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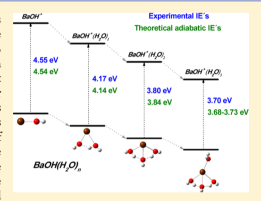


Hydration of Barium Monohydroxide in $(H_2O)_{1-3}$ Clusters: Theory and **Experiment**

Iván Cabanillas-Vidosa,† Maximiliano Rossa,‡ Gustavo A. Pino,‡ Juan C. Ferrero,*,‡ and Carlos I. Cobos*,

Supporting Information

ABSTRACT: The ionization energies (IE_e's) of small BaOH(H₂O)_m clusters (m = 1-3), as generated in a laser vaporization-supersonic expansion source have been determined by laser photoionization experiments over the 3.65-4.55 eV energy range. Complementary ab initio studies show that the IE 's are in good agreement with computed adiabatic ionization energies and that BaOH(H₂O)_m structures with a direct coordination of the Ba atom to water molecules are favored over those that are characterized by H-bonded networks involving H₂O molecules and the OH group of BaOH. Additional calculations have been performed on the hydration energies for the most stable isomers of the relevant $BaOH(H_2O)_{1-3}$ clusters. A comparison is made between the closed-shell title system and the results of related theoretical studies on the open-shell alkali monohydroxides, which allows for an interpretation of the opposite trends that are found in the cluster size dependence of the vertical



ionization energies for both series of systems, and highlights the role of the BaOH unpaired electron in its ionization process. Altogether, the present evidence suggests for the initial steps of the BaOH hydration process to be dominated by electrostatic and polarization interactions between the Ba+ and OH- ion cores, which become both increasingly solvated upon sequential addition of water molecules.

1. INTRODUCTION

Over the last 20 years, research related to the hydration of metal atoms and ions has largely benefitted from the increasing potentiality of gas phase spectroscopy and mass spectrometry to probe a variety of solvation phenomena and the related chemistry of metal-doped water clusters. 1-4 In particular, a considerable interest on group 2 of metals has been aroused due to their divalent nature that allows only one intermediate charge state, M+, between the stable neutral metal, M, and the doubly charged M²⁺ ions, which are present in solutions.³ Many aspects of the structure, energetics, and reactivity of M(H2O), $(M = Mg, Ca, Ba)^{5-7} M^{+}(H_{2}O)_{n} (M = Mg, Ca, Sr)^{8-11}$ and $M^{2+}(H_{2}O)_{n} (M = Mg, Ca, Sr, Ba)^{12,13}$ clusters have been explored at the molecular level from the experimental and theoretical viewpoints, leading to a case study of the stepwise microhydration of different metal oxidation states. In these studies, the formation of bare and hydrated MOH and MOH+ products has been reported, as a result of reactions that occur either during the cluster formation process^{6,8-11} or upon their subsequent photoionization,⁵ photodissociation,^{8–11} and collision-induced dissociation.¹² For the former case, the investigation of the hydration of a metal atom/ion or a MOH/ MOH⁺ radical in (H₂O), clusters through cluster size-resolved experiments is feasible. Recent results on the hydration and

reactions of neutral barium atoms in water clusters from this laboratory provide a case in point.⁶ A series of experiments and ab initio calculations has been conducted to show that, as a result of distinct cluster-growing processes involving groundstate and electronically excited state Ba atoms, $Ba(H_2O)_n$ (n = 1-4) and BaOH(H_2O)_m (m = 1-3) clusters are produced with similar abundances in a laser vaporization-supersonic expansion source. This was taken in the present study as an advantage point to investigate the stepwise hydration in water clusters of the BaOH radical.

The relevance of the title system relies on the open-shell nature of BaOH, which distinguishes it from closed-shell, stoichiometric $MOH(H_2O)_m$ (M = Li, Na, K, Rb, Cs)^{14–18} and $MOH^+(H_2O)_m$ (M = Mg, Ca)^{19,20} systems that have been studied previously. Owing to the neutral charge state of BaOH and the highly ionic character of its Ba-OH bond, it will be significant to contrast the present results with those obtained in studying the hydration process of the related MOH (M = alkaline metal) species, $^{14-18}$ in order to address the role of the

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[†]Instituto de Investigaciones Fisicoquímicas Teóricas y Aplicadas (INIFTA), Universidad Nacional de La Plata, Casilla de Correo 16, Sucursal 4, La Plata (1900), Argentina

[‡]Centro Láser de Ciencias Moleculares, INFIQC, Departamento de Fisicoquímica, Facultad de Ciencias Químicas, Universidad Nacional de Córdoba, Córdoba (X5000IUS), Argentina

BaOH unpaired electron on its stepwise hydration in small water clusters.

In this paper we report ionization energies for $BaOH(H_2O)_m$ (m=1-3) clusters determined by laser one-photon ionization experiments. Complementary high level ab initio calculations show that the experimental values are in good agreement with a chiefly adiabatic ionization process and allow for interpreting the nature and energetics of the BaOH hydration structures, which are formed under the prevailing experimental conditions. A comparison is made between the title system and the results of related theoretical studies on the alkali monohydroxides, $^{14-18}$ which allows for an interpretation of the opposite trends that are found in the cluster size dependence of the vertical ionization energies for both series of systems, as well as addressing the similarities and differences between the corresponding stepwise hydration behaviors.

2. EXPERIMENTAL DETAILS

The experimental setup has been fully described in previous works. 6,7,21 Briefly, neutral BaOH(H₂O)_m clusters were generated in a pick-up source combining 1064 nm laser vaporization [(0.7–1.4) J cm⁻² laser fluence] of a rotating Ba disk, with a supersonic expansion of a gaseous He:H₂O (0.988: 0.012) mixture through a pulsed solenoid valve (400 μ m diameter) at a stagnation pressure of 2 bar. The molecular beam of clusters was collimated with a 4.0 mm diameter skimmer placed 10 cm downstream from the nozzle, before entering into the ionization region of a differentially pumped Wiley–McLaren time-of-flight mass spectrometer (TOF-MS).

The one-photon ionization of the BaOH($\rm H_2O$)_m clusters was performed through the frequency-doubled output of a dye laser (0.04 cm⁻¹ bandwidth), which was pumped by the second harmonic (532 nm) of a pulsed Nd:YAG laser. The ionization energy was varied by scanning the dye laser in steps of 0.02 eV over the 272.5–340.0 nm (4.55–3.65 eV) range. Some experiments were also performed with the fourth harmonic (266 nm) of the same Nd:YAG laser as the ionization source.

