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# Semiempirical investigations on the stabilization energies and ionic hydrogen-bonded structures of $F^-(H_2O)_n$ and $CI^-(H_2O)_n$ (n = 1-4) clusters

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#### **Abstract**

Several semiempirical methods were utilized to analyze the structures and stabilities of  $X^-(H_2O)_n$  (X = F, CI; n = 1-4) clusters with respect to the number of water molecules through their comparison with *ab initio* molecular orbital calculations. Our results show that the recently developed PM6-DH+ semiempirical method can provide reasonable binding energies of hydrated fluoride and chloride ion clusters, which are consistent with the corresponding experimental results. For the optimized geometries of X = F, however, the semiempirical methods show that the global minima are close to  $HF(OH)^-(H_2O)_{n-1}$  structures, which are different from the *ab initio* calculations. Meanwhile, the topological characteristics for the global minima of X = CI obtained by semiempirical methods have the same symmetries with *ab initio* calculations. All calculation levels agree on the trend of decreasing ion-water interaction with the increasing number of water molecules. We also found a new structure of  $CI^-(H_2O)_4$  with a second hydration shell as a complement of previous studies. Those are very important data for our near-future study of on-the-fly semiempirical molecular dynamics (MD) or path integral MD simulation.

**Keywords:** Semiempirical potential, *Ab initio* molecular orbital calculations, Ionic hydrogen-bonded structures, Halide anion water clusters

PACS: 36.40.Mr, 31.15.bu, 31.15.A-

# **Background**

The ionic hydrogen bond, which has a stronger intermolecular interaction than the typical hydrogen bond, plays significant roles in various biological and chemical processes [1-4], such as protein folding [5] and molecular recognition [6,7]. Meanwhile, the ionic hydrogen-bonded structures provide a powerful lens through which the important phenomena of ion hydration could be further understood [8-11]. Intensive studies [12-15], with experimental techniques [8,16-22] and theoretical approaches [23-38], have been focused on the detailed structures of the hydrated halide ion clusters.

Experimentally, the formation enthalpies of small hydrated halide ion clusters in the gas phase have been studied using a pulsed electron beam high-pressure mass spectrometer [21]. Diffraction studies [22] have provided limited structural information on these clusters, for example, the average distance between fluoride/chloride and oxygen is 2.6 to approximately 2.9/3.1 to approximately 3.5 Å. Recently, vibrational spectroscopy using argon predissociation technique has provided more abundant structural information about the fundamentals and first two overtones for  $X^-(H_2O)_n$  (X = F, Cl; n = 1-4) clusters [8,16-18].

On the other hand, to understand ion hydration at the molecular level, intensive theoretical works have focused on the structures and binding energies of hydrated ion water clusters [23-26]. For example, Kim and coworkers reported various local minima and transition state structures for these clusters with n = 1 to 6 through high-level *ab initio* molecular orbital (MO) calculations [23-25]. Recently, Kamp et al. studied the solvation of fluoride and

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chloride ion water clusters with  $n \le 20$  using the effective fragment potential (EFP) method and Monte Carlo simulations [26]. In addition, other theoretical efforts, such as quantum mechanical/molecular mechanical (QM/MM) [28] and molecular dynamics (MD) with potential including polarization term [29,39], are also devoted to these clusters. The interaction potentials are essential to the theoretical studies and high-level *ab initio* calculations are considered to be the most reliable. However, due to the bottleneck of computational limitations, it is formidable to use such expensive potentials to deal with the energetics and geometries of these clusters with respect to a large number of water molecules.

As an alternative solution, semiempirical potentials have been successfully applied into various systems with extremely cheap computation cost [40,41]. Recently, the PM6 semiempirical potential [42,43] has been successfully applied into the quantum chemistry study of neutral hydrogen-bonded systems, such as glycine-water clusters [44] and hydrogen storage material MOF-5 [45]. The PM6 results show reasonable accurate structures and stabilization energies compared to high-level ab initio calculations. More recently, new semiempirical potentials of PM6-DH [46], PM6-DH+ [47], and PM6-DH2 [48], which improve PM6 by including two important corrections of dispersion energy and hydrogen bonding, have shown promising results for extended hydrogen-bonded complexes. To the best of our knowledge, however, there is still no report on ionic hydrogenbonded systems with PM6 or PM6-DH+ semiempirical potentials.

In this study, thus, we apply the semiempirical potentials to  $X^-(H_2O)_n$  (X = F, Cl; n = 1-4) clusters for investigating the various possible geometries. On the other hand, *ab initio* MO and density functional theory (DFT) methods are also utilized to examine the performance of the semiempirical potentials, with respect to the dependence of the number of water molecules on the energetics and geometries of the ionic hydrogen-bonded clusters. Those are very important data for our near-future study of on-the-fly semiempirical MD or path integral MD simulation.

