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Infrared spectroscopy of  $[H_2O-X_n]^+$  (n = 1-3, $X = N_2$ ,  $CO_2$ , CO, and  $N_2O$ ) radical cation clusters: competition between hydrogen bond and hemibond formation of the water radical cation\*

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The water radical cation H<sub>2</sub>O<sup>+</sup> is an important intermediate in radiation chemistry and radiobiology, and its role in radical reactions has recently attracted much attention. However, knowledge of intermolecular interactions of  $H_2O^+$  remains very limited due to its high reactivity. We focus on structures of  $[H_2O-X]^+$ , formed by  $H_2O^+$  with a counter molecule X, as a model for intermediates in reactions of  $H_2O^+$ . Such structural information provides the basis for understanding reaction processes of H<sub>2</sub>O<sup>+</sup>. Two structural motifs for  $[H_2O-X]^+$  have been known: hydrogen bond and hemibond, which are expected to have very different reactivities from each other. Due to the high acidity of H<sub>2</sub>O<sup>+</sup>, the H-bonded form is mostly considered to be preferred. However, it has recently been reported that the hemibonded form is preferred in some cases. We perform infrared photodissociation spectroscopy and quantum chemical calculations on  $[H_2O-X_n]^+$  (n=1-3,  $X=N_2$ ,  $CO_2$ ,  $CO_3$ , and  $N_2O_3$ ) to determine their structural motifs. The competition between the hydrogen bond and hemibond formation is systematically examined based on the firm structure information. The competition is interpreted in terms of the proton affinity (PA) and the ionization potential (IP) of X. The rough ranges of PA and IP for the priority of the hemibond motif are determined. The impact of other factors on the competition is also discussed.

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#### 1. Introduction

Water ionization and subsequent processes are central issues in the aqueous environmental chemistry and biology of various radiation-related processes.1 Water (in aqueous solutions) ionized upon the interaction with ionizing radiation initially forms the water radical cation H<sub>2</sub>O<sup>+</sup>, which is immediately followed by the formation of the active chemical species, the hydrated electron e and OH radical. The OH radical is well known as an important intermediate of radiation-induced damage in hydrated environments due to its high reactivity. In recent years, however, the role of H<sub>2</sub>O<sup>+</sup> itself as well as the OH radical has attracted much attention in radical chemistry of water induced by ionizing radiation.<sup>2-13</sup> Direct reactions with H<sub>2</sub>O<sup>+</sup> can be important when water molecules are ionized in close proximity to other molecules, e.g., at the interface between water and biomolecules. Otherwise, H<sub>2</sub>O<sup>+</sup> should undergo ultrafast proton transfer with surrounding water molecules to form the OH

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radical, as described above. Indeed, in the highly concentrated solutions, 2-9 where water molecules are in contact with other molecules, and in microdroplets, 10-13 it has been indicated that H<sub>2</sub>O<sup>+</sup> plays a key role in chemical reactions. Despite such importance, H<sub>2</sub>O<sup>+</sup> is still challenging to study, especially for direct measurements in the liquid phase, because of its high reactivity. Although there have been some reports, 14-20 knowledge of H<sub>2</sub>O<sup>+</sup> remains scarce. Gas-phase cluster studies are a powerful approach to such ultrashort-lived radical cation species, as their reactions are well controlled under isolated conditions. In the present study, we focus on structures of  $[H_2O-X]^+$ formed by H<sub>2</sub>O<sup>+</sup> and a counter molecule X as a model for reaction intermediates of H2O+. Such structural information is expected to provide the basis for understanding reaction processes of H<sub>2</sub>O<sup>+</sup>.

Structures of the  $[H_2O-X]^+$  (X = He, Ne, Ar, Kr, CO<sub>2</sub>, N<sub>2</sub>O, and H<sub>2</sub>O) radical cation clusters have been studied by infrared (IR) spectroscopy in the gas phase.<sup>21–31</sup> Two structural motifs are confirmed: hydrogen bond (H-bond) and hemibond. The H-bonded type has a structure in which X is H-bonded to one of the OH groups of  $H_2O^+$  (HOH···X). On the other hand, the hemibonded structure (H2O···X) has a non-classical (two center-three electron, 2c-3e) covalent bond formed by overlapping lone pair orbitals of H2O+ and X. The hemibond has

<sup>†</sup> Electronic supplementary information (ESI) available: Comparison of the observed and simulated spectra of  $[H_2O-X_n]^+$  (n =1-3, X = N<sub>2</sub>, CO<sub>2</sub>, CO, and N<sub>2</sub>O). See DOI: https://doi.org/10.1039/d3cp00753g

Paper

attracted strong attention in radiation chemistry, mainly of sulfur-containing systems. 32-47 Recently, it has been reported

that OH radicals form hemibonds in the aqueous phase. 48,49 However, roles of hemibonds of H<sub>2</sub>O<sup>+</sup> in aqueous systems have been scarcely evidenced so far. The formation of these two possible motifs, H-bond and hemibond, should compete with each other, and the preferential motif is expected to depend on X.

The X =  $H_2O$  case, i.e., the  $(H_2O)_n^+$  cluster, has been actively studied as the simplest model for ionization of water. 21-23,50-89 Some early theoretical calculations showed that the hemibond formation is preferred. However, many recent high-level theoretical computations agree well that the hemibonded form is less stable than the H-bonded form. In fact, only the H-bonded form has been observed by IR spectroscopic experiments of  $(H_2O)_n^{+21-23}$  Therefore, it can be concluded that the H-bond formation is favored for  $X = H_2O$ .

