COMPARATIVE INSIGHTS INTO HYDROGEN ABSTRACTION OF CCL-H BY SMALL OXYGEN-CONTAINING ANIONS MO- WITH M = BE, B, AND AL

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ABSTRACT

In the present work, the considered hydrogen abstraction (HAT) reactions of CClH molecule driven by three different small anions MO- (M = Be, B, and Al) have been investigated using electronic structure calculations. While full geometry optimizations were operated to locate all of the relevant stationary points using the DFT-BHandHLYP/aug-cc-pVTZ level, the potential-energy profiles were constructed using the coupled-cluster theory with extrapolation to complete basis set CCSD(T)/CBS. Our theoretical findings suggest that the most favored pathway determined for the HAT reactions mainly stems from the MO- type, namely, for facilitating the HAT pathway the B atom is predicted to be an inherent key in the BO- reaction whereas it becomes O atom in both BeO- and AlO- reactions. Of the three favored pathways obtained here, the activations of the CClH in the presence of both BeO- and AlO- anions are significantly efficient, in which the energy barrier for the cleavage of the C-H bond with the assistance of BeO- was to be relatively low. Again, through the transition state theory the rate constants at 298-1000 K are also evaluated for the most favored HAT reactions studied here, indicating the lower the temperature, the faster the BO- chemical reaction.

Keywords: Diatomic anion; H-atom abstraction; quantum chemical calculation.

1. INTRODUCTION

Nowadays, making widely use of efficient chemical process to achieve cleavage of the strong organic bonds is still a major issue for academic challenge. Of the many possible agents for organic bonds activation, the gas-phase anions, in particular, the diatomic anions, continues to attract the researcher’s attention, such as reactions of RCl (R = methyl, ethyl, isopropyl, and tert-butyl) with ClO⁻ anion, [1, 2] hence largely viewed as a prospective alternative for the expensive metal-based catalysts used for organic bonds activation. In such case, reproduction of the required diatomic anions is of very important. In this work we deal with a class of possible diatomic anions through replacing of the Cl atom in anion ClO⁻ using some available elements.

It is known that when neutral diatomic compounds provide wide variety of opportunities for investigations of the unusual bonding characteristics, the corresponding anions studies also have received considerable interest. [3-9] For example, addition of an electron for BeO forming BeO²- may generally exhibit a much slighter lengthening in the bond but a great larger decrease in the vibrational frequency. [10, 11] However, nature of the ionic bonding was apparently enhanced in the ground state AlO⁻ electron configuration. [12] As an insight for regarding periodic trends, although the B and Al atoms are definitely categorized as the same group, possessing almost identical s-p excitation energies, [13] the electronic properties of the BO⁻ anion were not yet available in experiments. [14] Prompted to these findings, together with reminiscence of the boron atom that possesses unique and diverse chemistries due to its electron deficiency [15], in this paper we set out a quantum chemical simulation to obtain a better understanding toward the characters of the BO⁻ anion with the aid of an investigation of the CClH + BO⁻ reaction, representing a continuation of our interest in boron chemistry.

Most of previous anions investigations on the bonds activation are primarily controlled by their reactions of bimolecular nucleophilic substitution (S₂,2); however, probability of the HAT mechanism for the type of X⁻ + CH₃Y reaction has typically reported as well, in which the HAT is usually endothermic for these reactions and does not occur at low collision energies. [16, 17] The importance of HAT at the low energies is evidently illustrated by studies of the OH⁻ + CH₃I reaction. [18, 19] Obviously, the resulting HAT should be an available pathway for the ongoing anionic reactions, which allows one to probe the intrinsic reactivity of the anions. Moreover, large amounts of studies on the reactions of carbene radical anions have already discovered occurrence of the remarkable HAT. [20-27] Therefore, we were strong curious about whether the hydrogen abstraction (HAT) is a general pathway for studies of the anionic reactions? Recently, we explicitely predicted reaction mechanism of CClH + XO⁻ (X = F, Cl, or Br) by means of the HAT pathway, proposing activation of C-H bond in CClH increases as Y goes from F to Br and decreases as X from F to Br. [28] Simultaneously, our investigations performed for the type reactions have reported potential energy surfaces (PESs) with a central barrier that separates the potential minima for the pre- and postreaction ion-dipole complexes, except of the FO⁻ reaction. An intriguing question is whether substituent X in the halogen-oxygen anion XO⁻ can be changed that the PES no longer retains two wells, then, the substituent-activity relationship controlling the reaction pathways has inevitably intrigued our interest.

