}{m_2 + m_3} \right)^2 + \frac{R^2}{d^2} - \frac{2m_2}{m_2 + m_3} rR \cos \theta \\ r_{12}^2 &= \left( \frac{m_3 dr}{m_2 + m_3} \right)^2 + \frac{R^2}{d^2} + \frac{2m_3}{m_2 + m_3} rR \cos \theta \end{aligned} \quad (24)$$

Thus, the phase-space integrals, which appear in the classical rovibrational partition function (without any correction) and the

hybrid LCP/QFH method, as well as on the corresponding moments, can be written in terms of internal mass-weighted Jacobi coordinates and the moduli of the conjugate momenta as follows:

$$\begin{aligned} I &= \frac{1}{h^6} \int \int_{\mathcal{B}} F(\mathbf{R}, \mathbf{r}, \mathbf{P}_R, \mathbf{P}_r) d\mathbf{R} d\mathbf{r} d\mathbf{P}_R d\mathbf{P}_r \\ &= \frac{128\pi^4}{h^6} \int \int_{\mathcal{B}} F(R, r, \theta, P_R, P_r) R^2 r^2 P_R^2 P_r^2 d\mathbf{R} d\mathbf{r} d \\ &\quad (\cos \theta) dP_R dP_r \end{aligned} \quad (25)$$

Such expressions can be obtained by adopting spherical polar coordinates to describe  $\mathbf{R}$ ,  $\mathbf{r}$ ,  $\mathbf{P}$ , and  $\mathbf{p}$  and performing analytically all of the integrals involving coordinates on which the Hamiltonian does not depend explicitly. Note that the multidimensional phase-space integral is then reduced from 12D to 5D. The function  $F(\dots)$  collects the integrands of eqs 2, 3, 7, 8, 9, and 10.

Moreover, for the hybrid LCP/QFH method, one requires the effective potential [eq 4] in terms of internal mass-weighted Jacobi coordinates. After some simple algebra, we can write the terms in  $\beta$  and  $\beta^2$  as

$$\begin{aligned} \nabla^2 V &= \nabla_R^2 V + \nabla_r^2 V \\ &= \frac{\partial^2 V}{\partial R^2} + \frac{2}{R} \frac{\partial V}{\partial R} + \frac{\partial^2 V}{\partial r^2} + \frac{2}{r} \frac{\partial V}{\partial r} + \\ &\quad \left( \frac{1}{R^2} + \frac{1}{r^2} \right) \left( \frac{\partial^2 V}{\partial \theta^2} + \cotg \theta \frac{\partial V}{\partial \theta} \right) \end{aligned} \quad (26)$$

and

$$\begin{aligned} (\nabla V)^2 &= (\nabla_R V)^2 + (\nabla_r V)^2 \\ &= \left( \frac{\partial V}{\partial R} \right)^2 + \left( \frac{\partial V}{\partial r} \right)^2 + \left( \frac{1}{R^2} + \frac{1}{r^2} \right) \left( \frac{\partial V}{\partial \theta} \right)^2 \end{aligned} \quad (27)$$

Finally, to perform the integrations involved in the rovibrational partition function and related thermodynamic quantities, we employ a Monte Carlo procedure, which will be described next.

**2.3. Monte Carlo Approach.** The methods generically classified as Monte Carlo offer one of the most powerful techniques to evaluate multidimensional integrals (e.g., see refs 38 and 39). Examples of their use in chemical physics are the determination of classical partition functions and density of states for molecular systems with realistic potential energy surfaces.<sup>40-49</sup> In previous work,<sup>15,23,26,29</sup> we have used such a method based on an adaptation of the Monte Carlo algorithm originally reported by Barker<sup>30</sup> within the context of transition-state theory. The spirit of such an algorithm is akin to the idea of importance sampling and consists of choosing a sampling domain that coincides as much as possible with the integration domain. Thus, the variables are not sampled independently of each other, but instead some kind of dependence is introduced. This leads to a normalized but nonuniform distribution and, hence, requires the use of appropriate weighting factors (see refs 29 and 30 for details).

However, as pointed out elsewhere,<sup>23</sup> the sampling of the configurational space for systems of which the potential energy surfaces possess two or more minima is not a trivial matter when using the Barker algorithm. In fact, the sampling becomes complicated and time-consuming, which led us to utilize here

the simpler crude Monte Carlo approach to sample the internal mass-weighted Jacobi coordinates ( $R, r, \theta$ ). However, we keep using Barker's algorithm to sample the modulus of the conjugate momenta ( $P_{\mathbf{R}}$  and  $P_{\mathbf{r}}$ ). Note that in the crude Monte Carlo integration the variables are sampled independently of each other using a sequence of pseudorandom numbers, which generate a uniform distribution over the configuration space. As already noted, all multidimensional integrals encountered here have the general form of eq 25, and hence, we summarize below the general procedure adopted to evaluate them. It involves the following steps:

(1) Define a minimum and maximum displacement for each of the internal mass-weighted Jacobi coordinates, namely,  $R^{\min}$ ,  $R^{\max}$ ,  $r^{\min}$ ,  $r^{\max}$ ,  $\theta^{\min}$ , and  $\theta^{\max}$ , so that the sampled hyperrectangular volume defined by these three coordinate intervals includes the true volume of integration (i.e.,  $V(R, r, \theta) \leq D_e$  for  $P_{\mathbf{R}} = P_{\mathbf{r}} = 0$ ) but is as much as possible close to it.

(2) Sample randomly  $R, r$ , and  $\cos(\theta)$  within their range to obtain the values  $R^S, r^S$ , and  $\theta^S$  according to

$$\begin{aligned} R^S &= R^{\min} + (R^{\max} - R^{\min})\xi \\ r^S &= r^{\min} + (r^{\max} - r^{\min})\xi \\ \cos(\theta^S) &= \cos(\theta^{\max}) + (\cos(\theta^{\min}) - \cos(\theta^{\max}))\xi \end{aligned} \quad (28)$$

where  $\xi$  is a random number in the range  $[0, 1]$ .