As for other molecules X, structural studies have been performed on rare gas atoms and triatomic molecules. For X = He, Ne, and Ar, it was concluded that the H-bond formation is preferred.<sup>24-28</sup> Inokuchi et al. also concluded that the H-bond formation is preferred over the hemibond formation for  $X = CO_2$ . Thus,  $H_2O^+$  seems to favor H-bond formation because of its high acidity. Recently, however, it was found that the hemibond formation is favored over the H-bond formation in the cases of  $X = N_2O$  and  $Kr.^{30,31}$  Here, a question arises; what factors govern the competition between H-bond and hemibond formation.

Plausible factors that determine the competition between H-bond and hemibond formation are the proton affinity (PA) and the ionization potential (IP) of X. When the PA of X is high, the H-bond strength would increase, resulting in the preference for H-bond formation, while when the IP of X is close to H<sub>2</sub>O, the strong charge resonance interaction would lead to hemibond formation. 34-36 In fact, the IPs of N2O and Kr are close to H<sub>2</sub>O, which would be a favorable condition for the hemibond formation. Then, we have an alternative question; what are the ranges of IP and PA of X for the preferential hemibond formation in [H2O-X]+. This range may provide a guide to judge the realization of hemibond formation of H<sub>2</sub>O<sup>+</sup> in the condensed phase.

The structural motifs of [H<sub>2</sub>O-X]<sup>+</sup> have been studied individually for each molecule X so far. However, this competition between H-bond and hemibond formation has not been fully systematically examined and is still poorly understood. In this study, we perform IR predissociation spectroscopy and quantum chemical calculations of  $[H_2O-X_n]^+$   $(n = 1-3, X = N_2, CO_2, C$ CO, and N2O) to determine their structural motif. Here, as a counterpart molecule X, we choose molecules that have IPs relatively close to H2O and are expected to have high hemibond strength comparable to H-bond strength. We should also note that the acidy of water is highly enhanced with ionization, and its H-bond is expected to be much stronger than typical H-bonds among neutral molecules, even though these X molecules are ordinarily categorized into very weak proton acceptors. Based on the results obtained in this study and those of the previous studies, the correlation between the competing H-bond/hemibond formation and the PA/IP values is investigated to determine the conditions for the preference of hemibond formation of H<sub>2</sub>O<sup>+</sup>.

For IR spectroscopy of cluster ions in the gas phase, we ordinately employ the infrared predissociation (IRPD) spectroscopic technique, 90,91 in which predissociation fragments upon vibrational excitation of parent cluster ions are detected as a measure of IR absorption. This is because the concentration of cluster ions is too low to detect their IR absorption directly. Here, we note that for n = 1 of  $[H_2O-X_n]^+$ , an IR spectrum of the most stable isomer may not be obtained by IRPD spectroscopy because its dissociation energy is estimated to be much higher than the IR photon energy. Moreover, considering dissociation energies, it is expected that higher energy (less stable) isomers are rather observed. Therefore, in the present study, the Ar-"tagging" technique  $^{91-94}$  is employed for n = 1 to observe low-energy isomers. In this technique, an Ar atom is attached to the cluster of interest. The Ar atom is easily released from the cluster with the IR vibrational excitation, and this is detected as a measure of the IR absorption in the scheme of the dissociation spectroscopy. We can safely suppose that the attachment of Ar is inert enough not to perturb the structural motif of the "bare" cluster (the smaller interaction energy of Ar than the X molecules will be shown later). Moreover, the Ar-tagged cluster becomes cold because only clusters of low internal (vibrational) energy can hold the Ar atom until the interaction region of the spectrometer (irradiation of the IR light). The structural motifs for  $X = CO_2$  and  $N_2O$  have already been reported. <sup>29,30</sup> However, in these previous studies, there remains some ambiguity in the structure determination of n = 1. For  $X = CO_2$ , a high-energy isomer could be observed for n = 1, as mentioned above. For  $N_2O$ , the IR spectrum was not measured for n = 1, and the structural motif was determined based on theoretical computations and the analogy to the  $n \ge 2$  clusters. The present measurements of the Ar-tagged clusters of n = 1 provide firm and direct experimental evidence for the most stable structural motif of  $[H_2O-X]^+$ .

# 2. Experimental and computational methods

Details of the experimental apparatus have been described elsewhere.<sup>93</sup> In brief,  $[H_2O-X_n]^+$  (n = 1-3,  $X = N_2$ ,  $CO_2$ ,  $CO_3$ and  $N_2O$ ) and their Ar-tagged clusters (n = 1 only) were generated by electron ionization in a supersonic expansion. A gaseous mixture of water, X, and Ar or He carrier (~8 MPa) was expanded through a high-pressure pulsed valve (Even-Lavie valve<sup>95</sup>) into a vacuum chamber. An electron beam accelerated to 200 V from an electron gun (Omegatron Corporation) was used for electron ionization. The produced ions were introduced into a tandem quadrupole mass spectrometer. The cluster ions of interest were mass-selected at the first stage of the mass spectrometer. The mass resolution was set higher than  $\Delta m/z \leq 1$ , and the corresponding protonated cluster, H<sub>3</sub>O<sup>+</sup>-X<sub>n</sub>, was carefully removed. The mass-selected ions were then introduced into the

octopole ion guide and were irradiated by IR light. The IR light was the output of an IR optical parametric oscillator/amplifier (LaserVision) pumped by the fundamental output of a Nd:YAG laser (Spectra Physics GCR 230). Fragment ions produced by vibrational predissociation were mass-selected in the second stage of the mass spectrometer and were detected by a channel electron multiplier. An IR spectrum was measured by monitoring the photofragment intensity as a function of the IR frequency. The loss channel of a single X molecule was monitored for the bare clusters, and the loss channel of Ar for the Ar-tagged clusters.

