LETTER TO THE EDITOR

Comment on “Exploring nature and predicting strength of hydrogen bonds: A correlation analysis between atoms-in-molecules descriptors, binding energies, and energy components of symmetry-adapted perturbation theory”

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Abstract
We evaluate the correlation between binding energy (BE) and electron density $\rho(r)$ at the bond critical point for 28 neutral hydrogen bonds, recently reported by Emamian and co-workers (J. Comput. Chem., 2019, 40, 2868). As an efficient tool, we use local stretching force constant $k^O_{\text{HB}}$ derived from the local vibrational mode theory of Konkoli and Cremer. We compare the physical nature of BE versus $k^O_{\text{HB}}$, and provide an important explanation for cases with significant deviation in the BE–$k^O_{\text{HB}}$ relation as well as in the BE–$\rho(r)$ correlation. We also show that care has to be taken when different hydrogen bond strength measures are compared. The BE is a cumulative hydrogen bond strength measure while $k^O_{\text{HB}}$ is a local measure of hydrogen bond strength covering different aspects of bonding. A simplified and unified description of hydrogen bonding is not always possible and needs an in-depth understanding of the systems involved.

KEYWORDS
binding energy, electron density, hydrogen bond, local hydrogen bond energy, local vibrational mode theory

In a recent article, Emamian and co-workers judiciously selected a diverse collection of 42 hydrogen-bonded dimers including 28 neutral and 14 charged complexes, in order to study the nature of hydrogen bonds with quantum chemistry. The geometry of these 42 dimer complexes was optimized at B3LYP-D3(BJ)/ma-TZVPP level$^{2-4}$ and the binding energies (BEs) characterizing the strength of H-bonding were calculated at CCSD(T)/jul-cc-pVTZ level$^{5,6}$ employing half of basis set superposition error correction.$^{7,8}$ The authors utilized symmetry-adapted perturbation theory (SAPT) at the SAPT2 + (3)iMP2/aug-cc-pVTZ level$^{9-11}$ to decompose the BEs into energy components and proposed a novel classification of hydrogen bonding into four categories: very weak, weak to medium, medium, and strong. Emamian and co-workers observed fairly strong linear correlation between the CCSD(T) BEs and the electron density $\rho(r)$ at the (3,−1) hydrogen bond critical point (BCP) for both the neutral ($R^2 = 0.9732$) and charged ($R^2 = 0.9644$) complexes. They advocated the use of this correlation between BE and $\rho(r)$ as a convenient way to predict in the molecular systems the BE of individual hydrogen bonds when these BEs are unattainable. We reassessed the correlation between BE and $\rho(r)$ for those hydrogen bonds in the 28 neutral complexes by utilizing the local vibrational mode theory$^{12,13}$ and obtained interesting insights which provide a different perspective complementing the work of Emamian and co-workers.

As shown in Figure 1(A), the correlation between BE and $\rho(r)$ at the BCP of 28 neutral hydrogen bonds contains four cases with significant deviation identified with a deviation criterion of 1.5$\sigma$ ($\sigma$ is the standard deviation of the residuals after fitting). However, Emamian and co-workers did not provide any remarks on possible reasons.

In order to better understand and assess the above correlation, we employed the local vibrational mode theory originally developed...
by Konkoli and Cremer to calculate the local stretching force constant \( k_{a}^{HB} \) of these 28 hydrogen bonds (collected in Table 1) based on the Hessian (second-order derivatives of energy) matrices of the equilibrium geometries optimized at the B3LYP-D3(BJ)/ma-TZVPP level.