(3) Calculate the potential at the sampled point,  $V^S = V(R^S, r^S, \theta^S)$ . If it represents a bound-state situation (i.e.,  $V^S \leq D_e$ ), move to the next step. Otherwise (i.e., the sampled point lies outside the true hypervolume of integration  $\mathcal{B}$ ), go to step 6.

(4) Following Barker's procedure, find the minimum and maximum displacements and the sampled value for each conjugate momenta  $P_{\mathbf{R}}$  and  $P_{\mathbf{r}}$  according to

$$\begin{aligned} P_{\mathbf{R}}^{\min} &= 0 \\ P_{\mathbf{R}}^{\max} &= \sqrt{2\mu(D_e - V^S)} \\ P_{\mathbf{R}}^S &= P_{\mathbf{R}}^{\max}\xi \\ P_{\mathbf{r}}^{\min} &= 0 \\ P_{\mathbf{r}}^{\max} &= \sqrt{2\mu(D_e - V^S) - (P_{\mathbf{R}}^S)^2} \\ P_{\mathbf{r}}^S &= P_{\mathbf{r}}^{\max}\xi \end{aligned} \quad (29)$$

The sampled point  $\mathbf{x}_g^S = (R^S, r^S, \theta^S, P_{\mathbf{R}}^S, P_{\mathbf{r}}^S)$  is therefore within the hypervolume of integration  $\mathcal{B}$ .

(5) Calculate the weight factor associated with the sampled point  $\mathbf{x}_g^S$  according to

$$w_g = \frac{128\pi^4}{h^6} W_R W_r W_{P_{\mathbf{R}}} W_{P_{\mathbf{r}}} \quad (30)$$

where

$$\begin{aligned} W_R &= (R^S)^2 (R^{\max} - R^{\min}) (\cos(\theta^{\min}) - \cos(\theta^{\max})) \\ W_r &= (r^S)^2 (r^{\max} - r^{\min}) \\ W_{P_{\mathbf{R}}} &= (P_{\mathbf{R}}^S)^2 P_{\mathbf{R}}^{\max} \\ W_{P_{\mathbf{r}}} &= (P_{\mathbf{r}}^S)^2 P_{\mathbf{r}}^{\max} \end{aligned}$$

which represents the hypervolume (divided by  $h^6$ ) associated with  $\mathbf{x}_g^S$ .

(6) Repeat  $N_T$  times the steps 2–5 to evaluate the integral of the eq 25 as

$$I \approx I^{N_T} = \frac{1}{N_T} \sum_{g=1}^{N_{\text{in}}} w_g F_g \quad (31)$$

where  $F_g = F(\mathbf{x}_g^S) \equiv F(R^S, r^S, \theta^S, P_{\mathbf{R}}^S, P_{\mathbf{r}}^S)$  is the function to be integrated,  $N_T$  is the total number of sampled points, and  $N_{\text{in}}$  is the total number of sampled points that are within the hypervolume of integration  $\mathcal{B}$ . The standard deviation associated with eq 31 assumes the form

$$\sigma^2 = \frac{1}{N_{\text{in}}(N_{\text{in}} - 1)} \sum_{g=1}^{N_{\text{in}}} \left( \frac{N_{\text{in}}}{N_T} w_g F_g - I^{N_T} \right)^2 \quad (32)$$

Note that the efficiency of this Monte Carlo procedure, which is defined by  $\epsilon = N_{\text{in}}/N_T$ , is not close to 1 as in Barker's algorithm but is certainly larger than for the crude Monte Carlo method because Barker's method is still employed to sample the momenta. Note further that such a fact does not imply that the present Monte Carlo procedure requires a larger computational effort than the one based on Barker's algorithm. The reason is that the determination of an integration domain close to the true hypervolume  $\mathcal{B}$  as required in Barker's algorithm can itself be time-consuming because of the necessity of calling the potential-energy surface routine many times.

### 3. Technical Details and Results

The rovibrational partition function and its first and second moments have been computed from the standard classical statistical mechanics expressions in eqs 2, 7, and 8 by using the Monte Carlo method described above. Calculations have also been carried out by using the hybrid LCP/QFH method (eqs 3, 9, and 10). Moreover, calculations of thermodynamic functions are reported from  $Q_{\text{rovib}}$ ,  $Q'_{\text{rovib}}$ , and  $Q''_{\text{rovib}}$  by employing eqs 11–14. All calculations considered a standard state pressure of  $p = 1$  bar ( $10^5$  Pa) as used in the JANAF<sup>10</sup> tables and by Vidler and Tennyson.<sup>32</sup> To describe the H<sub>2</sub>O molecule, we have employed the energy-switching (ES) potential energy surface reported by one of us.<sup>50</sup> This has been obtained by merging a modified form of the global many-body expansion (MBE) potential of Murrell and Carter<sup>51</sup> and a spectroscopically accurate polynomial form reported by Polyansky et al.<sup>52</sup> (known as PJT1). The classical dissociation energy of the ES potential energy surface is  $D_e = 0.199\,865\,54E_h$ . Besides being global and having spectroscopic accuracy where this is known, such a potential energy surface offers the advantage of its analytical first derivatives with respect to the internuclear distances having been obtained.<sup>53</sup> Thus, only its second derivatives are needed to be calculated numerically for the purpose of evaluating the effective potential (eq 4) according to the hybrid LCP/QFH method. They have been calculated numerically from the known analytical first-derivatives, thus avoiding errors inherent to the numerical estimation of high-order derivatives.