The photoions were detected by dual microchannel plates (MCP), and the MCP output signal was digitized and afterward processed in a personal computer. An average of 512 events was used to generate the mass spectra, which were further normalized to the ionization laser power, as recorded on a relative basis by a photomultiplier tube. It was observed that the integrated intensity of any given $\text{BaOH}(\text{H}_2\text{O})_{n+1}$ cluster remains almost constant at energies near the IE threshold of $\text{BaOH}(\text{H}_2\text{O})_n$. This fact was used to normalize the mass spectra of the corresponding $\text{BaOH}(\text{H}_2\text{O})_n$ cluster, as the $\text{BaOH}(\text{H}_2\text{O})_n/\text{BaOH}(\text{H}_2\text{O})_{n+1}$ ratio, to determine the ionization energy of $\text{BaOH}(\text{H}_2\text{O})_n$. Numerical integration of the mass signals corresponding to the $\text{BaOH}^+(\text{H}_2\text{O})_m$ (m=1-3) clusters then led to the relevant photoionization yield curves.

The electric field in the TOF-MS ionization region ($E_z = 192 \text{ V cm}^{-1}$) reduces the apparent ionization energy of the clusters by $2e(eE_z)^{1/2}$. Hence, a correction factor of 0.01 eV was applied to the experimental ionization energy (IE_e) values.

3. THEORETICAL METHODS

As shown in a previous study addressing to neutral and cationic barium atom—water clusters, 7 a good agreement between experimental and theoretical ionization energies was obtained by combining the mPW1PW91 method of the density functional theory $(DFT)^{22}$ to derive optimized structures and

harmonic vibrational frequencies and the high correlated coupled cluster singles and doubles excitations approach, including a perturbational estimate of the triples, CCSD-(T,Full), 23,24 to compute accurate total energies. In a similar way, in the present study the energetics of the neutral and cationic BaOH(H_2O) $_m$ (m=0-3) species was derived from single-point CCSD(T,Full) calculations on mPW1PW91 geometries, using for both methods the relativistic effective core potential (RECP) developed by Lim et al. 25 for Ba and the split-valence 6-311++G(d,p) basis sets for the O and H atoms. All of the calculations were performed by using the Gaussian 09 suite of programs. 26

To analyze the stability of the barium monohydroxide—water clusters, total binding energies $[\Delta E_0(m)]$ for the hydration processes $BaOH^x+m$ $H_2O \rightarrow BaOH^x(H_2O)_m$ (with x=0 for neutral and x=+1 for cationic clusters), were estimated as:

$$\Delta E_0(m) = E_0[BaOH^x(H_2O)_m] - E_0[BaOH^x] - mE_0[H_2O]$$
 (1)

where $E_0[{\rm BaOH^x(H_2O)}_m]$, $E_0[{\rm BaOH^x}]$ and $E_0[{\rm H_2O}]$ denote the CCSD(T,Full) total electronic energies of the relevant species, including zero-point vibrational energy ($E_{\rm ZPE}$) corrections based on the $m{\rm PW1PW91}$ calculations.

As usual, vertical ionization energies ($\rm IE_v$) were computed as the total electronic energy difference, including $E_{\rm ZPE}$ corrections, between the cationic and neutral species, taken both at the optimized geometries of the neutral clusters. In a similar way, adiabatic ionization energies ($\rm IE_a$) were derived from the optimized geometries of the cationic and the corresponding neutral clusters.

Additional calculations have been performed to evaluate the hydration energies ($E_{\rm hyd}$'s) of both Ba- and BaOH-doped water clusters, as defined by the reactions

$$Ba^{x}(H_{2}O)_{n-1} + H_{2}O \rightarrow Ba^{x}(H_{2}O)_{n}$$
 (2)

$$BaOHx(H2O)m-1 + H2O \rightarrow BaOHx(H2O)m$$
 (3)

and calculated according to the expressions:

$$E_{\text{hyd}}(n) = E_0[Ba^x(H_2O)_n] - E_0[Ba^x(H_2O)_{n-1}] - E_0[H_2O]$$
(4)

$$E_{\text{hyd}}(m) = E_0[\text{BaOH}^x(\text{H}_2\text{O})_m] - E_0[\text{BaOH}^x(\text{H}_2\text{O})_{m-1}] - E_0[\text{H}_2\text{O}]$$
(5)

where $E_0[\mathrm{Ba}^x(\mathrm{H_2O})_n]$ is the energy of $\mathrm{Ba}^x(\mathrm{H_2O})_n$ clusters. The optimized geometries for the most stable $\mathrm{Ba}(\mathrm{H_2O})_n$ (n=0-4) isomers were taken from ref 7.

4. EXPERIMENTAL RESULTS

The formation of $BaOH(H_2O)_m$ clusters in the reaction of electronically excited Ba atoms with water clusters (eq 6), has been recently demonstrated.⁶

$$Ba^* + (H_2O)_{m+1} \to BaOH(H_2O)_m + H$$
 (6)

The internal energy excess of the resulting barium monohydroxide—water clusters may be released in the early stages of the supersonic expansion, through collisions with He and by the following evaporative process:

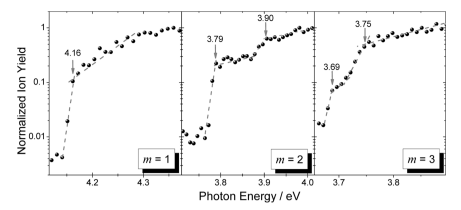


Figure 1. Photoionization yield curves for $BaOH(H_2O)_m$ clusters. The position of the lower breaks associated with the IE_e 's are indicated by arrows.

