Cage effects on conformational preference and photophysics in the host–guest complex of benzenediols with 18-Crown-6†

Fumiya Morishima, Ryoji Kusaka, Yoshiya Inokuchi, Takeharu Haino and Takayuki Ebata*

The conformational preference and modification of photophysics of benzenediols, namely hydroquinone (HQ), resorcinol (RE) and catechol (CA), upon host–guest complex formation with 18-Crown-6 (18C6) have been investigated, under supersonically jet-cooled conditions. Laser induced fluorescence (LIF) and UV–UV hole-burning spectra indicate the presence of two conformers for HQ and RE and one conformer for CA. On the other hand, the number of isomers is reduced to one in the 18C6-HQ and 18C6-RE complexes, while the 18C6-CA complex has three stable isomers. The IR spectra of the OH stretching vibration reveal that the two OH groups are H-bonded in 18C6-CA and 18C6-RE. In 18C6-RE, RE adopts the highest energy conformation in the bare form. In 18C6-HQ, the H-bonding of one OH group affects the orientation of the other OH group. The complex formation changes the photophysics of the S1 state of the benzenediols in a different manner. In our previous work, we reported a remarkable S1 lifetime elongation in 18C6-CA complexes; the S1 lifetime of CA is elongated more than 1000 times longer (8 ps → 10.3 ns) in 18C6-CA (F. Morishima et al., J. Phys. Chem. B, 2015, 119, 2557–2565), which we called the “cage effect”. In 18C6-RE, the increase of S1 lifetime is moderate: 4.0 ns (monomer) → 10.5 ns (complex). On the other hand, the S1 lifetime of HQ is shortened in 18C6-HQ: 2.6 ns (monomer) → 0.54 ns (complex). Density functional theory (DFT) calculations suggest that these behaviors are related to the S1 (\(\pi\pi^*\))–\(\pi\sigma^*\) energy gap, the character of the S2 state and the symmetry of benzenediol. These experimental results clearly show the potential ability of 18C6 to control the conformation and modification of the electronic structure of guest species.

1. Introduction

Crown ethers (CEs) have been widely known as functional molecules, especially as inclusion compounds in host-guest chemistry. CEs can hold cations in their cavity and are soluble in organic solvents as well as protic solvents. Owing to these characteristics, they are used as phase transfer catalysts. It is also known that CEs exhibit a preference to encapsulate cations in bulk, depending on the size matching between the cation and the CE cavity. Applications of CEs as molecular receptors, metal cation extraction agents, fluoroionophores and phase transfer catalytic media have been described in a number of studies.1–24 CEs can include not only metal ions but also various ionic and neutral species through non-covalent interactions.25–31

In our previous study, we investigated the structure of a cold 3\(n\)-Crown-\(n\)-phenol neutral complex (\(n = 5–8\))29 and found that 18-crown-6 (18C6) and phenol form a single substantially stable 18C6-phenol 1:1 host–guest complex. 18C6 modifies its shape to be nicely matched to phenol showing “induced fit”. The structure is strongly stabilized by collective intermolecular interactions, such as OH · · · O H-bonding and CH · · · \(\pi\) bonding. Furthermore, we recently studied the 18C6-catechol (CA) host–guest complex and found that 18C6 modifies not only the geometrical but also the excited state electronic structures of the guest species,31 that is the S1 (\(\pi\pi^*\))–\(\pi\sigma^*\) energy gap, the character of the S2 state and the symmetry of benzenediol. These experimental results clearly show the potential ability of 18C6 to control the conformation and modification of the electronic structure of guest species.
conformations of benzenediols and 18C6 are modified for forming the stable host–guest complex through “induced-fitting”. We then investigate how the complex formation changes the electronic structure of benzenediols by measuring their S₁ lifetime. We apply various laser spectroscopic methods for the supersonically cooled molecules and complexes to measure the electronic and IR spectra. The number of conformers and isomers is determined by UV–UV hole-burning spectroscopy, and the structure of the specific isomer is investigated by IR–UV double resonance spectroscopy and quantum chemical calculations at density functional theory. The S₁ state lifetime is obtained by deconvoluting the fluorescence decay profiles or the picosecond pump–probe time profiles. We will discuss how the induced-fit model can be applied for forming the most stable complex structures and how the complex formation affects the excited state electronic structure of guest benzenediols.

2. Experimental and computational

2.1 Experimental

Details of the experimental setup have been described elsewhere. The supersonic free jet of benzenediol and 18C6-benzenediol complexes was generated by the supersonic expansion of a gaseous mixture of 18C6 and benzenediols diluted in He carrier gas (4 bar). The solid 18C6 and benzenediol samples were separately heated at different sample housings (340–370 K for 18C6, and 348, 363 and 473 K for CA, RE and HQ respectively) to be vaporized. The gas mixture was expanded into a vacuum chamber through a 1 mm orifice of a pulsed nozzle. The S₁–S₀ electronic spectra were observed by laser induced fluorescence (LIF) spectroscopy. A tunable UV light obtained by frequency doubling the output of Nd<sup>3+</sup>:YAG laser pumped dye laser (Lambda Physik Scanmate/Continuum Surelite II) was introduced into the vacuum chamber to be crossed with the supersonic jet at ~30 mm downstream (x/D ≈ 30) of the orifice. The LIF spectra were obtained by detecting the total fluorescence as a function of UV frequency. The vibronic bands belonging to different isomers were discriminated by UV–UV hole-burning (HB) spectroscopy; the frequency of the probe UV laser was fixed to a certain vibronic band and the IR laser frequency was scanned. The depletion of the fluorescence induced by the IR pump laser was detected, giving fluorescence-dip IR spectra for the UV monitored species.

The S₁ lifetimes of 18C6-CA and 18C6-RE complexes were obtained by deconvoluting the time profiles of the fluorescence decay curves by assuming the laser pulse shape as a Gaussian function with a 4.5 ns pulse width. The S₁ lifetimes of benzenediol monomers and H₂O-HQ and 18C6-HQ complexes were too short to be determined from the fluorescence decay, so that they were obtained by pump–probe experiments using a picosecond laser system. In this experiment, the molecules or complexes in the molecular beam were ionized by stepwise two-photon ionization with two picosecond lasers. The setup of the picosecond laser system has also been described in detail elsewhere. The first picosecond pump pulse excites the molecules or complexes to the S₁ origin, and the second probe laser pulse ionizes the S₁ state species. The ions are mass-analyzed with a 50 cm time-of-flight tube and are detected using a channeltron (Burle 4900). The ion signals were processed by using a boxcar integrator (Par model 4401/4420) connected with a PC. By changing the delay time between the pump and probe lasers, the decay time profile of the S₁ state was obtained. The decay time constants were determined by deconvoluting the time profiles with the 12 ps laser pulses. All the decay curves were fitted as a single exponential decay. 18C6 and all benzenediols were purchased from Sigma-Aldrich and Nacalai Tesque, respectively, and used without further purification.

