The purpose of this study is to develop a method of calculating the vibration partition function of diatomic molecules for the Morse potential energy. After a brief introduction about the eigensolutions obtained for the problem in question, via the Euler-Maclaurin formula, we determine the thermal properties for four diatomic molecules, such as H$_2$, HCl, LiH, and CO. These different cases are exposed and explained by the appropriate plots of the thermal properties. In addition, we show that our method to calculate the thermal properties can be used to determine important thermodynamic quantities.

Keywords: Partition function; Bound states of three-dimensional Morse potential; the Euler-Maclaurin formula; diatomic molecules.

PACS: 03.60.-w; 03.60.Ge; 05.30.-d; 05.70.-a

1. Introduction

The molecules of polyatomic gases are considered to have internal degrees of freedom that are related to internal (rotational, vibrational, and electronic) energy states. At very low temperatures, the degrees of freedom are frozen. By increasing the temperature, the rotational modes are excited. The vibrational modes are excited at higher temperatures, where, normally, the rotational modes are completely excited. The electronic modes are excited at the very highest temperatures. A rational thermodynamic theory for a diatomic gas with one excited mode was developed at first by Muller and an extended thermodynamic theory by Kremer. Their approach is based on the equations of the balance of mass, momentum, and internal energy customary in thermodynamics and is supplemented by a general equation of balance for the vibrational energy. These equations of balance and simple constitutive equations give rise to field equations which a thermodynamic process must satisfy [1–5].

There has been an increasing interest in the study of the partition function and its derivatives of diatomic molecules because it is the essential link between the coordinates of microscopic systems and the thermodynamic properties. In the atomic and molecular physics, the interaction between atoms in diatomic and even in polyatomic molecules is usually described by the Morse potential. The Morse potential gives an excellent description of the interaction between the two atoms in a diatomic molecule. The several studies show that it yields an exactly solvent Schrödinger problem. Besides this potential is the most simple and realistic anharmonic potential model, which has been widely used in the description of the vibrational motion of diatomic molecules. Finally, the exact solutions of the Schrodinger equation with Morse potential based on the Pekeris approximation have been obtained [6, 7]: In the last decade a large community of researchers has been involved a search of approximate solutions for wave equations (non-relativistic or relativistic) including the centrifugal term and subject to different potential functions $V(r)$. The main characteristic of these solutions lies in the substitution of the centrifugal term by an approximation so that one can obtain an equation, normally hypergeometric, which is solvable. Pekeris [8] is the pioneer in the study. He managed to obtain analytic solutions for the radial Schrodinger equation with the Morse potential [9], through expansion for the centrifugal term.

Taking advantage of the exact energy spectrum of the Morse potential, we could analytically obtain the individual partition function, and consequently the thermodynamic functions can be deduced: analytical representations of thermodynamic functions of gases over the whole temperature range from zero to the thermal dissociation limit have aroused much interest in dealing with diatomic and polyatomic systems. Through the exact form of its spectrum of energy, the vibration partition function, which is of great importance to many issues in chemical physics and engineering, can be obtained. Following its definition, the molecular vibrational partition function can be calculated by direct summation over all possible vibrational energy levels. Many efforts have been made to acquire explicit expressions of partition function for molecular potential energy models in diatomic and polyatomic molecules: in this context, and also in the non-relativistic case, the investigation of thermodynamic functions of some type of potentials such as Morse and improved Manning-Rosen potentials, improved Rosen-Morse and Tietz oscillators, through a partition function and its derivatives with respect to temperature, were an important field of research in the literature [9–14, 18–25].
Strekalov [15,16], derived a simple analytical formula for the partition function of Morse oscillators. With the cumulant expansion method, the approximation equally suitable within the whole range of temperatures, where a molecule exists as a bound system, has been obtained. Based on the Poisson summation formula (see Appendix A) and in order to reduce the errors with the experimental results, he proposed an accurate closed-form expression for the partition function in the case of the Morse oscillator. His approach becomes a most useful method used to calculate the thermal properties of some diatomic molecules for a different type of potentials (see following Refs [9–14, 18–25]). Recently, in the same context, another approach has been proposed: based on the Euler-Maclaurin formula (see Appendix A), the author has studied the case of the one-dimensional Morse potential in its q-deformed version: the results found are promising [26].

Now, before to give the principal aims of our works, we are in obligation to make some remarks about the difference between both methods described above.