Before we present the results, we should define the  $R, r$ , and  $\theta$  displacement intervals, of which the importance has been highlighted in the previous section. Thus, to improve the efficiency of the Monte Carlo procedure, we should establish the smallest hyperrectangle that contains the domain of integration  $\mathcal{B}$ . This can be defined by the minimum values of  $R$  and  $r$  when varied independently from each other, while the maximum values of  $R$  and  $r$  are fixed at the asymptotic region of the potential. Using such a procedure, we have obtained  $R^{\min}$

**TABLE 1: Convergence of Rovibrational Partition Function Calculations for Water Using the Hybrid LCP/QFH Method**

$T, K$	run 1	run 2	run 3	all
	$N_T = 4 \times 10^8$ $N_{in} = 25\ 992\ 144$	$N_T = 4 \times 10^8$ $N_{in} = 25\ 999\ 790$	$N_T = 4 \times 10^8$ $N_{in} = 25\ 991\ 985$	$N_T = 1.2 \times 10^9$ $N_{in} = 77\ 983\ 919$
1000	336.0 ± 7.9	335.9 ± 8.0	320.8 ± 7.8	330.9 ± 4.6
2000	1411.3 ± 10.0	1407.1 ± 10.0	1396.4 ± 9.9	1404.9 ± 5.8
3000	4158.8 ± 16.0	4156.5 ± 15.9	4160.8 ± 15.9	4158.7 ± 9.2
4000	10 226.3 ± 25.6	10 228.8 ± 25.6	10 257.6 ± 25.6	10 237.5 ± 14.8
5000	22 070.6 ± 39.6	22 083.0 ± 40.0	22 143.1 ± 39.7	22 098.9 ± 22.9
6000	42 842.8 ± 58.7	42 876.3 ± 58.7	42 968.5 ± 58.9	42 895.9 ± 33.9

**TABLE 2: Rovibrational Partition Function of H<sub>2</sub>O as a Function of Temperature**

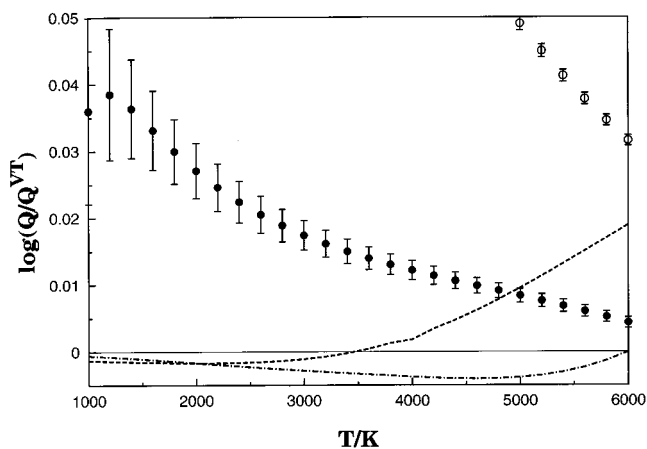
$T, K$	$Q_{rovib}^{CM}$ <sup>a</sup>	$Q_{rovib}^{LCP/QFH}$ <sup>b</sup>	$Q_{rovib}^{VT}$ <sup>c</sup>	$Q_{rovib}^{HVMT}$ <sup>d</sup>	$Q_{rovib}^{Irwin}$ <sup>e</sup>
500	12 474.3 ± 2703	56.8 ± 3.6	96.583 33	96.4132	
1000	3263.3 ± 28.9	330.9 ± 4.6	304.580	303.670	304.172
1200	2417.7 ± 16.9	469.1 ± 4.6	429.315	427.918	428.560
1400	2178.6 ± 12.5	637.2 ± 4.7	586.027	583.987	584.696
1600	2188.9 ± 10.5	843.4 ± 5.0	781.478	778.608	779.298
1800	2343.7 ± 9.7	1096.4 ± 5.3	1023.29	1019.39	1019.89
2000	2607.3 ± 9.4	1404.9 ± 5.8	1320.00	1314.85	1314.87
2200	2968.9 ± 9.4	1778.9 ± 6.3	1681.04	1674.49	1673.58
2400	3428.6 ± 9.7	2229.0 ± 6.9	2116.92	2108.87	2106.37
2600	3992.0 ± 10.1	2766.8 ± 7.6	2639.20	2629.76	2624.68
2800	4668.6 ± 10.7	3405.4 ± 8.3	3260.65	3250.27	3241.08
3000	5470.1 ± 11.4	4158.7 ± 9.2	3995.27	3984.98	3969.40
3200	6410.6 ± 12.2	5042.3 ± 10.1	4858.42	4850.13	4824.74
3400	7505.9 ± 13.2	6072.9 ± 11.2	5866.89	5863.76	5823.60
3600	8773.6 ± 14.2	7269.0 ± 12.3	7038.94	7045.89	6983.93
3800	10 232.9 ± 15.4	8650.1 ± 13.5	8394.37	8418.59	8325.24
4000	11 904.7 ± 16.6	10 237.5 ± 14.8	9954.54	9996.1	9868.67
4200	13 811.2 ± 18.0	12 053.8 ± 16.2	11 742.4	11 834.7	11 637.1
4400	15 975.8 ± 19.5	14 122.6 ± 17.7	13 782.2	13 932.9	13 655.3
4600	18 423.2 ± 21.1	16 469.1 ± 19.3	16 099.8	16 331.0	15 949.9
4800	21 179.0 ± 22.8	19 119.0 ± 21.0	18 722.1	19 061.2	18 549.8
5000	24 269.5 ± 24.6	22 098.9 ± 22.9	21 677.3	22 157.1	21 485.8
5200	27 721.2 ± 26.6	25 435.8 ± 24.8	24 994.1	25 653.4	24 791.4
5400	31 561.1 ± 28.7	29 157.1 ± 26.9	28 702.2	29 586.0	28 502.3
5600	35 815.9 ± 30.9	33 289.8 ± 29.1	32 831.2	33 990.8	32 656.9
5800	40 511.9 ± 33.2	37 860.6 ± 31.5	37 411.2	38 904.3	37 296.7
6000	45 674.9 ± 35.7	42 895.9 ± 33.9	42 471.8	44 362.4	42 465.8

<sup>a</sup> Standard classical results from eq 2; this work. <sup>b</sup> Hybrid LCP/QFH results from eq 3; this work. <sup>c</sup> Reference 32. <sup>d</sup> Reference 8. <sup>e</sup> Reference 13.

$= 1.244\ 44a_0$ ,  $R^{\max} = 10.0a_0$ ,  $r^{\min} = 1.230\ 90a_0$ , and  $r^{\max} = 10.0a_0$ . For  $\theta$ , we have taken  $\theta^{\min} = 0.0$  rad and  $\theta^{\max} = \pi$  rad.