2.2 Computational

The initial structures of 18C6-benzenediol complexes were obtained by Monte Carlo simulation by mixed torsional search with low-mode sampling in MacroModel V.9.1 with the MMFF94s force field. The geometries were optimized by the PRCG algorithm with a convergence threshold of 0.05 kJ mol<sup>−1</sup>. From this calculation, 165 isomers for 18C6-HQ, 46 isomers for 18C6-RE and 193 isomers for 18C6-CA were obtained, within 20 kJ mol<sup>−1</sup> of the most stable one. All of these isomers were geometry-optimized by density functional theory (DFT) calculations at the M05-2X/6-31+G* level with loose optimization criteria. This calculation yielded 5, 19, and 61 isomers for 18C6-HQ, 18C6-RE and 18C6-CA, respectively, within 20 kJ mol<sup>−1</sup>. These loose-optimized isomers were further optimized at the oB97X-D/6-31++G** level with tight optimization criteria and ultrafine grid. The total energies were corrected by nonscaled zero-point vibrational energy (ZPE). The interaction energy (E<sub>int</sub>) of 18C6-benzenediol complexes was computed with counterpoise correction to remove the basis set superposition error (BSSE),

\[
E_{\text{int}}(18C6 \cdot \cdot \cdot \text{benzenediol}) = E(18C6 \cdot \cdot \cdot \text{benzenediol}) - E(18C6) - E_{\text{int}}(\text{benzenediol}) + E_{\text{BSSE}}(18C6 \cdot \cdot \cdot \text{benzenediol}),
\]

where conformations of 18C6 and benzenediol fragments were the same as those in the 18C6-benzenediol complex. The IR spectra were obtained by vibrational analysis, and the vertical electronic transition energies were obtained by TD-DFT calculations.
at the ωB97X-D/6-31++G** level. All the DFT calculations were performed using the Gaussian 09 package, revision D.01. The OH stretching frequencies were scaled by 0.9346, 0.9325 and 0.9325 for HQ, RE, and CA, respectively, to reproduce the observed OH stretching vibration of each monomer. Though the calculated electronic energies are the vertical excitation energies, they are scaled by 0.8645, 0.8603 and 0.8598, for HQ, RE and CA, respectively, to reproduce the observed S1–S0 band origin energy of each monomer.

3. Results
3.1 S1–S0 spectra of benzenediols and their complexes
Fig. 1 shows the LIF and UV–UV HB spectra of the S1–S0 transition of jet cooled HQ, RE, and CA and their complexes with 18C6. Fig. 1(a) shows the LIF spectrum of HQ in the S1–S0 band origin region. Two strong bands at 33 515 and 33 550 cm⁻¹ are already assigned to the 0,0 band of two conformers of HQ, namely trans- and cis-HQ, respectively (see Scheme 1). Weak bands at 33 164 and 33 201 cm⁻¹ are the band origins of the H2O-HQ 1:1 complex due to the residual water in the sample. Fig. 1(b) shows the LIF spectrum measured by adding 18C6 vapor to the gas mixture. New bands appearing around 33 100–33 200 cm⁻¹ can be assigned to the 18C6-HQ 1:1 complex. The UV–UV HB spectrum obtained by monitoring band AHQ is shown in Fig. 1(c). This result clearly shows that the 18C6-HQ 1:1 complex has a single isomer in spite of the fact that the HQ monomer has two conformers with almost equal abundance. The ratio of the band intensity for cis-HQ/trans-HQ is 0.86 without 18C6 vapor, while it becomes 0.57 with 18C6 vapor, indicating that the relative abundance of cis-HQ is reduced by the addition of 18C6 and that the cis-HQ conformer prefers the complex formation with 18C6 more than trans-HQ.

Fig. 1(d) and (e) exhibit the LIF spectra of RE without and with 18C6 vapor, respectively. The RE monomer has three
possible conformations (see Scheme 1, conformers I, II and III). Hereafter we call these conformers RE(I), RE(II) and RE(III), respectively. The bands at 35 965 and 36 217 cm\(^{-1}\) are assigned to the 0,0 band of RE(I) and RE(II), respectively.\(^{42}\) Thus, under the present jet-cooled conditions, only RE(I) and RE(II) are observed in the LIF spectrum. The total energy of RE(III) is 8.86 kJ mol\(^{-1}\) higher than that of RE(I) at the present level of the calculation, and the electronic transition of RE(III) appears only under warmer jet conditions.\(^{43}\) By the addition of 18C6 vapor, many bands due to the 18C6/CA complexes in the S\(_0\) state will be much stronger. The UV–UV HB spectra, not shown here, indicate that the new bands in Fig. 1(h) are attributed to vibronic bands belonging to either of three species labeled ACA–CCA.\(^3\) Thus, the 18C6–CA complex has three stable isomers, although bare CA has only one conformation. This is in contrast to the other two benzenediols, HQ and RE. The structures of the 18C6-benzenediol 1:1 complexes will be described in a later section. The frequencies of the 0,0 bands of the S\(_1\)–S\(_0\) transition for the observed species are listed in Table 1.