Quantum chemical computations of  $[H_2O-X_n]^+$  were performed using the Gaussian 16 program package. 96 Energyoptimized structure search and harmonic vibrational simulations were performed at the MP2/aug-cc-pVTZ level for  $X = N_2$ ,  $CO_2$ , and CO. Only for  $X = N_2O$ , the B3LYP/aug-cc-pVTZ level was employed. This is because, for  $X = N_2O$ , the SCF calculations for the MP2 calculation did not converge. The calculated frequencies were scaled by a factor of 0.955 for the MP2 calculations and 0.958 for the B3LYP calculations. Only for the n = 1 clusters, we also performed second order vibrational perturbation theory (VPT2) calculations for anharmonic analyses of minor combination and overtone bands.

### 3. Results and discussion

#### 3.1. $[H_2O-(N_2)_n]^+$ radical cation clusters

Fig. 1 shows the observed and simulated IR spectra of [H<sub>2</sub>O-N<sub>2</sub>]<sup>+</sup>-Ar. As expected, the H-bonded and hemibonded isomers were obtained for the calculated stable structures.

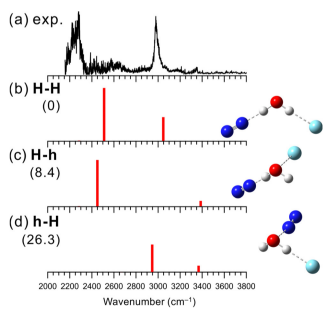


Fig. 1 Comparison of (a) observed IRPD spectrum of  $[H_2O-N_2]^+$ -Ar and (b-d) simulated spectra of the stable isomers calculated at MP2/aug-ccpVTZ. The simulated spectra were scaled by a factor of 0.955. The intensities of the simulated spectra of the isomers are plotted in the same scale. The schematic structures of the isomers are also shown. Numbers in parentheses are ZPE-corrected energies (in kJ mol<sup>-1</sup>).

Two labels H and h represent the H-bond and hemibond in the isomers, respectively. For [H<sub>2</sub>O-X]<sup>+</sup>-Ar, the first label corresponds to the bonding type formed by H2O+ and X, and the second label to that of Ar  $(e.g., in [H_2O-N_2]^+-Ar, isomer h-H has$ H<sub>2</sub>O<sup>+</sup> forming a hemibond with N<sub>2</sub> and a H-bond with Ar).

In the observed spectrum, strong bands were seen at 2990 and around 2250 cm<sup>-1</sup>, which are attributed to the Ar-bound and N<sub>2</sub>-bound OH stretching vibrations of H<sub>2</sub>O<sup>+</sup>, respectively. This indicates that in  $[H_2O-N_2]^+$ -Ar, both the OH groups of the H<sub>2</sub>O<sup>+</sup> moiety are H-bonded to the N<sub>2</sub> and Ar, respectively. Moreover, there are no remarkable bands in the 3300-3500 cm<sup>-1</sup> region, where free OH stretching bands of H<sub>2</sub>O<sup>+</sup> are expected. These spectral features mean that an isomer without free OH should be the major carrier of the spectrum. Therefore, it is concluded that H-bond formation is preferred over hemibond formation for  $X = N_2$ . Some minor bands are observed at  $\sim$  2600, 3160 and 3350 cm<sup>-1</sup>. The former two bands could be a combination bands of the N<sub>2</sub> (or Ar)-bound OH stretching and the intermolecular stretching vibration between H2O+ and H-bonded N<sub>2</sub> (or Ar), respectively. The latter could be attributed to contributions from other isomers (see below).

Three stable isomers are found in the theoretical calculations of [H<sub>2</sub>O-N<sub>2</sub>]<sup>+</sup>-Ar at the MP2/aug-cc-pVTZ level. Their schematic structures, relative energies at the zero-point energy (ZPE) level, and harmonic vibrational spectra are shown in Fig. 1(b)-(d). The most stable structure of  $[H_2O-N_2]^+$ -Ar is isomer H-H, in which both N2 and Ar are H-bonded to the OH groups of H<sub>2</sub>O<sup>+</sup>. Isomer **H-h** has a structure in which a water molecule is H-bonded to N2, and the oxygen atom of water and Ar form a hemibond, while h-H is the reversed one: N<sub>2</sub> forms a hemibond, and Ar forms a H-bond. These computational results are consistent with the experimental result. Isomer H-H, in which both OH groups are H-bonded, is calculated to be the most stable, and its simulated spectrum well reproduces the major features of the observed spectrum. In addition, as shown in Fig. S1 in the ESI,† the VPT2 anharmonic vibrational calculations for isomer H-H reproduce the two minor bands, the combination bands at  $\sim 2600$  and 3160 cm<sup>-1</sup>. Therefore, isomer **H-H** is assigned to the major carrier of the observed spectrum. This means that N2 favors the H-bond formation with H<sub>2</sub>O<sup>+</sup> over the hemibond formation. This is also consistent with the MP2/aug-cc-pVTZ level computations of the "bare" [H<sub>2</sub>O-N<sub>2</sub>]<sup>+</sup> cluster shown Fig. S2 in ESI;† the H-bonded (H) isomer is calculated to be more stable  $(-32.1 \text{ kJ mol}^{-1})$  than the hemibonded (**h**) isomer.

