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Interactions between the chloride anion and aromatic molecules: infrared spectra of the Cl-\_C6H5CH3, Cl-\_C6H5NH2 and Cl-\_C6H5OH complexes

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Abstract
The Cl$^-$-$\text{C}_6\text{H}_5\text{CH}_3$, Cl$^-$-$\text{C}_6\text{H}_5\text{NH}_2$, and Cl−-$\text{C}_6\text{H}_5\text{OH}$ anion complexes are investigated using infrared photodissociation spectroscopy and ab initio calculations at the MP2/aug-cc-pVDZ level. The results indicate that for Cl$^-$-$\text{C}_6\text{H}_5\text{NH}_2$ and Cl$^-$-$\text{C}_6\text{H}_5\text{OH}$, the Cl$^-$ anion is attached to the substituent group by a single near-linear hydrogen bond. For Cl$^-$-$\text{C}_6\text{H}_5\text{CH}_3$, the Cl$^-$ is attached to an ortho-hydrogen atom on the aromatic ring and to a hydrogen atom on the methyl group by a weaker hydrogen bond. The principal spectroscopic consequence of the hydrogen-bonding interaction in the three complexes is a red-shift and intensity increase for the CH, NH, and OH stretching modes. Complexities in the infrared spectra in the region of the hydrogen-bonded XH stretch band are associated with Fermi resonances between the hydrogen-stretching vibrational modes and bending overtone and combination levels. There are notable correlations between the vibrational red-shift, the elongation of the H-bonded XH group, and the proton affinity of the aromatic molecule's conjugate base.

Keywords
cl, spectra, infrared, molecules, aromatic, anion, chloride, between, interactions, complexes, _c6h5nh2, _c6h5oh, _c6h5ch3, GeoQUEST

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Interactions between the Chloride Anion and Aromatic Molecules: Infrared Spectra of the Cl$^-$–C$_6$H$_5$CH$_3$, Cl$^-$–C$_6$H$_5$NH$_2$ and Cl$^-$–C$_6$H$_5$OH Complexes

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The Cl$^-$–C$_6$H$_5$CH$_3$·Ar, Cl$^-$–C$_6$H$_5$NH$_2$·Ar, and Cl$^-$–C$_6$H$_5$OH·Ar anion complexes are investigated using infrared photodissociation spectroscopy and ab initio calculations at the MP2/aug-cc-pVDZ level. The results indicate that for Cl$^-$–C$_6$H$_5$NH$_2$ and Cl$^-$–C$_6$H$_5$OH, the Cl$^-$ anion is attached to the substituent group by a single near-linear hydrogen bond. For Cl$^-$–C$_6$H$_5$CH$_3$, the Cl$^-$ is attached to an ortho-hydrogen atom on the aromatic ring and to a hydrogen atom on the methyl group by a weaker hydrogen bond. The principal spectroscopic consequence of the hydrogen-bonding interaction in the three complexes is a red-shift and intensity increase for the CH, NH, and OH stretching modes. Complexities in the infrared spectra in the region of the hydrogen-bonded XH stretch band are associated with Fermi resonances between the hydrogen-stretching vibrational modes and bending overtone and combination levels. There are notable correlations between the vibrational red-shift, the elongation of the H-bonded XH group, and the proton affinity of the aromatic molecule’s conjugate base.

1. Introduction

This paper is concerned with the properties of the Cl$^-$–C$_6$H$_5$CH$_3$, Cl$^-$–C$_6$H$_5$NH$_2$, and Cl$^-$–C$_6$H$_5$OH complexes as revealed through their infrared spectra in the hydrogen-stretching region and through ab initio calculations. Motivation for investigating these systems stems in part from the importance of interactions between simple ions such as Na$^+$, K$^+$, and Cl$^-$ and the phenyl, alkyl, amino, and hydroxyl functional groups in different biological contexts, including ion transport in membrane channels. Prompted by this significance, Lisy and co-workers have investigated the interactions between simple metal cations (M$^+$ = Na$^+$, K$^+$) and aromatic molecules (L = C$_6$H$_6$, C$_6$H$_5$OH) through spectroscopic investigations of mixed M$^+$–L$_n$–(H$_2$O)$_m$ clusters. It was found that whereas Na$^+$ ions are preferentially solvated by H$_2$O rather than by C$_6$H$_6$, the reverse is true for K$^+$. This situation is consistent with the preferential transfer of K$^+$ ions in certain membrane ion channels, where cation–π interactions play an important role. There have also been spectroscopic investigations of complexes involving benzene and other metal cations, including Al$^+$, V$^+$, Fe$^+$, Co$^+$, and N$^+$, in the 500–1500 cm$^{-1}$ range using light generated by a free electron laser$^{6,7}$ and in the benzene CH stretch region (2700–3300 cm$^{-1}$) using light from an OPO. $^{7,9}$ These investigations confirm that the metal cation prefers to bind to the π cloud of the benzene ring.