All results from the present work have been calculated using three distinct Monte Carlo sequences of random numbers obtained from different seeds for the pseudorandom number generator. In particular, we have used the ran2 subroutine from Numerical Recipes.<sup>54</sup> Each sequence has been computed from a total of  $4 \times 10^8$  sampled points, totaling  $N_T = 1.2 \times 10^9$ . Table 1 shows some of the values calculated for the rovibrational partition function  $Q_{rovib}^{LCP/QFH}$  using the hybrid LCP/QFH method, together with the corresponding statistical uncertainties. As expected, the various calculations coincide within their statistical uncertainties. Moreover, the Monte Carlo error of the final results is smaller than 1.4% for temperatures above 1000 K and decreases with  $T$  (e.g., for  $T = 6000$  K, it is only 0.08%). Note that the Monte Carlo efficiency ( $\epsilon = N_{in}/N_T$ ) of our procedure is about 6.5%.

**3.1. Rovibrational Partition Function.** The calculated rovibrational partition functions of H<sub>2</sub>O based on the standard classical procedure ( $Q_{rovib}^{CM}$ ) and the hybrid LCP/QFH method ( $Q_{rovib}^{LCP/QFH}$ ) are reported in Table 2. For comparison, we also give in this table the results of Vidler and Tennyson<sup>32</sup> ( $Q_{rovib}^{VT}$ ), which were obtained by performing an explicit summation over experimental and theoretical rovibrational energy levels, and of Harris et al.<sup>8</sup> ( $Q_{rovib}^{HVMT}$ ), which were determined from a summation over theoretical energy levels. Also included is the



**Figure 1.** Logarithm of ratio of the rovibrational partition function with respect to that calculated by Vidler and Tennyson<sup>32</sup> ( $Q_{rovib}^{VT}$ ) as function of temperature: (○) standard classical ( $Q_{rovib}^{CM}$ ) results with error bars from eq 2; (●) hybrid LCP/QFH ( $Q_{rovib}^{LCP/QFH}$ ) results with error bars from eq 3; (---) Harris et al.<sup>8</sup> ( $Q_{rovib}^{HVMT}$ ); (-·-) Irwin<sup>13</sup> ( $Q_{rovib}^{Irwin}$ ).

Irwin fit<sup>13</sup> ( $Q_{rovib}^{Irwin}$ ) to the partition function data from the JANAF thermochemical tables.<sup>10</sup> Note that the rovibrational energy levels used in the Vidler and Tennyson calculations have been obtained from three separate sources: experiment where available,<sup>55</sup> computations from the spectroscopically determined PJT2<sup>56</sup> potential energy surface for levels with total angular momentum  $J \leq 42$  and  $E_i \leq 30\ 000\ \text{cm}^{-1}$ , and the assumption that vibrational and rotational motions can be separated for higher energies up to dissociation. In this case, the vibrational levels were computed by Mussa and Tennyson<sup>57</sup> using an ab initio potential energy surface,<sup>58</sup> while the rotational levels were estimated using the Padé approximant model of Polyansky.<sup>59</sup> Regarding the calculations by Harris et al., the computed rovibrational levels were obtained from the spectroscopically determined PJT2<sup>56</sup> potential energy surface for  $J \leq 35$  and  $E_i \leq 30\ 000\ \text{cm}^{-1}$ , while a procedure similar to that employed by Vidler and Tennyson was employed for higher energy levels. Assuming the results of Vidler and Tennyson as reference, we observe that our  $Q_{rovib}^{LCP/QFH}$  values are more accurate than  $Q_{rovib}^{CM}$  over the whole range of temperatures, as already found in a previous study<sup>26</sup> for diatomics. For example, at  $T = 2000$  K, the error relative to  $Q_{rovib}^{VT}$  [defined as  $\Delta Q_{rovib} = (Q_{rovib} - Q_{rovib}^{VT})/Q_{rovib}^{VT}$ ] in  $Q_{rovib}^{LCP/QFH}$  is  $\sim 6.5\%$  while that of  $Q_{rovib}^{CM}$  is  $\sim 98\%$ . In turn, for  $T = 6000$  K, one observes a deviation of 1% in  $Q_{rovib}^{LCP/QFH}$  and 7.5% in  $Q_{rovib}^{CM}$ . Moreover, for temperatures above 4900 K, the hybrid LCP/QFH approach gives results in better agreement with the Vidler and Tennyson ones than those computed by Harris et al.<sup>8</sup> This can also be seen from Figure 1, in which the logarithm of the ratios  $Q_{rovib}^{CM}/Q_{rovib}^{VT}$ ,  $Q_{rovib}^{LCP/QFH}/Q_{rovib}^{VT}$ ,  $Q_{rovib}^{HVMT}/Q_{rovib}^{VT}$ , and  $Q_{rovib}^{Irwin}/Q_{rovib}^{VT}$  are plotted as a function of temperature. It is important to point out that the comparison between our results and previous ones (mainly those of Vidler and Tennyson<sup>32</sup> and Harris et al.<sup>8</sup>) is somewhat arbitrary because of the different potential energy surfaces that have been employed for the calculations. In any case, the agreement between the hybrid LCP/QFH rovibrational partition function with previous results ( $Q_{rovib}^{VT}$ ,  $Q_{rovib}^{HVMT}$ , and  $Q_{rovib}^{Irwin}$ ) is quite satisfactory at low temperatures ( $T \approx 500$  K) and good at moderate and high temperatures.

The other two sets of results evaluated directly by using the Monte Carlo multidimensional integration are presented in Tables 3 and 4 (respectively, the first and second moments of