#### **Methods**

First, we have explored the optimized energies and geometries of  $X^-(H_2O)_n$  (X = F, Cl; n = 1-4) clusters by semiempirical PM6. Then two modified semiempirical methods, PM6-DH+ and PM6-DH2, were also applied to these clusters. PM6-DH+ was applied to examine both the energies and geometries of the clusters. However, due to the limitation on geometry optimizations of PM6-DH2 [48], we examined the interaction energies of PM6-DH2 on the basis of the optimized geometries

from PM6. All the semiempirical calculations were performed using the MOPAC2009 (SCC, Colorado Springs, CO, USA) program package [49].

On the other hand, we have examined the above-obtained semiempirical results by applying *ab initio* MO and DFT calculations for such ionic hydrogen-bonded systems. The *ab initio* calculations were performed with Hartree-Fock (HF) and second-order Møller-Plesset perturbation theory (MP2) [50] including electron correlations. The DFT scheme utilized Becke's three parameter hybrid method using the Lee-Yang-Parr correlation function (B3LYP) [51,52]. The aug-cc-pVDZ basis set was used for the comparison of all methods. Of the above-mentioned theoretical levels, MP2 is usually considered to be the most reliable method, while it requires the most computational cost. The computations were performed using the Gaussian 09 (Gaussian, Inc., Wallingford, CT, USA) program package [53].

## Results and discussion

The schematic illustrations of optimized structures for  $X^-(H_2O)_{1-4}$  are shown in Figure 1. Here, X refers to F or Cl. These topological structures obtained from semiempirical calculations are consistent with those of the *ab initio* calculations. In addition, the global minimum structures of fluoride clusters, which are optimized by only semiempirical methods, are shown in Figure 2. The binding energies for fluoride and chloride anion water clusters are listed in Tables 1 and 2, respectively. The binding energy is defined as follows:

$$\Delta E = (E_{x-} + nE_{(H2O)}) - E_{x-(H2O)n}, \tag{1}$$

where  $E_Z$  refers to the optimized energies of species  $Z = X^-$ ,  $H_2O$ , and  $X^-(H_2O)_m$  respectively. The selected bond lengths and bond angles, which represent the ionic hydrogen bond strengths of  $X^-(H_2O)_{1-4}$  clusters, are shown in

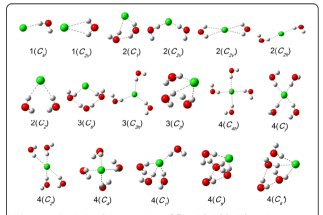


Figure 1 Optimized geometries of fluoride/chloride anion water clusters at MP2/aug-cc-pVDZ level. The colors of green, red, and white represent fluoride/chloride, oxygen, and hydrogen atoms, respectively.

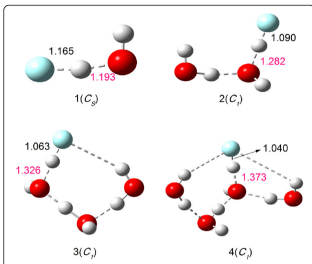


Figure 2 Optimized global minima of fluoride anion water clusters at PM6-DH+ level. The F...H and H...O distances in each cluster are shown in black and pink color, respectively. The optimized F...H and H...O distances in isolated HF and H $_2$ O molecules are 0.966 and 0.949 Å at the same calculation level, respectively.

detail Tables S1 and S2 in Additional file 1 for X = F and X = Cl, respectively. In addition, the values and directions of the imaginary frequencies of the corresponding transition state structures are shown in Table S3 and Figure S1, respectively, in Additional file 1.

We would focus on the performance of the semiempirical methods on the optimized geometries and binding energies of  $X^-(H_2O)_n$  clusters with respect to the increasing water number through comparison with the *ab initio* and DFT calculations.

# Fluoride anion water clusters

In the  $F^-(H_2O)$  cluster, the stable structure takes the  $C_s$ symmetry, and the transition state takes the  $C_{2v}$  symmetry for all calculation methods. The potential barrier height between the transition state and stable structure is 7.7 kcal/mol at PM6 level with zero-point vibrational energy (ZPE) correction, which is exactly the same as that at MP2 level. We note here that PM6-DH+ and PM6-DH2 show the same stabilization energies as PM6. The ionic hydrogen-bonded structures of  $F^-(H_2O)_n$  are exhibited through  $r(F...H_{near})$ , r(F...O), and  $\theta(F...H-O)$ . In comparison with the  $C_{2v}$  structure, the semiempirical results show that the  $r(F...H_{near})$  of  $C_s$  structure is shorter and  $\theta(F...H-O)$  is much larger, which agree well with other calculation levels (see Table S1 in Additional file 1 for details). The results indicate that the ionic hydrogen bond strength in the stable structure is stronger than that in the transition state structure. However, r(F...O) of the  $C_{\rm s}$  structure is slightly longer for the semiempirical methods, which is just on the contrary at higher calculation