Table 1. Experimental and Calculated Vertical and Adiabatic Ionization Energies for BaOH $(H_2O)_m$ $(m = 0-3)^a$

		ionization energy (eV)						
			IE_{v}			IE_a		
$BaOH(H_2O)_m$	ΙE _e	${\it neutral\ structure}^b$	DFT	CCSD(T)	ionic structure ^c	DFT	CCSD(T)	
m = 0	4.55 ± 0.03^d	BaOH	4.60	4.58	$BaOH^+$	4.55	4.54	
m = 1	4.17 ± 0.05	1U01	4.38	4.36	1U01 ⁺	4.28	4.28	
		1U10	4.13	4.05	1U11 ⁺	4.09	4.01	
		1U11	4.35	4.32	1U11 ⁺	4.20	4.14	
m = 2	3.80 ± 0.05	2U02	4.81	4.77	2U01+	4.37	4.38	
		2U20	3.95	3.80	2U21 ⁺	3.78	3.63	
	3.91 ± 0.05	2U11	4.25	4.21	2U21 ⁺	4.00	3.87	
		2U22	4.10	4.06	2U22 ⁺	3.94	3.84	
m = 3	3.70 ± 0.05	3U02	4.72	4.70	3U01 _b ⁺	4.47	4.50	
		3U23	4.02	3.98	3U23 ⁺	3.75	3.68	
		3U12 _a	4.04	4.01	3U22 _a +	3.81	3.70	
		3U20	4.00	3.80	3U21 _b ⁺	3.47	3.31	
	3.76 ± 0.05	3U11	4.20	4.17	3U21 _a ⁺	3.73	3.50	
		3U12 _b	4.00	3.96	3U32 _a +	3.62	3.55	
		3U22	4.00	3.95	3U22 _b ⁺	3.80	3.73	
		3D33	3.97	3.95	3U33 ⁺	3.70	3.54	
		3U32	3.89	3.79	3U32 _b +	3.66	3.53	

^aSelected hydration structures are highlighted (see text for details). ^bSee neutral clusters in Figure 2. ^cSee ionic clusters in Figure 3. ^dExtracted from ref 21.

$$Ba^* + (H_2O)_{m+1} \to BaOH(H_2O)_{m-p} + H + pH_2O$$
 (7)

Since the pick-up process occurs mostly in the postexpansion region, where the probability of multiple collisions with the buffer gas is low, the evaporation of water molecules is likely the dominant stabilization process for neutral $BaOH(H_2O)_m$ clusters.

All of the observed $BaOH^+(H_2O)_m$ ions arise from onephoton ionization of the corresponding neutral clusters that are entrained in the molecular beam, that is,

$$BaOH(H2O)m + h\nu \rightarrow BaOH+(H2O)m$$
 (8)

Considering that the calculated energies for the $BaOH^+(H_2O)_m \rightarrow BaOH^+(H_2O)_{m-1} + H_2O$ process are 1.03, 0.93, and 0.88 eV for m=1, 2, and 3, respectively, the evaporative photoionization of neutral $BaOH(H_2O)_m$ species can be neglected so that fragmentation-free photoionization is assumed under the prevailing, near-threshold ionization conditions.

Figure 1 shows the photoionization yield curves for the $BaOH(H_2O)_{1-3}$ clusters. As in the case of $Ba(H_2O)_{1-4}$, the lowest post-threshold breaks in the logarithmic plot of the ion

yield as a function of the photon energy (Watanabe-type plots) were assigned to the ${\rm IE_e}$'s for the various ${\rm BaOH}({\rm H_2O})_m$ clusters. The resulting values, which represent the average of three individual measurements, are listed in Table 1. The associated uncertainty of 0.05 eV accounts for the deviation of these measurements and includes an estimate of the error arising from the fact that the photoionization energy was varied in steps of 0.02 eV. Higher-energy breaks can be also observed in Figure 1 for ${\rm BaOH^+}({\rm H_2O})_{2,3}$. The corresponding ${\rm IE_e}$ values are listed in Table 1.

As can be seen in Table 1, all of the $\rm IE_e$'s for $\rm BaOH(H_2O)_m$ (m=1-3) are smaller than the value of (4.55 \pm 0.03) eV that was previously determined for bare $\rm BaOH$. In addition, these $\rm IE_e$'s decrease systematically with the increasing size m of the cluster. This behavior is quite similar to that observed previously for $\rm Ba(H_2O)_{1-4}$ clusters. Yet, the observation of two lowest and close $\rm IE_e$'s for both $\rm BaOH(H_2O)_2$ and $\rm BaOH(H_2O)_3$ species might be rationalized in a number of ways, including the presence in the molecular beam of either nearly isoenergetic isomers, or two low-lying vibrational states of a given isomer for such clusters. To elucidate this, ab initio

calculations of stable structures, binding energies, and vertical and adiabatic ionization energies were performed.

5. THEORETICAL RESULTS

5.1. BaOH Radical. The calculated geometry of the BaOH radical agrees very well with previous experimental and theoretical values. The linear structure is well-reproduced, and the computed Ba–O bond distance of 2.204 Å is in excellent agreement with the experimental value of 2.201 Å. The similarity of the neutral and cationic BaOH geometries leads to similar vertical and adiabatic IE's of 4.58 and 4.54 eV, respectively, which are also in very good agreement with the recent experimental determination of $(4.55 \pm 0.03 \text{ eV})$.

Figures 2 and 3 show the optimized structures of BaOH and BaOH⁺, respectively, along with the corresponding charge

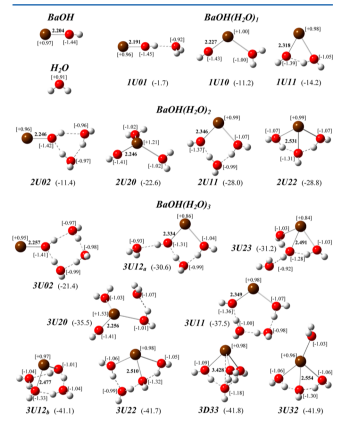


Figure 2. Optimized structures of neutral $BaOH(H_2O)_m$ (m=0-3) clusters calculated at the mPW1PW91 level. Binding energies (in kcal mol^{-1}) calculated at the CCSD(T,Full) level are given under each structure. The Ba-OH lengths (R_0) are given in angstroms, and the natural population analysis evaluated at the DFT level is indicated between brackets.

distributions, as estimated employing a natural population analysis treatment. It is evident from these results that both species are strongly polarized as $Ba^+(OH)^-$ and $Ba^{2+}(OH)^-$.