**TABLE 3: Calculated First Moment of Internal Partition Function of H<sub>2</sub>O as a Function of Temperature**

<i>T</i> , K	$Q_{\text{rovib}}^{\text{CM}^a}$	$Q_{\text{rovib}}^{\text{LCP/QFH}^b}$	$Q_{\text{rovib}}^{\text{VT}^c}$
500	-1 131 384 ± 28 161	239.4 ± 9.5	149.5118
1000	-7279.7 ± 108	631.1 ± 7.5	553.9156
1200	-2643.1 ± 45.7	910.4 ± 7.1	838.0108
1400	-606.3 ± 25.3	1300.3 ± 7.2	1224.177
1600	734.6 ± 16.9	1825.1 ± 7.6	1738.323
1800	1911.3 ± 13.3	2512.8 ± 8.3	2409.863
2000	3132.8 ± 12.1	3395.9 ± 9.4	3272.146
2200	4510.6 ± 12.2	4511.8 ± 10.7	4362.889
2400	6123.0 ± 13.1	5902.9 ± 12.2	5724.670
2600	8036.6 ± 14.6	7617.3 ± 13.9	7405.270
2800	10 316.0 ± 16.5	9708.5 ± 15.9	9458.213
3000	13 027.6 ± 18.7	12 236.5 ± 18.2	11 943.2
3200	16 241.7 ± 21.2	15 267.2 ± 20.6	14 926.3
3400	20 033.5 ± 23.9	18 873.1 ± 23.4	18 480.5
3600	24 483.7 ± 27.0	23 133.2 ± 26.4	22 685.6
3800	29 677.9 ± 30.3	28 132.0 ± 29.7	27 628.2
4000	35 706.0 ± 33.9	33 959.0 ± 33.3	33 401.0
4200	42 661.6 ± 37.9	40 708.0 ± 37.2	40 102.3
4400	50 640.2 ± 42.1	48 475.1 ± 41.4	47 834.7
4600	59 737.6 ± 46.7	57 357.1 ± 45.9	56 703.6
4800	70 048.8 ± 51.6	67 450.2 ± 50.8	66 816.0
5000	81 665.5 ± 56.9	78 847.8 ± 56.1	78 278.4
5200	94 675.1 ± 62.6	91 638.9 ± 61.7	91 195.1
5400	109 158.5 ± 68.6	105 906.4 ± 67.7	105 666.5
5600	125 189.5 ± 75.0	121 725.8 ± 74.1	121 787.3
5800	142 833.0 ± 81.9	139 164.0 ± 80.9	139 645.3
6000	162 144.5 ± 89.1	158 278.2 ± 88.1	159 320.0

<sup>a</sup> Standard classical results from eq 7; this work. <sup>b</sup> Hybrid LCP/QFH results from eq 9; this work. <sup>c</sup> Reference 32.

**TABLE 4: Calculated Second Moment of Internal Partition Function of H<sub>2</sub>O as a Function of Temperature**

<i>T</i> , K	$Q_{\text{rovib}}^{\text{CM}^a}$	$Q_{\text{rovib}}^{\text{LCP/QFH}^b}$	$Q_{\text{rovib}}^{\text{VT}^c}$
500	10 835 142 ± 300 896	690.9 ± 44.9	399.183
1000	31 221 ± 469	1787.7 ± 21.7	1758.363
1200	14 014 ± 154	2895.6 ± 19.2	2824.526
1400	10 237.9 ± 69.2	4471.4 ± 19.1	4343.094
1600	10 411.6 ± 39.8	6634.5 ± 20.7	6439.341
1800	12 492.6 ± 30.2	9529.0 ± 23.6	9258.888
2000	15 982.4 ± 29.5	13 324.5 ± 27.5	12 970.58
2200	20 828.6 ± 32.8	18 217.9 ± 32.4	17 769.12
2400	27 149.6 ± 38.2	24 434.5 ± 38.1	23 877.75
2600	35 150.5 ± 44.7	32 230.9 ± 44.8	31 550.88
2800	45 094.0 ± 52.4	41 895.9 ± 52.3	41 076.76
3000	57 288.5 ± 61.0	53 752.8 ± 60.8	52 779.7
3200	72 081.8 ± 70.6	68 158.7 ± 70.3	67 021.0
3400	89 855.7 ± 81.3	85 502.7 ± 80.9	84 199.0
3600	111 019.0 ± 93.0	106 200.5 ± 92.5	104 745.8
3800	135 998 ± 106	130 686 ± 105	129 121.6
4000	165 227 ± 120	159 403 ± 119	157 806.7
4200	199 134 ± 135	192 787 ± 135	191 290.4
4400	238 125 ± 152	231 259 ± 151	230 058.2
4600	282 578 ± 170	275 206 ± 169	274 578.9
4800	332 824 ± 190	324 973 ± 189	325 290.7
5000	389 142 ± 211	380 847 ± 209	382 588.9
5200	451 748 ± 233	443 056 ± 232	446 815.3
5400	520 790 ± 257	511 758 ± 256	518 249.1
5600	596 350 ± 282	587 041 ± 281	597 100.7
5800	678 437 ± 310	668 921 ± 308	683 507.8
6000	766 993 ± 338	757 343 ± 337	777 534.1

<sup>a</sup> Standard classical results from eq 8; this work. <sup>b</sup> Hybrid LCP/QFH results from eq 10; this work. <sup>c</sup> Ref 32.

the H<sub>2</sub>O rovibrational partition function) and are compared with the accurate results of Vidler and Tennyson.<sup>32</sup> It is clear that, for both cases, the hybrid LCP/QFH method shows an improved agreement with respect to the standard classical statistical mechanics results over the whole range of temperatures. In particular, the standard classical approach is seen to give values

**TABLE 5: The Gibbs Enthalpy Function (in J K<sup>-1</sup> mol<sup>-1</sup>) of H<sub>2</sub>O as a Function of Temperature**