### 3.2 IR spectra of benzenediols and 18C6-benzenediol complexes in the S\(_0\) state

Fig. 2(a)–(i) display the IR–UV DR spectra of HQ, RE, CA and their complexes with 18C6 in the OH stretching frequency region obtained by monitoring their 0,0 bands. For the CA monomer, the position of the IR bands reported in a previous study.\(^{10}\) The band labeled CA in Fig. 1(g) at 35 695 cm\(^{-1}\) is the 0,0 band of the CA monomer. In the spectrum, the bands of the H\(_2\)O–CA complex also appeared. This is due to the presence of residual water in the sample. In CA, two adjacent OH groups form an intramolecular H-bond, so that it has a stable single conformer as seen in Scheme 1. The intensity of band CA in Fig. 1(g) is very weak due to the fast internal conversion (IC) to the \(^1\)\(^\pi\)E* state. On the other hand, the band intensities of the 18C6–CA 1:1 complex in Fig. 1(h) are much stronger. The UV–UV HB spectra, not shown here, indicate that the new bands in Fig. 1(h) are attributed to vibronic bands belonging to either of three species labeled ACA–CCA.\(^3\) Thus, the 18C6–CA complex has three stable isomers, although bare CA has only one conformation. This is in contrast to the other two benzenediols, HQ and RE. The structures of the 18C6-benzenediol 1:1 complexes will be described in a later section. The frequencies of the 0,0 bands of the S\(_1\)–S\(_0\) transition for the observed species are listed in Table 1.

### Table 1

| Band labeling | Obs. | Calc. | S\(_1\)–S\(_0\) transition energy [cm\(^{-1}\) [eV]] | \(\Delta E_{2\rightarrow1}\) [eV] | S\(_1\) lifetime [ns] | Symmetry in S\(_0\) | Symmetry in S\(_1\) |
|---------------|------|-------|---------------------------------|-----------------|-----------------|----------------|----------------|
| cis-HQ        | 3663 | 3662, 3663 | 33 350 (4.160) | 4.649(0.0000) | 0.488 | 2.6 | C\(_{2h}\) | C\(_{2h}\) |
| trans-HQ      | 3662 | 3663 | 33 515 (4.156) | 4.726(0.0000) | 0.571 | 1.2 | C\(_{2v}\) | C\(_{2v}\) |
| H\(_2\)O-cis-HQ | —    | — | 33 201 (4.117) | 4.555(0.0000) | 0.343 | 0.402 | — |
| H\(_2\)O-trans-HQ | —    | — | 33 164 (4.106) | 4.534(0.0000) | 0.251 | 0.802 | — |
| 18C6-cis-HQ-I | A\(_{HQ}\) | 3444, 3657 | 3420, 3653 | 33 115 (4.106) | 4.398(0.0376) | 0.541 | — |
| 18C6-cis-HQ-II | —    | 3410, 3628 | — | 4.347(0.0422) | 0.496 | 0.54 | — |
| RE(I)         | 3658 | 3656, 3658 | 35 965 (4.460) | 4.320(0.0513) | 0.343 | 1.2 | C\(_{1}\) | — |
| RE(II)        | 3658 | 3659 | 36 217 (4.491) | 4.347(0.0422) | 0.496 | 0.54 | — |
| RE(III)       | —    | — | 33 201 (4.117) | 4.555(0.0000) | 0.343 | 0.402 | — |
| 18C6-RE(III)-I | A\(_{RE}\) | 3465 | 3371, 3396 | 35 339 (4.382) | 4.320(0.0422) | 0.496 | 0.54 | — |
| 18C6-RE(III)-II | —    | — | 3359, 3401 | — | 4.347(0.0422) | 0.496 | 0.54 | — |
| 18C6-RE(III)-III | —    | — | 3391, 3429 | — | 4.347(0.0422) | 0.496 | 0.54 | — |
| 18C6-RE(III)-IV | —    | — | 3358, 3369 | — | 4.347(0.0422) | 0.496 | 0.54 | — |
| 18C6-RE(III)-V | —    | 3412, 3462 | — | 4.347(0.0422) | 0.496 | 0.54 | — |
| CA            | 3611, 3673 | 3612, 3672 | 35 695 (4.426) | 4.682(0.0004) | 0.256 | 0.008 | C\(_{1}\) | C\(_{1}\) |
| 18C6-CA-A1   | C\(_{CA}\) | 3281, 3507 | 3311, 3584 | 35 397 (4.389) | 4.895(0.0020) | 0.548 | — |
| 18C6-CA-E1   | A\(_{CA}\) | 3385, 3407 | 3372, 3429 | 35 230 (4.369) | 5.041(0.0241) | 0.721 | 10.3 | C\(_{1}\) |
| 18C6-CA-E4   | B\(_{CA}\) | 3424 | 3386, 3396 | 35 548 (4.408) | 4.993(0.0181) | 0.639 | 10.3 | — |

\(^a\) Calculated energies are scaled by 0.8645, 0.8603 and 0.8598 for HQ, RE and CA, respectively, so as to reproduce the observed S\(_1\)–S\(_0\) transition of each benzenediol monomer. \(^b\) Ref. 53.
Table 1 also lists the observed frequencies of the OH stretching vibrations. The OH bands of trans- and cis-HQ appear at 3662 and 3663 cm\(^{-1}\), respectively. In HQ, the two OH groups are symmetrically equivalent in the two conformers. So, the observed IR band is assigned to the antisymmetric OH stretching vibration in trans-HQ, while the symmetric and antisymmetric OH stretching vibrations are thought to be overlapped in cis-HQ. The IR spectrum of the 18C6-HQ complex (A\(_{HQ}\)) is shown in Fig. 2(c). The bands at 3444 and 3657 cm\(^{-1}\) are assigned to the H-bonded OH stretch and the free OH stretch, respectively. Several bands appearing at the higher frequency side of the H-bonded OH are assigned to the combination bands with low frequency vibrations, probably intermolecular vibrations. Thus, in the 18C6-HQ complex, one OH group is H-bonded and the other is free from the H-bond.

Fig. 2(d) and (e) show the IR spectra of the RE(I) and RE(II) monomers, respectively. In RE(II), the antisymmetric OH stretch band has a stronger IR intensity, while in RE(I) two OH bands may have similar intensity because of the lower molecular symmetry. Actually, in both RE(I) and RE(II), only...
one OH band appeared at the same frequency at 3658 cm\(^{-1}\) in the observed spectra.\(^\text{45}\) This result indicates that the coupling between the two OH groups is very weak, which is also supported in the calculated spectra of Fig. 2(k). In the IR–UV spectrum of the 18C6-RE complex (A\(_{\text{RE}}\)) (Fig. 2(f)), no free OH band is seen at \(\sim 3650\) cm\(^{-1}\) and three bands appear in the 3460–3500 cm\(^{-1}\) region, which are assigned to the H-bonded OH. Among them, the intensity of the lowest frequency band at 3465 cm\(^{-1}\) is the strongest. Since RE has only two OH groups, the appearance of three bands means that one of the two higher frequency bands is the combination band with a low frequency intermolecular vibration.