5.2. Solvation of BaOH in Water Clusters. Figure 2 shows the mPW1PW91 optimized conformations of BaOH- $(H_2O)_m$ (m=0-3) and the computed ΔE_0 values at the CCSD(T,Full) level of theory. As expected, the number of minimum energy structures increases significantly as the number of water molecules increases.

In a previous study on $Ba(H_2O)_n$ clusters,⁷ the different isomers were classified according to the number of water molecules present in the first, second, third, and fourth

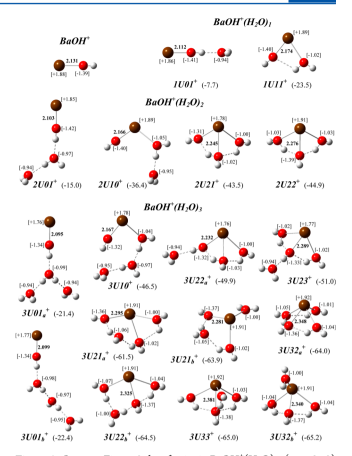


Figure 3. Same as Figure 2, but for ionic $BaOH^+(H_2O)_m$ (m=0-3) clusters.

hydration shells of the Ba atom. For the present system, it is necessary instead to distinguish between the water molecules that are bonded to the Ba atom from those bonded to the OH group of BaOH. Hence, we used here the notations " $mUwy_i$ " and " $mDwy_i$ " proposed by Kim et al., ^{14–17} where m is the total number of water molecules, w and y are the hydration numbers of Ba and OH, respectively, U/D indicates the undissociated/ dissociated state of Ba—OH, and the subscript i distinguishes among different conformers of the same isomer.

Three stable structures were found for $BaOH(H_2O)_1$. The most stable isomer, 1U11, presents both H₂O and OH bounded to Ba and has a binding energy of $\Delta E_0 = -14.2$ kcal mol⁻¹. The O-Ba-O frame is bent (60.5°), and the water molecule is H-bonded to the OH group. The 1U01 isomer is the less stable of the three $(-1.7 \text{ kcal mol}^{-1})$ and lacks direct Ba-water interactions, and the oxygen atom of the water molecule is H-bonded to the OH group. The 1U10 isomer is intermediate in energy $(-11.2 \text{ kcal mol}^{-1})$. The 1U10 structure lacks for H-bonds and the O-atom of water interacts directly with the Ba atom $[<(O-Ba-O) = 110.8^{\circ}]$. The Ba-OH(H₂O)₁ distance gradually increases in the order 1U01, 1U10, and 1U11 (2.19, 2.23, and 2.32 Å, respectively), and it is found that hydration structures of the HOBa-OH2 type are much more stable than those of the BaOH-OH₂ type. The latter is also observed in higher order clusters (m > 1), which determines that the corresponding analogues to 1U01 having the BaOH-OH₂ structure are generally much less stable than the remaining stable isomers.

Four stable structures have been located on the potential energy surface (PES) of BaOH(H₂O)₂. The most stable

corresponds to the 2U22 isomer $(-28.8 \text{ kcal mol}^{-1})$ with both water molecules directly bonded to the Ba atom and H-bonded to the OH group. In addition, a cyclic isomer, 2U11, with an energy difference of only 0.8 kcal mol^{-1} with respect to the 2U22 isomer was found. This isomer exhibits only one Ba— $O(H_2)$ interaction, and the second water molecule is bonded to both the former and the OH group by two H-bonds. The 2U20 isomer $(-22.6 \text{ kcal mol}^{-1})$ has both water molecules directly bonded to the Ba atom and lacks for H-bonds. The corresponding $(H_2)O-Ba-OH$ and $(H_2)O-Ba-O(H_2)$ angles are 120.8° and 95.5° , respectively, with the Ba and the three O atoms in a nearly pyramidal geometry. The 2U02 isomer has a six-membered ring structure, and it lacks direct metal—water interactions. This fact probably accounts for its observed minor relative stability $(-11.4 \text{ kcal mol}^{-1})$.

Nine stable structures have been characterized for the BaOH(H₂O)₃ cluster. The four more stable isomers (3U12_b, 3U22, 3D33, and 3U32) are essentially iso-energetic, with a relative energy difference <0.8 kcal mol⁻¹. Out of them, the two lowest-energy isomers, namely, 3D33 and 3U32, have the three water molecules bonded to barium atom and at least two Hbonds to the OH group. A higher-energy 3U11 isomer (-37.5)kcal mol⁻¹) has a cyclic structure with one Ba-O(H₂) interaction. The 3U20 isomer presents two water molecules bonded to the Ba atom, that is, one more than 3U11. The former isomer is 2 kcal mol⁻¹ less stable than the latter, probably because the third H-bonded water molecule in 3U20 can be effectively considered in the second hydration shell of Ba. The $3U12_a$ (-30.6 kcal mol⁻¹) and 3U23 (-31.2 kcal mol⁻¹) isomers are related to 2U11 and 2U22, respectively. Both add a water molecule that is H-bonded to the OH group. The 3U02 isomer was found to be the highest in energy (-21.4)kcal mol⁻¹), with a H-bonded ring structure between the OH group and the water molecules. As for the 2U02 isomer of BaOH(H₂O)₂, the lack of direct metal-water interactions in 3U02 could explain its lower stability with regard to the remaining $BaOH(H_2O)_3$ isomers.

The above results indicate that the sequential addition of water molecules to BaOH leads to increases in (i) the number of H-bonding interactions within the BaOH($\rm H_2O$) $_m$ cluster, in (ii) the coordination number of both the Ba atom and the OH group, and in (iii) the Ba—OH bond distance that, compared to the value of 2.20 Å for bare BaOH, is 2.318 Å for 1U11 and 2.554 Å for 3U32.

Such effects are accompanied by an increasing trend for the generation of stable dissociated isomers with increasing hydration. Whereas all of the lowest-energy isomers of $BaOH(H_2O)_{1,2}$ clusters are undissociated, one stable lowenergy isomer of $BaOH(H_2O)_3$ (3D33) is found to be dissociated (Ba–OH bond distance of 3.43 Å).

A larger number of stable dissociated species has been found for the most stable isomers of $BaOH(H_2O)_{4,5}$ at the mPW1PW91 level of theory (Figure 4). For these species the Ba-OH bond distance also increases with the hydration number, for example, 3.726 Å for 4D33 and 3.938 Å for $5D33_a$. In addition, the undissociated 4U22 and 5U22 isomers exhibit binding energies very close to those of the nearly iso-energetic, dissociated isomers. Seemingly, five solvent molecules do not lead to BaOH full dissociation.