<i>T</i> , K	gef <sup>CM</sup> <sup>a</sup>	gef <sup>LCP/QFH</sup> <sup>b</sup>	gef <sup>VT</sup> <sup>c</sup>	gef <sup>HVMT</sup> <sup>d</sup>	gef <sup>JANAF</sup> <sup>e</sup>
500	252.243 ± 0.180	188.270 ± 0.526	192.681	192.53	192.68
1000	226.452 ± 0.074	207.423 ± 0.115	206.734	206.58	206.73
1200	226.098 ± 0.058	212.465 ± 0.081	211.727	211.58	211.73
1400	227.257 ± 0.048	217.035 ± 0.062	216.340	216.19	216.34
1600	229.188 ± 0.040	221.258 ± 0.049	220.624	220.47	220.62
1800	231.516 ± 0.034	225.199 ± 0.040	224.626	224.47	224.62
2000	234.042 ± 0.030	228.901 ± 0.034	228.383	228.23	228.37
2200	236.653 ± 0.026	232.395 ± 0.029	231.924	231.77	231.90
2400	239.284 ± 0.023	235.703 ± 0.026	235.274	235.12	235.25
2600	241.895 ± 0.021	238.847 ± 0.023	238.454	238.30	238.42
2800	244.465 ± 0.019	241.842 ± 0.020	241.481	241.33	241.44
3000	246.981 ± 0.017	244.702 ± 0.018	244.368	244.23	244.32
3200	249.435 ± 0.016	247.439 ± 0.017	247.130	247.00	247.07
3400	251.825 ± 0.015	250.063 ± 0.015	249.776	249.65	249.70
3600	254.148 ± 0.013	252.584 ± 0.014	252.317	252.21	252.23
3800	256.407 ± 0.012	255.010 ± 0.013	254.760	254.66	254.66
4000	258.601 ± 0.012	257.346 ± 0.012	257.113	257.04	256.99
4200	260.732 ± 0.011	259.601 ± 0.011	259.383	259.33	259.25
4400	262.802 ± 0.010	261.777 ± 0.010	261.574	261.55	261.42
4600	264.814 ± 0.010	263.881 ± 0.010	263.693	263.69	263.52
4800	266.768 ± 0.009	265.917 ± 0.009	265.743	265.77	265.56
5000	268.666 ± 0.008	267.887 ± 0.009	267.727	267.79	267.53
5200	270.511 ± 0.008	269.796 ± 0.008	269.650	269.75	269.44
5400	272.303 ± 0.008	271.645 ± 0.008	271.514	271.65	271.29
5600	274.045 ± 0.007	273.437 ± 0.007	273.322	273.49	273.09
5800	275.738 ± 0.007	275.175 ± 0.007	275.076	275.28	274.84
6000	277.383 ± 0.006	276.861 ± 0.007	276.779	277.02	276.54

<sup>a</sup> Standard classical results; this work. <sup>b</sup> Hybrid LCP/QFH results; this work. <sup>c</sup> Reference 32. <sup>d</sup> Reference 8. <sup>e</sup> Reference 10.

of the correct magnitude only at  $T \leq 1600$  K, while the hybrid LCP/QFH method gives acceptable results from  $T = 500$  K upward. The only exception is for the second moment of the internal partition function in which, unexpectedly,  $Q_{\text{rovib}}^{\text{CM}}$  lies closer to  $Q_{\text{rovib}}^{\text{VT}}$  than does  $Q_{\text{rovib}}^{\text{LCP/QFH}}$  at temperatures above 5300 K. However, as noted above, the results of Vidler and Tennyson<sup>32</sup> employ (for high temperatures) calculated high-energy rovibrational levels and a model for the highest rotationally excited states, and hence, such a behavior can partly be attributed to the use of different potential energy surfaces.

**3.2. Thermodynamic Quantities.** The values reported in Tables 2–4 have been used to obtain the Gibbs enthalpy function (gef), the Helmholtz function (hcf), the entropy ( $S$ ), and the specific heat capacity at constant pressure ( $C_p$ ). The results, with their associated errors obtained from the standard error propagation formulas, are given in Tables 5–8, respectively. For comparison, we also give the results from Vidler and Tennyson (VT)<sup>32</sup> and Harris et al. (HVMT),<sup>8</sup> as well as those from the JANAF thermochemical tables.<sup>10</sup> As expected, the hybrid LCP/QFH results are more accurate than those computed from standard classical statistical mechanics. In general, the thermodynamic quantities calculated with the hybrid LCP/QFH method are in good agreement with previous studies (i.e., deviations of about 1%) at temperatures above 1000 K, while those obtained using the standard classical procedure show a similar agreement only at much higher temperatures. Moreover, assuming as reference the results of Vidler and Tennyson,<sup>32</sup> Tables 5–7 show that the hybrid LCP/QFH results look better than the HVMT and JANAF ones over some range of temperatures. For example, gef<sup>LCP/QFH</sup> lies closer to gef<sup>VT</sup> than gef<sup>HVMT</sup> and gef<sup>JANAF</sup> for  $T \leq 5300$  K and  $T \leq 4800$  K, respectively. In turn, for the Helmholtz function, such a pattern is observed at temperatures above 3800 K when comparing to the Harris et al.<sup>8</sup> results and at  $4000 \leq T \leq 5000$  K in relation to the JANAF<sup>10</sup> values. For the entropy, we observe a better agreement between  $S^{\text{LCP/QFH}}$  and  $S^{\text{VT}}$  for temperatures above 1500 K when compar-

**TABLE 6: The Helmholtz Function (in J mol<sup>-1</sup>) of H<sub>2</sub>O as a Function of Temperature**

T, K	hcf <sup>CM</sup> <sup>a</sup>	hcf <sup>LCP/QFH</sup> <sup>b</sup>	hcf <sup>VT</sup> <sup>c</sup>	hcf <sup>HVMT</sup> <sup>d</sup>	hcf <sup>JANAF</sup> <sup>e</sup>
500	-37 216 ± 121	18 006 ± 1800	6925	6925	6925
1000	-7666 ± 111	26 741 ± 408	26 003	26 000	26 000
1200	4132 ± 112	34 402 ± 340	34 515	34 509	34 506
1400	15 957 ± 117	42 952 ± 307	43 513	43 504	43 493
1600	27 819 ± 124	52 141 ± 289	52 946	52 936	52 908
1800	39 716 ± 135	61 812 ± 281	62 756	62 750	62 693
2000	51 649 ± 149	71 863 ± 276	72 890	72 891	72 790
2200	63 616 ± 163	82 218 ± 274	83 300	83 317	83 153
2400	75 619 ± 177	92 829 ± 273	93 946	93 992	93 741
2600	87 660 ± 189	103 655 ± 272	104 797	104 892	104 520
2800	99 740 ± 200	114 669 ± 272	115 828	115 998	115 464
3000	111 860 ± 209	125 848 ± 271	127 019	127 302	126 549
3200	124 022 ± 216	137 172 ± 271	138 353	138 795	137 757
3400	136 222 ± 223	148 623 ± 270	149 816	150 471	149 073
3600	148 456 ± 227	160 184 ± 269	161 394	162 321	160 485
3800	160 717 ± 231	171 838 ± 269	173 072	174 334	171 980
4000	172 992 ± 234	183 562 ± 267	184 883	186 490	183 552
4200	185 266 ± 236	195 334 ± 266	196 660	198 766	195 191
4400	197 519 ± 238	207 127 ± 264	208 529	211 130	206 892
4600	209 729 ± 239	218 915 ± 263	220 418	223 545	218 650
4800	221 870 ± 239	230 668 ± 261	232 301	235 973	230 458
5000	233 917 ± 239	242 356 ± 259	244 149	248 371	242 313
5200	245 845 ± 239	253 951 ± 257	255 936	260 697	254 215
5400	257 629 ± 239	265 425 ± 255	267 634	272 911	266 164
5600	269 247 ± 238	276 753 ± 253	279 218	284 978	278 161
5800	280 680 ± 237	287 913 ± 250	290 663	296 864	290 204
6000	291 911 ± 236	298 888 ± 248	301 850	308 544	302 295