Fig. 2(g)–(i) display the IR–UV spectra (red curves) of 18C6-CA by monitoring the intensity of bands C\(_{\text{CA}}\), A\(_{\text{CA}}\) and B\(_{\text{CA}}\), and the calculated IR spectra (black curves) of the corresponding structures. Since the IR spectrum of monomer CA could not be obtained because of its weak fluorescence intensity, we adopted the reported values of the frequencies of IR bands.\(^\text{44}\) The IR spectrum of species C\(_{\text{CA}}\) shows a broad OH band centered at 3281 cm\(^{-1}\) and a sharp one at 3507 cm\(^{-1}\). The former band is assigned to the H-bonded OH with 18C6 and the latter to the intramolecular H-bonded OH band. The IR spectra of species A\(_{\text{CA}}\) and B\(_{\text{CA}}\) exhibit the OH bands at the 3350–3450 cm\(^{-1}\) region, indicating that the two OH groups are H-bonded with 18C6 with almost equal strength. The assignments of the OH vibrations by comparison with the calculated ones will be given in a later section.

3.3 \(S_1\) lifetimes of the benzenediol monomers and their complexes

We then measured the \(S_1\) lifetime of each monomer and complexes. Fig. 3 shows the picosecond pump–probe decay profiles of (a) cis-HQ, (d) RE(I), (e) RE(II) and (g) CA monomers, respectively. They are observed by exciting at the 0,0 band of \(S_1\), giving the lifetime of 2.6 ± 0.2 ns, 4.2 ± 0.3 ns, 4.5 ± 0.5 ns, and 8.0 ps, respectively. The very short \(S_1\) lifetime of CA was investigated by Livingstone \textit{et al.}\(^\text{46}\) and Chatterley \textit{et al.}\(^\text{47}\) and is attributed to the fast IC to the nearby *π*-π* state, leading to an atomic hydrogen elimination of the acceptor OH (\(b\text{OH}\) in Scheme 1). Fig. 3(c), (f), (h) and (i) display the decay profiles of the 18C6 complexes. The change of the \(S_1\) lifetime upon complexation is quite different for different benzenediols; in the case of HQ, the \(S_1\) lifetime of the 18C6-HQ complex (0.54 ns, Fig. 3(c)) is shorter by a factor of 5 than that of the monomer. In contrast, the lifetime of the 18C6-RE complex (10.5 ns) is 2.5 times longer than that of the RE monomer, and the lifetime of the 18C6-CA complex (10.3 ns) is 1280 times longer than that of the CA monomer.

4. Discussion

4.1 Structure of 18C6-benzenediol 1 : 1 complexes

4.1.1 18C6-HQ complex. As shown in Fig. 1(c), only one isomer appears for the 18C6-HQ complex. The fact that the...
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An interesting point is that HQ adopts the cis-form in the complex can be explained by the fact that the complex is largely stabilized by the dipole–dipole interaction between cis-HQ and 18C6; cis-HQ has a dipole moment of 2.789 D, while that of trans-HQ is almost zero. In addition, though the most stable 18C6 has a planar structure (C₃ symmetry) in the bare form, it changes the structure so that it has a dipole moment to include HQ. Thus, both the molecules cooperatively change the conformation to have dipole moments and stabilize during the process of the host–guest complex formation. Thus, the complex formation mechanism can be called as ′induced-fitting′.

Fig. 2(j) shows the calculated IR spectra of bare HQ and the 18C6-cis-HQ-I complex of Fig. 4(a). Both trans- and cis-HQ in the bare form show the free OH stretching bands at 3663 cm⁻¹. In the complex, the calculated IR spectrum of 18C6-cis-HQ-I shows the H-bonded (3420 cm⁻¹) and free (3653 cm⁻¹) OH stretching vibrations, which nicely reproduces the observed spectrum of species A₁HQ (Fig. 2(c)). It should be noted that 18C6-cis-HQ-I, -II and 18C6-trans-HQ show very similar calculated IR spectra with each other (see Fig. S1, ESI†). Based on the relative energies and IR spectra, it is concluded that the structure of species A₁HQ is 18C6-cis-HQ-I. This complex is stabilized through multiple interactions such as the OH⋯O H-bond, CH⋯π, and dipole–dipole interactions. This synergistic effect is very important for the stabilization and conformation preference of this host–guest complex.

4.1.2 18C6-RE complex. Fig. 5(a)–(e) show the stable isomers of the 18C6-RE complex within the energy of 10 kJ mol⁻¹. As shown in Fig. 5, RE takes the RE(III) conformation in all isomers, which did not appear in the monomer form, actually, the calculated energies of R(II) and R(III) are 0.40 and 8.86 kJ mol⁻¹ higher than R(I) in the monomer. In the complex, however, the lowest energy isomer having RE(I) conformation (18C6-RE(I)) is 27.1 kJ mol⁻¹ higher than 18C6-RE(III)-I (see Fig. S2(a), ESI†),

Table 2. Relative total energies (ΔE [kJ mol⁻¹]) of the complexes, interaction energies (E_int [kJ mol⁻¹]), relative energies of the conformers of the 18C6 part (ΔE_CE [kJ mol⁻¹]) and the benzenediol part (ΔE_HQ,RE or CA [kJ mol⁻¹]) in the complexes, and dihedral angles of the OH groups of benzenediol in the complexes. The species with bold letters are those experimentally observed. The calculations are performed at the b97X-D/6-31++G** level.

| 18C6-cis-HQ | 18C6-RE(III) | 18C6-CA |
|---|---|---|
| | Dihedral angle | | Dihedral angle | | Dihedral angle |
| | | a | b | OH | OH | | | a | b | OH | OH | | | a | b | OH | OH |
| I | 0.00 | 94.3 | I | 0.00 | 113 | 20.0 | 11.4 | 178 | 171 | | A1 | 0.00 | 92.1 | 10.2 | 2.28 | 179 | 179 |
| II | 9.10 | 88.7 | 9.26 | 2.11 | 168 | 178 | | | | | E1 | 3.94 | 93.2 | 9.34 | 7.98 | 151 | 160 |
| — | — | — | — | | — | — | — | — | — | E4 | 8.66 | 93.9 | 7.01 | 16.2 | 170 | 140 |
| — | — | — | 1.97 | 9.12 | 12.6 | 167 | 163 | | | | | | | |
| III | 3.34 | 93.7 | 2.83 | 10.6 | 179 | 179 | | | | | | | | | | | | | |
| IV | 8.75 | 91.3 | 4.13 | 12.9 | 158 | 154 | | | | | | | | | | | | | |
| — | — | — | — | — | — | — | | | | | | | | | | | | | |