levels. Here, we have to mention that the r (F. . .  $H_{near}$ ) predicted by PM6-DH+ (1.165 Å) is much shorter than that of MP2 (1.413 Å). The optimized F. . . H and H. . . O distances in isolated HF and  $H_2O$  molecules are 0.966 and 0.949 Å at PM6-DH+ level, respectively. Thus, the global minimum would be close to the HF(OH $^-$ ) structure (see Figure 2).

In the case of F<sup>-</sup>(H<sub>2</sub>O)<sub>2</sub>, ab initio calculations show that stable and transition state structures take the  $C_2$ and C<sub>2v</sub> symmetries, respectively. The potential barrier height between these two structures is 0.2 kcal/mol at the MP2 level with ZPE correction. The energy barrier is guite small in comparison with the F-(H<sub>2</sub>O) cluster, which indicates the frequent changes of cluster geometry under certain thermal fluctuations. The semiempirical methods can also provide such small energy difference (not more than 0.1 kcal/mol) between  $C_2$  and  $C_{2v}$  structures. However, the preferred global minimum structure by semiempirical methods takes the  $C_1$  symmetry, rather than the  $C_2$  symmetry by ab initio calculations. The results indicate that the water-water interaction in F-(H<sub>2</sub>O)<sub>2</sub> is not well estimated by semiempirical methods. In addition,  $2(C_{2v'})$  of the  $F^-(H_2O)_2$  structure is shown to be a saddle point with the smallest binding energies by semiempirical as well as ab initio and DFT calculations. The  $2(C_{2h})$  structure was optimized as a saddle point by MP2/6-311++G\*\* and MP2/TZ (2df, pd)++ in [24]; however, using a larger basis set of aug-cc-pVDZ in this work, it is shown to be a transition state at the MP2 level and to be a local minimum at the HF and B3LYP levels. The semiempirical methods support that it is a saddle point structure. On the other hand, the geometry differences of the  $C_2$ ,  $C_{2v}$ , and  $C_{2h}$ structures, such as  $r(F...H_{near})$ , r(F...O), and  $\theta(F...H-O)$ , are very close for both semiempirical and ab initio calculations. The results imply that the ionic hydrogen bond strengths in these geometries are very close. Similar to the case of n = 1, short  $r(F...H_{near})$ is predicted by PM6-DH+ (1.087 Å), the  $C_1$  structure is probably HF(OH<sup>-</sup>)H<sub>2</sub>O.

In the case of  $F^-(H_2O)_3$ , *ab initio* calculations show that stable structures take the  $C_3$  and  $C_s$  symmetries, and the  $C_3$  structure is more stable. It is interesting from the *ab initio* results that when ZPE corrections are included, the energy of the transition state with the  $C_{3h}$  symmetry could be even lower than that with the  $C_3$  symmetry. Unfortunately, this important indication could not be well supported by PM6 or PM6-DH+. Instead, the most stable  $C_1$  structure of  $F^-(H_2O)_3$  optimized by semiempirical methods takes a planar structure similar to  $C_{3h}$ . What differs with the  $C_{3h}$  symmetry is that the fluoride ions are located outside rather than inside the cluster. The semiempirical results show that the binding energy of the  $C_s$  structure is higher than