<sup>a</sup> Standard classical results; this work. <sup>b</sup> Hybrid LCP/QFH results; this work. <sup>c</sup> Reference 32. <sup>d</sup> Reference 8. <sup>e</sup> Reference 10.

**TABLE 7: The Entropy (in J K<sup>-1</sup> mol<sup>-1</sup>) of H<sub>2</sub>O as a Function of Temperature**

T, K	S <sup>CM</sup> <sup>a</sup>	S <sup>LCP/QFH</sup> <sup>b</sup>	S <sup>VT</sup> <sup>c</sup>	S <sup>HVMT</sup> <sup>d</sup>	S <sup>JANAF</sup> <sup>e</sup>
500	177.81 ± 0.42	224.28 ± 4.13	206.53	206.38	206.53
1000	218.79 ± 0.18	234.16 ± 0.52	232.74	232.58	232.73
1200	229.54 ± 0.15	241.13 ± 0.36	240.49	240.33	240.48
1400	238.66 ± 0.13	247.72 ± 0.28	247.42	247.26	247.40
1600	246.57 ± 0.12	253.85 ± 0.23	253.71	253.55	253.69
1800	253.58 ± 0.11	259.54 ± 0.20	259.49	259.33	259.45
2000	259.87 ± 0.10	264.83 ± 0.17	264.83	264.67	264.76
2200	265.57 ± 0.10	269.77 ± 0.15	269.79	269.64	269.70
2400	270.79 ± 0.10	274.38 ± 0.14	274.42	274.28	274.31
2600	275.61 ± 0.09	278.71 ± 0.13	278.76	278.64	278.62
2800	280.09 ± 0.09	282.80 ± 0.12	282.85	282.76	282.68
3000	284.27 ± 0.09	286.65 ± 0.11	286.71	286.66	286.50
3200	288.19 ± 0.08	290.30 ± 0.10	290.36	290.36	290.12
3400	291.89 ± 0.08	293.78 ± 0.09	293.84	293.90	293.55
3600	295.39 ± 0.08	297.08 ± 0.09	297.15	297.29	296.81
3800	298.70 ± 0.07	300.23 ± 0.08	300.30	300.54	299.91
4000	301.85 ± 0.07	303.24 ± 0.08	303.32	303.66	302.88
4200	304.84 ± 0.07	306.11 ± 0.07	306.21	306.65	305.72
4400	307.69 ± 0.06	308.85 ± 0.07	308.97	309.53	308.44
4600	310.41 ± 0.06	311.47 ± 0.07	311.61	312.28	311.06
4800	312.99 ± 0.06	313.97 ± 0.06	314.14	314.93	313.57
5000	315.45 ± 0.06	316.36 ± 0.06	316.56	317.46	315.99
5200	317.79 ± 0.05	318.63 ± 0.06	318.87	319.88	318.32
5400	320.01 ± 0.05	320.80 ± 0.05	321.08	322.18	320.58
5600	322.13 ± 0.05	322.86 ± 0.05	323.18	324.38	322.76
5800	324.13 ± 0.05	324.82 ± 0.05	325.19	326.46	324.87
6000	326.04 ± 0.05	326.68 ± 0.05	327.10	328.44	326.92

<sup>a</sup> Standard classical results; this work. <sup>b</sup> Hybrid LCP/QFH results; this work. <sup>c</sup> Reference 32. <sup>d</sup> Reference 8. <sup>e</sup> Reference 10.

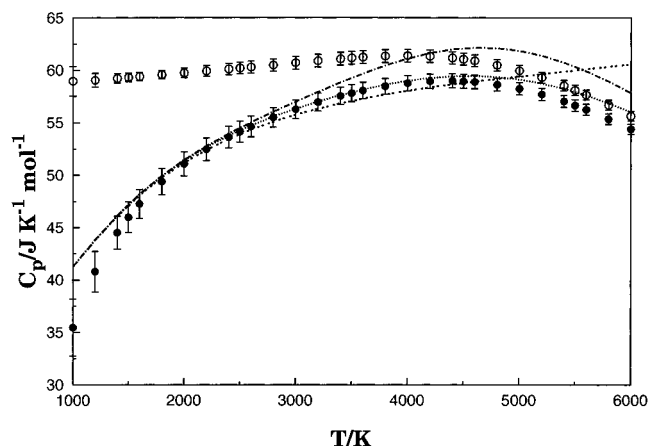
ing to S<sup>HVMT</sup> and for temperatures over the range 2000 ≤ T ≤ 5700 K when considering S<sup>JANAF</sup>.