Infrared spectroscopic investigations of the corresponding anion-aromatic complexes and clusters are scarcer. In one of the few previous studies, complexes of benzene with Cl$^-$, Br$^-$, and I$^-$ were characterized using infrared photodissociation (IRPD) spectroscopy and through ab initio calculations. The infrared spectra and quantum chemical calculations suggest that the complexes have planar structures in which the F$^-$ anion is attached to the benzene by a single H-bond, whereas the larger halides (Cl$^-$, Br$^-$, I$^-$) are attached by twin H-bonds. In all cases, the configuration with the halide anion sitting atop the aromatic ring was found to be unstable, presumably due to repulsive interactions between the halide and the ring π cloud.

In the current study, we extend spectroscopic investigations to complexes consisting of a chloride anion attached to substituted benzene (SBz) molecules (toluene, aniline, phenol). We aim to identify the complexes’ structures by following changes to the XH stretch vibrational frequencies of the SBz subunit caused by interaction with the chloride anion. Although there have been no previous infrared spectroscopic studies of halide–SBz complexes, thermochemical data (binding enthalpies and entropies) have been measured for several systems including Cl$^-$–C$_6$H$_5$OH, Br$^-$–C$_6$H$_5$CH$_3$, and Br$^-$–C$_6$H$_5$NH$_2$. Additionally, Continetti and co-workers recently studied the dissociative photodetachment dynamics of the I$^-$–C$_6$H$_5$NH$_2$ complex using photoelectron–photofragment coincidence spectroscopy. Density functional calculations accompanying the experimental work predicted that the anion complex has a structure in which the I$^-$ anion is attached by a single hydrogen bond to the amine group.

2. Experimental and Theoretical Methods

2.1. Experimental Technique. The vibrational predissociation spectra of the Cl$^-$–SBz complexes were recorded in a low-energy guided ion beam apparatus. The machine consists of a cluster ion source, a primary quadrupole mass filter, an octopole ion guide, a secondary quadrupole mass filter, and a microchannel plate detector. The complexes were synthesized in an electron beam crossed supersonic expansion of argon (6 bar) seeded with traces of CCl$_4$ (as a Cl$^-$ precursor) and the SBz molecule of interest.

In all three cases, the Cl$^-$–SBz binding energy is likely to exceed the IR photon energy. For example, the binding enthalpy of Cl$^-$–C$_6$H$_5$OH has been determined as 109.0 ± 8.4 kJ mol$^{-1}$ ($\sim$9100 cm$^{-1}$). Additional to our knowledge, the binding enthalpies of Cl$^-$–C$_6$H$_5$CH$_3$ and Cl$^-$–C$_6$H$_5$NH$_2$ have not been reported, they almost certainly exceed those of Br$^-$–C$_6$H$_5$CH$_3$.

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and Br–C₆H₅CH₃ (36. ± 7.5 and 61.1 ± 7.5 kJ mol⁻¹, respectively³). So that we could monitor single IR photon absorption by cold complexes, we have probed argon-solvated species (Cl–SBz·Ar), monitoring Ar atom loss as a function of the infrared wavelength. The presence of a weakly attached Ar atom has been shown to have a minimal effect on the IR spectra of other anion complexes, the main consequence being a narrowing and slight shifting of the vibrational bands.¹⁹,²⁰

The parent Cl–SBz·Ar clusters were mass-selected by a quadrupole mass filter, turned through 90° by a quadrupole bender, and injected into the octapole ion guide. Here, the clusters were irradiated with the counter-propagating output of a Nd:YAG-pumped OPO/OPA system (Continuum Mirage 3000). Resulting fragment ions were selected by a second quadrupole mass filter and sensed by a microchannel plate/scintillator/multiphoton multiplier detector. Wavelength calibration was achieved by measuring the output from the first stage of the OPO and the 532 nm output from the Nd:YAG laser using a pulsed wavemeter (HighFinesse WS-7).