Konkoli and Cremer derived in 1998 the local vibrational modes associated with individual internal coordinates \( q_{n} \) (e.g., H\_D–H/C1/C1/C1 \_Ahydrogen bond length in D–H–D hydrogen bonding) directly from the normal vibrational modes (within the harmonic approximation) by solving the mass-decoupled Euler-Lagrange equations. The local vibrational mode associated with a bond stretching can be considered as the motion being obtained via an infinitesimal change in the bond length followed by the relaxation of all other atoms in the molecular system. Each local mode has its corresponding local mode frequency \( \omega_{a}^{n} \) and local mode force constant \( k_{a}^{n} \) (subscript \( n \) represents the target internal coordinate parameter \( q_{n} \)). The local mode force constant is independent of atomic masses and characterizes pure electronic structure effects. The underlying physical nature of local mode force constant associated with a chemical bond is the curvature of the Born–Oppenheimer potential energy surface (PES) in the direction of the bond (i.e., diatomic) stretching. Therefore, the local stretching force constant has been extensively used to quantify the...
The linear correlation between BE and local stretching force constant $k_{HB}^a$ (see Figure 3(A)) is in the same range as the correlation between BE and $\rho(\mathbf{r})$. On one hand, Emamian and co-workers carefully picked small-sized molecules for their set of 28 neutral hydrogen bonded dimers with marginal secondary interactions, i.e. additional interactions between the two monomers besides the target hydrogen bonding, so that in this particular case the BE predominantly reflects the pure hydrogen bonding strength. On the other hand, the local stretching force constant $k_{HB}^a$ with its physical nature as the curvature of PES in the direction of hydrogen bond stretching has been widely recognized as an intrinsic bond strength descriptor derived from vibrational spectroscopy. Therefore, it is of interest to check in more detail to what extent BE and $k_{HB}^a$ as two different hydrogen bond strength descriptors are compatible with each other.

We got particularly interested in the five cases with significant deviation in the BE versus $k_{HB}^a$ correlation identified with the deviation criterion of $1.5/\sigma$ (see Figure 3(A)) because four of them overlap with the significantly deviated points in the BE versus $\rho(\mathbf{r})$ correlation.

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The intrinsic strength of both covalent bonds \(^{14-16}\) and noncovalent interactions including hydrogen\(^{17-20}\), halogen\(^{21-24}\), chalcogen\(^{25-27}\), pnicogen\(^{28-30}\), and tetrel bonding\(^{31}\). Concerning the detailed mathematical derivation of local vibrational mode theory, interested readers are referred to a recent review article.\(^{13}\)

We observed a rather strong correlation between the local stretching force constant $k_{HB}^a$ and $\rho(\mathbf{r})$ at the BCP for the 28 neutral hydrogen bonds (see Figure 2(A)) and this correlation is marginally stronger than that between BE and $\rho(\mathbf{r})$ according to the coefficient of determination ($R^2$). For hydrogen bonding denoted as D–H···A (D: donor atom/group; A: acceptor atom/group), the (3,–1) BCP of hydrogen bond H···A is a point between two bonding atoms (H and A) where the first derivative of electron density vanish (i.e., $\nabla_\mathbf{r} \rho(\mathbf{r}) = 0$) while the Hessian of dimension (3×3) for $\rho(\mathbf{r})$ has two negative eigenvalues and one positive eigenvalue.\(^{22}\) Furthermore, the local stretching force constant $k_{HB}^a$ of the H···A hydrogen bond is the curvature of the PES in the direction of H···A stretching. Both $\rho(\mathbf{r})$ and $k_{HB}^a$ are local second-order response properties associated with the H···A bond. This explains the strong correlation and also corroborates the fact that the local mode force constant characterizes the pure electronic structure effects.

The linear correlation between BE and local stretching force constant $k_{HB}^a$ (see Figure 3(A)) is in the same range as the correlation between BE and $\rho(\mathbf{r})$. On one hand, Emamian and co-workers carefully picked small-sized molecules for their set of 28 neutral hydrogen bonded dimers with marginal secondary interactions, i.e. additional interactions between the two monomers besides the target hydrogen bonding, so that in this particular case the BE predominantly reflects the pure hydrogen bonding strength. On the other hand, the local stretching force constant $k_{HB}^a$ with its physical nature as the curvature of PES in the direction of hydrogen bond stretching has been widely recognized as an intrinsic bond strength descriptor derived from vibrational spectroscopy. Therefore, it is of interest to check in more detail to what extent BE and $k_{HB}^a$ as two different hydrogen bond strength descriptors are compatible with each other.

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comparison with local stretching force constants. Emamian and co-workers claimed that the BE is best to exhibit intrinsic binding strength of the monomers without considering the structural relaxation of the monomers. In other words, the BE includes all energy contributions collectively upon complexation of the monomers. As we sketch in Figure 4, the local stretching force constant $k_{HB}^a$ for H--A bond covers the pure hydrogen bond interaction connecting the two monomers, it does not cover additional interactions within the monomers caused by hydrogen bonding. Based on this analysis, we can explain the cases with significant deviation in the correlation between BE and $k_{HB}^a$ shown in Figure 3(A).