For the specific heat capacity at constant pressure, the agreement between the various results is worse than for the other thermodynamic quantities. In fact, C<sub>p</sub> is particularly sensitive to convergence of the partition function, because it is determined from the difference between the second and the square of the

**TABLE 8: The Specific Heat Capacity at Constant Pressure (in J K<sup>-1</sup> mol<sup>-1</sup>) of H<sub>2</sub>O as a Function of Temperature**

T, K	C <sub>p</sub> <sup>CM</sup> <sup>a</sup>	C <sub>p</sub> <sup>LCP/QFH</sup> <sup>b</sup>	C <sub>p</sub> <sup>VT</sup> <sup>c</sup>	C <sub>p</sub> <sup>HVMT</sup> <sup>d</sup>	C <sub>p</sub> <sup>JANAF</sup> <sup>e</sup>
1000	58.96 ± 1.40	35.46 ± 2.72	41.287	41.278	41.268
1200	59.05 ± 0.66	40.79 ± 1.94	43.809	43.795	43.768
1400	59.21 ± 0.44	44.51 ± 1.58	46.124	46.114	46.054
1600	59.40 ± 0.39	47.26 ± 1.37	48.157	48.160	48.050
1800	59.58 ± 0.41	49.38 ± 1.24	49.904	49.929	49.749
2000	59.75 ± 0.46	51.07 ± 1.15	51.394	51.452	51.180
2200	59.93 ± 0.50	52.45 ± 1.08	52.668	52.776	52.408
2400	60.11 ± 0.54	53.62 ± 1.03	53.766	53.953	53.444
2600	60.30 ± 0.57	54.62 ± 0.98	54.724	55.026	54.329
2800	60.50 ± 0.59	55.50 ± 0.93	55.571	56.033	55.089
3000	60.70 ± 0.61	56.27 ± 0.89	56.326	56.996	55.748
3200	60.91 ± 0.61	56.95 ± 0.85	57.005	57.928	56.323
3400	61.09 ± 0.61	57.55 ± 0.82	57.614	58.824	56.828
3600	61.25 ± 0.61	58.05 ± 0.79	58.152	59.671	57.276
3800	61.35 ± 0.61	58.46 ± 0.76	58.613	60.441	57.675
4000	61.39 ± 0.60	58.76 ± 0.73	58.986	61.104	58.033
4200	61.34 ± 0.59	58.94 ± 0.70	59.259	61.627	58.357
4400	61.18 ± 0.57	58.98 ± 0.67	59.418	61.981	58.650
4600	60.90 ± 0.56	58.88 ± 0.65	59.451	62.143	58.918
4800	60.49 ± 0.55	58.63 ± 0.62	59.350	62.098	59.164
5000	59.96 ± 0.53	58.23 ± 0.60	59.111	61.844	59.390
5200	59.30 ± 0.51	57.69 ± 0.57	58.734	61.384	59.628
5400	58.52 ± 0.50	57.02 ± 0.55	58.255	60.732	59.864
5600	57.64 ± 0.48	56.24 ± 0.53	57.591	59.907	60.100
5800	56.67 ± 0.47	55.35 ± 0.51	56.846	58.934	60.335
6000	55.63 ± 0.45	54.38 ± 0.49	56.003	57.838	60.571

<sup>a</sup> Standard classical results; this work. <sup>b</sup> Hybrid LCP/QFH results; this work. <sup>c</sup> Reference 32. <sup>d</sup> Reference 8. <sup>e</sup> Reference 10.



**Figure 2.** Specific heat at constant pressure, C<sub>p</sub>, as function of temperature: (○) standard classical results (C<sub>p</sub><sup>CM</sup>) with error bars, this work; (●) hybrid LCP/QFH results (C<sub>p</sub><sup>LCP/QFH</sup>) with error bars, this work; (···) C<sub>p</sub><sup>VT</sup> from ref 32; (- · -) C<sub>p</sub><sup>HVMT</sup> from ref 8; (- - -) C<sub>p</sub><sup>JANAF</sup> from ref 13.

first moments. In particular, previous workers<sup>8,32</sup> have called attention to the fact that it is difficult to obtain reliable results for C<sub>p</sub> at high temperatures. In our calculations, this can be quantified from the size of the Monte Carlo error associated with the C<sub>p</sub> results, which is proportionally higher than the Monte Carlo error for the other thermodynamic quantities. However, conversely to previous calculations, the uncertainties in both sets of calculations reported in the present work decrease with temperature. This is particularly clear in Figure 2, which shows C<sub>p</sub> as a function of temperature. In any case, C<sub>p</sub><sup>LCP/QFH</sup> follows much better the general behavior of the Vidler and Tennyson,<sup>32</sup> Harris et al.,<sup>8</sup> and JANAF<sup>10</sup> results than C<sub>p</sub><sup>CM</sup>, specially for T ≤ 4000 K. Moreover, if we assume C<sub>p</sub><sup>VT</sup> as reference, then we can claim that the hybrid LCP/QFH results are more accurate than C<sub>p</sub><sup>HVMT</sup> at temperatures above 2400 K

and  $C_p^{\text{JANAF}}$  at  $T \geq 2200$  K. Finally, we comment on an interesting feature that is observed at temperatures above 4000 K. While the JANAF specific heat value continues to increase with temperature, both our results (based on standard classical statistical mechanics and the hybrid LCP/QFH method) and those of Vidler and Tennyson<sup>32</sup> and Harris et al.<sup>8</sup> show a maximum at about 4500 K before decreasing for higher temperatures. Such a feature may be attributed to a saturation of the energy levels of water in its ground electronic state, given that the phase-space hypervolume  $\mathcal{B}$  associated with a bound-state regime is finite. A similar explanation was suggested by Vidler and Tennyson<sup>32</sup> on the basis of the fact that the number of rovibrational energy levels is finite; see also ref 60. Such a feature is absent from the JANAF results because the results for  $T \geq 4000$  K have been obtained using a linear extrapolation.

#### 4. Conclusions

Through the use of the standard classical statistical mechanics and hybrid LCP/QFH methods, calculations of the rovibrational partition function of water and related thermodynamic quantities have been reported as a function of temperature. The hybrid LCP/QFH results are found to be rather more accurate than the standard classical ones, both for the partition function (as shown previously for diatomic systems<sup>26</sup>) and for thermodynamic properties (Gibbs enthalpy function, Helmholtz function, entropy, and specific heat at constant pressure). Moreover, the hybrid LCP/QFH results have been found to be in good agreement with previous calculations based on experimental or theoretical rovibrational energy levels or both<sup>8,32</sup> and an approximate compilation<sup>10</sup> for temperatures between 1000 and 6000 K. In summary, our hybrid LCP/QFH method can provide accurate values of the partition function and related thermodynamic properties of polyatomic molecules described by realistic potential energy surfaces at moderate- and high-temperature regimes in which the exact sum-over-states quantum mechanical treatment is unaffordable.

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