5.3. Ionization Energies of BaOH(H_2O)_m **Clusters.** To calculate the adiabatic ionization energies of $BaOH(H_2O)_m$ clusters, the corresponding cationic structures were optimized at the mPW1PW91 level of theory. The derived geometries and

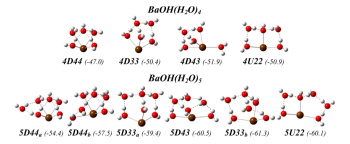


Figure 4. Same as Figure 2, but for selected most stable $BaOH(H_2O)_m$ (m = 4 and 5) clusters. The binding energy (in kcal mol^{-1}) next to the label of each structure was calculated at mPW1PW91 level. The results show an increase in the number of dissociated stable geometries as a function of m.

CCSD(T,Full) binding energies are given in Figure 3. As for neutral clusters, the hydratation in the $BaOH^+(H_2O)_m$ species is dominated by the direct coordination of the water molecules to the Ba atom center over formation of H-bonded networks involving the OH group. For example, the $1U11^+$, $2U22^+$, and $3U32^+$ isomers, which exhibit water molecules bonded by Ba^+ O interactions, are more stable than the $1U01^+$, $2U01^+$, and $3U01^+$ isomers, which have a H_2O molecule H-bonded to the OH group.

Table 1 lists the vertical and adiabatic ionization energies for the BaOH($\rm H_2O$) $_m$ (m=0-3) clusters derived from the DFT and the CCSD(T,Full) methods. As it can be seen, both levels of theory lead to very close $\rm IE_v$ and $\rm IE_a$ values for all m. A good agreement between the experimental and calculated $\rm IE$'s is apparent, and the systematic decrease of $\rm IE_e$'s with increasing size of the cluster is well-reproduced. Such a noteworthy trend is opposite to the reported behavior in previous theoretical studies of the $\rm MOH(H_2O)_m$ ($\rm M=Li, Na, K, Rb, Cs$) systems, $\rm ^{14-17}$ for which the $\rm IE_v$'s, as derived from the MP2 method were found to increase consistently with the number of water molecules. Unfortunately, no experimental data seem to be available to confirm the findings for the hydrated alkaline hydroxides.

A comparison between the experimental and predicted ionization energies for $BaOH(H_2O)_m$ (m=0-3) is relevant at this time. In fact, except for the 1U10 and 2U20 isomers, the IE_v 's for the mono-, di-, and trihydrated clusters are larger than the IE_e values. Instead, the calculated adiabatic ionization energies for the most stable $BaOH(H_2O)_{1,2}$ clusters are very similar to the IE_e 's. The same holds for $BaOH(H_2O)_3$ species except for their lowest-energy isomers.

The adiabatic ionization process for the less stable clusters involves a substantial geometry rearrangement of the ionic species. For example, the *ring* structure is broken upon the $2\text{U}02 + h\nu \rightarrow 2\text{U}01^+$ or $3\text{U}02 + h\nu \rightarrow 3\text{U}01_b^+$ processes. As a result, the adiabatic process for these clusters is likely to be spectroscopically unfavorable. Considering that the actual ionization process probably involves a structural relaxation of the ion to some extent, the computed IE_v's can be seen as upper theoretical limits.

6. DISCUSSION

6.1. Comparison of Experimental and Theoretical Results. As above-mentioned, the theoretical analysis shows that cluster structures with Ba—OH₂ bonds are mostly favored. The quantum chemical results of Table 1 also indicate that the more energetic isomers probably do not account for the

experimental data. In fact, the calculated $\rm IE_a$'s for the 1U01, 2U02, and 3U02 configurations (4.28, 4.38, and 4.50 eV, respectively) are systematically higher than the $\rm IE_e$'s (4.17, 3.91, and 3.76 eV).

Regarding the most stable isomers of the whole BaOH- $(H_2O)_m$ (m=1-3) set, the CCSD(T,Full) values for IE $_v$ and IE $_a$ are close to the experimentally measured values. This is the case of the 1U11 isomer, for which IE $_e$ = 4.17 \pm 0.05 eV lies between the calculated IE $_v$ (4.32 eV) and IE $_a$ (4.14 eV). On the contrary, the computed IE $_v$ and IE $_a$ for the 1U10 isomer of 4.05 and 4.01 eV, respectively, are somewhat lower than the photoionization threshold for BaOH⁺(H₂O) $_1$ ions at 4.17 eV. On this basis, it is suggested that the 1U11 isomer is the most populated BaOH(H₂O) $_1$ cluster of this series in the molecular beam.

The lowest IE_e for $BaOH(H_2O)_2$ at 3.80 eV could be certainly assigned to the adiabatic photoionization of the 2U22 isomer. While the lowest IE_e also coincides with the CCSD(T,Full) IE_v (3.80 eV) of the 2U20 isomer, the present calculations show that the corresponding $2U20^+$ structure is unstable. Besides, the expected population of the 2U20 isomer in the molecular beam is much lower than that of the most stable isomer 2U22. Even if the 2U20 isomer is present in the molecular beam, its vertical photoionization would imply a strong reorganization in the formed cationic cluster, thereby turning unlikely the whole processes. This fact, along with the lack of observation for an IE_e feature similar to the IE_a of 2U20 (3.63 eV), suggests that this isomer may be at least a minor component in the molecular beam.

A rationalization of the higher-energy IE_e for $BaOH(H_2O)_2$ at 3.91 eV should be addressed. Whereas this is in agreement with the IE_a value for the 2U11 isomer, alternative assignments cannot be excluded at present. The latter include the presence in the molecular beam of two low-lying vibrational states of a given isomer for BaOH(H2O)2, as well as the promotion of a given neutral isomer in its vibrational ground-state to two different low-lying vibrational states of its cationic form upon photoionization. Indeed, the energy difference of (0.11 ± 0.07) eV $[(887 \pm 565) \text{ cm}^{-1}]$ between the two lowest IE_e's for BaOH(H₂O)₂ lies within the energy range of vibrational quanta, which are associated to 7-8 normal modes for all of the isomers involved in the $2U22 + h\nu \rightarrow 2U22^+$ and $2U11 + h\nu \rightarrow$ 2U21⁺ processes, as derived from the present calculations. Hence, the discussion hereafter will rely on considering the lowest IE_e for BaOH(H₂O)₂ alone which, as mentioned above, is in agreement with the IE_a value of the 2U22 isomer.