2.2. Theoretical Methods. Optimized geometries and vibrational frequencies for the Cl–C₆H₅CH₃, Cl–C₆H₅NH₂, and Cl–C₆H₅OH complexes were calculated out at the Møller–Plesset second-order (MP2) level of theory, using Dunning’s aug-cc-pVDZ basis set.²¹ This method has previously been demonstrated as suitable for describing the structures and energetics of hydrogen-bonded complexes and clusters containing halide anions.²²,²³ Calculations utilized the Gaussian 03 suite of programs.²⁴

Simulated infrared spectra of the complexes were generated by scaling the calculated harmonic frequencies using a factor determined from the experimental and calculated harmonic hydrogen stretch frequencies of the SBz monomer. Experimental vibrational frequencies for toluene, aniline, and phenol were taken from refs 25–27. The resulting scaling factors were 0.9503 for toluene, 0.9583 and 0.9482 for the NH and CH stretches of aniline, and 0.9602 and 0.9514 for the OH and CH stretches of phenol. To generate the simulated IR spectra, the calculated transitions were convoluted by Gaussians having widths of 6 cm⁻¹. Scaled vibrational frequencies, intensities, and assignments for the three complexes are shown in Table 1.

3. Results

3.1. Cl–C₆H₅CH₃, Theoretical Results. (a) Conformations and Dissociation Energy. A Cl– anion can interact with a toluene molecule either through aliphatic or aromatic hydrogen atoms, making it an interesting system to study the competition between different types of hydrogen bonding. Different conformations of the Cl–C₆H₅CH₃ complex, corresponding to stationary points on the potential energy surface (PES), were calculated at the MP2/aug-cc-pVDZ level of theory and are shown in Figure 1.

The global minimum energy structure (structure 1) has the Cl– attached to toluene by a double contact H-bond, through an ortho-hydrogen atom on the aromatic ring as well as a hydrogen atom on the methyl group. The dissociation energy corrected for vibrational zero-point energy (ZPE) is 45.4 kJ mol⁻¹; correction for basis set superposition error by the method of Boys and Bernardi²⁸ reduces the dissociation energy to 38.2 kJ mol⁻¹. Interestingly, it appears that the H-bond to the aromatic ring is stronger than the one to the methyl group. Evidence for this is that the distance between the ortho-H atom and the Cl– ion is predicted to be 2.504 Å, whereas the distance between the methyl group H atom and the Cl– ion is 2.628 Å. Furthermore, the H-bonded ortho-CH bond on the ring is lengthened by 0.5%, whereas the H-bonded CH on the methyl group is lengthened by only 0.2%. Double-contact H-bonds between halides and solvent molecules have also been observed for the I–HOCON complex, in which both the OH and CH groups are attached to the I–.²⁹ and in the Cl–C₆H₆, Br–C₆H₆, and I–C₆H₆ complexes where the halide anions are attached to two neighboring hydrogen atoms.¹⁰

There are several higher-energy stationary points for the Cl–C₆H₅CH₃ system; these are discussed in order of increasing energy. Starting with structure 1, rotation of the methyl group by 60° leads to structure 2, which is a first-order saddle point lying 3.2 kJ mol⁻¹ higher in energy. Structure 3, in which the Cl– anion is attached to the ortho- and meta-hydrogen atoms of the aromatic ring, is a stable minimum lying 8.2 kJ mol⁻¹ above structure 1. For this structure, rotation of the methyl group by 60° changes the energy by <0.1 kJ mol⁻¹. Structure 4, in which the Cl– is attached to the meta- and para-hydrogen atoms, is a stable geometry lying 10.4 kJ mol⁻¹ above structure 1. Structure 5, in which the Cl– is bound by a single H-bond to the para-hydrogen, lies 12.1 kJ mol⁻¹ above structure 1. Finally, for structure 6, the Cl– ion is attached to the methyl group by three equivalent hydrogen bonds. This configuration is a second-

### Table 1: Scaled Hydrogen Stretch Frequencies for Cl–C₆H₅CH₃, Cl–C₆H₅NH₂, and Cl–C₆H₅OH (in cm⁻¹) with Infrared Intensities (km mol⁻¹) Given in Brackets

<table>
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<tr>
<th></th>
<th>Cl–C₆H₅CH₃</th>
<th>Cl–C₆H₅NH₂</th>
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<td>νOH</td>
<td>νOH</td>
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<td>3005 (18)</td>
<td>2815 (2528)</td>
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</tr>
<tr>
<td>νCH₃CAM</td>
<td>3050 (43)</td>
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</tbody>
</table>

* For Cl–C₆H₅CH₃, superscripts al and ar denote localization of the vibration on the aliphatic or aromatic CH groups, respectively. Asymmetric and symmetric vibrations are indicated by a and s, respectively.