Table 1 Binding energies (kcal/mol) of fluoride anion water clusters

F <sup>-</sup> (H <sub>2</sub> O) <sub>n</sub>	Experiment	PM6			PM6-DH+			ı	PM6-DH2			HF/aug-cc-pVDZ			B3LYP/aug-cc-pVDZ			MP2/aug-cc-pVDZ		
		ΔΕ	$\Delta E_{ZPE}$	IF	ΔΕ	$\Delta E_{\text{ZPE}}$	IF	ΔΕ	$\Delta E_{ZPE}$	IF	ΔΕ	$\Delta E_{ZPE}$	IF	ΔΕ	Δ <i>E</i> <sub>ZPE</sub>	IF	ΔΕ	Δ <i>E</i> <sub>ZPE</sub>	IF	
1(C <sub>s</sub> )	23.3	24.5	23.0	0	24.6	23.0	0	24.6	23.0	0	23.6	22.2	0	27.3	26.5	0	26.8	26.0	0	
1(C <sub>2v</sub> )		16.8	15.3	1	16.8	15.3		16.8	15.3	1	18.0	16.7	1	19.6	18.6	1	19.4	18.3	1	
2(C <sub>2</sub> )	42.5	36.0	32.8	1	36.1	33.0	1	36.2	33.1	1	42.9	39.4	0	48.1	45.3	0	48.1	45.0	0	
2(C <sub>2v</sub> )		35.2	32.9	1	35.3	32.9	0	35.3	33.0	0	42.7	39.4	1	47.7	45.2	1	47.5	44.8	1	
$2(C_{2v'})$		31.1	28.1	3	31.2	28.2	3	31.2	28.2	3	33.7	31.1	3	36.1	33.9	3	36.1	33.9	3	
$2(C_{2h})$		35.3	33.1	3	35.4	33.2	2	35.4	33.2	3	42.9	39.6	0	47.8	45.2	0	47.6	44.9	1	
2(C <sub>1</sub> )		43.5	39.3	0	46.5	41.8	0	49.8	45.0	0	-	-	-	-	-	-	-	-	-	
3(C <sub>3</sub> )	57.8	48.5	42.5	2	51.2	44.2	2	50.7	44.3	2	59.0	52.7	0	64.8	59.7	0	66.9	60.4	0	
3(C <sub>s</sub> )		51.1	45.9	1	56.1	48.1	1	54.2	48.7	1	56.5	49.9	0	64.1	58.1	0	65.4	59.3	0	
$3(C_{3h})$		44.8	41.5	3	45.2	41.9	3	45.2	42.0	3	58.9	53.5	1	64.8	60.0	1	65.3	60.4	1	
3(C <sub>1</sub> )		57.2	50.7	0	66.6	57.1	0	65.7	58.7	0	-	20	-	-	-	-	-	-	-	
4(C <sub>1</sub> )	71.7	70.6	62.2	0	80.0	70.0	0	84.5	74.6	0	72.8	64.3	0	79.1	70.9	0	82.2	73.7	0	
4(C <sub>2</sub> )		61.8	54.8	0	66.6	58.8	2	65.0	57.3	3	72.5	64.3	0	78.6	70.9	0	81.1	73.2	0	
4(C <sub>4</sub> )		59.7	51.1	3	66.2	55.3	3	64.0	55.0	3	72.0	63.1	0	78.0	69.2	0	82.2	72.8	0	
4(C <sub>s</sub> )		63.4	55.8	1	71.6	61.8	1	68.6	59.8	1	71.5	61.9	0	79.9	70.7	0	83.5	74.1	0	
4(C <sub>i</sub> )		60.7	53.9	1	66.0	58.2	1	64.3	57.0	1	72.0	63.7	0	77.6	69.9	1	80.1	72.2	1	
4(C <sub>4h</sub> )		52.3	47.8	6	53.2	48.8	5	53.3	48.8	5	71.6	63.9	2	77.4	70.4	2	79.7	72.8	5	

 $\Delta E$  and  $\Delta E_{ZPE}$  represent the binding energies without and with zero-point vibrational energy (ZPE) corrections, respectively. IF represents the number of imaginary frequencies. The experiment values are taken from [21].

Cl <sup>-</sup> (H <sub>2</sub> O) <sub>n</sub>	Experiment	PM6			PM6-DH+			PM6-DH2			HF/aug-cc-pVDZ			B3LYP/aug-cc-pVDZ			MP2/aug-cc-pVDZ		
		ΔΕ	$\Delta E_{ZPE}$	IF	ΔΕ	$\Delta E_{\text{ZPE}}$	IF	ΔΕ	$\Delta E_{ZPE}$	IF	ΔΕ	∆ <i>E</i> <sub>ZPE</sub>	IF	ΔΕ	Δ <i>E</i> <sub>ZPE</sub>	IF	ΔΕ	Δ <i>E</i> <sub>ZPE</sub>	IF
1(C <sub>s</sub> )	14.7	15.3	14.2	0	15.3	14.2	0	15.3	14.2	0	11.8	10.5	0	14.3	13.2	0	14.7	13.5	0
1(C <sub>2v</sub> )		15.4	13.8	0	15.5	13.9	0	15.4	13.8	0	11.0	10.0	1	12.5	11.6	1	13.1	12.2	1
2(C <sub>1</sub> )	27.7	30.1	26.5	0	32.2	28.1	0	31.5	27.7	0	23.5	19.8	0	28.0	24.5	0	29.8	26.2	0
2(C <sub>2</sub> )		29.1	25.6	1	29.5	25.9	1	29.7	26.0	1	23.1	19.7	1	27.5	24.4	1	29.3	26.1	1
2(C <sub>2h</sub> )		-	-	-	-	-	-	-		-	22.4	19.8	2	26.6	24.2	2	27.6	25.2	3
2(C <sub>2v</sub> )		-	-	-	-	-	-	-		<u> </u>	22.3	19.8	3	26.6	24.2	2	27.7	25.2	2
2(C <sub>2v′</sub> )		29.6	26.3	0	29.7	26.2	0	29.7	26.4	0	21.0	19.0	4	23.4	21.6	5	24.9	23.1	4
3(C <sub>3</sub> )	39.5	44.2	37.6	0	48.8	40.8	0	47.2	40.4	0	35.4	28.8	0	42.0	35.4	0	46.1	39.4	0
3(C <sub>s</sub> )		41.6	35.4	0	47.0	36.8	1	44.3	37.0	1	33.9	28.0	0	40.9	35.2	0	43.5	37.6	0
3(C <sub>3h</sub> )		41.7	36.7	3	42.0	37.1	3	42.0	37.1	3	32.1	28.2	1	37.8	34.2	1	39.6	36.0	1
4(C <sub>2</sub> )	51.1	54.2	46.6	0	58.2	47.8	2	56.3	48.2	2	43.1	35.8	0	50.2	43.0	0	54.5	47.2	0
4(C <sub>4</sub> )		57.9	48.3	0	67.7	52.5	0	63.6	53.7	0	46.1	36.9	0	54.6	45.1	0	60.8	51.1	0
4(C <sub>s'</sub> )		54.0	46.0	0	63.7	52.5	0	60.5	51.7	0	44.0	35.7	0	52.4	44.1	0	57.5	49.1	0
4(C <sub>s</sub> )		53.1	45.9	2	60.9	49.8	2	57.8	49.6	2	44.9	36.2	1	54.0	45.3	1	59.3	50.4	1