- The Poisson summation is an equation that relates the Fourier series coefficients of the periodic summation of a function to values of the function’s continuous Fourier transform. For appropriate functions \( f \), the Poisson summation formula may be stated as [17–19]:

\[
\sum_{n=-\infty}^{+\infty} f(n) = \sum_{n=-\infty}^{+\infty} \hat{f}(n) \tag{1}
\]

where \( \hat{f} \) is the Fourier transform of \( f \) and \( n \in \mathbb{Z} \).

- The Euler-Maclaurin formula provides a powerful connection between integrals and sums. It provides expressions for the difference between the sum and the integral in terms of the higher derivatives \( f \) evaluated at the endpoints of the interval. It has the following definition [17–19]:

\[
\sum_{n=0}^{\infty} f(n) = \frac{1}{2} f(0) + \int_0^\infty f(x) \, dx - \sum_{p=1}^{\infty} \frac{B_{2p}}{(2p)!} f^{(2p-1)}(0), \tag{2}
\]

where \( B_{2p} \) are the Bernoulli numbers, \( f^{(2p-1)} \) is the derivative of order \( 2p - 1 \). Both methods are basic examples of deeper ideas. But at their core, the underlying ideas are very different.

- The Euler-Maclaurin formulas are used to give high accuracy estimates of integrals in standard numerical analysis methods, as discrete sums with understandable error terms are computable and estimable. Thus, this method is the most important method for the summation of infinite series. Also, this formula is a bit tedious to use, and the amount of work depends on the derivatives.

- The Poisson summation is very different in nature. It concerns Fourier series, or more fundamentally, the Fourier transform. This looks superficially similar to the integrals in Euler-Maclaurin summation, but the underlying ideas are substantively different. So the Poisson formula usually does not give an immediate answer but is a transform allowing other procedures to be applied.

Thus, the purpose of the present work is to give a simple analytical expression of the partition function of the Morse potential via the well-known Euler-Maclaurin method. This approach also enables one to estimate the influence of rotation-vibration interaction effects on the thermal properties of diatomic molecules. Precisely, we obtain the basic thermodynamic functions, such as the partition function and vibrational specific heat, are obtained. Section 4 will be a conclusion.

### 2. The behavior of the eigenvalues of the Morse potential in three dimensions

The time-independent Schrödinger equation for an arbitrary potential \( V(r) \) is given by

\[
-\frac{\hbar^2}{2\mu} \Delta \psi(r) = \{ E - V(r) \} \psi(r), \tag{3}
\]

with the total wave function and the three-dimensional Morse potential are written as follows: [5, 6]

\[
\psi(r) = \frac{1}{r} R_{nl}(r) Y_{lm}^n(\theta, \phi) \tag{4}
\]

\[
V(r) = D \left\{ e^{-\alpha(r-r_e)} - 2e^{-\alpha(r-r_e)} \right\}, \quad D > 0, \quad \alpha > 0, \tag{5}
\]

where \( D \) is dissociation energy, \( r_e \) is the equilibrium internuclear distance and \( \alpha \) is the parameter controlling the width of the potential well. In order to solve Eq. (3), we first put \( x = (r - r_e)/r_e \) and then rewrite the potential function in terms of \( x \) as follows

\[
V(x) = D \left( e^{-2\alpha x} - 2e^{-\alpha x} \right), \quad \alpha = \alpha r_e. \tag{6}
\]

Now, inserting Eqs. (6) and (4) into (3) we obtain

\[
\left\{ -\frac{d^2}{dx^2} - \frac{2\mu \alpha}{\hbar^2} \left[ D \left( e^{-2\alpha x} - 2e^{-\alpha x} \right) - E_{nl} \right] + \frac{l(l+1)}{(r+1)^2} R_{nl}(r) \right\} = 0, \tag{7}
\]

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where \( n \) and \( l \) are the vibration-rotation quantum numbers, \( \mu \) is the reduced mass of the diatomic molecule, and \( E_{nl} \) is the appropriate energy eigenvalue. Now adopting the following approximation [7, 8]

\[
\frac{1}{(1 + x)^2} \approx C_0 + C_1 e^{-\alpha r} + C_2 e^{-2\alpha r},
\]

(8)

with

\[
C_0 = \frac{3}{\alpha} + \frac{3}{\alpha^2},
\]

(9)

\[
C_1 = \frac{4}{\alpha} + \frac{6}{\alpha^2},
\]