However, to the best of our knowledge, there are currently less available works for the CClH HAT reactions induced by the modeling MO⁻ anions with different active sites, where M is metal atom. More important, the associated metal atom in the family MO⁻ lying positively charges can overcome their dangling bond, [29] thus dramatically changing corresponding electronic and chemical features, which are not observed in the halogen-oxygen anions. For the reason stated above, a comparative HAT reaction model, MO⁻ + CCIH → CCI⁻ + MOH and H-MO (M = Be, B, and Al), is deservedly considered for our present studying system, as they yield good insights regarding periodic trends. Our computed results shed new light on the reactivity abilities of the novel amphiprotic-elements-containing anions to achieve the C-H bond activation, and thereby suggest directions for the search for the anions as potential catalysts for dehydrogenation processes.

2. METHODOLOGY

All electronic structure computations are fully conducted using both density functional theory (DFT) and wave-function methods implemented in the Gaussian 03 suite of program. [30] Noted that the popular DFT-BHandHLYP functional [31] belongs to a better functional for calculations of the H-atom transfer pattern. [28, 32] Moreover, this BHandHLYP is previously proved to be very efficient for the estimations of the molecular energies and geometries. [33-35] Although the recent studies show that it was fairly advantageous to use DFT-B3LYP wave function as the reference set for the BeO⁻ higher-order calculation, [36] our present BHandHLYP/aug-cc-pVTZ computation gives a 1.235 Å bond length for the ground state of BO⁻ (Σ⁻), which is in excellent agreement with the B3LYP/aug-cc-pVTZ result (1.234 Å) [37], especially the experimental measurement 1.236 ± 0.001 Å [38]. With the above exposures, the hybrid BHandHLYP functional should be very appropriate and reliable for investigations of the CI⁻ + MO⁻ systems and thus used to optimize the geometries of all stationary structures studied in this paper, in conjunction with Dunning’s all-electron basis set [aug-cc-pVTZ] [39]. Meanwhile, this DFT method has also been used to characterize the distinctive nature of the stationary points on PES, in which the minima were properly confirmed with positive frequencies and the transition state structure had single imaginary frequency. The intrinsic reaction coordinate (IRC) was often carried out to track the minimum energy paths from transition state to the corresponding minima [40]. The natural population analysis (NPA) has been expressly made with the natural bond orbital (NBO) method [41] to obtain significant insights into the bonding properties. Furthermore, much more accurate CCSD(T)/CBS quality energies are reasonably extrapolated by using the open-source Psi4 program [42], in which there is a driver that does the extrapolation procedure automatically.

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Rate coefficients over the temperature range 298-1000 K were typically calculated for the favorable pathways found in the title reactions using the conventional transition-state theory:[43, 44].

\[ k(T) = \kappa(T)k_{\text{TST}}(T) = \frac{k_B T}{h} e^{-\frac{\Delta G^*}{R T}} \]  

(2)

where \( k_B \) and \( h \) are the Boltzmann and Planck constants respectively; the transmission coefficient \( \kappa(T) \) was effectively evaluated through the Wigner method [45]. \( \kappa^W(T) \) is given by:

\[ \kappa^W(T) = 1 + \frac{1}{24} \left( \frac{\hbar m (v^*)^2}{k_B T} \right) \]  

(3)

where \( v^* \) is the TS imaginary frequency.

3. RESULTS AND DISCUSSION

With regard to the title reactions presented here, Fig. 1 illustrates clearly the BHandHLYP optimized geometrical shapes marked with some key parameters, in which precursor complex, transition state, product complex, and final product are artifically denoted by the labels IM1-Y, TS1-Y, IM2-Y, and P-Y, respectively, where Y stands for either OM or MO, indicating the C-H bond activation of CCl\(_3\)H by the MO anions can be directly induced by its O or M site. Moreover, the predicted PESs together with relevant geometries were vividly described in Fig. 2 on the basis of the CCSD(T)/CBS relative energies. Estimation of the exothermic and endothermic reactions was understandably characterized by employing enthalpies (\( \Delta H \)) of formation of the products relative to the corresponding reactants. All mentioned energy values respectively summarized in TABLE S1 (Supporting Information). The previous observation shows that electron transfer from K atom to CCl\(_3\)H molecule produces mainly the Cl\(_2\) ion, [46] indicating that the \( \sigma^*(\text{C-H}) \) antibonding orbital is not activated. However, in this paper following the acceptor ability of the \( \sigma^*(\text{C-H}) \) and \( \sigma^*(\text{Cl-Cl}) \) orbitals (cf. Table 1), in the IM1-Y complexes the NBO second-order perturbation energy, \( E_2^{(2)} \), for LP(O/M) \( \rightarrow \sigma^*(\text{C-H}) \) is quite sensitive, proposing the greater \( E_2^{(2)} \) values with respect to LP(O/M) \( \rightarrow \sigma^*(\text{Cl-Cl}) \) case, to thus yield the severe H-activation in the CCl\(_3\)H molecule. In this regard, pathway of the H-atom abstraction from CCl\(_3\)H is more favorable than that of the C-Cl bond breaking and consequently the most stable complex prefers a HAT result over the C-Cl cleavage. Additionally, highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) energies of the predicted precursor complexes (IM1-Y) were successively checked as shown visibly in Fig. 3, as well as corresponding pictures. We can see that formation of the IM1-OAl complex is preferred in thermodynamics by 3.68 eV \( E_{\text{comp}} \) (energy gap). Indeed, as clearly shown in Fig. 2, formation of the IM1-OBe should be very predominant owing to the largest complexation energies (\( \Delta E_{\text{comp}} \)) among all IM1-Y complexes, in which \( \Delta E_{\text{comp}} \) is defined by the relative energy of the “precursor” complex in regard to the total energy of isolated ones. Why. The more relevant information will be elucidated in detail in the following sections. Here, to make the discussions more clearly, in this paper the studied CCl\(_3\)H + MO reactions were designedly divided into the three: (1) Reaction of CCl\(_3\)H + BeO; (2) Reaction of CCl\(_3\)H + BO; (3) Reaction of CCl\(_3\)H + AlO.