The five points with significant deviation can be grouped into two parts including three above the regression line as Group 1+ and two below the regression line as Group 2+. Additionally, Group 0 refers to the standard model obtained after excluding Group 1+ and Group 2+, which will be discussed in detail later. We find that Group 1+ complexes share the similarity that (1) the donor molecule is hydrogen fluoride and (2) the acceptor atom A has more than one lone pair available to delocalize into the $\sigma^*(F-H)$ antibonding orbital. Since $F-H$ as its hydrogen bond donor, it belongs to Group 1+. The fluorine atom is extremely electronegative and therefore tends to attract $F-H$ bonding electron density creating a partially positive charge on hydrogen atom. When hydrogen fluoride and the acceptor monomer is hydrogen bonded, the lone pair electrons from acceptor atom A delocalize into the $\sigma^*(F-H)$ antibonding orbital. The delocalization is further strengthened by the pulling of the...
electronegative fluorine atom. This provides the hydrogen atom in F–H with excess electron density leading to repulsive interaction with the electron density of the fluorine atom. Such a repulsion counting as a destabilizing factor toward the total BE is not reflected in the local stretching force constant $k_{\text{HB}}^a$ explaining why these four data points are located above the regression line.

For Group 2 complexes, the donor monomers are hydrogen azide (HN$_3$) and hydrogen fluoride (HF), both of which are weak acids while the acceptor monomer is ammonia (NH$_3$) as weak base. The association of ammonia and weak acid molecules reminds us of the formation of ammonium salt consisting of an ammonium cation (NH$_4^+$) and an anion. Therefore, in the case of hydrogen bonding between ammonia (NH$_3$) and a weak acid (D–H), the partially ionic character of ammonia salt arising from the delocalization of ammonia lone pair into the $\sigma^*$ (D–H) antibonding orbital (covalent character of H–N) leads to an electrostatic attraction between the anion (D–) and the ammonium cation (H$^+$–NH$_3$)$_3$. Such an ionic attraction counts as a stabilizing factor toward the total BE but is not reflected by the local stretching force constant $k_{\text{HB}}^a$ of H–N explaining why these two cases with significant deviation are found below the regression line.

Furthermore, to demonstrate that the observed deviations to the BE and $k_{\text{HB}}^a$ model are not artifacts of the statistical method used here, but instead have a sound physical basis, we calculated BE, $\rho(r)$ (at the same levels of theory as in the ref. 1), and $k_{\text{HB}}^a$ for a set of newly designed dimers (see Table 2). These were designed to have similar hydrogen bonding environment as in Group 1 or Group 2 dimers, which deviate from the simple hydrogen bond situation due to the presence of additional (de)stabilization. The new dimers contain as their acceptor or donor at least one of the monomers incorporated in the original 28 dimers. For a systematic comparison of how the presence of complexes that deviate from simple hydrogen bond situation can affect the quality of the BE and $k_{\text{HB}}^a$ model, we did the following. First, we removed all the systems belonging to and as characterized by Group 1 and Group 2 dimer classes (i.e., Group 1: (20), (22), (25), (27), and Group 2: (26), (28), respectively) and only considered the rest of the 22 dimers (denoted as Group 0). These dimer systems have the simple situation in their hydrogen bonding nature and the corresponding linear regression line is considered as a standard/simple model between BE and $k_{\text{HB}}^a$, which has $R^2 = 0.9871$ with RMSE = 0.1937 (see Figure 3(B)). Then, with the inclusion of the seven new dimers (denoted by Group 1+ and Group 2+) we did linear regression for the extended data set with 35 dimers, and the corresponding linear regression line between BE and $k_{\text{HB}}^a$ has $R^2 = 0.9413$ with RMSE = 1.0356 (see Figure 3(C)). In comparison to the standard model between BE and $k_{\text{HB}}^a$, we could still see a fairly high correlation for the extended data set, but the deviation (given by RMSE) is much more pronounced now. This is a clear indication that in the presence of complexes that diverge from the simple hydrogen bond situation, the quality of the model is affected and as a caveat we like to point out that careful consideration should be taken when presenting a unified picture of hydrogen bonding under such circumstances. Also, this analysis further confirmed, the identification of the deviations defined as Group 1 and Group 2, is not merely based on the points produced by the statistical method we used here, but rather guided by our chemical intuition. A similar kind of evaluation was also carried out for BE and $\rho(r)$, and $k_{\text{HB}}^a$ and $\rho(r)$ relationships as shown in Figures 1(B), (C) and 2(B), (C), respectively.