*ΔE_CE are relative values to that of the most stable 18C6 conformer in the monomer. bΔE_{HQ,RE or CA} are relative values to that of the most stable conformer in each benzenediol monomer. cDihedral angles of ⁴OH and ⁵OH are defined in Scheme 1.*
and the 18C6·RE(II) isomer is not obtained as a stable structure even up to 40 kJ mol\(^{-1}\). The stability of the 18C6·RE(III) isomer mostly comes from the formation of two OH·–O H-bonds as seen in Fig. 5. Fig. 2(k) shows the calculated IR spectra of RE(I), RE(II), 18C6·RE(III)-I and 18C6·RE(II). Similar to HQ, bare RE shows only a free OH stretch band. RE(I) and RE(II) have \(C_1\) and \(C_{2v}\) symmetry, respectively. The two free OH bands are overlapped with each other with equal intensity in RE(I), but in RE(II) the anti-symmetric OH stretch band appears much stronger than the symmetric OH stretch. In any case, both the symmetric and anti-symmetric OH stretches have almost the same frequency. The IR spectrum of 18C6·RE(III)-I shows two H-bonded OH stretching vibrations at 3371 and 3396 cm\(^{-1}\), which reproduces the observed IR spectra of A\(_{RE}\). The IR spectrum of the 18C6·RE(I) isomer exhibits a free OH stretching vibration at \(\sim3655\) cm\(^{-1}\), which does not reproduce the observed IR spectrum of A\(_{RE}\). Thus, the observed species A\(_{RE}\) is assigned to the 18C6·RE(III) isomer.

Similar to the case of HQ, multiple conformers coexist in the bare form in RE, but the complexation with 18C6 controls the conformation of RE to the highest energy one (RE(III)) to form the most stable complex. In addition, 18C6 also changes its conformation so as to fit the guest species in its cavity. The complex is finally stabilized by two OH·–O H-bonds and CH·–π interactions, and this synergetic effect is also essential for the stabilization of the host-guest complex of crown ether.

**4.1.3 18C6·CA complex.** The difference between CA and the other benzenediols (HQ and RE) is the presence of intramolecular H-bonding in bare CA, which leads to a unique feature of CA, such as the very short \(S_1\) lifetime compared to HQ or RE. Complexation with 18C6 largely affects this intramolecular H-bonding. Fig. 6 shows the calculated structures of 18C6·CA assignable to the observed three species, A\(_{CA}\), B\(_{CA}\) and C\(_{CA}\), in our previous study. \(^31\) In that paper, species A\(_{CA}\) and B\(_{CA}\) were assigned to 18C6·CA-E1 (Fig. 6(b)) and 18C6·CA-E4 (Fig. 6(e)), respectively, in which the intramolecular H-bonding of CA is broken and the two OH groups are independently H-bonded to the oxygen atoms of 18C6 (we call it Type-E). On the other hand, in species C\(_{CA}\) (structure 18C6·CA-A1), CA retains its intramolecular H-bonding and its acceptor OH is H-bonded to the oxygen of 18C6 (we call it Type-A). Other possible isomers of the 18C6·CA complex are shown in Fig. S2b (ESI\(^†\)). The calculated IR spectra (black curves) of the three isomers of 18C6·CA are shown below the observed ones. The calculated IR spectrum of 18C6·CA-E1 shows H-bonded OH stretches at 3372 and 3429 cm\(^{-1}\), and that of 18C6·CA-E4 at 3386 and 3396 cm\(^{-1}\). In the IR spectrum of 18C6·CA-A1, an intermolecular H-bonded OH appears at 3325 cm\(^{-1}\) and intramolecular H-bonded OH at 3584 cm\(^{-1}\). Though isomer 18C6·CA-A1, assigned to species C\(_{CA}\), is the most stable structure, the fluorescence intensity of species C\(_{CA}\) is rather weak compared to A\(_{CA}\) or B\(_{CA}\). One of the possibilities of the weakness of this band is a smaller fluorescence quantum yield of this species than that of A\(_{CA}\) and B\(_{CA}\). A measurement of the fluorescence lifetime of band C\(_{CA}\) was unsuccessful because of its weakness, so that it is difficult to discuss the relative abundance of the complexes.

Based on the results of IR measurements, it is concluded that the 18C6·CA complex has two types of isomers having different H-bonding structures. An important point is that 18C6 has an ability to break the intramolecular H-bond of CA,
which is connected to the drastic changes of the excited state electronic structure of guest CA, as will be discussed later.

4.2 Energetics of 18C6-benzenediol 1:1 complexes

As described above, 18C6 and benzenediols form the most stable host–guest complexes at the expense of changing each structure to higher energy ones. In this section, we compare the energies of the bare form in the most stable conformations, and those of the distorted one for forming the complex, and that of the complex. Fig. 7(a) and (b) show the schematic energy diagram of the 18C6-HQ and 18C6-RE systems: the most stable structures of benzenediols and 18C6 (left), distorted structures (middle), and the stable 18C6-benzenediol complexes (right). It should be noted that in the figures the horizontal axis does not mean the complex formation coordinate. In Fig. 7, $\Delta E_{CE}$ is the energy difference of 18C6 between the most stable structure in the bare form and that in the complex. $\Delta E_{HQ}$, $\Delta E_{RE}$ or $\Delta E_{CA}$ is also the energy difference of the benzenediol part. $E_{int}$ is the interaction energy of the complex, which is obtained by using eqn (1) described in the Computational section. In the 18C6-HQ-I complex, $E_{int}$ is obtained to be 94.3 kJ mol$^{-1}$. In the case of HQ (Fig. 7(a)), the most stable structure of bare HQ has the trans form. The energy difference between “most stable trans-HQ monomer + most stable 18C6” and 18C6-HQ-I complex corresponds to the binding energy of the 18C6-HQ-I complex. In the complex, the 18C6 and HQ components take higher energy structures, (18C6)$^+$ and (HQ)$^-$ (middle in Fig. 7(a)), to form the stable complex (right in Fig. 7(a)). The energy necessary for this distortion, ($\Delta E_{CE} + \Delta E_{HQ}$), is 10.6 kJ mol$^{-1}$. Hence, the interaction energy $E_{int}$ is represented by using the binding energy and ($\Delta E_{CE} + \Delta E_{HQ}$), as follows:

$$E_{int} = \text{Binding energy} + (\Delta E_{CE} + \Delta E_{HQ}) \quad (2)$$

The BSSE corrected binding energy of the complex is obtained to be 83.7 kJ mol$^{-1}$. This value leads $E_{int}$ to be 94.3 kJ mol$^{-1}$ by using eqn (2), which is equal to that obtained by using eqn (1). Fig. 7(b) shows the schematic energy diagram for the 18C6-RE complex. The energy difference between the most stable monomers and distorted ones, ($\Delta E_{CE} + \Delta E_{RE}$), is 31.4 kJ mol$^{-1}$,
the binding energy of the complex is 81.6 kJ mol\(^{-1}\), and the interaction energy \(E_{\text{int}}\) is 113 kJ mol\(^{-1}\).

Fig. 8 shows a similar energy diagram for the 18C6-CA complex system. In this case, two different types of stable complexes are formed, 18C6-CA-A1 and 18C6-E1, as described previously. The value of \((\Delta E_{\text{CE}} + \Delta E_{\text{CA}})\) is 12.5 kJ mol\(^{-1}\) for 18C6-CA-A1 and 17.3 kJ mol\(^{-1}\) for 18C6-CA-E1. The binding energy of the complex is 79.7 for 18C6-CA-A1 and 75.8 kJ mol\(^{-1}\) for 18C6-CA-E1, and \(E_{\text{int}}\) is 92.1 kJ mol\(^{-1}\) for 18C6-CA-A1 and 93.2 kJ mol\(^{-1}\) for 18C6-CA-E1.

Table 2 lists the relative energies of the complexes \((\Delta E)\), \(E_{\text{int}}, \Delta E_{\text{CE}}\), \(\Delta E_{\text{CA}}\), \(\Delta E_{\text{H}2\text{O}}, \Delta E_{\text{RE}}\) and \(\Delta E_{\text{CA}}\) and the dihedral angles of OH groups \((\text{OH}, \text{OH})\) with respect to the benzene plane. For the benzenediol part, \(\Delta E_{\text{H}2\text{O}}\) is 1.64–2.11 kJ mol\(^{-1}\), \(\Delta E_{\text{RE}}\) is 10.6–12.9 kJ mol\(^{-1}\), and \(\Delta E_{\text{CA}}\) is 2.28–16.2 kJ mol\(^{-1}\). These values are quite different for different species and conformations. As seen in Fig. 7 and 8 and Table 2, all the necessary energies for forming the higher energy complexes are easily compensated by the large interaction energy of 90–110 kJ mol\(^{-1}\) or the binding energies of 83.7 kJ mol\(^{-1}\) for 18C6-cis-HQ-I, 81.6 kJ mol\(^{-1}\) for 18C6-RE(III)-I and 70.7–79.6 kJ mol\(^{-1}\) for 18C6-CA complexes. These binding energies are roughly 4 times larger than the normal OH···O H-bond energy (for example the H-bond energy of the water dimer is 19–20 kJ mol\(^{-1}\))\(^\text{,52}\) Such a large binding energy comes from the multiple interactions between benzenediols and 18C6, such as the OH···O H-bond, CH···π and dipole–dipole interactions. These interactions are actually seen in the calculated structures in Fig. 4–6. In these figures, the number of OH···O and CH···π bonding are also shown; the most stable isomers have the largest number of intermolecular bonding between the host and guest molecules. These results strongly indicate that the 18C6-benzenediol host–guest complex formation is not described as a simple rigid “lock-and-key” model, but as a flexible “induced-fit” model, like an enzyme–substrate fitting.