Figure 1. Low-energy conformations of Cl–C₆H₅CH₃ calculated at the MP2/aug-cc-pVDZ level. Relative energies (in kJ mol⁻¹) are given for each conformer.
order saddle point (with a doubly degenerate vibrational mode with imaginary frequency). The predicted infrared spectra associated with the different stable minima are discussed below in relation to the experimental spectrum.

(b) Rotation of the Methyl Group. One can anticipate that the methyl torsional potential for Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\) will differ profoundly from that of C\(_6\)H\(_5\)CH\(_3\) because the Cl\(^{-}\) anion is H-bonded to both the methyl group and the aromatic ring. To investigate the influence of the attached Cl\(^{-}\), we calculated the methyl torsional potentials for C\(_6\)H\(_5\)CH\(_3\) and Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\) at the MP2/aug-cc-pVDZ level in relaxed scans along the methyl torsional angle \(\phi\). The resulting torsional potential energy curves are shown in Figure 2. The potential energy curve for the toluene monomer exhibits a very small barrier for methyl internal rotation (13 cm\(^{-1}\)) and a parallel minimum energy configuration (\(\phi = 0^\circ\)). Experimentally, toluene is found to have a 4.9 cm\(^{-1}\) barrier for internal rotation and a perpendicular minimum energy configuration (\(\phi = 90^\circ\)).\(^{30,31}\) As the torsional potential curve for C\(_6\)H\(_5\)CH\(_3\) is extremely shallow and depends on rather subtle interactions, it is perhaps not surprising that the MP2/aug-cc-pVDZ calculations fail to predict the correct configuration. As expected, the torsional potential is significantly modified for Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\) because of the dual H-bonds between Cl\(^{-}\) and H atoms on the methyl group and the ring; the torsional barrier increases to 266 cm\(^{-1}\) and the hindering potential exhibits 3 equivalent minima.

Experimental Results. The photodissociation IR spectrum of Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\){\_}Ar is displayed in Figure 3, along with the scaled MP2/aug-cc-pVDZ vibrational frequencies and intensities for the four stable structures (structures 1 and 3\textendash}5 in Figure 1). The experimental spectrum is dominated by an intense broad feature centered near 2970 cm\(^{-1}\), with weaker features occurring above 3000 cm\(^{-1}\) and below 2940 cm\(^{-1}\). The Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\){\_}Ar spectrum differs significantly from the spectrum of the jet-cooled C\(_6\)H\(_5\)CH\(_3\) monomer recorded by Fuji et al.\(^{32}\) For Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\), the CH stretch band structure spans a range (2850\textendash}3075 cm\(^{-1}\)) that is slightly lower in frequency than for the toluene monomer (2870\textendash}3100 cm\(^{-1}\)). Most noticeably, the dominant feature in the Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\){\_}Ar spectrum occurs around 65 cm\(^{-1}\) lower in energy than the dominant peak in the toluene spectrum (3035 cm\(^{-1}\)). The differences are presumably due to the influence of the ionic H-bonds on the CH stretch vibrational frequencies.

Evidence for the structure of Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\) can be obtained by comparing the experimental and simulated infrared spectra in the CH stretch region. Inspection of Figure 3 shows that the

![Figure 2](image_url) Torsional potential for the methyl group in the Cl\(^{-}\)-C\(_6\)H\(_5\)-CH\(_3\) complex (●) and bare toluene (▲) calculated at the MP2/aug-cc-pVDZ level.

![Figure 3](image_url) (a) Experimental infrared spectrum of Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\){\_}Ar, and (b\textendash}e) simulated spectra based on calculations at the MP2/aug-cc-pVDZ level of theory for the 4 local minima shown in Figure 1. Trace (b) corresponds to the global minimum (structure 1), (c) to structure 3, (d) to structure 4, and (e) to structure 5. Scaled vibrational frequencies, intensities, and assignments for the global minimum structure are given in Table 1.

Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\) experimental spectrum bears a reasonable resemblance to the simulated spectrum associated with global minimum energy structure (structure 1 in Figure 1). In particular, the broad, intense central feature is reproduced in the simulated spectrum by a band that is the superposition of two transitions associated with the symmetric and asymmetric stretch vibration of the two H-bonded hydrogen atoms. The weaker features above 3000 cm\(^{-1}\) correspond to aromatic CH stretch vibrational modes, whereas features below 2940 cm\(^{-1}\) are attributable to the methyl CH stretch vibrational modes.

Aside from the broad central feature, it is difficult to make a direct correspondence between the peaks in the experimental and simulated spectrum for structure 1; the experimental spectrum contains 8\textendash}10 distinct bands, whereas the simulated spectrum exhibits eight bands. Additional bands may arise from Fermi resonances between the IR bright CH stretch states and near resonance overtone and combination levels associated with the in-plane bend modes; such resonances also play a significant role in the spectral structure of the toluene and benzene monomers.\(^{32,33}\)

The simulated spectra of the higher-energy Cl\(^{-}\)-C\(_6\)H\(_5\)CH\(_3\){\_}Ar structures (structures 3\textendash}5 in Figure 3) show less resemblance to the experimental spectrum; in each case, the intense central feature is absent from the simulated spectrum. Structures 3 and
4 would be expected to show a prominent methyl CH stretch band at 2885 cm\(^{-1}\), whereas structure 5 should be dominated by a band at 2935 cm\(^{-1}\) associated with stretching vibration of the H-bonded para-hydrogen atom. Although most of the ions probed experimentally probably have structure 1, it is possible that minor fractions of the population have the higher-energy structures 3–5. Therefore, it is conceivable that additional peaks in the experimental spectrum are associated with these less stable forms.

3.2. \(\text{Cl}^- - \text{C}_6\text{H}_5\text{NH}_2\). Theoretical Results. At the MP2/aug-cc-pVDZ level, \(\text{Cl}^- - \text{C}_6\text{H}_5\text{NH}_2\) is found to have the three stable minima shown in Figure 4. The lowest-energy structure features a near-linear H-bond between the \(\text{Cl}^-\) and one amine hydrogen (structure 1; bond angle of 173.2°), similar to the calculated structure of the \(\Gamma^- - \text{C}_6\text{H}_5\text{NH}_2\) complex.\(^{15}\) This conformer has a ZPE corrected dissociation energy \(D_0 = 75.6\) kJ mol\(^{-1}\), with BSSE correction reducing this to 68.5 kJ mol\(^{-1}\). Although the association enthalpy of \(\text{Cl}^- - \text{C}_6\text{H}_5\text{NH}_2\) has not been measured, the calculated \(D_0\) is bracketed by the experimental association enthalpies of \(\text{F}^- - \text{C}_6\text{H}_5\text{NH}_2\) (131.0 ± 8.4 kJ mol\(^{-1}\)) and \(\text{Br}^- - \text{C}_6\text{H}_5\text{NH}_2\) (61.1 ± 7.5 kJ mol\(^{-1}\)). There are two higher-energy structures. In the first (structure 2), which lies higher in energy than the global minimum by 44.1 kJ mol\(^{-1}\), the \(\text{Cl}^-\) is H-bonded to the meta- and para-hydrogens. In the second (structure 3), which lies an additional 2.1 kJ mol\(^{-1}\) higher in energy, the \(\text{Cl}^-\) is bound linearly to the para-hydrogen of the aromatic ring. Barriers for interconversion between structures 2 and 1 and between structures 3 and 2 are 0.5 kJ mol\(^{-1}\) and 2.1 kJ mol\(^{-1}\), respectively. Because of the much greater stability of structure 1 and low interconversion barriers it is unlikely that structures 2 or 3 would be formed in the ion source.

