Relationship between the 2-body Energy of the Biswas-Hamann and the Murrell-Mottram Potential Functions

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Z. Naturforsch. 59a, 116 – 118 (2004); received December 30, 2003

The two-body interactions in the Biswas-Hamann (BH) and Murrell-Mottram (MM) potential functions are analytically related in this paper by equating the zeroth to second differentials at equilibrium bond length. By invoking the Maclaurin series expansion for the exponential term, the MM potential function could be expressed in a manner that enables comparison of repulsive and attractive terms. Approximate and refined sets of scaling factors were obtained upon comparing the indices and coefficients, respectively. Finally, the suitability for each set of scaling functions is discussed in terms of the “softness” of the bonds.

Key words: 2-body Energy; Empirical Potential Functions; Parametric Conversion.

1. Introduction

Notwithstanding the lack of fundamental basis in comparison to the quantum mechanical method that requires solving the Schrödinger equation, empirical potential energy functions find their worth in computation of large-scale many-body condensed matter systems coupled with large distortion, due to their convenient execution. With various potential functions proposed, parameters in these functions were parametrized for a number of elements in crystalline, microcluster and other structural arrangements. An issue encountered for these empirical potential functions is that each of these functions requires different sets of parameters. Thus parametrization – which includes experimentation and curve-fitting – can be costly and time consuming. A more economical and rapid approach for obtaining a set of potential function parameters from another potential function is by analytical conversion of their parameters. Recently parametric relationships have been developed for converting parameters among bond-stretching potential functions [1], bond-bending potential functions [2], bond-twisting potential functions [3], and van der Waals potential functions [4–6]. Upon refinements [7], a prototype molecular potential function converter has been developed in a spreadsheet format [8]. However, a majority of these relationships are useful only to the chemistry community, as the parameters that were related were based on potential functions used in physical/computational chemistry softwares. Within the context of condensed matter, Stoneham et al. [9] performed a comparison of eight valence-force potentials for silicon. However, valence-force potentials are useful only for describing small distortions from equilibrium. A group of six potential functions was compared in detail by Balamane et al. [10], but these functions were studied without analytical conversion amongst the potential functions. In this paper, a set of analytical relationships is developed for converting parameters among the 2-body interactions of the Biswas-Hamann (BH) and the Murrell-Mottram (MM) potential functions. BH and MM for parametric connection was selected because these two potential functions were developed for silicon [11–13]. Another reason for their selection was that both BH and MM belong to a class of potential functions where the total energy of a system is the sum of the energy in the 2-body and 3-body interactions:

$$\Phi = \phi_{\text{2-body}} + \phi_{\text{3-body}} = \sum_{i<j} U_{ij} + \sum_{i<j<k} W_{ijk}.$$  \hfill (1)

Generally speaking, (1) is a subset of the Born-Oppenheimer approximation. However higher moments (e.g. 4-body and higher) are neglected due to quick convergence [14]. Specifically, the 2-body interactions for the BH and MM potential functions were given as

$$U_{\text{BH}} = A_1 \exp(-\lambda_1 r) + A_2 \exp(-\lambda_2 r)$$  \hfill (2)

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we have the following 3 independent equations:

\[ U_{\text{BH}} = U_{\text{MM}}(D, a, R, r) \]
\[ U_{\text{BH}}(A_i, A_2, \lambda_1, \lambda_2, r) \]
\[ \lambda_1 = \frac{\xi_1}{\xi_2} \]
\[ \lambda_2 = \frac{\xi_2}{\xi_2} \]

and

\[ U_{\text{MM}} = -D \left[ 1 + a \left( \frac{r}{R} - 1 \right) \right] \exp \left( -a \left( \frac{r}{R} - 1 \right) \right), \quad (3) \]

respectively. The relationship between BH parameters \((A_i, \lambda_i \text{ for } i=1,2)\) and MM parameters \((D, a, R)\) is developed in the next section.

2. Analysis

Equating the 2-body BH and MM potential functions from the zeroth to the second differential with respect to the interatomic distance, \(r\), at the equilibrium bond length, \(R\),

\[ \left( \frac{d^2 U_{\text{BH}}}{dr^2} \right)_{r=R} = \left( \frac{d^2 U_{\text{MM}}}{dr^2} \right)_{r=R}; \quad n = 0, 1, 2, (4) \]

we have the following 3 independent equations:

\[ A_1 \exp(-\xi_1) + A_2 \exp(-\xi_2) = -D, \quad (5) \]
\[ \xi_1 A_i \exp(-\xi_1) + \xi_2 A_2 \exp(-\xi_2) = 0, \quad (6) \]
\[ \xi_1^2 A_1 \exp(-\xi_1) + \xi_2^2 A_2 \exp(-\xi_2) = a^2 D, \quad (7) \]

with scaling factors

\[ \xi_2 = \lambda_i R; \quad (i = 1, 2). \quad (8) \]