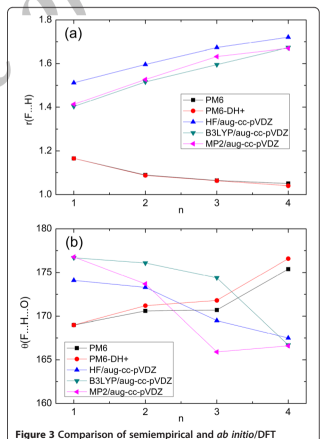
 $\Delta E$  and  $\Delta E_{ZPE}$  represent for the binding energies without and with zero-point vibrational energy (ZPE) corrections, respectively. IF represents the number of imaginary frequencies. The experiment values are taken from [21].

that of the  $C_3$  structure, which may be the result of the overestimation of water-water interactions. On the other hand, in agreement with the ab initio results, semiempirical methods show that  $r(F...H_{near})$  of the  $C_s$ structure is the shortest in the F-(H2O)3 clusters, which is followed by  $C_{3h}$  and  $C_3$  symmetries. The largest r(F...O) also appears in the  $C_{3h}$  structure at PM6 level, while other higher calculation levels favor the  $C_3$  structure. The largest  $\theta(F...H-O)$  appears in the  $C_s$  structure at the PM6 level, while other higher calculation levels prefer the  $C_{3h}$  symmetry. The  $C_1$  structure, which is optimized through the PM6-DH+ method, has a very short F...H distance (1.063 Å). The results indicate that the structure of HF(OH<sup>-</sup>)(H<sub>2</sub>O)<sub>2</sub> is probably formed instead of F<sup>-</sup>(H<sub>2</sub>O)<sub>3</sub>. However, the *ab initio* calculations do not support these results. In addition, both the semiempirical and ab initio calculations show that due to the formation of the second hydration shell in the  $C_s$  structure, a shorter  $r(F...H_{near})$  and a larger  $\theta(F...H-O)$  are obtained in comparison with the  $C_3$  symmetry. The results indicate that the ionic hydrogen bond strength in the structures with the second hydration shell is stronger than that in single hydrogen shell structures, when the water molecule number included in the cluster is the same. More interestingly, the ionic hydrogen bond strength in the  $C_s$  structure of  $F^-(H_2O)_3$  is stronger than that in the  $C_2$  structure of  $F^-(H_2O)_2$ . The result indicates that the ionic hydrogen bond strength is strengthened when additional water molecules are connected via water-water interaction.