Here, we discuss the contributions of the other isomers in the observed IR spectrum. The weak band observed at  $3350 \text{ cm}^{-1}$ can be attributed to the free OH stretching vibrational band of higher energy isomers H-h or h-H, of which the population would be much less than isomer H-H. It is difficult at the present stage to clearly identify the spectral carrier of this band. The calculated relative energies show that isomer H-h, where Ar forms a hemibond, is favored. However, the contribution of isomer h-H cannot be ruled out since we have obtained an IR spectrum attributed to the N<sub>2</sub>-hemibonded (h) form in the measurement of the "bare" [H<sub>2</sub>O-N<sub>2</sub>]<sup>+</sup> cluster, in which higher energy isomers tend to appear

because predissociation following IR excitation can occur only with the assistance of internal energy (see Fig. S2 in ESI† for

Similarly, IRPD spectroscopy and quantum chemical calculations were performed for n = 2 and 3 of  $[H_2O-(N_2)_n]^+$ . The results are summarized in Fig. S3 and S4 in ESI,† respectively. For n = 2, a strong band attributed to the N<sub>2</sub>-bound OH stretching vibrations appears, while no obvious bands corresponding to free OH stretching vibrations are seen, as in the case of [H2O-N2]+-Ar. In addition, isomer H-H is the most stable one in the theoretical calculations. Therefore, both the N<sub>2</sub> molecules are preferentially bonded to the OH groups of  $H_2O^+$  for n=2. This also indicates again that the H-bond formation is favored for  $X = N_2$  over the hemibond formation. For n = 3, an IR spectrum similar to that of n = 2 was obtained. Based on the experimental and calculational results, the spectrum was attributed to the structure in which two N2 molecules are H-bonded to the OH groups and the third one is hemibonded to the O atom of  $H_2O^+$ .

#### 3.2. $[H_2O-(CO_2)_n]^+$ radical cation clusters

The IR photodissociation spectra of bare  $[H_2O-(CO_2)_n]^+$  (n =1-7) have been reported by Inokuchi et al., and they have concluded that the H-bond formation is favored over the hemibond formation.<sup>29</sup> In n = 1, however, because of its large dissociation energy, the dissociation efficiency decreases as the excitation energy decreases. As a result, a remarkable disagreement of the vibrational band intensities between the observed and calculated spectra was seen, which leaves the ambiguity of the spectral and structural assignment. In the present study, we measured an IR spectrum of n = 1 using Ar-tagging for clearer assignments.

Fig. 2 shows the IR spectrum obtained in the present experiment of [H<sub>2</sub>O-CO<sub>2</sub>]<sup>+</sup>-Ar and the calculated vibrational spectra of the stable isomers. In the observed spectrum, a strong band is seen at 3070 cm<sup>-1</sup>, which is attributed to the Ar-bound OH stretching vibration. The bands at 2390, 3560, and 3740 cm<sup>-1</sup> are attributed to the antisymmetric CO stretching vibration  $(v_3)$  and the  $v_1 + v_3$  and  $2v_2 + v_3$  combination bands of CO<sub>2</sub>, respectively, as proposed in the previous studies.<sup>29,97</sup> Here,  $v_1$  and  $v_2$  represent the symmetric stretching and bending vibrations of CO2, respectively. The other weak peaks around 3230 and 3370 cm<sup>-1</sup> are attributed to a combination band between Ar-bound OH stretching and intermolecular stretching vibrations and to contributions of higher energy isomers, respectively, as in the case of  $[H_2O-N_2]^+$ -Ar shown in the previous subsection. It is noted that no remarkable free OH band is seen in the observed spectrum of [H<sub>2</sub>O-CO<sub>2</sub>]<sup>+</sup>-Ar. Therefore, we also conclude that the H-bond formation is preferred over the hemibond formation for  $X = CO_2$ , as Inokuchi et al. have shown. The band of the CO2-bound OH stretching vibration is not clearly seen in the observed spectrum; however, the rising of the signal below 2300 cm<sup>-1</sup> would correspond to the tail of the CO2-bound OH stretching band. CO<sub>2</sub> has a larger proton affinity than N<sub>2</sub> (541 and 494 kJ mol<sup>-1</sup>, respectively<sup>98</sup>). Therefore, the CO<sub>2</sub>-bound OH stretching

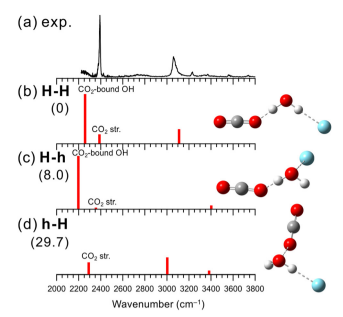


Fig. 2 Comparison of (a) observed IRPD spectrum of [H<sub>2</sub>O-CO<sub>2</sub>]<sup>+</sup>-Ar and (b-d) simulated spectra of the stable isomers calculated at MP2/augcc-pVTZ. The simulated spectra were scaled by a factor of 0.955. The intensities of the simulated spectra of the isomers are plotted in the same scale. The schematic structures of the isomers are also shown. Numbers in parentheses are ZPE-corrected energies (in kJ mol<sup>-1</sup>).

vibration of [H<sub>2</sub>O-CO<sub>2</sub>]<sup>+</sup>-Ar is expected to show a lower frequency than that of the N<sub>2</sub>-bound OH stretching vibration ( $\sim 2250 \text{ cm}^{-1}$ ) of [H<sub>2</sub>O-N<sub>2</sub>]<sup>+</sup>-Ar, and the band would be shifted to out of the measured frequency range. Inokuchi et al. have attributed a weak and broad band appearing at  $\sim 2600 \text{ cm}^{-1}$  to CO<sub>2</sub>-bound OH, which was not obtained in the present measurement. Therefore, the present Ar-tagged measurement provides the firm structural and spectral assignments of [H<sub>2</sub>O-CO<sub>2</sub>]<sup>+</sup>.