![Figure 1](image_url.png)

**Figure 1.** Optimized geometries for the HAT reactions of CCl\(_3\)H + MO (M = Be, B, and Al) at the BHandHLYP/aug-cc-pVTZ level of theory, with bond distances Å and angles in degree. (The symbol “—” denotes nonexistent).
Figure 2. Energy profiles of the MO reactions (M = Be, B, and Al). Relative energies are taken from the CCSD(T)/CBS energies summarized in TABLE S1 (Supporting Information).

Figure 3. Frontier molecular orbitals of the IM1-MO and IM1-OM complexes, where M = Be, B, and Al.
3.1. Reaction of CCl₃H + BeO⁻

3.1.1 Precursor complex

It should be noted that the initial possible C-H bond activation of CCl₃H molecule mediated by the BeO⁻ anion experiences firstly two alternative precursor complexes, CCl₃H · BeO⁻ (IM1-BeO) and CCl₃H · BeO⁻ (IM1-BeO⁻), along the reactions’ collision, as outlined distinctly in Fig. 1 together with Fig. 2.

The most noteworthy feature depicted here is that the two loose ion-molecular species on PES are lying lower in energy than the separated reactants, in which the formation of the IM1-BeO⁻ is more favorable in thermodynamics thanks to its larger \( \Delta E_{\text{comp}} \) when comparing the IM1-BeO case, such as -96.72 vs. -38.90 kJ mol⁻¹. Besides, as demonstrated clearly in Fig. 3, the generating intermediate IM1-BeO⁻ is calculated to be ~0.92 ev larger in \( E_{\text{comp}} \) than that of the IM1-BeO, implying more stabilization for the IM1-BeO⁻, consisting with ~90.00 kJ mol⁻¹ contribution of the interaction LP(O)–σ*(C-H) exists in the IM1-BeO⁻ structure from our NBO analysis (see Table 1). Here, we failed to locate the interaction of LP(Be)–σ*(C-H) in the IM1-BeO case, despite our extensive attempt. Thus, we will mainly discuss the characters of complex IM1-BeO⁻ and its corresponding reaction, as will be shown below.

**Table 1.** NBO analysis obtained at the BHandHLYP/aug-cc-pVTZ level for the precursor complexes of the reactions of CCl₃H + MO⁻ (M = Be, B, and Al).

<table>
<thead>
<tr>
<th>Species</th>
<th>Donor</th>
<th>Acceptor</th>
<th>( E^{(2)} ) (kJ mol⁻¹)</th>
<th>( \Delta E ) (a.u.)</th>
<th>( F_{ij} ) (a.u.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IM1-BeO⁻</td>
<td>LP(O)</td>
<td>σ*(C-H)</td>
<td>90.00</td>
<td>0.94</td>
<td>0.179</td>
</tr>
<tr>
<td></td>
<td>LP(O)</td>
<td>σ*(C-CI)</td>
<td>0.79</td>
<td>0.60</td>
<td>0.014</td>
</tr>
<tr>
<td>IM1-BeO</td>
<td>LP(Be)</td>
<td>σ*(C-H)</td>
<td>61.42</td>
<td>0.97</td>
<td>0.107</td>
</tr>
<tr>
<td></td>
<td>LP(Be)</td>
<td>σ*(C-CI)</td>
<td>0.79</td>
<td>0.61</td>
<td>0.010</td>
</tr>
<tr>
<td>IM1-OB</td>
<td>LP(O)</td>
<td>σ*(C-H)</td>
<td>243.97</td>
<td>0.47</td>
<td>0.153</td>
</tr>
<tr>
<td></td>
<td>LP(B)</td>
<td>σ*(C-CI)</td>
<td>2.34</td>
<td>0.14</td>
<td>0.009</td>
</tr>
<tr>
<td>IM1-BO</td>
<td>LP(O)</td>
<td>σ*(C-H)</td>
<td>119.54</td>
<td>1.14</td>
<td>0.162</td>
</tr>
<tr>
<td></td>
<td>LP(O)</td>
<td>σ*(C-CI)</td>
<td>1.13</td>
<td>0.80</td>
<td>0.013</td>
</tr>
<tr>
<td>IM1-AIO</td>
<td>LP(AI)</td>
<td>σ*(C-H)</td>
<td>87.32</td>
<td>0.53</td>
<td>0.094</td>
</tr>
<tr>
<td></td>
<td>LP(AI)</td>
<td>σ*(C-CI)</td>
<td>1.09</td>
<td>0.23</td>
<td>0.007</td>
</tr>
</tbody>
</table>