**4.3 The S\(_1\) lifetime of 18C6-benzenediol complexes**

The observed \(S_1\) lifetimes of benzenediols and 18C6-benzenediol complexes are also listed in Table 1. As described in the Introduction section, the major non-radiative pathway of the \(S_1\) state of benzenediols is thought to be IC to \(S_2\) \(\pi\pi^*\) via the conical intersection, followed by the O–H bond fission along the repulsive \(S_2\) potential curve or the further IC to \(S_0\) via the \(S_2\)-\(S_0\) conical intersection. Anomalous short lifetime of bare CA (8 ps) is described by the low barrier height of the \(S_1\)-\(S_2\) conical intersection due to the small \(S_1\)/\(S_2\) energy gap \((\Delta E_{2–1})\) and low symmetry \((C_1)\) in the \(S_1\) state.\(^\text{53,54}\) So, we calculated the vertical excitation energies to the \(S_1\) and \(S_2\) states with the fixed \(S_0\) geometry and obtained the \(S_1\)-\(S_2\) energy gap \((\Delta E_{2–1})\), which is also listed in Table 1. Since we do not calculate the potential energy curves along the O–H coordinate, \(\Delta E_{2–1}\) is not equal to the barrier height at the \(S_1\)-\(S_2\) \(\pi\pi^*\) conical intersection. Even if it is so, we consider that this value is roughly proportional to the barrier height. The calculated \(\Delta E_{2–1}\) of bare HQ and RE is 0.439–0.571 eV, while that of CA is much smaller, 0.256 eV. This difference is in good agreement with the shorter \(S_1\) lifetime of CA than HQ or RE. The complex formation with 18C6 greatly affects the nonradiative process of each benzenediol in a different manner. First, in the case of HQ, the \(S_1\) lifetime is shortened by a factor of five (2.6 ns \(\rightarrow\) 0.54 ns) in the complex, while the lifetime is 2.5 times lengthened in the RE complex (4.0 ns \(\rightarrow\) 10.5 ns) and 1280 times in the Type-E complex of CA (8.0 ps \(\rightarrow\) 10.3 ns). Thus, the 18C6-HQ complex shows an opposite trend of the \(S_1\) lifetime behavior compared to that of 18C6-CA.
18C6-RE and 18C6-CA. This difference is attributed to the presence of free OH in 18C6-HQ but not in 18C6-RE and 18C6-CA. As seen in Table 1, the presence of free OH does not affect strongly the $\Delta E_{2-1}$ value: 0.471 eV for RE[I] changes to 0.512 eV for 18C6-RE[III]-I and 0.488 eV for cis-HQ changes to 0.496 eV for 18C6-HQ-II. Only CA shows a large change in the $\Delta E_{2-1}$ value upon complex formation: 0.256 eV for CA $\rightarrow$ 0.721 eV for 18C6-CA-E1. However, the difference can be seen clearly in the molecular orbitals of the electronic states between 18C6-HQ, 18C6-RE and 18C6-CA. The molecular orbitals responsible for the $S_1$ and $S_2$ electronic states are shown in Fig. S3 [ESI†]. In the figure, it is clear that the $\sigma^*$ character dominates in the $S_2$ state in all the monomer species. On the other hand, for the 18C6 complexes, the $\sigma^*$ character can be seen only at the free OH site of 18C6-HQ [Fig. S3(c), ESI†]. In addition, the dihedral angle defined in Scheme 1 of cis-HQ becomes 168–178 degree, indicating that the molecular symmetry of cis-HQ becomes lower from $C_{2h}$ to $C_1$ by complexation. Hence, this lower molecular symmetry causes the shorter $S_1$ lifetime of HQ upon complexation. In the case of 18C6-RE and 18C6-CA, the cage of 18C6 diminishes the $\sigma^*$ character in the $S_2$ state, leading to the suppression of the OH cleavage. Rizzo and co-workers observed a similar lifetime elongation in a protonated triphenyl-($H_2O)_2$ complex.55 The protonation to triphenyl lowers the energy of the $\pi^*$ ($\sigma^*$ of N–H) state, facilitating a mixing between the $\pi^*$ and indole $\pi \pi^*$ states. Thus, the protonated triphenyl monomer exhibits a very broad electronic spectrum. In contrast, the hydration to the $NH_3^+$ group by two water molecules shifts the $\sigma^*$ of N–H to higher energy, resulting in the decoupling of $\pi^*$ from $\pi \pi^*$. In other words, two water molecules “block” the N–H bond fission of protonated triphenyl in the excited state, resulting in the sharp electronic spectrum. In the case of 18C6-HQ, the $\pi^*$ character is still retained in the free OH group in the $S_2$ state as seen in Fig. S3(e) [ESI†].

5. Conclusion

In the present study, we investigated the formation of host–guest complexes between 18C6 and benzenediols in the gas phase under supersonically jet-cooled conditions. We found that 18C6 and benzenediols cooperatively change their conformations from the stable bare forms to higher energy ones for the best-fitted stable complexes. The complexes are stabilized synergistically through multiple interactions. Especially, the RE part in 18C6-RE takes the characteristic conformation (RE[III]), which is the highest energy one in the monomer form. In the 18C6-CA Type-E complex, CA breaks its intramolecular H-bond to form two intermolecular H-bonds with 18C6. It is also found that complex formation controls the direction of the OH group remote from the direct contact. All the complexes are stabilized by synergetic intermolecular interactions, which are produced by the geometry changes of the host and guest species via a typical “induced-fit” mechanism.

It was also found that the host–guest complexation with crown ethers drastically changes the electronic excited state of guest species. In the case of RE and CA, the $S_1$ lifetime is lengthened by a factor of 2.5 and 1280, respectively. This drastic change is described partly by an increase of the $S_1/S_2$ energy gap and mostly by the destruction of the $\pi^*$ character of the $S_2$ state. In contrast, in HQ, the complexation shortens the $S_1$ lifetime of HQ by a factor of five, thus accelerating the nonradiative decay rate. The present study has shown that 18C6, a simple crown ether, can exert isomeric preference and modify the excited state dynamics of guest species through multiple interactions. In future work, we will extend this work to more flexible molecules as guest molecules and elucidate the potential ability of crown ether to control the conformational preference as well as the excited state dynamics of the guest.