The structures, energies, and simulated spectra of three isomers of [H2O-CO2]+-Ar obtained by the MP2 calculations are shown in Fig. 2(b)-(d). The obtained stable structures have the same structural motifs as for  $X = N_2$ . The most stable structure is isomer H-H. For CO2, forming a H-bond with H<sub>2</sub>O<sup>+</sup> is 29.7 kJ mol<sup>-1</sup> more stable than forming a hemibond (isomer h-H). Considering the missing of the largely red-shifted CO2-bound OH stretching vibration, all the major features of the observed spectrum are consistent well with the calculated spectrum of isomer H-H. The anharmonic calculation results shown in Fig. S5 (ESI†) also demonstrate that the most stable structure, isomer H-H, well reproduces the features of the observed spectrum.

Finally, we note that the weak band near 3370  $\text{cm}^{-1}$  in the observed IR spectrum of [H2O-CO2]+-Ar can be attributed to the minor population of the Ar-hemibonded isomer, H-h, or the CO<sub>2</sub>-hemibonded isomer, h-H. Isomer H-h is energetically more stable than **h-H**, while the spectrum of bare  $[H_2O-CO_2]^+$ measured in the present study suggested the hemibonded (h) isomer formation with CO<sub>2</sub> (see Fig. S6 in ESI† for details), as in the case of  $[H_2O-N_2]^+$ . Therefore, it is difficult to identify which isomer is the spectral carrier of this minor band.

We also re-measured the IR spectra of n = 2 and 3 and performed their theoretical computations. The results are summarized in Fig. S7 and S8 in ESI,† and they provide the same conclusion (the superiority of the H-bond formation over the hemibond formation between H<sub>2</sub>O<sup>+</sup> and CO<sub>2</sub>) as reported in the previous study by Inokuchi et al.29

#### 3.3. $[H_2O-(CO)_n]^+$ radical cation clusters

Fig. 3 shows the observed IR spectrum of  $[H_2O-CO]^+$ -Ar and the simulated vibrational spectra of the stable isomers. The peaks at 3180 and 3300 cm<sup>-1</sup> in the observed spectrum can be attributed to the Ar-bound OH stretching vibration and its combination band with the intermolecular stretching vibration, respectively. In addition, the shoulder around 3070 cm<sup>-1</sup> could be a bending overtone of  $H_2O^+$ . Unlike the cases of  $X = N_2$ and CO<sub>2</sub> mentioned above, a strong band is seen at 3470 cm<sup>-1</sup>, which should be attributed to a free OH stretching vibration. The presence of the free OH stretching vibrational band indicates a preference for a different structural motif from the H-bonded form for X = CO.

The structures, energies, and spectra of the four stable isomers of [H<sub>2</sub>O-CO]<sup>+</sup>-Ar obtained by the MP2 calculations are shown in Fig. 3(b)-(e). The most stable structural motif is

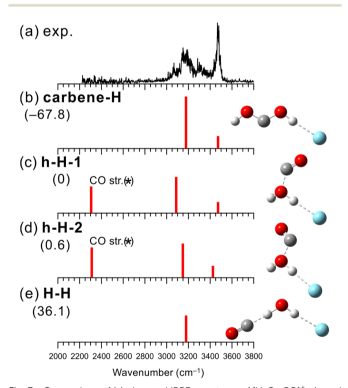


Fig. 3 Comparison of (a) observed IRPD spectrum of [H<sub>2</sub>O-CO]<sup>+</sup>-Ar and (b-e) simulated spectra of the stable isomers calculated at MP2/aug-ccpVTZ. The simulated spectra were scaled by a factor of 0.955. The intensities of the simulated spectra of the isomers are plotted in the same scale. The schematic structures of the isomers are also shown. Numbers in parentheses are ZPE-corrected energies (in kJ mol<sup>-1</sup>). In calculated spectra (c) and (d), the frequency of the CO stretching vibration (asterisked band) could be overestimated with the MP2 calculation (see text for details.)

the carbene-type isomer (carbene-H), which is formed by the rearrangement of the covalent bonds between the two molecules. The hemibonded isomers (h-H-1 and h-H-2) are more stable than the H-bonded isomer (H-H) by 36 kJ mol<sup>-1</sup> while 68 kJ mol<sup>-1</sup> more unstable than the carbene-type isomer. It would be straightforward to assume that the carbene-type isomer is dominantly generated. However, the previous computational studies showed that the isomerization barrier between the hemibonded and carbene-type isomers is calculated to be quite high ( $\sim 170 \text{ kJ mol}^{-1}$ ), <sup>99,100</sup> and the isomerization of all the produced clusters seems unlikely to occur.