\( E^{(2)} \) is the perturbative analysis hyperconjugative energy, \( \Delta E \) is the energy difference between the \( r \) and \( r^* \) NBOs, and \( F_{ij} \) is the Fock matrix element between the NBOs \( i \) and \( j \). The symbol “—” denotes nonexistent.

**Complex IM1-BeO⁻**

Let us first review the electronic structure of BeO⁻ anion. As the point of interest for BeO⁻, the corresponding additional electron resides in an antibonding orbital that is primarily a mixture of Be 2s/p, and O 2s/p, hybrid atomic orbitals, giving rise to a \( \Sigma^2 \) ground state. [47] Then, and compared to the quartet counterpart the doublet open-shell structure of BeO⁻ was found to be much more favored and thus used for their CCSD(T) and DFT computations in this paper. When the BeO⁻ anion collinearly approaching H-CCl₃, there is very strong interaction between O (BeO⁻) and H (CCl₃H), completely favoring the molecular complex IM1-BeO⁻ formation, by considering the net charges population on Be and O atoms of the BeO⁻. As given clearly in Fig. 4(a), the Be atom has an impressive positive NBO charge, whereas the net charge of the corresponding O atom is negative. Obviously, the O atom being high electronegativity tends to attract the positively charged H atom from the CCl₃H substrate. Such gives rise to charges transfer operating from BeO⁻ to CCl₃H, thus resulting in the CCl₃H activation. As shown in Fig. 1, in the IM1-BeO the distance of C-CI bond is elongated by 0.024 Å at the same time, the C-H bond is remarkably elongated by 0.083 Å, with respect to the isolated CCl₃H reactant. The PES profile associated with IM1-BeO⁻ formation found for the interaction energy between BeO⁻ and CCl₃H is -96.72 kJ mol⁻¹, namely, the step starting from the associated reactants (BeO⁻ + CCl₃H) can be very accessible without the need of extra energy. Indeed, as can be seen in discussion of 3.1.2 Section in this paper, the forming intermediate IM1-BeO located below the zero energy reference strongly facilitates the occurrence of the subsequent steps.

**Fig. 4.** NBO charges population on the anions MO⁻. (a) O-Be; (b) O-B; (c) O-Al.

3.1.2 Mechanism

Fig. 2 vividly plots PES profiles illustrating the pathways of H-atom transfer from CCl₃H to BeO⁻. Very clearly, there are two different HAT pathways found for the reaction of CCl₃H + BeO⁻. The first pathway, with very striking double-well-potential character, initiated by the O atom of BeO⁻ via TS1-BeO drasticaly undergoes dissociation of the C-H bond in CCl₃H and following formation of the BeOH product, with a very tiny barrier of 5.6 kJ mol⁻¹. Here, the energy barrier is defined in this paper by energy of the least-stable transition state on PES in regard to that of the corresponding precursor complex.

Moreover, the calculated energy profiles (Fig. 2) visibly display the energy difference between IM1-BeO and IM2-BeO is very closer each other, only ~6.2 kJ mol⁻¹, however, the possible formed P-BeO product, with respect to complex IM2-BeO, is deliberately characterized by activation energy of 59.55 kJ mol⁻¹. Seemingly, the isomerization between IM1-BeO and IM2-BeO configurations takes place instead of yielding available BeOH product. This is very similar to results of the anionic Sc²⁻ reactions that illustrate the important role of resonances. [48-53]

In fact, the P-BeO is located by 43.37 kJ mol⁻¹ below the zero-energy reference, in such a way the product BeOH appears that can be spontaneously obtained without additional energy requirement. Alternatively, another pathway related to H-atom abstraction from the CCl₃H molecule, going forward by Be atom of the BeO⁻, may be predicted as shown in Fig. 1 clearly. The corresponding reaction-energy profile (Fig. 2) markedly indicates a single-well-potential curve path, in which the most remarkable feature is absence of the corresponding transition state. Meanwhile, a considerable endothermicity of 91.43 kJ mol⁻¹ computed for forming of the P-BeO product. Thereby, formation of the H-BeO molecule happens in a difficult way, even though IM1-BeO is still located below the zero-energy level. Overall, of the two HAT pathways studied, the studied HTA reaction opened by O atom of BeO⁻ is the most favorable; that is to say, the product H-BeO is less detected in experiment.