(10)

\[
C_2 = -\frac{1}{\alpha} + \frac{3}{\alpha^2},
\]

(11)

and substituting (8) into (7), equation (7) is transformed into

\[
\left( \frac{d^2}{dx^2} - \eta^2 e^{-2\alpha x} + 2\zeta^2 e^{-\alpha x} - \beta_1^2 \right) R_{nl}(x) = 0,
\]

(12)

with

\[
\beta_1^2 = -\frac{2\mu r^2}{\hbar^2} E_{nl} + l(l + 1) C_0,
\]

(13)

\[
\zeta^2 = \frac{2\mu r^2 \hbar^2}{D} \frac{l(l + 1)}{2} C_1,
\]

(14)

\[
\eta^2 = \frac{2\mu r^2 \hbar^2}{D} + l(l + 1) C_2.
\]

(15)

Solving Eq. (12), we found that the rotation-vibrational energy levels of the Morse potential for diatomic molecules are given by [7]

\[
E_{nl} = \frac{\hbar^2}{2\mu r^2} \left\{ l(l + 1) C_0 - \alpha^2 \left( n + \frac{1}{2} - \frac{\zeta^2}{\eta \alpha} \right)^2 \right\},
\]

\[
n = 0, 1, 2, \ldots n_{\text{max}},
\]

(16)

with \( n_{\text{max}} \) denotes the upper bound. In this stage, two remarks seem important to notice:

- Equation (16) is obtained by using the Pekeris approximation. The validity of this approximation depends on the magnitude of the rotational quantum number \( l \): so it is not reliable for higher values of \( l \) [27].

- The closest to the maximum value allowed can be determined by putting the following condition \( dE/\!\!\!d\eta = 0 \) [20–26]: following this condition, we obtain that

\[
n_{\text{max}} = \frac{\zeta^2}{\alpha \eta} - \frac{1}{2},
\]

(17)

with \( \zeta \) and \( \eta \) are defined by Eqs. (14) and (15).

---

**Table I. Spectroscopic parameters of selected molecules used in the present calculation.**

<table>
<thead>
<tr>
<th>Molecule</th>
<th>( \mu )(amu)</th>
<th>( D )(eV)</th>
<th>( a )(Å(^{-1}))</th>
<th>( r_e )</th>
</tr>
</thead>
<tbody>
<tr>
<td>HCl</td>
<td>0.9801045</td>
<td>4.61907</td>
<td>2.38057</td>
<td>1.2746</td>
</tr>
<tr>
<td>H(_2)</td>
<td>0.50391</td>
<td>4.7446</td>
<td>1.440558</td>
<td>0.7416</td>
</tr>
<tr>
<td>CO</td>
<td>6.8606719</td>
<td>11.2256</td>
<td>2.59441</td>
<td>1.1283</td>
</tr>
<tr>
<td>LiH</td>
<td>0.8801221</td>
<td>2.515287</td>
<td>1.7998368</td>
<td>1.5956</td>
</tr>
</tbody>
</table>

Now, we are ready (i) to compute the ro-vibrational energy levels of diatomic molecules and (ii) to discuss our results about the variation of the energies in different situations. For convenience, we choose four molecules, \( H_2 \), LiH, HCl, and CO, which have been most widely studied in the literature. The spectroscopic parameters of selected molecules are shown in Table I [27].

In Fig. 1, representative plots are given, with respect to the vibrational quantum number \( l \) at four selected values of \( n = 0, 5, 10, 50 \) for \( H_2 \), LiH, HCl and CO molecules. In the other hand, Fig. 2 correspondings to the energy changes for rotational quantum number \( n \) at four selected \( l \) values, namely 0, 10, 20, 30. Following these figures, some remarks can be made [28]:

- The choice of a fixed value of \( n \) is a consequence of the fact that this potential supports a limited number of bound states for these molecules; the estimated \( n_{\text{max}} \) for our molecules under consideration are shown in Table II; these values are obtained by using that \( \hbar c = 1973.269 \) eV, Å and 1amu = \( 931.5 \times 10^6 \) eV (Å\(^{-1}\)). Following this table, contrary to the case of \( H_2 \), the number of quantum levels are finite and decrease with the quantum number \( l \).

- These figures allow us to obtain the allowed values of \( l \) leaving our spectrum of energy negative (bound states): the positive values of energy mean that the state is in continuum.