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References

1 C. J. Pedersen, J. Am. Chem. Soc., 1967, 89, 7017–7036.
2 C. J. Pedersen and H. K. Frensdorff, Angew. Chem., Int. Ed. Engl., 1972, 11, 16–25.
3 C. J. Pedersen, Science, 1988, 241, 536–540.
4 R. M. Izatt, J. H. Rytting, D. P. Nelson, B. L. Hayamore and J. J. Christensen, Science, 1969, 164, 443–444.
5 R. M. Izatt, D. P. Nelson, J. H. Rytting and J. J. Christensen, J. Am. Chem. Soc., 1971, 93, 1619–1623.
6 R. M. Izatt, R. E. Terry, B. L. Haymore, L. D. Hansen, N. K. Dalley, A. G. Avondet and J. J. Christensen, J. Am. Chem. Soc., 1976, 98, 7620–7626.
7 R. M. Izatt, R. E. Terry, D. P. Nelson, Y. Chan, D. J. Eatough, J. S. Bradshaw, L. D. Hansen and J. J. Christensen, J. Am. Chem. Soc., 1976, 98, 7626–7630.
8 J. D. Lamb, R. M. Izatt, C. S. Swain and J. J. Christensen, J. Am. Chem. Soc., 1980, 102, 475–479.
9 H. Zhang, J. H. Chu, S. Leming and D. V. Dearden, J. Am. Chem. Soc., 1991, 113, 7415–7417.
10 H. Zhang and D. V. Dearden, J. Am. Chem. Soc., 1992, 114, 2754–2755.
11 S. Maleknia and J. Brodbelt, J. Am. Chem. Soc., 1992, 114, 4295–4298.
12 I. H. Chu, H. Zhang and D. V. Dearden, J. Am. Chem. Soc., 1993, 115, 5736–5744.
13 L. X. Dang, J. Am. Chem. Soc., 1995, 117, 6954–6960.
14 P. D. J. Grootenhuis and P. A. Kollman, J. Am. Chem. Soc., 1989, 111, 2152–2158.
15 D. Ray, D. Feller, M. B. More, E. D. Glendening and P. B. Armentrout, J. Phys. Chem., 1996, 100, 16116–16125.
16 F. Sobott, W. Kleinekofort and B. Brutschy, Anal. Chem., 1997, 69, 3587–3594.
17 M. B. More, D. Ray and P. B. Armentrout, J. Phys. Chem. A, 1997, 101, 4254–4262.
18. M. B. More, D. Ray and P. B. Armentrout, J. Phys. Chem. A, 1997, 101, 7007–7017.
19. M. B. More, D. Ray and P. B. Armentrout, J. Am. Chem. Soc., 1999, 121, 417–423.
20. Y. Inokuchi, O. V. Boyarkin, R. Kusaka, T. Haino, T. Ebata and T. R. Rizzo, J. Am. Chem. Soc., 2011, 133, 12256–12263.
21. Y. Inokuchi, O. V. Boyarkin, R. Kusaka, T. Haino, T. Ebata and T. R. Rizzo, J. Phys. Chem. A, 2012, 116, 4057–4068.
22. Y. Inokuchi, R. Kusaka, T. Ebata, O. V. Boyarkin and T. R. Rizzo, ChemPhysChem, 2013, 14, 649–660.
23. Y. Inokuchi, T. Mizuuchi, T. Ikeda, T. Haino, T. Kimura, H. Guo and Y. Furutani, J. Am. Chem. Soc., 2014, 136, 90–95.
24. Y. Inokuchi, T. Ebata, T. R. Rizzo and O. V. Boyarkin, Chem. Phys. Lett., 2014, 136, 1815–1824.
25. R. Kusaka, Y. Inokuchi and T. Ebata, Phys. Chem. Chem. Phys., 2007, 9, 4452–4459.
26. R. Kusaka, Y. Inokuchi and T. Ebata, Phys. Chem. Chem. Phys., 2008, 10, 6238–6244.
27. R. Kusaka, Y. Inokuchi and T. Ebata, Phys. Chem. Chem. Phys., 2009, 11, 9132–9140.
28. S. Kokubo, R. Kusaka, Y. Inokuchi, T. Haino and T. Ebata, Phys. Chem. Chem. Phys., 2010, 12, 3559–3565.
29. R. Kusaka, Y. Inokuchi, T. Haino and T. Ebata, J. Phys. Chem. Lett., 2012, 3, 1414–1420.
30. Y. Inokuchi, O. V. Boyarkin, T. Ebata and T. R. Rizzo, Phys. Chem. Chem. Phys., 2012, 14, 4457–4462.
31. F. Morishima, R. Kusaka, Y. Inokuchi, T. Haino and T. Ebata, J. Phys. Chem. B, 2015, 119, 2557–2565.
32. Y. Inokuchi, Y. Kobayashi, T. Ito and T. Ebata, J. Phys. Chem. A, 2007, 111, 3209–3215.
33. R. J. Lipert and S. D. Colson, J. Phys. Chem., 1989, 93, 3894–3896.
34. Y. Yamada, N. Mikami and T. Ebata, Proc. Natl. Acad. Sci. U. S. A., 2008, 105, 12690–12695.
35. R. Kusaka and T. Ebata, Angew. Chem., Int. Ed., 2010, 49, 6989–6992.
36. I. Kolossváry and W. C. Guida, J. Am. Chem. Soc., 1996, 118, 5011–5019.
37. MacroModel, version 9.1, Schrödinger, LLC, New York, 2005.
38. T. A. Halgren, J. Comput. Chem., 1999, 20, 730–748.
39. M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, Ö. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, Gaussian 09, Revision D.01, Gaussian, Inc., Wallingford CT, 2009.
40. S. J. Humphrey and D. W. Pratt, J. Chem. Phys., 1993, 99, 5078 (J. Chem. Phys., 1996, 104, 2752).
41. P. S. Meenakshi, N. Biswas and S. Waregaonkar, Phys. Chem. Chem. Phys., 2003, 5, 294–299.
42. M. Gerhards, W. Perl and K. Kleinermanns, Chem. Phys. Lett., 1995, 240, 506–512.
43. S. Melandri, G. Maccaferr, W. Caminati and P. G. Favero, Chem. Phys. Lett., 1996, 256, 513–517.
44. M. Gerhards, C. Unterberg and K. Kleinermanns, Phys. Chem. Chem. Phys., 2000, 2, 5538–5544.
45. K. Hattori and S. Ishiuchi, J. Phys. Chem. A, 2007, 111, 6028–6033.
46. R. A. Livingstone, J. O. F. Thompson, M. Iljina, R. J. Donaldson, B. J. Sussman, M. J. Paterson and D. Townsend, J. Chem. Phys., 2012, 137, 184304.
47. A. S. Chatterley, J. D. Young, D. Townsend, J. M. Żurek, M. J. Paterson, G. M. Roberts and V. G. Stavros, Phys. Chem. Chem. Phys., 2013, 15, 6879–6892.
48. G. Wipff, P. Weiner and P. Kollman, J. Am. Chem. Soc., 1982, 104, 3249–3258.
49. Y. Sun and P. A. Kollman, J. Chem. Phys., 1992, 97, 5108.
50. E. D. Glendening, D. Feller and M. A. Thompson, J. Am. Chem. Soc., 1994, 116, 10657–10669.
51. M. J. Bovill, D. J. Chadwick, I. O. Sutherland and D. Watkin, J. Chem. Soc., Perkin Trans. 2, 1980, 1529–1543.
52. M. W. Feueriesen, D. Feller and D. A. Dixon, J. Phys. Chem., 1996, 100, 2993–2997.
53. (a) M. Gerhards, C. Unterberg and S. Schumm, J. Chem. Phys., 1999, 111, 7966; (b) W. B. Tseng, K. Narayanan, C. Y. Hsieh and C. C. Tung, Spectrochim. Acta, Part A, 1997, 53, 2595–2604; (c) S. J. Humphrey and D. W. Pratt, J. Chem. Phys., 1993, 99, 5078; (d) G. A. King, T. A. A. Oliver, R. N. Dixon and M. N. R. Ashfold, Phys. Chem. Chem. Phys., 2012, 14, 3338–3345.
54. M. Weiler, M. Miyazaki, G. Féraud, S. Ishiuchi, C. Dedonder, C. Jouvet and M. Fujii, J. Phys. Chem. Lett., 2013, 4, 3819–3823.
55. S. R. Mercier, O. V. Boyarkin, A. Kamariotis, M. Guglielmi, I. Tavernelli, M. Cascella, U. Rothlisberger and T. R. Rizzo, J. Am. Chem. Soc., 2006, 128, 16938–16943.