In the case of F<sup>-</sup>(H<sub>2</sub>O)<sub>4</sub>, PM6 and PM6-DH+ show that the most stable structure takes the  $C_1$  symmetry, and  $F^-$  is located at the surface of the clusters. However, in the global minimum optimized by HF and B3LYP, F<sup>-</sup> is in the middle of the cluster with the  $C_1$  symmetry. Besides, MP2 favors the C<sub>s</sub> symmetry with a 0.4-kcal/mol binding energy higher than the  $C_1$  symmetry. The binding energy of the  $C_1$  structure by PM6-DH+ is closer to MP2 in comparison with PM6. However, the binding energy differences between  $C_1$  and other symmetries by PM6 or PM6-DH+ are greater than 6 kcal/mol, as compared with the differences of less than 2 kcal/mol in the MP2 level. Since the energy differences between the stable configurations are quite small at the MP2 level, the cluster tends to be delocalized and has the coexistence of several configurations. However, the semiempirical results indicate a more localized structure of the  $C_1$  symmetry. In this structure, the F...H distance (1.040 Å) at the PM6-DH+ level is close to the isolated HF bond distance (0.966 Å). Thus, the structure of HF(OH<sup>-</sup>)(H<sub>2</sub>O)<sub>3</sub> is probably formed. From the results of ab initio and DFT calculations, the longest  $r(F...H_{near})$  and r(F...O) appear in the transition state structure with the  $C_i$  symmetry. The largest and smallest  $\theta(F...H-O)$  appear in the  $C_{4h}$  symmetry and  $C_{1}$ 

structure, respectively. However, the semiempirical results show that the longest  $r(F...H_{near})$  and r(F...O) appear in the  $C_1$  and  $C_s$  structures, respectively, while the largest and smallest  $\theta(F...H-O)$  both appear in the  $C_2$  structure. Similar to the case of n=3, the *ab initio* and DFT calculations agree that the shortest  $r(F...H_{near})$  and r(F...O) appear in the  $C_s$  structure, which indicates that the strongest hydrogen bond exits in a second hydration shell structure.

As the water molecules in the  $F^-(H_2O)_n$  clusters increase from 1 to 4, the semiempirical methods represent reasonable binding energies of the global minima. However, PM6-DH+ and PM6 probably favor an optimized global minimum structure of  $HF(OH^-)(H_2O)_{n-1}$ . As n increases from 1 to 4, the F...H distance in the global minimum structure is getting closer to the HF bond distance (see Figure 2). Similar structures have been optimized by the EFP method [26] as local minima, while they are not global minima on the potential energy surface. However, the  $HF(OH^-)(H_2O)_{n-1}$  structures are not optimized by ab initio or DFT calculations. As shown in Figure 3, the ab initio and DFT calculations



calculations on ionic hydrogen-bonded structure of fluoride anion water clusters. The horizontal axis represents the water numbers and the vertical axis represents (**a**) F...H distance and (**b**) F...H...O angle.

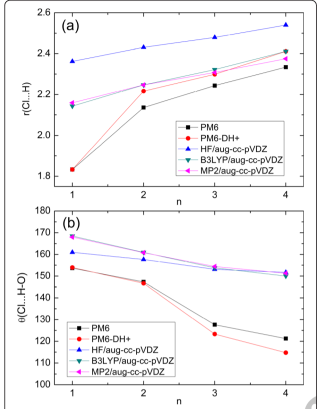


Figure 4 Comparison of semiempirical and *ab initio/DFT* calculations on ionic hydrogen-bonded structure of chloride anion water clusters. The horizontal axis represents the water numbers and the vertical axis represents (a) Cl...H distance and (b) Cl...H-O angle.

show that as n increases from 1 to 4,  $r(F...H_{near})$  and r(F...O) distances of the stable configurations with only one hydration shell increase, and the  $\theta(F...H-O)$  angle decreases. The results imply that the ionic hydrogen bond strength decreases with the increase of cluster size. However, the opposite trend emerged for the semiempirical methods. The semiempirical methods are sufficient to show that the ionic hydrogen bond strength is enhanced when the second hydration shell forms, compared with the structure without the additional water molecules. However, the semiempirical methods are not accurate enough to study the structures of small fluoride ion water clusters, where strong ionic hydrogen bond exists.

#### Chloride anion water clusters

In the Cl<sup>-</sup>( $H_2O$ ) cluster, all calculations show that the most stable structure takes the  $C_s$  symmetry. The semiempirical methods also show the  $C_{2v}$  symmetry to be a local minimum, though it is supposed to be a transition state by *ab initio* and DFT calculations. The energy difference between the  $C_s$  and  $C_{2v}$  symmetries is 0.4 kcal/mol at the

PM6-DH2 level with ZPE correction, which is slightly lower than 1.3 kcal/mol at the MP2 level. The small energy barrier indicates that the cluster geometry is very flexible. In comparison with the  $C_{2v}$  structure, the  $C_s$  structure shows shorter  $r(\text{Cl...H}_{near})$  and r(Cl...O) and a much larger  $\theta(\text{Cl...H-O})$  by all calculation levels (see Table S2 in Additional file 1 for details). The results indicate that the ionic hydrogen bond of  $\text{Cl}^-(\text{H}_2\text{O})$  is stronger in the asymmetric  $C_{2v}$  structure.