The calculated barrier for the umbrella inversion of the amine group in \(\text{Cl}^- - \text{C}_6\text{H}_5\text{NH}_2\) is 2.8 kJ mol\(^{-1}\), compared to 6.0 kJ mol\(^{-1}\) for bare aniline (experiment: 6.3 kJ mol\(^{-1}\)).\(^{35,36}\) The rather low barrier for umbrella inversion in the complex is unexpected given that MP2/aug-cc-pVTZ calculations predicted a larger inversion barrier for \(\text{Cl}^- - \text{NH}_2\) compared to bare \(\text{NH}_2\).\(^{37}\) A possible explanation for the low inversion barrier in \(\text{Cl}^- - \text{C}_6\text{H}_5\text{NH}_2\) is that the \(\text{Cl}^-\)–H cloud repulsion is minimized for the planar configuration and the opportunity for a weak secondary H-bond between the ortho-CH group and the \(\text{Cl}^-\) is maximized.

The barrier for transferring the \(\text{Cl}^-\) from one H to the other on the \(\text{NH}_2\) group is 17.2 kJ mol\(^{-1}\), considerably higher than the 6.4 kJ mol\(^{-1}\) barrier for \(\text{Cl}^- - \text{NH}_2\).\(^{37}\) This may be a consequence of the stronger intermolecular H-bond in the \(\text{Cl}^- - \text{C}_6\text{H}_5\text{NH}_2\) complex compared to \(\text{Cl}^- - \text{NH}_2\).

Experimental Results. The IR spectrum of the jet-cooled aniline monomer contains a series of aromatic CH stretch bands in the 3000–3120 cm\(^{-1}\) region and symmetric and asymmetric NH2 stretch bands at 3423 and 3509 cm\(^{-1}\), respectively.\(^{19}\) The aromatic CH stretch bands and the NH2 bands are of comparable intensity. In contrast, the IR spectrum for \(\text{Cl}^- - \text{C}_6\text{H}_5\text{NH}_2\), shown in Figure 5, is dominated by a strong peak at 2920 cm\(^{-1}\) that can be assigned to a red-shifted H-bonded NH stretch mode for structure 1 shown in Figure 4. The free NH stretch band appears as a small but unmistakable peak at 3475 cm\(^{-1}\), approximately midway between the symmetric and asymmetric NH stretch bands of the \(\text{C}_6\text{H}_5\text{NH}_2\) monomer (indicated by arrows in Figure 5). The red-shift for the H-bonded NH stretch band for \(\text{Cl}^- - \text{C}_6\text{H}_5\text{NH}_2\) with respect to the average of the symmetric and asymmetric NH2 stretch bands of the \(\text{C}_6\text{H}_5\text{NH}_2\) monomer (3466 cm\(^{-1}\)) is 546 cm\(^{-1}\).

Several much weaker bands appear in the aromatic CH stretch region (3000–3100 cm\(^{-1}\)). The feature centered at 2970 cm\(^{-1}\) does not correspond to any band in the simulated spectrum and may represent a combination or overtone band deriving intensity through Fermi resonance with the strongly red-shifted NH stretch mode.

Comparison of the experimental spectrum and the simulated spectrum for structure 1 suggests that the scaled harmonic frequency for the H-bonded NH stretch mode (3073 cm\(^{-1}\)) overestimates the experimental value (2920 cm\(^{-1}\)). Presumably, this is because the scaling does not adequately account for the extreme anharmonicity of the H-bonded NH stretch coordinate. This issue is addressed more thoroughly in the Discussion section.

3.3. \(\text{Cl}^- - \text{C}_6\text{H}_5\text{OH}\). Theoretical Results. The MP2/aug-cc-pVDZ calculations suggest that \(\text{Cl}^-\) prefers to bind to the hydroxy group of the \(\text{C}_6\text{H}_5\text{OH}\) molecule (structure 1, Figure 6). This structure has a ZPE-corrected dissociation energy of \(D_0 = 107.1\) kJ mol\(^{-1}\), with BSSE correction reducing this to 99.3 kJ mol\(^{-1}\). The calculated value compares reasonably well with the measured association enthalpy (109.0 ± 8.4 kJ mol\(^{-1}\)).\(^{15}\) The OH bond is elongated from 0.968 Å for bare phenol to 1.013 Å for the complex. This significant elongation, which can be viewed as reflecting incipient proton transfer to the \(\text{Cl}^-\) anion, is also reflected in a large red-shift of 841 cm\(^{-1}\) in the scaled harmonic stretch frequency. The \(\text{Cl}^-\)···H intermolecular bond length (1.969 Å) is significantly shorter than that for \(\text{Cl}^- - \text{C}_6\text{H}_5\text{CH}_3\) (2.628 and 2.504 Å) and \(\text{Cl}^- - \text{C}_6\text{H}_5\text{NH}_2\) (2.166 Å).
phenol monomer recorded by Yamada et al. contains an OH formed in the ion source. For interconversion, it is unlikely that structures 2-
- bands occurring in the 3000-
-1 region (Figure 7). As
-70.2
-(3)
-(4) 72.6

Figure 7. (a) Experimental infrared spectrum of Cl$^-$C$_6$H$_5$OH:Ar, and (b) a simulated spectrum based on calculations at the MP2/aug-
-cc-pVDZ level of theory. Scaled vibrational frequencies, intensities, and assignments are given in Table 1.