3.2. Reaction of CCl₃H + BO⁻

3.2.1 Precursor complex

Upon replacement of Be in the BeO⁻ anion by B atom, the generated BO⁻ ion is a close-shell molecule with valence electronic configuration of \( (1\sigma)^2(2\sigma)^2(1\pi)^4(1\pi')^2(3\sigma)^2 \), where a single electron occupies a \( \pi \)-type molecular orbital which is mainly of B 2s character. Thus, the \( \Sigma^2 \) singlet state is normally considered as its ground structure. Our results point out that, as demonstrated distinctly in Fig. 2, two more favored intermediates IM1-OB and IM1-BO from the initial separated reactants (BO⁻ + CCl₃H) are spontaneously obtained without additional activation energy. Because there are substantial \( \Delta E_{\text{comp}} \) found for formation of the two precursor complexes, 63.26 (IM1-OB) and 65.51 (IM1-BO) kJ mol⁻¹, which is large enough to compensate in keeping a favorable profile, discussions of the initial reactive species might be very interesting and important to provide a wealth of insights into characters of the BO⁻-reaction.

As a comparison of stabilization difference between the two containing-boron complexes we can clearly see that (Fig. 3) a smaller \( E_{\text{comp}} \) resides preferably in the IM1-OB (2.71 ev) relative to the IM1-BO (3.22 ev), namely, the IM1-BO is more stable than the IM1-OB, which is in good agreement with our \( \Delta E_{\text{comp}} \) calculations. Otherwise, it should be noted that (Table 1) the contribution of the donor of the nonbonding electrons to the \( \sigma^*(C-H) \) antibonding orbital in the IM1-BO structure (LP(B)–\( \sigma^*(C-H) \)) is fairly remarkable, suggesting by \(-182.55 \) kJ mol⁻¹ larger than the IM1-OB case (LP(O)–\( \sigma^*(C-H) \)), which may make HAT more easily in IM1-BO and so to support more favorable H-abstraction pathway. This result is in complete agreement with that in the BO + H₂ reaction [54], in which the most favorable mechanism is described using the abstraction of H by the B atom. In the following sections, we will discuss in detail the above-mentioned predictions.
Complex IM1-OB

An ion-molecular complex IM1-OB can be firstly obtained through activation of CCI$_3$H by the O atom of BO$^{-}$ and the corresponding geometrical shape is visibly illustrated in Fig. 1. Unlike formation of IM1-OBs found in the reaction of BeO$^-$, as shown in Fig. 1, the C-Cl and C-H bonds (Å) described in the IM1-OB are slightly elongated by 0.012 and 0.024, respectively, in regard to the isolated CCI$_3$H molecule. Actually, although the NBO population analysis illustrates distinctively the presence of the notable negative charges (-1.193 e) on the O atom of BO (see Fig. 4(b), known as the B atom having deficient electrons, the HOMO (3e) of BO$^{-}$ is basically a 2s lone pair with slight BO antibonding character, [37] whose weak antibonding character is expected to partially cancel the bonding character. Such action does not facilitate the reaction of CCI$_3$H + BO$^{-}$ to proceed continually its following process, even though the precursor complex IM1-OB having considerable $\Delta E_{\text{comp}}$ of 63.26 kJ mol$^{-1}$.

Complex IM1-BO

As just predicted above, captivity of the B atom in BO$^{-}$ for the CCI$_3$H activation is much stronger than that of the corresponding O atom, thus the possible IM1-BO complex for the reaction of BO$^{-}$ with CCI$_3$H that is identified rationally. As shown clearly in Fig. 1, formation of the IM1-BO complex entirely stems from a near-linear approach of B atom (in BO$^{-}$) to the H atom of the CCI$_3$H substrate. However, the computed distance between B and H atoms in the IM1-BO is approximately 0.364 Å longer than that between O and H atoms in the IM1-OB.