In the case of  $Cl^{-}(H_2O)_2$ , semiempirical methods show that the stable and transition state structures take the  $C_1$ and C<sub>2</sub> symmetries, respectively. The results are compatible with the ab initio and DFT calculations. The potential barrier height is 1.7 kcal/mol at the PM6-DH2 level with ZPE correction, which is higher than 0.1 kcal/mol for the MP2 level. Such small energy difference indicates that the delocalized cluster geometry varies easily under certain thermal fluctuations. The  $2(C_{2v})$  and  $2(C_{2h})$  for Cl<sup>-</sup>(H<sub>2</sub>O)<sub>2</sub> structures are not obtained by semiempirical calculations. In addition, the  $2(C_{2v'})$  structure is optimized as a local minimum by semiempirical calculations, while it is a saddle point structure for ab initio and DFT methods. Despite the underestimation of  $\theta(Cl...H-O)$ for  $C_2$  structure, the semiempirical methods could present the same trend as the ab initio and DFT methods, i.e., the bond lengths of r(Cl...H<sub>near</sub>) and r(C1...O) in the  $C_2$  structure are close to the average of two asymmetric bond lengths in the  $C_1$  structure. The results imply that the ionic hydrogen bond strength of the transition state  $C_2$  structure is between the two ionic hydrogen bond strengths of the local minimum  $C_1$ structure.

In the case of Cl<sup>-</sup>(H<sub>2</sub>O)<sub>3</sub>, all calculations show that the most stable structure takes the  $C_3$  symmetry. By including ZPE correction, it is found that PM6-DH+ and PM6-DH2 could present binding energies very close to those of MP2 as well as experimental results. From the geometry point of view, the semiempirical PM6 and PM6-DH+ show that the shortest r(Cl...O) in the  $Cl^{-}(H_2O)_3$  clusters is in the  $C_s$  structure, which is consistent with the ab initio and DFT methods. The higher calculations also show that the shortest  $r(Cl...H_{near})$  and largest  $\theta$ (Cl...H-O) are in the  $C_s$  structure, while it takes the  $C_3$  symmetry by semiempirical method. The higher calculation results indicate that the ionic bond strength is weakened in the sequence of  $C_s$ ,  $C_{3h}$ , and  $C_3$  structures. However, the semiempirical method PM6 or PM6-DH+ do not show a clear tendency.

In the case of  $Cl^-(H_2O)_4$ , the semiempirical methods show that the most stable structure takes the  $C_4$  structure, which is consistent with the higher calculation results without ZPE corrections. When considering ZPE corrections, B3LYP shows that the  $C_s$  structure becomes

the most stable, while HF and MP2 still favor the  $C_4$ structure. The PM6 and PM6-DH+ methods show that the shortest  $r(Cl...H_{near})$  and r(Cl...O) and largest  $\theta$ (Cl...H-O) come up in the second hydration shell's  $C_{\rm s}'$  structure. The results are consistent with the other calculations and indicate that the strongest ionic hydrogen bond exits in the second hydration shell structure of  $Cl^{-}(H_2O)_4$ . The longest  $r(Cl...H_{near})$  and r(Cl...O) appear in the  $C_s$  structure for semiempirical methods and in the  $C_2$  structure for higher calculations, respectively. The smallest  $\theta(Cl...H-O)$  is obtained in the stable structure with  $C_2$  symmetry for all calculations. We note that the  $C'_s$  structure of  $Cl^-(H_2O)_4$  is reported here for the first time. It is a different configuration from the  $C_s$ structure found previously. In the  $C'_s$  structure, it is interesting that the strength of the single ionic hydrogen bond is stronger than that of the other two symmetric ionic hydrogen bonds, while it is contrary to the case of the  $C_s$  structure. This new local minimum obtained from PM6 and PM6-DH+ methods could be verified through ab initio and DFT results.

The binding energies and topological characteristics of the global minimum structures presented by PM6 and PM6-DH+ are in good agreement with those of other ab initio and DFT results. Thus, it is reasonable to apply semiempirical methods on Cl<sup>-</sup>(H<sub>2</sub>O)<sub>n</sub> clusters for on-the-fly MD studies from both energetic and geometric considerations. Due to the complex coupling of ion-water and water-water interactions, the ionic hydrogen bond strength is strengthened by the formation of the second hydrogen shell. As the number of water molecules in the Cl<sup>-</sup>(H<sub>2</sub>O)<sub>n</sub> cluster increases from 1 to 4, the tendency of weakening the ionic hydrogen bond strength with the increasing number of water molecules could be obtained both from semiempirical and ab initio and DFT calculations (see Figure 4). We note that the mean values of r(Cl...H) and  $\theta$ (Cl...H-O) are adopted for cases of structures with more than one ionic hydrogen bonds. Especially for Cl...H distances, PM6-DH+ shows better results than PM6 as n increases in comparison with MP2.