The above considerations hold true for BaOH(H₂O)₃ as well, on the basis of comparing the (0.06 \pm 0.06) eV [(484 \pm 484) cm⁻¹] energy difference between its two lowest IE_e's and the values of the calculated harmonic vibrational frequencies for the relevant isomers (see Supporting Information). Hence, only the lowest IE_e for BaOH(H₂O)₃ will be considered hereafter. The CCSD(T,Full) computed IE_v and IE_a values for three out of nine BaOH(H_2O)₃ isomers agree with the lowest IE_e at 3.70 eV. Because the estimated IE_a (4.50 eV) of the highest-energy 3U02 species is 0.8 eV higher than the IEe's, this isomer could be, in principle, discarded. It is interesting to focus now on the group of the most stable, near iso-energetic isomers ($\Delta E_0 \approx$ -42 kcal mol⁻¹): 3U12_b, 3U22, 3D33, and 3U32. Whereas the adiabatic IE (3.73 eV) of the 3U22 isomer agrees with the lowest IE_e within the assigned uncertainty (0.05 eV), the IE_a's for the remaining isomers are very similar, ~3.54 eV, and in all cases somewhat lower than relevant IEe. Regarding the

remaining four isomers, that is, $3U12_a$, 3U23, 3U20, and 3U11, only $3U12_a$ and 3U23 have IE_a values of 3.70 and 3.68 eV, respectively, in agreement with the relevant IE_e . Therefore, an unambiguous assignment of the lowest IE_e for $BaOH(H_2O)_3$ clusters to an adiabatic or vertical photoionization process of any predicted isomer does not appear to be possible.

Figure 5 compares the lowest IE_e 's with the theoretical IE_v 's and IE_a 's values of the relevant $BaOH(H_2O)_m$ clusters, as

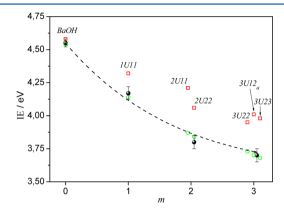


Figure 5. Experimental and calculated IE's for $BaOH(H_2O)_m$ (m=0-3) clusters as a function of m. \bullet , lowest experimental IE_e ; red \square , vertical IE_v ; green \square , adiabatic IE_a . The theoretical values correspond to the structures highlighted in Table 1.

derived from the CCSD(T,Full) method: overall, there is a good agreement between the IE_e 's and the calculated IE_a 's for such isomers.

6.2. BaOH Solvation in $(H_2O)_m$ (m=1-5) Clusters. The present results indicate that the sequential attachment of water molecules to BaOH changes its linear geometry to some extent (see Figure 2) as well as the Ba-OH bond length. This is a consequence of the changes in the local charges of the Ba and O atoms in BaOH, and thus to the polarization of the Ba-OH bond, as induced by the water molecules.

The strong Ba⁺(OH)⁻ ionic character of the BaOH radical is observed in all of the $BaOH(H_2O)_m$ clusters. To obtain reliable estimates of the charge distribution on the Ba and O atoms of the BaOH radical, a natural population analysis on the characterized clusters was carried out. The resulting values are listed in Figure 2. It is apparent that the charges change slightly when the O atom of the water molecule is H-bonded to OH group. For the weakly bonded 1U01 complex, a Ba-OH bond length decrease of 0.013 Å with respect to bare BaOH is observed. Instead, the O atom charge in BaOH becomes less negative when the OH group acts as a proton acceptor. This is the case for the 1U11 isomer of BaOH(H₂O)₁. As a result of strong HO-HOH and Ba-OH2 interactions, 1U11 is most stable ($\Delta E_0 = -14.2 \text{ kcal mol}^{-1}$) than 1U01 and displays a Ba-OH bond polarization decrease and a Ba-OH bond distance increase of 0.114 Å.

Figure 6 shows relaxed PES scans performed at the mPW1PW91 level for selected neutral BaOH(H₂O) $_m$ species as a function of the Ba-OH bond distance. As expected, the dissociation of bare BaOH is a barrierless process having a well depth (D_e) of 125.9 kcal mol $^{-1}$ (Figure 6a). On the other hand, for BaOH(H₂O) $_m$ (m=1-5) clusters, a H-transfer from one water molecule to the OH group leads to a H₂O elimination channel, that is, BaOH(H₂O) $_{m-1}$ + H₂O, instead of the competitive OH elimination Ba(H₂O) $_m$ + OH. This is

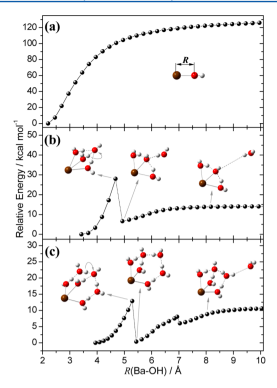


Figure 6. Potential energy curves calculated at the mPW1PW91 level, for the ground neutral state of BaOH(H₂O)_m as a function of the BaOH bond length (from the equilibrium distance, R_0). (a) m = 0, BaOH; (b) m = 3, 3D33 isomer; (c) m = 5, 5D33_a isomer.

illustrated in Figure 6b and c for the 3D33 and 5D33_a isomers of the BaOH($\rm H_2O$)₃ and BaOH($\rm H_2O$)₅ clusters. Upon $\rm H_2O$ elimination, the resulting species, BaOH($\rm H_2O$)₂ and BaOH-($\rm H_2O$)₄ are stabilized by 14.0 and 2.3 kcal mol⁻¹ with respect to the highest energy point along the reaction coordinate. These results indicate that, despite the Ba–OH bond distance increases with m, five solvent molecules are not enough to fully dissociate the BaOH radical. Overall, the present experimental and theoretical results for BaOH($\rm H_2O$)_m together with results of previous theoretical studies related to the $\rm MOH(\rm H_2O)_m$ ($\rm M=Li, Na, K, Rb, Cs$) systems, $^{14-17}$ allow estimates in the number of water molecules which seem to be required for full dissociation of the relevant metal monohydroxides: m > 5 for BaOH, 7 for LiOH, 16 6 for NaOH and KOH, 17 5 for RbOH, 15 and 4 for CsOH. 14