When we compare the observed and simulated spectra in Fig. 3, we note that in the calculated spectra of h-H-1 and h-H-2, a band corresponding to the CO stretching vibration is predicted at ~2300 cm<sup>-1</sup>, though the experimental spectrum shows no band in this region. However, the stretching vibration of bare neutral CO is 2147 cm<sup>-1</sup>, 101 suggesting that the MP2 calculation overestimates the CO stretching frequency. Indeed, the harmonic vibrational analysis at the MP2/aug-cc-pVTZ level of CO and CO+ calculated their frequencies to be 2015 and 2745 cm<sup>-1</sup>, respectively, while the experimental values were 2147 and 2188 cm<sup>-1</sup>, respectively. 101 For CO<sup>+</sup>, the extremely large overestimation occurs. The CO site is partially charged in the hemibonded isomer, and this would lead to an overestimation of the frequency of the CO site. The CO stretching vibration of the hemibonded isomers is expected out of the measured spectral range. Therefore, the main features of the observed spectra are in good agreement with both the calculated spectra of the carbene-type and hemibonded isomers. However, the shoulder band around 3070 cm<sup>-1</sup>, which is not predicted in both the harmonic vibrational simulations, could be attributed to a bending overtone of H<sub>2</sub>O<sup>+</sup>. Since isomer carbene-H has no H<sub>2</sub>O subunit, this could be a marker band for the presence of the hemibonded isomer. The results of the anharmonic vibrational calculations are shown in Fig. S9 in ESI.† For isomer carbene-H, only the Ar-bound OH stretching band is predicted in the 3000-3200 cm<sup>-1</sup> region, while for the hemibonded isomers, a water bending overtone is calculated in this region in addition to the Ar-bound OH stretching band (the Fermi resonance occurs). Therefore, the calculated anharmonic spectra of the hemibonded isomers (especially h-H-2) well reproduce the observed spectrum with its shoulder band around 3070 cm<sup>-1</sup>. To summarize for [H<sub>2</sub>O-CO]<sup>+</sup>-Ar, the contribution of the most stable structure, the carbene-type isomer, to the observed spectrum is highly plausible. However, because the isomerization barrier to the carbene form is quite high, some contribution of the hemibonded isomers could occur, as evidenced by the shoulder band around 3070 cm<sup>-1</sup>. Therefore, it is concluded that the coexistence of the carbene-type and hemibonded isomers are plausible. Finally, we emphasize that the experimental spectral features clearly exclude the dominance of the H-bonded isomer (H-H). In addition, the H-bonded isomer is calculated to be much less stable than the hemibonded isomers. In other words, both the observed spectral features and the calculated energies indicate that hemibond formation is preferred over H-bond formation for X = CO.

Experimental measurements and theoretical calculations were also performed for n = 2-4. The results are summarized in Fig. S10–S12 in ESI.† For  $n \ge 2$ , it has been calculated that CO acts as a catalyst and significantly reduces the isomerization barrier between the hemibonded and carbene-type isomers  $(\sim 0.8 \text{ kJ mol}^{-1})$ . Therefore, it is likely that the carbenetype isomer is predominantly formed for  $n \ge 2$  since the carbene form is energetically much more stable than the hemibonded form.

#### 3.4. $[H_2O-(N_2O)_n]^+$ radical cation clusters

Structures of  $[H_2O-(N_2O)_n]^+$  have been investigated by collision induced dissociation (CID), photodissociation, and IR spectroscopic experiments. 30,102-105 Matsushima et al. have measured the IR predissociation spectra of  $[H_2O-(N_2O)_n]^+$  at n = 2-7 and have performed their theoretical computations.<sup>30</sup> They have shown that N<sub>2</sub>O favors the hemibond formation with H<sub>2</sub>O<sup>+</sup> over the H-bond formation. However, the experimental spectrum of n = 1 was not obtained because of its large dissociation energy, and the structure of [H<sub>2</sub>O-N<sub>2</sub>O]<sup>+</sup> was inferred by the theoretical computations and analogy with the larger clusters  $(n \ge 2)$ . Therefore, in the present study, we measured an IR spectrum of [H<sub>2</sub>O-N<sub>2</sub>O]<sup>+</sup> by using Ar-tagging to determine the binding motif unequivocally.

Fig. 4 shows the observed IR spectrum of  $[H_2O-N_2O]^+$ -Ar and the simulated vibrational spectra of its stable isomers. The observed spectrum has similar spectral features to those of  $[H_2O-CO]^+$ -Ar. The peaks at 3240, 3390, and 3460 cm<sup>-1</sup> in the observed spectrum can be assigned to an Ar-bound OH stretching band, its combination band with intermolecular stretching,

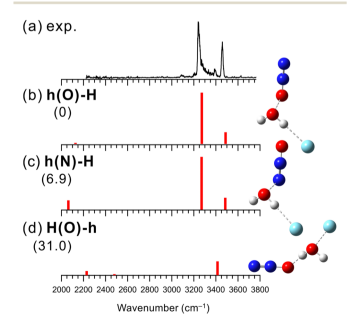


Fig. 4 Comparison of (a) observed IRPD spectrum of  $[H_2O-N_2O]^+$ -Ar and (b-d) simulated spectra of the stable isomers calculated at B3LYP/ aug-cc-pVTZ. The simulated spectra were scaled by a factor of 0.958. The intensities of the simulated spectra of the isomers are plotted in the same scale. The schematic structures of the isomers are also shown. Numbers in parentheses are ZPE-corrected energies (in kJ mol<sup>-1</sup>).

and a free OH stretching band, respectively. In addition, the very weak peak at 3090 cm<sup>-1</sup> could be a bending overtone of H<sub>2</sub>O<sup>+</sup>. The presence of the free OH stretching band indicates that the hemibond formation is preferred over the H-bond formation for  $X = N_2O$ .

Fig. 4(b)-(d) show the three lowest energy stable isomer structures, relative energies, and vibrational spectra of [H<sub>2</sub>O-N<sub>2</sub>O]<sup>+</sup>-Ar. Here, the B3LYP/aug-cc-pVTZ level of theory was used for the calculations. This is because the SCF calculations for the MP2 calculations did not converge in this system, and the alternative choice of B3LYP/aug-cc-pVTZ follows the theory employed in the previous study by Matsushima et al.30 The calculated relative energies show that hemibonded isomer h(O)-H is more stable than H-bonded isomer H(O)-h by 31 kJ mol<sup>-1</sup>, which supports the present experimental result. Also, this preference for the hemibond over the H-bond is consistent with the previous theoretical results.30,106 Since isomers h(O)-H and h(N)-H exhibit the calculated IR spectra similar to each other, it is difficult to distinguish the contribution of each isomer to the observed spectrum. However, h(O)-H is more stable than h(N)-H by 6.9 kJ mol<sup>-1</sup> and could be predominantly present. As shown in Fig. S13 in ESI,† the anharmonic vibrational calculations for the hemibonded isomers reproduce the observed spectrum including the minor bands, combination band between the Ar-bound OH stretching and the intermolecular stretching vibration and bending overtone of  $H_2O^+$ . We can conclude that  $X = N_2O$  favors the hemibond formation in [H<sub>2</sub>O-N<sub>2</sub>O]<sup>+</sup>-Ar. This conclusion is consistent with the previous study of bare  $[H_2O-(N_2O)_n]^+$ .  $^{30,106}$ However, the present study provided firm experimental evidence of the hemibond formation in  $[H_2O-N_2O]^+$ .