# Comparison of fluoride and chloride clusters

For both fluoride and chloride clusters, PM6-DH+ could substantially improve the binding energies of PM6 in comparison with experiments as well as MP2 results. The semiempirical methods are sufficient to show the well-known results that the binding energy of fluoride cluster is larger than that of chloride with similar geometries.

From the geometric point of view, the optimized geometries by PM6 and by PM6-DH+ show little distinction for both X = F and X = Cl. The results show that the HF(OH)<sup>-</sup>(H<sub>2</sub>O)<sub>n-1</sub> structures are probable to be global minima for X = F. However, the *ab initio* and DFT results all show that F prefers to be located inside

the small  $F^-(H_2O)_n$  clusters. The strong ion-water interactions are dominated in these clusters, while they are not well estimated by PM6 or PM6-DH+. On the other hand, the semiempirical results show that it is quite ubiquitous to find the cooperation of ion-water and waterwater interactions in  $Cl^-(H_2O)_n$  clusters. The results are compatible with those of the *ab initio* and DFT calculations and that the global minima take the same topological characteristics.

#### **Conclusions**

We have performed semiempirical calculations to analyze the stabilization energies and ionic hydrogen-bonded structures of  $X^-(H_2O)_n$  (X = F, Cl; n = 1-4). The results were also compared with the high-level *ab initio* and DFT calculations.

For fluoride and chloride clusters, the stabilization energies by PM-DH+ are better than those by PM6 in comparison with those by MP2 as well as of the experimental results. On the other hand, PM6-DH+ presents tiny differences in optimized geometries from PM6. In the case of X = F, the optimized global minima are more probable to take the HF(OH)<sup>-</sup>(H<sub>2</sub>O)<sub>n-1</sub> structures, which are not supported by the *ab initio* or DFT calculations. The results may be due to the fact that the strong ion-water interactions are not well estimated by PM6 or PM6-DH+ methods. However, the optimized global minima for X = Cl present the same topological characteristics as MP2, which take  $C_s$ ,  $C_1$ ,  $C_3$ , and  $C_4$  symmetries for n = 1 to 4, respectively.

As the number of water molecules increases, the ionic hydrogen bond strength becomes weaker for  $X^-(H_2O)_n$  clusters. The ionic hydrogen bond strength of fluoride cluster is stronger than that of the chloride cluster with similar geometry. It is interesting to find that the emergence of the second hydration shell enhances the ionic hydrogen bond strengths, as compared with the structure without the additional water molecular. Meanwhile, we have reported a new stable structure of  $Cl^-(H_2O)_4$  with  $C_s'$  symmetry with the second hydration shell. The aforementioned conclusions obtained from semiempirical methods have been verified by *ab initio* and DFT calculations. The study of on-the-fly semiempirical MD or path integral MD simulation with PM6 or PM6-DH+ potentials is in progress.

### **Additional file**

**Additional file 1: Table S1:** selected ionic hydrogen-bonded structures of r (F. . . H<sub>near</sub>), r (F. . . O), and  $\theta$  (F. . . H-O). **Table S2:** selected ionic hydrogen-bonded structures of r (Cl. . . H<sub>near</sub>), r (Cl. . . O), and  $\theta$  (Cl. . . . H-O). **Table S3:** the values of the imaginary frequencies of the transition state structures. **Figure S1:** Schematic illustration of directions of the imaginary frequencies of the transition state structures.

#### Competing interests

The authors declare that they have no competing interests.

#### Authors' contributions

QW carried out the calculations and drafted the manuscript. KS conducted the *ab initio* and DFT calculations. UN, MT, and SY provided guidance at various stages of the study and reviewed the manuscript. All the authors read and approved the final manuscript.

#### Acknowledgements

Qi Wang thanks the financial support from the program of China Scholarships Council. Masanori Tachikawa thanks Grant-in-Aid for Scientific Research and for the priority area by the Ministry of Education, Culture, Sports, Science and Technology, Japan. Umpei Nagashima and Shiwei Yan acknowledge the support from the National Natural Science foundation of China under grant number 10975019.

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#### Received: 25 July 2012 Accepted: 30 January 2013 Published: 12 February 2013

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#### doi:10.1186/2251-7235-7-7

Cite this article as: Wang et al.: Semiempirical investigations on the stabilization energies and ionic hydrogen-bonded structures of  $F^-(H_2O)_n$  and  $CI^-(H_2O)_n$  (n=1-4) clusters. Journal of Theoretical and Applied Physics 2013 7:7.