Indeed, as can be observed from Fig. 2, the IM1-BO is only -2.25 kJ mol$^{-1}$ more stable than the IM1-OB, namely, they are quite close in $\Delta E_{\text{comp}}$, which assume existence of a rather free isomerization process from IM1-OB to IM1-BO and the detailed discussions are well-rounded described in Section 3.2.2. Again, the C-H bond activation of the CCI$_3$H by B atom of the BO$^{-}$ is found to be very predominant, as displayed in Fig. 1 clearly, C-H bond in IM1-BO is acutely elongated by 0.047 Å compared to the isolated CCI$_3$H case. Obviously, the complex IM1-BO plays a role as a key precursor complex for HAT from CCI$_3$H to BO$^{-}$, which favors the subsequent step to happen.

3.2.2 Mechanism

Schematic PES profile associated with possible HAT pathways for the reaction of CCI$_3$H + BO$^{-}$ is very clearly displayed in Fig. 2. As expected, there are two types of pathways found for the considered reactions, i.e., CCI$_3$H→OB$^{-}$ and CCI$_3$H→BO$^{-}$. Of the two cases for the C-H bond scission of CCI$_3$H molecule, the former pathway is graphically characterized by a single-well reaction-energy profile, whereas the latter becomes the double-well-potential pathway. Again, formation of the product P-OB derived from the former pathway is approved to be less preferred in energies, being an enormous endothermicity of 181.57 kJ mol$^{-1}$, whereas in the latter the corresponding product P-BO can readily be created with an energy release of 48.44 kJ mol$^{-1}$.

Those are in good accordance with some predicted studies that the energy difference between the two isomers, H-BO (in P-BO) and BOH (in P-OB), is in a range of 214-244 kJ mol$^{-1}$, [55-61] Although the former reaction can be achieved with considerable endothermic, the C-H bond activation with the help of the O atom of BO$^{-}$ stills takes an advantage over the isolated CCI$_3$H case, owing to favored generation of the stable precursor complex IM1-OB lying 63.26 kJ mol$^{-1}$ below the zero-energy level. In most case, a rather free isomerization process from IM1-OB to IM1-BO via transition state TS-OB/BO may likely occur. Such behavior can be elucidated by lower barrier heights, being around ~18.48 kJ mol$^{-1}$, particularly, the TS-OB/BO is located beneath the zero-energy level as is clearly observed from Fig. 2. In other words, the HAT reaction of CCI$_3$H with BO$^{-}$ happens mostly upon abstraction of the B atom of BO$^{-}$ forward H atom of CCI$_3$H, leading to the obtained the H-BO product in preference over the final BOH product.

Hence, we only focus here on the pathway starting from the IM1-BO complex. As the most favored channel for reaction of CCI$_3$H + BO$^{-}$, the pathway involving TS1-BO yielding product P-BO is reliably characterized to be an exothermic process. Despite there is a high energy barrier of 14.7 kJ mol$^{-1}$ for the HAT reaction of CCI$_3$H + BO$^{-}$, the related TS1-BO lies now 50.81 kJ mol$^{-1}$ below the separated reactants. Apparently, the HAT process via TS1-BO is unquestionably spontaneous with absence of an overall energy barrier and thus turned out to be the most favored for the reaction of CCI$_3$H with BO$^{-}$.

3.3. Reaction of CCl$_3$H + AlO$^-$

3.3.1 Precursor complex

Analogous to cases of the BeO$^-$ reaction reported in our present study (Section 3.1), there are also two alternative pathways found for activation of the substrate CCl$_3$H when AlO$^-$ attacks, emerging to have two stable precursor complexes IM1-OAl and IM1-AIO, respectively. As is observably shown in Fig. 2, the two complexes are not only located beneath the zero-energy level, but also characterized by a relatively strong interaction, such as the predicted $\Delta E_{\text{comp}}$ being -94.27 kJ mol$^{-1}$ for the IM1-OAI and -40.41 kJ mol$^{-1}$ for the IM1-AIO. The calculated $E_{\text{TS}}$ (see Fig. 3) that the IM1-OAI is 1.24 eV larger than the IM1-AIO, so favoring the IM1-OAI more stabilization. In other words, there exists a strong orbital interaction between the p orbital of O (AlO$^-$) and the s of H (CCI$_3$H), which is very beneficial for activation of CCI$_3$H C-H σ bond. Hence, differing from the channel coming from IM1-AIO to form P-AIO (H-AlO + CCl$_3$) product that is endothermic, the pathway starting from IM1-OAI to yield product P-OAI (AlOH + CCl$_3$) is found to be exothermic, which is in good agreement with the result that the formation of the AlOH more stable than that of the H-AlO. [62] As a result, the HAT reaction of CCI$_3$H mediated by the O atom of AlO$^-$, similar to the initial step of the reaction of AlO$^-$ + H$_2$O, [63] is more favored than that by the corresponding Al atom and we only report here some important properties of the IM1-OAI complex and its corresponding mechanism.