- Also, it is seen that the energies \( E_{nl} \) versus \( n \) plots for different \( l \) become more closely spaced, with \( H_2 \) showing maximum sparsity. The rate of increase in energy, in general, increases as one move toward \( H_2 \rightarrow CO 

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Finally, in order to understand well these observations, we have plotted the vibrational energy level vs both vibrational quantum numbers \((n, l)\) in Figs. 3 and 4, following these figures, we observe that the molecule CO shows a linear behavior, and both molecules HCl and LiH have a similar variation. To our knowledge, such detailed energy plots have not been presented, and we hope these results would be helpful for future studies: precisely, to show the interval when the spectrum of energies becomes positive: this condition can be used as a rule of the selection of allowed values of \(l\) and \(n\) and consequently, used them for calculating the thermal properties of these molecules.

3. Thermal properties of the Morse potential for some Diatomic molecules in three dimensions

3.1. Vibrational partition function

As we know, all thermodynamic quantities can be obtained from the partition function \(Z\); therefore, the partition function of the system is the starting point to derive all thermal properties of the system in question. This function can be calculated by direct summation over all possible vibrational energy levels available to the system. Given the energy spectrum, the partition function \(Z\) at finite temperature \(T\) is obtained through the Boltzmann factor is

\[
Z = \sum_{n=0}^{\infty} e^{-\beta(E_n - E_0)},
\]

with \(\beta = 1/k_B T\), where \(k_B\) is the Boltzmann constant.

Now in order to calculate the \(Z\) function, we use the Euler-MacLaurin formula: according to this approach, the sum transforms to the integral as follows (see Appendix A):

\[
\sum_{n=0}^{\infty} f(x) = \frac{1}{2} f(0) + \int_{0}^{\infty} f(x) \, dx - \sum_{p=1}^{\infty} \frac{B_{2p}}{(2p)!} f^{(2p-1)}(0),
\]

where \(B_{2p}\) are the Bernoulli numbers, and \(f^{(2p-1)}\) is the derivative of order \((2p - 1)\). Up to \(p = 3\), the partition function \(Z\) is written as

\[
Z = \sum_{n=0}^{\infty} e^{-\beta(E_n - E_0)},
\]

Figure 1. The ro-vibrational energy level to vibrational quantum number \(l\) for different values of vibrational quantum number \(n\).
Figure 2. The ro-vibrational energy level versus vibrational quantum number \( n \) for different values of vibrational quantum number \( l \).

\[
Z = \sum_{n=0}^{n_{\text{max}}} e^{-\beta(E_{nl}-E_0)} = \frac{1}{2} + \int_0^{n_{\text{max}}} f(n) \, dn - \sum_{p=1}^{\infty} \frac{B_{2p}}{(2p)!} f^{(2p-1)}(0), \tag{20}
\]

with \( B_2 = 1/6 \) and \( B_4 = -1/30 \) and

\[
f(x) = e^{-\beta(E_{nl}-E_0)}. \tag{21}
\]

In our case, we focus only on the study on the partition function \( Z \) and the specific heat \( C_v \). The influence of the quantum parameter \( l \) on these functions will be also examined.

### 3.2. Applications for some diatomic molecules

In the present work, we choose four diatomic molecules: \( \text{H}_2 \), HCl, LiH, and CO, which have been most widely studied in the literature. The typical values of molecular constants for the electronic state of these molecules are given in Table I.

\[
F = -\frac{1}{\beta} \ln Z, \tag{22}
\]

\[
U = -\frac{d \ln Z}{d \beta}, \tag{23}
\]

\[
S = \ln Z - \beta \frac{d \ln Z}{d \beta}, \tag{24}
\]

\[
C_v = \beta^2 \frac{d^2 \ln Z}{d \beta^2}. \tag{25}
\]
With the aid of these values, the vibrational partition function $Z$ is determinate; via this function, the thermal properties of these molecules can be found easily. These thermodynamic functions are represented versus the inverse of temperature $\beta = 1/k_B T$ and the quantum number $l$. The Fig. 4(d) show some thermal quantities for the following diatomic molecules, $\mathrm{H}_2$, $\mathrm{HCl}$, $\mathrm{LiH}$, and $\mathrm{CO}$: from this figure, some remarks can be made:

- The Fig. 4(a) shows both the partition function $Z$ and the reduced specific heat $C_v/k_B$ of $\mathrm{H}_2$ versus $\beta$ for different values of vibrational quantum number $l$: it reveals that for increasing inverse temperature $\beta$ the par-
Figure 5. Thermal properties for the following Diatomic molecules, H₂, H Cl, LiH, and CO. (a) Both the vibrational partition function \( Z \) and the reduced specific heat \( C/k_B \) of H₂ as a function of \( \beta \) for different values of vibrational quantum number \( l \). (b) Both the vibrational partition function \( Z \) and the reduced specific heat \( C/k_B \) of CO as a function of \( \beta \) for different values of vibrational quantum number \( l \). (c) Both the vibrational partition function \( Z \) and the reduced specific heat \( C/k_B \) of HCl as a function of \( \beta \) for different values of vibrational quantum number \( l \). (d) Both the vibrational partition function \( Z \) and the reduced specific heat \( C/k_B \) of LiH as a function of \( \beta \) for different values of vibrational quantum number \( l \).
The partition function is decreased as well. The variation of the reduced specific heat versus $\beta$ for various values of $l$ in the interval $0 \leq \beta \leq 5$ is shown in Fig. 5(a). We observe that the $C_v/k_B$ first increases with the increasing $\beta$ until it reaches the maximum value $\beta_C = 1/k_B T_C$ and then decreases with it: we can see that this maximum value, $T_C$, decreases when $l$ increases (see Table II).

- The Fig. 5(b) shows both vibrational partition function $Z$ and the reduced specific heat $C_v/k_B$ of CO as a function of $\beta$ for different values of vibrational quantum number $l$: from these figures, we observe that both the partition function and $C_v/k_B$ are independent of the parameter $l$: so, we can state that all states of this molecule are identical.

- Finally, Figs. 5(c) and 5(d) show both the vibrational partition function $Z$ and the reduced specific heat $C_v/k_B$ of HCl and LiH as a function of $\beta$ for different values of the quantum number $l$: from these figures, we observe that the behavior of both molecules is identical. This comportment is because both molecules have similar spectroscopic parameters (see Table I).

In Fig. 6 we show the behavior of the reduced specific heat of the diatomic molecules $H_2$, HCl, LiH, and CO for two values of $l$: from this figure, we observe that the specific heat of $H_2$, HCl is approximately the same, contrarily to the case of the molecules LiH and CO where the difference is well observed.

In Table III we show some values of the critical temperature $T_C = 1/k_B \beta_C$, and its variation with the parameter $l$: as explained above, the condition of having the bound states (negative spectrum of energy) leads to the limitation of the quantum number $l$: following this, the relative value $T_C$ for each molecule, is $T_C^{H_2} = 8788$ K ($l \leq 10$), $T_C^{HCl} = 9355$, $T_C^{LiH} = 8923$, $T_C^{CO} = 20715$ K. The case of the molecule CO requires some attention: all curves are identical and independent of the quantum number $l$. It means that all states of the molecule CO can be considered as in the $S$ state ($l = 0$).

Also, we can see that the reduced vibrational specific heat $C_v/k_B$ is more sensitive to $n_{max}$ than the vibrational partition function. The reason of this is twofold: (i) the specific heat depends upon the second derivative of the partition function and (ii) the expansion of the specific heat as a function of $l$ is very clear in the figure for the different types of molecules: this enlargement can be explained by the decreases in the number of energy levels $n_{max}$ when $l$ decreases except to the case of $H_2$. Thus, for a system composed of diatomic molecules, a critical temperature value $T_C$ appears. This temperature means that the system becomes saturated and can no longer absorb more energy because all its excited states are occupied. We note here that this temperature was at first mentioned by [26,29]. It corresponds to the maximum in the specific heat curves. These curves show the anomalous behavior of the specific heat when diatomic systems interact under the Morse potential.
Finally, the specific heat is an important physical quantity for testing the existence of a transition phase, as well as its nature (first or second-order). Recently, predictions of entropy for diatomic molecules and gaseous substances have been the subject of two recent studies [14, 15]. The authors have obtained the exact expression of the vibrational entropy for (i) the improved Tietz oscillator and (ii) the Morse and Manning-Rosen oscillators. According to the results obtained by these authors, two remarks can be made: firstly, the diatomic molecules are treated as rigid rotors, and the interaction between two molecules is neglected. On the other hand, the specific heat values derived from experimental measurements are a combination of the translational, rotational, and vibrational specific heat. Thus, in order to compare our theoretical vibrational specific heat, it must have the experimental data of the total specific heat at first. But, we stress, though, as far as we know, we have not any data concerning the experimental values of the total specific heat that helps us to make a comparison with our theoretical results.