6.3. Solvation of BaOH vs Ba and MOH (M = Li, Na, K, Rb, Cs) in Water Clusters. A theoretical energy level diagram of the hydration energies for the relevant BaOH- and Ba-doped water clusters, for both neutral and cationic species is depicted in Figure 7. It is apparent that the hydration energies of BaOH- $(H_2O)_m$ clusters are larger than those of Ba- $(H_2O)_n$ species for m = n, for example, the E_{hvd} 's for BaOH-H₂O (14.2 kcal mol⁻¹) and BaOH(H₂O)-H₂O (14.5 kcal mol⁻¹) are larger than those for Ba- H_2O (11.2 kcal mol⁻¹) and Ba(H_2O)- H_2O (12.3 kcal mol⁻¹). A similar trend is observed by comparing the hydration energies of $BaOH^+(H_2O)_m$ and $Ba^+(H_2O)_n$ clusters, which are found to be about twice those for the corresponding neutral species. All findings can be rationalized on the basis of the results of the natural population analysis performed here for neutral and cationic BaOH $(H_2O)_m$ clusters and previously for the corresponding $Ba(H_2O)_n$ species. Indeed, Figures 2 and 3 show that the BaOH/BaOH⁺ cores of all neutral/cationic clusters are significantly polarized, with effective charges of

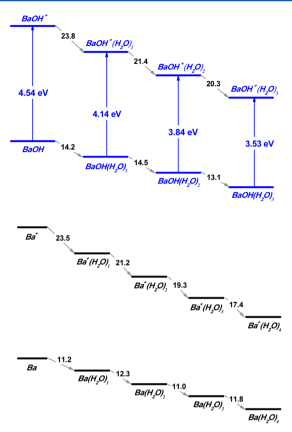


Figure 7. Theoretical energy level diagram performed at the CCSD(T,Full)//mPW1PW91 level, for hydration of BaOH (upper plot) and Ba (lower plot; extracted from ref 7), in the neutral and cationic states. Gray diagonal arrows are the hydration energies ($E_{\rm hyd}$) in kcal mol $^{-1}$, and blue vertical arrows are the adiabatic ionization energies ($E_{\rm hyd}$) in eV. The most stable isomer for each cluster was used.

-(1.2-1.4) on the O atom of the OH group and of +(0.9-1.2-1.4)(1.0)/+(1.8-1.9) on the Ba atom of BaOH(H₂O)_m/ BaOH⁺(H₂O)_m clusters. This situation contrasts with the relatively low Ba atom charges of +(0.03-0.20) in Ba $(H_2O)_n$ (n = 1-4) clusters, and of around +(1.0) in $Ba^+(H_2O)_n$. The relatively strong polarization of the BaOH and BaOH⁺ cores strengthens the bonding with water molecules as compared to the Ba and Ba⁺ cases, which in turn determines that the E_{hyd} 's of all $BaOH(H_2O)_m/BaOH^+(H_2O)_m$ clusters are much larger than those of $Ba(H_2O)_n/Ba^+(H_2O)_n$ for m = n. Similar trends have been found by Watanabe et al. ^{19,20} for the reactions between Mg⁺ and Ca⁺ ions with (H₂O)_n clusters. These theroretical studies were aimed to explain previous experimental observations that $M^+(H_2O)_n$ species are predominantly produced for n ≤ 5 and 4 for Mg⁺ and Ca⁺, respectively, whereas the $MOH^+(H_2O)_{n-1}$ products dominate in the ranges of 6–14 and 5-13, respectively.

A further insight into the stepwise hydration behavior of the BaOH radical could be gained from comparing the cluster size dependence of the theoretical IE_v 's for the most stable isomers of $BaOH(H_2O)_m$ (m=0-3) clusters, with the corresponding values for the $MOH(H_2O)_m$ (M=Li, Na, K, Rb, Cs) systems, as derived from MP2 calculations by Kim and co-workers. ^{14–17} Despite that the lack of CCSD(T,Full) IE_v 's for the hydrated alkali monohydroxides does not allow for a straightforward comparison to be made with the corresponding values for the title system, it was found that the IE_v 's for the most stable $MOH(H_2O)_m$ (M= alkaline metal) isomers generally increase

with the number of water molecules, which is opposite to the trends found for $BaOH(H_2O)_m$. The differing behaviors may be rationalized considering the nature of the photoionization process in bare BaOH and MOH first. Owing to the highly ionic character of the Ba-OH and M-OH bonds, ionization in such species can be envisaged as electron removal either from the Ba⁺/M⁺ or the OH⁻ moieties. The former is true in the case of BaOH, for which removal of the unpaired electron in its ground state occurs from a nonbonding orbital located primarily on the Ba+ ion core and polarized away from the OH ion, 21 whereas ionization of MOH corresponds essentially to removal of the electron from the OH⁻ moiety.²⁸ This picture is likely to hold for BaOH(H2O)_m clusters considering the effective charges on the Ba atom for BaOH(H2O)_m and $BaOH^+(H_2O)_m$ (Figures 2 and 3), i.e., the ionization process can be considered as $[Ba^+OH^-(H_2O)_m] + h\nu \rightarrow$ $[Ba^{2+}OH^{-}(H_2O)_m]^+ + e^-$. As the number m of water molecules increases, the solvation of the neutral BaOH $(H_2O)_m$ clusters is not as strong as for the corresponding ions BaOH+(H2O),,, because of the stronger charge-charge and charge-dipole interactions that are at play between the Ba²⁺ and OH⁻(H₂O)_m cores in the ionic clusters. In turn, this will lead to a decrease in the ionization energy of BaOH(H2O),, with increasing cluster size. The opposite picture holds in the case of MOH(H₂O)_m clusters, for which the ionization process can be considered as $[M^+OH^-(H_2O)_m] + h\nu \rightarrow [M^+OH(H_2O)_m]^+ + e^-$, thereby resulting in stronger charge—charge and charge—dipole interactions at play between the M⁺ and OH⁻(H₂O)_m cores in the neutral clusters than in cationic clusters, and leading to an increase in the $MOH(H_2O)_m$ ionization energy with increasing cluster size.