We also measured the IR spectra of the bare clusters of n = 1-3 and performed their theoretical computations. The results are summarized in Fig. S14-S16 in ESI.† In the present study, we could observe the spectrum of bare n = 1. However, its fragment signal intensity was very weak, and this suggests that we might observe very hot components, in which internal energy assists photodissociation. For n = 2 and 3, we have essentially the same results and conclusions as those in the previous study by Matsushima et al.30

#### 3.5. Correlation between the competing H-bond/hemibond formation and the PA/IP

The results obtained in this study (Sections 3.1-3.4) and the previous studies21-31 reveal the preferred binding motif of  $[H_2O-X]^+$  for the various X molecules (X = He, Ne, Ar, Kr, N<sub>2</sub>, CO<sub>2</sub>, CO, N<sub>2</sub>O, and H<sub>2</sub>O). Fig. 5 shows the PA-IP plot of the various X molecules, 98,107 color-coded with the preferred binding motifs (hemibond or H-bond) of  $[H_2O-X]^+$ . In this plot, points of X favoring the H-bond formation are colored in blue, and those favoring the hemibond formation are in red. This plot shows the ranges of PA and IP of X for the preference of the hemibond formation with H<sub>2</sub>O<sup>+</sup>. The preference for hemibond formation is limited only to X of which IP is in the narrow range within  $\sim$  1.5 eV from that of H<sub>2</sub>O (12.6 eV). This trend is consistent with the previous theoretical studies<sup>34-36</sup> that the hemibond strength

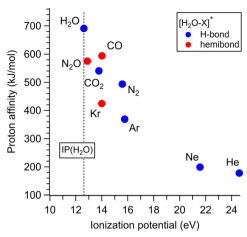


Fig. 5 PA-IP plot of the various X molecules color-coded with the preferred binding motifs of [H<sub>2</sub>O-X]<sup>+</sup>. Points of X favoring the H-bond formation are colored in blue and those favoring the hemibond formation are in red. This plot is based on the preferential structure determinations in the present and previous studies. <sup>21–31</sup> The dotted line shows the IP of H<sub>2</sub>O.

decreases exponentially with increases of  $\Delta$ IP, the difference in the ionization potentials of the two units. When X is H<sub>2</sub>O  $([H_2O-H_2O]^+)$ ,  $\Delta IP$  is 0, and therefore, the strongest hemibond formation is expected. However, it has been known that the H-bond formation is actually favored in  $[H_2O-H_2O]^+$ . This means that the PA of water (691 kJ mol<sup>-1</sup>) is high enough to form the H-bond stronger than the hemibond. Therefore, based on the plot in Fig. 5, we conclude that the hemibond formation in  $[H_2O-X]^+$  can be superior to the H-bond formation only when the IP of X is within  $\sim 1.5$  eV from that of water (12.6 eV) and the PA is lower than that of water (691 kJ mol<sup>-1</sup>).

However, the PA-IP plot does not fully explain the competition between the hemibond and H-bond formation. The point of  $X = CO_2$  seems exceptional; though one would expect the preference of the hemibond formation for CO2 because its IP is closer to H2O than CO and its PA is also lower than CO, [H<sub>2</sub>O-CO<sub>2</sub>]<sup>+</sup> actually favors the H-bond formation. Therefore, further consideration of the competition is required.

To investigate this correlation between the competing bond formations and PA/IP in more detail, the correlation between the H-bond strength and PA of X and that between the hemibond strength and IP was evaluated using binding energy  $D_0$ , respectively. In the following, all computations were performed at the CCSD/aug-cc-pVTZ level. This is because we frequently had failure of MP2 calculations for the homodimer cations, which are requested in the later discussion on hemibonds. Note that no stable hemibonded structure was obtained for X = Ne and He (only a hemibond-like structure was obtained; however, the charge is almost localized on H<sub>2</sub>O<sup>+</sup>, suggesting that its hemibond nature is weak.), and the SCF calculation did not converge for  $X = N_2O$ . The calculation results are shown in Fig. 6. Fig. 6 (top) shows the correlation between the binding energy of the H-bonded isomer and the PA value of X. The binding energy increases monotonically with increasing PA. This indicates that, as expected, the H-bond strength becomes

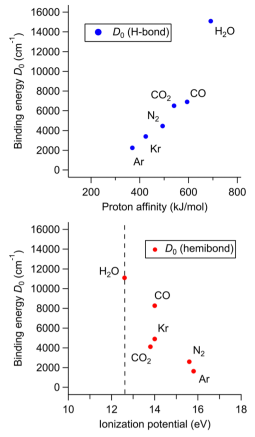


Fig. 6 (top) Correlation between  $D_0$  of the H-bonded isomers of  $[H_2O-X]^+$  and the PA of X. (bottom) Correlation between  $D_0$  of the hemibonded isomers and the IP of X. In both the plots, the  $D_0$  values were calculated at CCSD/aug-cc-pVTZ.

higher as the PA of X increases. Fig. 6 (bottom) shows the correlation between the binding energy of the hemibonded isomer and the IP value of X. The binding energy tends to increase as the IP approaches that of H<sub>2</sub>O. However, the point of  $X = CO_2$  seems exceptional; the binding energy of  $X = CO_2$  is lower than those of X = Kr and CO, even though the IP of CO<sub>2</sub> is lower than those of Kr and CO and closer to that of H2O. Therefore, the hemibond strength cannot be simply correlated only to IP (or  $\Delta$ IP).