Eliminating the terms \(A_i \exp(-\xi_i)\) for \(i = 1, 2\) from (5)–(7) and solving simultaneously leads to the upper and lower solutions

\[ \left\{ \begin{array}{l} \xi_1^{\text{up}} \\ \xi_1^{\text{low}} \end{array} \right\} = \left\{ \begin{array}{l} \xi_2 \\ (a^2 / \xi_2) \end{array} \right\} \quad (9) \]

By applying (6) and (8), the original 2-body BH potential given by (2) can be expressed as

\[ \frac{U_{\text{BH}}}{A_1} = \exp \left( -\xi_1 \frac{r}{R} \right) - \xi_1 \exp \left[ -\xi_1 + \xi_2 \left( 1 - \frac{r}{R} \right) \right], \quad (10) \]

Eliminating the term \(A_2 \exp(-\xi_2)\) from (5) and (6) gives

\[ A_1 = D \left( \frac{\xi_2}{\xi_1 - \xi_2} \right) \exp(\xi_1) \quad (11) \]

which, upon substitution into (10), leads to the loose form of (2)

\[ \frac{U_{\text{BH}}}{D} = \left( \frac{\xi_2}{\xi_1 - \xi_2} \right) \exp \left[ \xi_1 \left( 1 - \frac{r}{R} \right) \right] - \left( \frac{\xi_1}{\xi_1 - \xi_2} \right) \exp \left[ \xi_2 \left( 1 - \frac{r}{R} \right) \right]. \quad (12) \]

As such, (12) shows that the upper solution of (9) is invalid. In paving a way for relating parameters of these potential functions, we rewrite the MM potential in (3) as

\[ \frac{U_{\text{MM}}}{D} = \left( a \left( 1 - \frac{r}{R} \right) - 1 \right) \exp \left[ a \left( 1 - \frac{r}{R} \right) \right]. \quad (13) \]

Upon recalling the Maclaurin series expansion of the exponential term

\[ \exp \left[ a \left( 1 - \frac{r}{R} \right) \right] = \sum_{n=0}^{\infty} \frac{a^n \left( 1 - \frac{r}{R} \right)^n}{n!}, \quad (14) \]

we have the approximation

\[ a \left( 1 - \frac{r}{R} \right) \approx \exp \left[ a \left( 1 - \frac{r}{R} \right) \right] - 1 \quad (15) \]

which, upon substitution into (3), gives the approximate MM function

\[ \frac{U_{\text{MM}}^{\text{approx}}}{D} = \exp \left[ 2a \left( 1 - \frac{r}{R} \right) \right] - 2 \exp \left[ a \left( 1 - \frac{r}{R} \right) \right] \quad (16) \]

Now, comparison of the exponential indices of (12) and (16) gives \(\xi_1 = 2a\) and \(\xi_2 = a\). On the other hand, comparing the coefficients to the exponential terms these equations generally indicate that

\[ \xi_1 = 2\xi_2. \quad (17) \]
Hence substitution of (17) into the lower solution of (9) leads to $\xi_1 = \sqrt{2}a$ and $\xi_2 = a/\sqrt{2}$. A summary of the parametric relationships for the 2-body interactions of the BH and MM potential functions is furnished in Table 1.

3. Results and Discussion

For verification of the developed parametric relationships, curves of non-dimensional 2-body BH energy were plotted against the non-dimensional interatomic distance based on MM parameters. Parametrized values of $a$ are 5.5 for silicon crystals [12, 13], and 6.5 for silicon microclusters [15]. We select the average value of $a = 6$ for convenient illustration. Figure 1 shows a plot of the 2-body MM curve, and the corresponding BH curves based on parametric conversion from MM parameters. We observe an overestimation when the BH curve is plotted based on the approximate conversion from MM parameters. When the refined relationship is used, the BH curve gives an excellent correlation to the MM curve. The higher slope at the inflexion point corresponding to the approximate BH curve suggests its usefulness in curve-fitting “hard” bonds, which exhibit a steep rise in energy from equilibrium, followed by an abrupt leveling off of the 2-body interaction energy. The lower slope at the inflexion point corresponding to the refined BH curve indicates its usefulness in curve-fitting of “soft” bonds, characterized by a more gradual rise in 2-body energy from the equilibrium point. Discrepancies may well be attributed to the approximation adopted in (15) but, as evident from Fig. 1, the discrepancy is insignificant when the refined relation is used.

4. Conclusions and Suggestion

A method for theoretically relating the 2-body potential energy functions in the BH and MM potentials has been described and graphically verified. Essentially, both potential functions were equated at the equilibrium bond length for the zeroth, first and second differentials. Comparison of the repulsive and attractive terms gives two sets of scaling factors. An approximate and a refined set of scaling factors is obtained when the indices and the coefficients are equated, respectively. The approximate relations give a higher estimation of the BH curve based on the MM parameters – thereby more suited for curve-fitting “hard” bonds, while the refined relation gives excellent correlation. It is suggested that 3-body potentials in the BH and MM potential functions are considered in future work.