4. Conclusion

In this work, by using the vibrational energies obtained in the three-dimensional Morse potential, we have carried out a calculation of the vibrational partition function of the Morse potential for some diatomic molecules via the Euler-Maclaurin approach. Via this function, we have derived explicit expressions for the thermodynamic functions such as specific heat $C_v$. We have analyzed the behavior of the specific heat: this analysis shows the existence of a critical temperature $T_C$ in the curves of the specific heat: this temperature decreases when $l$ increases.

Appendix

A. Euler-Maclaurin formula and Poisson summation

A.1 Euler-Maclaurin Formula

In mathematics, the Euler-Maclaurin formula [31,32] is a formula for the difference between an integral and a closely related sum. It can be used to approximate integrals by finite sums, or conversely to evaluate finite sums and infinite series using integrals and the machinery of calculus.

If $m$ and $n$ are natural numbers and $f(x)$ is a complex or real, valued continuous function for real numbers $x$ in the interval $[m, n]$, then the integral

$$I = \int_{m}^{n} f(x) \, dx,$$

(A.1)

can be approximated by the sum (rectangle method)

$$S = f(m + 1) + \ldots + f(n - 1) + f(n).$$

(A.2)

The Euler-Maclaurin formula provides expressions for the difference between the sum and the integral in terms of the higher derivatives $f^{(k)}(x)$ evaluated at the endpoints of the interval, that is to say when $x = m$ and $x = n$. Thus, for $p$ a positive integer and a function $f(x)$ that is $p$ times continuously differentiable in the interval $[m, n]$, we have

$$S - I = \sum_{k=1}^{p} \frac{B_k}{k!} \left\{ f^{(k-1)}(n) - f^{(k-1)}(m) \right\} + R_p,$$

(A.3)

where $B_k$ is the $k$th Bernoulli number and $R_p$ is an error term which depends on $n, m, p$, and $f$ and is usually small for suitable values of $p$.

The formula is often written with the subscript taking only even values since the odd Bernoulli numbers are zero except for $B_1$. In this case, we have

$$\sum_{i=m}^{n} f(i) = \int_{m}^{n} f(x) \, dx + \frac{f(n) + f(m)}{2} + \sum_{k=1}^{p} \frac{B_{2k}}{(2k)!} \left\{ f^{(2k-1)}(n) - f^{(2k-1)}(m) \right\} + R_p.$$

(A.4)

Finally, in the context of computing asymptotic expansions of sums and series, usually the most useful form of the Euler-Maclaurin formula is

$$\sum_{n=a}^{b} f(n) \sim \int_{a}^{b} f(x) \, dx + \frac{f(b) + f(a)}{2} + \sum_{k=1}^{\infty} \frac{B_{2k}}{(2k)!} \left\{ f^{(2k-1)}(b) - f^{(2k-1)}(a) \right\}.$$

(A.5)

Now, when $a = 0$ and $b = \infty$, we have

$$\sum_{n=0}^{\infty} f(n) \sim \int_{0}^{\infty} f(x) \, dx + \frac{f(0)}{2} - \sum_{k=1}^{\infty} \frac{B_{2k}}{(2k)!} f^{(2k-1)}(0).$$

(A.6)

A.2 Poisson summation Formula

For appropriate functions $f$, the Poisson summation formula may be stated as:

$$\sum_{n=-\infty}^{+\infty} f(n) = \sum_{n=-\infty}^{+\infty} \hat{f}(n),$$

(A.7)

where $\hat{f}(n)$ is the Fourier transform of $f$; that is $\hat{f}(n) = \mathcal{F}\{f(n)\}$.

Consider the function

$$F(x) = \sum_{n \in \mathbb{Z}} f(x + n).$$

This is a periodic function of period 1: therefore, we can take its Fourier series expansion

$$F(x) = \sum_{n \in \mathbb{Z}} a_n e^{2\pi i n x},$$

(A.8)

where
B.2 Case of \( l \)

\[
a_n = \int_0^1 F(x) e^{-2\pi inx} dx = \int_0^1 \sum_{m \in \mathbb{Z}} f(x + m) e^{-2\pi inx} dx = \sum_{m \in \mathbb{Z}} \int f(x + m) e^{-2\pi inx} dx
\]

\[
= \sum_{m \in \mathbb{Z}} \int f(x + m) e^{-2\pi in(x + m)} dx (x + m) = \int f(t) e^{-2\pi int} dt = \hat{f}(n).
\]

Therefore:

\[
\sum_{n = -\infty}^{+\infty} f(x + n) = F(x) = \sum_{k = -\infty}^{+\infty} \hat{f}(n) e^{2\pi inx}.
\]

and the result follows by evaluating at \( x = 0 \).