According to the previous theoretical studies, 34-36 the binding energy  $D_{AB}$  of a hemibond is given by:

$$D_{\rm AB} \cong (D_{\rm AA} + D_{\rm BB})/2 \times \exp(-\Delta {\rm IP}/2\sqrt{D_{\rm AA}D_{\rm BB}})$$
 (1)

where  $D_{AA/BB}$  is the binding energy of the hemibonded homodimer cation and  $\Delta$ IP is the difference in IP between molecules A and B. Therefore, not only  $\Delta$ IP but also the binding energy of the homodimer cation is essential for the hemibond strength. Thus, we calculated the binding energy  $D_{AA}$  of the homodimer cation for each X. The results are shown in Table 1. We found that  $X = CO_2$  exhibits a much smaller binding energy  $D_{AA}$ (4198 cm<sup>-1</sup>) than others. This small binding energy of the homodimer cation would be reflected in the low hemibond strength of [H<sub>2</sub>O-CO<sub>2</sub>]<sup>+</sup>. To confirm this interpretation, the binding

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**Table 1** Binding energies of the hemibonded structure of  $[H_2O-X]^+$  and homodimer cation  $X_2^+$  for various molecules X.  $D_{AA}$  and  $D_0$  represent the binding energies calculated at the CCSD/aug-cc-pVTZ level for  $X_2^+$  and hemibonded  $[H_2O-X]^+$ , respectively.  $D_{AB}$  of  $[H_2O-X]^+$  was obtained by substituting  $D_{AA}$  and IP values for each X into eqn (1)

X	IP (eV)	$D_{\mathrm{AA}} \left( \mathrm{cm}^{-1} \right)$	$D_{\mathrm{AB}} \left( \mathrm{cm}^{-1} \right)$	$D_0 \left( \mathrm{cm}^{-1} \right)$
Ar	15.8	9703	3071	1626
$N_2$	15.6	9725	3300	2589
Kr	14.0	9769	6118	4904
CO	14.0	16 942	9308	8264
$CO_2$	13.8	4198	3873	4116
$H_2O$	12.6	11 100	11 100	11 100

energy  $D_{AB}$  of  $[H_2O-X]^+$  was evaluated by substituting the calculated binding energies  $D_{AA}/D_{BB}$  of the homodimer cations into eqn (1). The results and the hemibond binding energy  $D_0$ obtained from the CCSD calculation are also shown in Table 1. The evaluated binding energy  $D_{AB}$  shows a good correlation to  $D_0$ , as demonstrated for the various systems by Clark. <sup>36</sup>  $D_{AB}$  well reproduces the exceptional behavior of X = CO<sub>2</sub>, its particularly smaller value. This indicates that the small  $D_{AA}$  of  $(CO_2)_2^+$ contributes to the weak hemibond strength of [H<sub>2</sub>O-CO<sub>2</sub>]<sup>+</sup>. This is the reason why the H-bond formation overcomes the hemibond formation for  $X = CO_2$ . Thus, the competition between hemibond and H-bond formation in  $[H_2O-X]^+$  is basically governed by the IP and PA values of X. However, the binding energy of the hemibonded X<sub>2</sub><sup>+</sup> homodimer cation, which may reflect the spatial overlap of the lone pair electron orbitals, is also required to consider in some cases.

## 4. Conclusions

photodissociation spectroscopy and quantum chemical calculations of  $[H_2O-X_n]^+$  (n = 1-3,  $X = N_2$ ,  $CO_2$ ,  $CO_3$ and N<sub>2</sub>O) were performed to systematically investigate the competition between the hydrogen (H-) bond and hemibond formation of H<sub>2</sub>O<sup>+</sup> and its correlation with the proton affinity (PA) and ionization potential (IP) values of counter molecule X in the intermolecular bond formation. We found that the H-bond formation is dominant for  $X = N_2$  and  $CO_2$ , while the hemibond formation is preferred for X = CO and  $N_2O$ . The correlation of the preferred binding motif (hemibond or H-bond) with the PA/IP values was examined on the basis of the most stable isomer structures of [H<sub>2</sub>O-X]<sup>+</sup> obtained in this study and the previous studies. It was concluded that this competition was basically explained by the magnitudes of the PA and IP, which essentially reflect the magnitudes of the H-bond and hemibond, respectively. We roughly estimated the ranges of the PA and IP of X for the preferential hemibond formation with  $H_2O^+$ ; the hemibond formation in  $[H_2O-X]^+$  can overcome the competitive H-bond formation only when the IP of X is within 1.5 eV from that of water and the PA is lower than that of water. Therefore, when water is ionized in close proximity to molecules whose IP is within 1.5 eV from that of water, not only H-bond but also hemibond formation with H<sub>2</sub>O<sup>+</sup> can be considered. As has been pointed out theoretically, however,

the magnitude of the hemibond between  $H_2O^{\dagger}$  and X would also depend on some other properties of X, such as orbital overlap. Thus, some exceptional H-bond preference can occur even in the PA/IP ranges for the hemibond preference. The present results are hoped to advance understanding of the detailed processes of ionizing radiation reactions in aqueous solutions.

### Author contributions

M.K.: investigation, calculations, data analysis, and writing. A.F.: conceptualization, supervision, funding acquisition, and writing – review and editing.

### Conflicts of interest

There are no conflicts to declare.

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