The conclusions of our work can be summarized as follows.

- Those cases with significant deviation in the $\rho(r)$ versus BE correlation are difficult to explain in reminiscence of the Hohenberg–Kohn theorem which relates energy to electron density, but so far the actual functional between the two is still not known. The local stretching force constant $k_{\text{HB}}^a$ of hydrogen bond H–A characterizing the hydrogen bond strength between the donor and acceptor monomers is an effective tool for assessing cases with significant deviation in the BE–$k_{\text{HB}}^a$ and BE–$\rho(r)$ correlations.

- For 28 neutral hydrogen-bonded dimer complexes investigated by Emamian and co-workers, both BE and $k_{\text{HB}}^a$ are reasonable descriptors for quantifying the hydrogen bond strength and they provide in general consistent ordering of bond strength for these 28 complexes. We show that both BE and $k_{\text{HB}}^a$ provide similar measures of hydrogen bond strength based on the judicious selection of donor/acceptor monomers forming 28 simple dimer complexes made by Emamian and co-workers. However, the local stretching force constant $k_{\text{HB}}^a$ avoids the twist of borrowing additional local descriptors to indirectly quantify the hydrogen bond strength (like the use of $\rho(r)$ to predict BE). Therefore, the use of $k_{\text{HB}}^a$ is straightforward.

### Table 2

<table>
<thead>
<tr>
<th>Group</th>
<th>Complex</th>
<th>Structure</th>
<th>BE</th>
<th>$\rho(r)$</th>
<th>$k_{\text{HB}}^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group 1+</td>
<td>m1</td>
<td>FH·NCH</td>
<td>−7.40</td>
<td>0.0399</td>
<td>0.251</td>
</tr>
<tr>
<td></td>
<td>m2</td>
<td>FH·OC</td>
<td>−1.54</td>
<td>0.0160</td>
<td>0.083</td>
</tr>
<tr>
<td></td>
<td>m3</td>
<td>FH·FCCH</td>
<td>−1.01</td>
<td>0.0133</td>
<td>0.053</td>
</tr>
<tr>
<td>Group 2+</td>
<td>m4</td>
<td>N$_3$H·N(Me)$_3$</td>
<td>−9.47</td>
<td>0.0438</td>
<td>0.231</td>
</tr>
<tr>
<td></td>
<td>m5</td>
<td>FH·N(Me)$_3$</td>
<td>−16.38</td>
<td>0.0763</td>
<td>0.406</td>
</tr>
<tr>
<td></td>
<td>m6</td>
<td>HCOOH·NH$_3$</td>
<td>−12.02</td>
<td>0.0508</td>
<td>0.285</td>
</tr>
<tr>
<td></td>
<td>m7</td>
<td>HCO$_3$H$^+$·NH$_3$</td>
<td>−14.08</td>
<td>0.0557</td>
<td>0.309</td>
</tr>
</tbody>
</table>

*Peroxyl acid.
when measuring the hydrogen bond strength and generally applicable also to complex systems like water clusters or proteins.

- Our work provides a different point of view into the nature of hydrogen bonding by emphasizing the interactions within the donor/acceptor monomer which are not reflected by local electronic structure descriptors of H - A such as ρ(r) and k_HB but contribute marginally to the total BE of the 28 neutral hydrogen-bonded complexes. Therefore, care has to be taken when different hydrogen bond strength measures are compared. The BE is a cumulative hydrogen bond strength measure while k_HB is a local measure of hydrogen bond strength.

- As a caveat we want to highlight, any attempt to present a simplified and unified description of hydrogen bonding as done in the work of Emamian and co-workers via their BE and ρ(r) model, can be clouded by the fact that many systems tend to diverge from the simple hydrogen bond situation due to other factors that come into play. Thus, careful consideration should be taken under such situations.

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**DATA AVAILABILITY STATEMENT**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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**REFERENCES**


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