Complex IM1-OAI

Formation of the examined complex named IM1-OAI attained AlO$^-$ with electronic X$^{-}$` ground state has considerably large $\Delta E_{\text{comp}}$ of -94.27 kJ mol$^{-1}$ (cf. Fig. 2), thus the process is spontaneous with absence of an overall energy barrier. Specifically, character of a substantial bonds activation found for the CCl$_3$H moiety appears. As shown clearly in Fig. 1, comparing with the isolated CCI$_3$H molecule, the distances (Å) of C-Cl and C-H bonds in the IM1-OAI are elongated by 0.023 and 0.075, respectively. Such result that may mainly originate from the striking LP(O)-→σ*(C-H) hyperconjugative effect in the IM1-OAI structure, displaying -119.54 kJ mol$^{-1}$ interaction from NBO analysis collected in Table 1. Therefore, similar to the BeO$^-$ case, the O atom of AlO$^-$ tends to priority attraction of the positively charged H atom from CCI$_3$H molecule, as the result of the notable electronegativity difference between the Al and O atoms (see Fig. 4(c)).

3.3.2 Mechanism

As demonstrated distinctly in Fig. 2, the HAT pathway coming from the complex IM1-OAI takes place via transition state TS1-OAI, briefly describing C-H bond breaking and O-H bond formation, to yield another much favored intermediate IM2-OAI. This process needs to overcome very tiny energy barrier of 7.49 kJ mol$^{-1}$. If the pathway starts from IM2-OAI to IM1-OAI, an only barrier of 6.75 kJ mol$^{-1}$ is preferential. Meanwhile, the transition state TS1-OAI is located well below the zero-energy level and associated with overall barrier of -86.78 kJ mol$^{-1}$ with respect to the separated reactants.

Those strongly suggest the long-lived properties for the IM1-OAI and IM2-OAIs complexes and much free isomerization between them, [64] in such a way that the formation of product P-OAI happens in a difficult way. In fact, formation of the product P-OAI (AlOH + CCl$_3$) in this channel is energetically favored with an overall exothermic energy of 36.62 kJ mol$^{-1}$, in turn, the AlOH molecule can be easily obtained without the need of extra energy.

3.4. Comparisons of the most favored BeO$^-$, AlO$^-$, and BO$^{-}$-reactions

The most favored possible HAT pathways obtained for BeO$^-$, AlO$^-$, and BO$^{-}$-reactions are identified as the channels emanating from complexes IM1-OBe, IM1-OAI, and IM1-BO, respectively, in which both IM1-OBe and IM1-OAI are much more stable than the IM1-BO and the H-atom abstractions via the former two cases appear to be dominant in particular. Our MO analysis (Fig. 3) shows that the BO moiety in the HOMO of IM1-BO is primarily composed of B 2s AO with slight antibonding character between the B 2s and O 2p AO’s, whose the antibonding character is expected to partially weaken the bonding character. In contrast, the BeO fragment in the HOMO of IM1-OBe presents a σ bonding MO composed of O 2p and Be 2s AO’s, which involves significant charges back-donation from O to Be and therefore it is of strong ionic
character; This analogous phenomenon also found expectedly for the AlO moiety in the HOMO of IM1-OAl. Those might further suggest that both Al-O and Be-O bonding are primarily ionic while B-O bonding is more covalent, which is in good agreement with the experimental observation [37].