The above considerations bring out the significance of the open-shell nature of BaOH in determining the hydration structures and electronic properties of small BaOH-doped water clusters, as compared to the closed-shell $\mathrm{MOH}(\mathrm{H_2O})_m$ (M = Li, Na, K, Rb, Cs)^{14–18} and $\mathrm{MOH^+(H_2O)}_m$ (M = Mg, Ca)^{19,20} systems. For the latter, electrostatic interactions dominate the bonding of $\mathrm{MOH/MOH^+}$ to water molecules. This is revealed by the finding for hydrated alkali monohydroxides that a fully dissociated conformation is usually attained as the number of solvating water molecules is large enough, so as to tricoordinate the $\mathrm{OH^-}$ moiety with water molecules. Instead, the required hydration number for the series of isovalent $\mathrm{M^+}$ moieties, having a closed-shell spherical electron distribution, depends chiefly on their charge-to-radius ratio and generally decreases with the corresponding atomic mass. ^{14–18}

As pointed out above, electrostatic interactions contribute significantly to bonding between the strongly polarized BaOH radical and the water molecules as well. Notwithstanding, inductive effects on the BaOH unpaired electron distribution should be considered in order to address the different dissociation behavior in (H2O)_m clusters with respect to the alkali monohydroxides, $^{14-18}$ especially CsOH that seems to require a smaller number of water molecules for full dissociation (m = 4), as compared to BaOH (m > 5). The difference might be explained in terms of the effect that the successive binding of water molecules to BaOH is expected to have on the electron distribution around its Ba+ moiety. In bare ground-state BaOH, a polarized electron density on the opposite side of the barium atom from the OH moiety is predicted to result from repulsion between the unpaired electron and the electron cloud of the OH⁻ moiety. In analogy with the alkaline earth monohalides, ²⁹ such a polarization is achieved through $6s\sigma$ – $6p\sigma$ orbital mixing on the barium atom, which produces a nonbonding orbital where the BaOH's valence (unpaired) electron resides. This is corroborated by additional calculations on the electron density distribution of the singly occupied molecular orbital (SOMO) for bare BaOH (Figure 8). Within this picture, a fully dissociated conformation

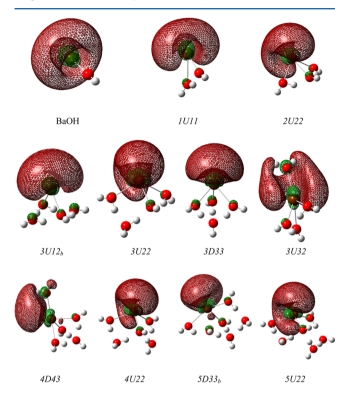


Figure 8. Electronic density distribution of the SOMO in BaOH- $(H_2O)_m$ clusters (m=0-5). The iso-density surfaces correspond to 0.03 Å⁻³.

in BaOH(H2O), clusters is expected to develop upon attainment of spatial charge polarization¹⁸ of the hydrated Ba+ and OH- moieties, along with delocalization to some extent of the corresponding unpaired electron over the surrounding water molecules. Figure 8 shows theoretical results on the SOMO's for the most stable isomers of BaOH $(H_2O)_{1-5}$ clusters as well. These SOMO's extend in a large vacant space around the barium atom and in a direction perpendicular to the Ba-OH bond, which indicates that they retain a nonbonding character upon sequential solvation by less than five water molecules. Most significantly, these unpaired electron density distributions are chiefly polarized away from the electron clouds on the oxygen atoms of both the water molecules and the OHmoieties, which for all cluster sizes are found to be more negative than the oxygen atom of bare H₂O (Figure 2). This is especially true for the oxygen atoms of the OH moieties, which in turn suggests that, despite the fact that Ba-OH bond distance increases with the cluster size (see Section 6.2), the OH⁻ ion exerts a major influence on the Ba⁺ counterion, to the extent that the valence electron of the most stable BaOH-(H₂O)₁₋₅ isomers do not effectively delocalize over such small water clusters. As a result, these hydrated OH moieties might be thought of interacting with nearby, closed-shell Ba2+ ion cores that are partially shielded by their (valence) electron clouds. The latter is expected to lead to stronger charge-dipole interactions with water molecules, as compared to the Cs⁺ ion

case, which in turn would explain the relatively large hydration numbers that are required for dissociating the BaOH radical into spatially polarized Ba⁺ and OH⁻ moieties.

7. CONCLUDING REMARKS

The present work reports the determination in laser onephoton ionization experiments of the ionization energies of $BaOH(H_2O)_m$ (m=1-3) clusters. Complementary ab initio calculations show that the experimental values are in good agreement with a chiefly adiabatic ionization process, while they allow for an assessment of the most likely BaOH hydration structures under the prevailing experimental conditions. Both the experimental and theoretical IE's for the open-shell BaOH(H₂O)_m clusters decrease with the number of water molecules, which is opposite to the trends reported for the IE,'s of the closed-shell $MOH(H_2O)_m$ (M = Li, Na, K, Rb, Cs) systems. On the basis of calculated hydration energies, atomcharge and valence-electron distributions for the most stable isomers of BaOH $(H_2O)_m$ clusters with up to m = 5, it was shown that the opposite behaviors is a consequence of the different electrons that are removed upon photoionization: The unpaired electron on the Ba+ moiety for BaOH and an electron of the OH⁻ moiety for closed-shell MOHs. Overall, the present evidence is compatible with an increasing solvation of the Ba⁺ and OH- ion cores upon sequential addition of a small number of water molecules to BaOH, and it is also suggestive for the initial steps of the BaOH hydration process to be dominated by Ba⁺-OH⁻ electrostatic and polarization interactions.

ASSOCIATED CONTENT

S Supporting Information

Calculated harmonic vibrational frequencies for the normal modes of all of the relevant neutral and cationic $BaOH(H_2O)_2$ and $BaOH(H_2O)_3$ isomers. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

Corresponding Author

*J.C.F.: Tel.: +54 351 4334169/80. Fax: +54 351 4334188. E-mail: jferrero@fcq.unc.edu.ar. C.J.C.: Tel.: +54 221 4257291/7430. Fax: +54 221 4254642. E-mail: cobos@inifta.unlp.edu.ar.

Notes

The authors declare no competing financial interest.

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