**B. Vibrational partition function for some values of vibrational quantum number \( l \)**

**B.1 Case of \( l = 0 \)**

\[
Z_{\text{H}_2} = \frac{1}{2} + \frac{1}{720} (0.00626265\beta^3 - 0.020674\beta^2) + 0.0330932\beta
\]

\[
+ \frac{e^{-5.45387\beta} (9.51412\text{erfi}(2.13164\sqrt{\beta}) - 9.51412\text{erfi}(0.0823662\sqrt{\beta}))}{\sqrt{\beta}}.
\]

(B.1)

\[
Z_{\text{C}_2} = \frac{1}{2} + \frac{1}{720} (0.0273855\beta^3 - 0.00370934\beta^2) + 0.0251184\beta
\]

\[
+ \frac{e^{-11.0743\beta} (19.5686\text{erfi}(3.3278\sqrt{\beta}) - 19.5686\text{erfi}(0.0217579\sqrt{\beta}))}{\sqrt{\beta}}.
\]

(B.2)

\[
Z_{\text{HCl}} = \frac{1}{2} + \frac{1}{720} (0.0976538\beta^3 - 0.0333989\beta^2) + 0.038375\beta
\]

\[
+ \frac{e^{-3.858\beta} (8.06064\text{erfi}(2.09423\sqrt{\beta}) - 8.06064\text{erfi}(0.00527475\sqrt{\beta}))}{\sqrt{\beta}}.
\]

(B.3)

\[
Z_{\text{LiH}} = \frac{1}{2} + \frac{1}{720} (0.019802\beta^3 - 0.0124903\beta^2)
\]

\[
+ 0.0225453\beta + \frac{e^{-2.37809\beta} (10.1031\text{erfi}(1.54211\sqrt{\beta}) - 10.1031\text{erfi}(0.0508877\sqrt{\beta}))}{\sqrt{\beta}}.
\]

(B.4)

**B.2 Case of \( l = 30 \)**

\[
Z_{\text{H}_2} = \frac{1}{2} + \frac{1}{720} (0.0919104\beta^3 - 0.0234941\beta^2) + 0.0376074\beta
\]

\[
+ \frac{e^{-5.86809\beta} (9.51412\text{erfi}(2.42241\sqrt{\beta}) - 9.51412\text{erfi}(0.000550072\sqrt{\beta}))}{\sqrt{\beta}}.
\]

(B.5)

\[
Z_{\text{C}_2} = \frac{1}{2} + \frac{1}{720} (0.0268359\beta^3 - 0.00368436\beta^2) + 0.0249493\beta
\]

\[
+ \frac{e^{-10.9256\beta} (19.5686\text{erfi}(3.30539\sqrt{\beta}) - 19.5686\text{erfi}(0.0446334\sqrt{\beta}))}{\sqrt{\beta}}.
\]

(B.6)
\[ Z_{\text{HCl}} = \frac{1}{2} + \frac{1}{720} \left( -0.0735999\beta^3 - 0.0303945\beta^2 + 0.034923\beta \right) 
+ \frac{e^{-3.63224/\beta}}{\sqrt{\beta}} \left( 8.06064\text{erfi} \left( 1.90584\sqrt{\beta} \right) - 8.06064\text{erfi} \left( 0.0367777\sqrt{\beta} \right) \right), \quad (B.7) \]

\[ Z_{\text{LiH}} = \frac{1}{2} + \frac{1}{720} \left( -0.0134738\beta^3 - 0.0109858\beta^2 + 0.0198297\beta \right) 
+ \frac{e^{-1.83971/\beta}}{\sqrt{\beta}} \left( 10.1031\text{erfi} \left( 1.35636\sqrt{\beta} \right) - 10.1031\text{erfi} \left( 0.0405763\sqrt{\beta} \right) \right), \quad (B.8) \]

with the imaginary error function, denoted \text{erfi}, is defined as [32]

\[ \text{erfi}(x) = -i\text{erf}(ix) = \frac{2}{\sqrt{\pi}} \int_0^x e^{t^2} dt. \quad (B.9) \]

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