Noted that in our presented MO\(^-{*}\) anions when the M is fixed as same group, along with the rising atomic number, such as from B to Al, it is found to be of different for the MO\(^-{*}\) to interplay with CCl\(_1\)H to complete corresponding most favored HAT reaction. As shown remarkably in Fig. 2, the most favored HAT in the BO\(^-{*}\) -reaction is promoted by the B atom, whereas it becomes O atom in the AlO\(^-{*}\)-reaction. Again, compared to ability of the BO\(^-{*}\) toward the C-H activation, the AlO\(^-{*}\) anion induces a larger reduction in energy barrier, which amounts now to \(-7.49\) kJ mol\(^{-1}\). In addition, difference between the energy barriers mainly arises from that between their stability ordering. As displayed in Fig. 1, in the TS1-BO the C-H bond length elongates to about 0.06 Å compared to the TS1-OAl case and thus leads to higher barrier. Obviously, the larger the radical of M atom is, the more easily the MO\(^-{*}\) induced HAT reaction occurs, for example, the HAT with assistance of the AlO\(^-{*}\) has taken place rather more efficiently than that with the BO\(^-{*}\) (see Fig. 2). Nevertheless, although the higher energy barrier for C-H cleavage pathway is revealed in the BO\(^-{*}\) -reaction, the most remarkable feature here is the absence of the overall energy barrier for the C-H breaking. It means that this process can also be more favored and the corresponding reaction may take place quickly (see the estimations of rate constant below). On the other hand, when comparing the reactions achieved by the nearly same size MO\(^-*\) anions, such as the BeO\(^-*\) and AlO\(^-*\) reactions, the favored HAT found for the route is consistently induced by the O atom of MO\(^-*\). Moreover, the energy barrier for the C-H dissociation in the AlO\(^-*\) reaction is slightly higher than that in the BeO\(^-*\) reaction, with a maximum difference of only \(-1.89\) kJ mol\(^{-1}\). Consequently, the H-abstraction presented here takes place last along the reaction path for TS1-OAl rather than TS1-OB. This can be well understood by considering their individual geometric characteristics such as the calculated bond orders (%(C-H)\(^{11}\), 25% (TS1-OB)) < 27% (TS1-OAl), according to Eqs reported by Ren [65]. In contrast, the newly formed O-H bonds in the TS are obviously strengthened relative to their final equilibrium values by ca. 28% for the BeO\(^-*\) abstraction and ca. 22% for the AlO\(^-*\) abstraction. These features indicate that the BeO\(^-*\) HAT reaction reaches the TS relatively early, whereas the AlO\(^-*\) case arrives at the TS relatively late. Thus, the reaction of CCl\(_1\)H + BeO\(^-*\) may have the highest exothermicity among all presented favored reactions. In summary, the various selections of the elements M may result in MO\(^-*\) to possess different activated ability, namely, the nearly same size MO\(^-*\) can induce stronger C-H bond activation than the MO\(^-*\) with same group M does.

Results of the rate constants at the BHandHLYP/aug-cc-pVTZ level of theory for the most favored HAT reactions of CCl\(_1\)H + MO\(^-*\) (M = Be, B, and Al) are systematically compiled in Table 2. For the transmission coefficient \(k^w\), there exists a very striking feature, as is shown clearly in Table 2, the \(k^w\) gradually decreased with the increasing in temperature, indicating that the entropic effects became less significant for the rate constant, and thus decrease in probability of the tunneling. Also, it may be noted that contribution of the temperature for increasing rate constants of \(k(T)\) (L mol\(^{-1}\) s\(^{-1}\)) after \(k^w\) correction is very appreciable for both BeO\(^-*\) and AlO\(^-*\) reactions with respect to BO\(^-*\) reaction, showing the higher the temperature, larger the \(k(T)\) value, the faster the reaction. In contrast, under the lower conditions, chemical rates of both BeO\(^-*\) and AlO\(^-*\) reactions are much smaller than that of the BO\(^-*\) reaction. In other words, reaction of BO\(^-*\) + CCl\(_1\)H occurs strongly faster at the lower temperature. In fact, for the BO\(^-*\) reaction the high HAT frequency value \((1124\) cm\(^{-1}\)) calculated from BHandHLYP level indicates the dominating contribution of the proton movement in the reaction coordinates and thus implies nonclassical effects, in particular the tunneling.

Table 2. Rate constants at the BHandHLYP/aug-cc-pVTZ level of theory for the most favored HAT reactions of CCl\(_1\)H + MO\(^-*\) (M = Be, B, and Al) along with the tunneling transmission coefficients as a function of temperature.

<table>
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<th>System</th>
<th>(T/K)</th>
<th>(\Delta G^\ddagger) (kJ mol(^{-1}))</th>
<th>(k^\text{TST}) (L mol(^{-1}) s(^{-1}))</th>
<th>(k^w) (L mol(^{-1}) s(^{-1}))</th>
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\(\Delta G^\ddagger\): TS free energy relative to its corresponding reactants, \(k^\text{TST}\): TST rate constant, and \(k^w\): Wigner transmission coefficient.
4. CONCLUSIONS

In summary, from the analysis in the present experiments, we are confident in predicting that, for the HAT reactions of CClH + MO (M = Be, B, and Al), (1) the most favored pathway determined in the BO -reaction is promoted by the B atom, whereas it becomes the O atom in both BeO - and AlO -reaction. (2) of the three favored pathways obtained here, there is a very striking double-well potential character in the channel described consistently by an exothermicity, (3) comparing with BO case, the CCCH activation in the presence of both BeO - and AlO - becomes very predominant, in which the energy barrier studied for the C=H bond dissociation with the assistance of BeO - was found to be low relatively. (4) of the rate of the favored HAT reactions presented here is governed by the temperature, in which the lower the temperature, the faster the HAT reaction of BO -.

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SUPPORTING INFORMATION

Energies for the HAT reactions of CClH + MO - (M = Be, B, and Al) stationary points.

CONFICT OF INTEREST

The author has no conflict of interest.

REFERENCES