There are several higher-energy structures in which the anion is attached to two hydrogen atoms on the aromatic ring (structure 2-4, Figure 6). In all cases, both the hydroxy group and the anion lie in the plane of the ring, so that the complexes have C$_s$ symmetry. The barrier to interconversion between structures 2 and 1 is only 0.2 kJ mol$^{-1}$, that between Structures 3 and 2 is 1.4 kJ mol$^{-1}$, and that between structures 4 and 3 is 0.5 kJ mol$^{-1}$. Given the relative stability of structure 1 and low barriers for interconversion, it is unlikely that structures 2-4 were formed in the ion source.

4. Discussion

The calculations and infrared spectra described in the preceding section suggest that the Cl$^-$C$_6$H$_5$CH$_3$, Cl$^-$C$_6$H$_5$NH$_2$, and Cl$^-$C$_6$H$_5$OH complexes all possess structures in which the Cl$^-$ is hydrogen bonded to a single hydrogen atom on the substituent group. In the case of Cl$^-$C$_6$H$_5$CH$_3$, there is also a second, stronger H-bond with the ortho-hydrogen atom on the aromatic ring. Although for all three complexes, higher-lying isomers in which the Cl$^-$ anion is H-bonded to one or two hydrogen atoms on the phenyl ring are predicted, the barriers for conversion to the lowest-energy form are generally found to be small. For the lowest-energy form of all three Cl$^-$SBz complexes, the principal spectroscopic consequence of the hydrogen-bonding interaction is a red-shift and intensity increase for the CH, NH, and OH stretching modes. Unfortunately, straightforward comparisons between the experimental and simulated spectra are complicated by Fermi resonances between the H-bonded XH groups.

The Cl$^-$SBz complexes can be considered as intermediates for the proton-transfer reaction

$$\text{Cl}^- + \text{HA} \rightarrow \text{HCl} + \text{A}^-$$

Because the proton affinity (PA) of Cl$^-$ (14.46 eV$^{39}$) is less than the PAs of C$_6$H$_5$CH$_3^-$, C$_6$H$_5$NH$_2^-$, and C$_6$H$_5$O$^-$ (16.45, 15.89, and 15.16 eV, respectively$^{39}$), the complexes consist essentially of a Cl$^-$ anion attached to a perturbed SBz molecule. Nevertheless, there is a degree of proton transfer from the SBz to the chloride anion, the extent of which is determined by the difference between the proton affinities (APA) of the Cl$^-$ and the A$^-$ base. The degree of proton transfer is reflected in the magnitudes of the vibrational red-shift for the H-bonded XH stretch mode and bending overtones and combinations, particularly for Cl$^-$C$_6$H$_5$CH$_3$ and Cl$^-$C$_6$H$_5$OH. Nevertheless, in all three cases, there is clearly a systematic red-shift and intensity increase in vibrational modes associated with the H-bonded XH groups.

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For Cl levels are superimposed on the potential curves in Figure 8. The effective mass of the H-bonded hydrogen atom. The effective mass of the Cl–C6H5NH2, C6H5OH, and Cl–C6H5OH. In each case, the lowest three energy levels and associated wave functions are shown.

A likely source of the discrepancy is anharmonicity in the H-bonded XH stretch coordinate not fully accounted for by frequency scaling. To investigate this possibility, we calculated one-dimensional potential energy curves (at the MP2/aug-cc-pVDZ level) for the H-bonded XH stretch motion (with remaining coordinates fixed). Plots for both the free SBz molecule and the corresponding Cl–SBz complex appear alongside one another in Figure 8, highlighting the influence of the attached Cl–ion. The distortion of the potential energy curves for Cl–C6H5NH2 and Cl–C6H5OH is quite evident, illustrating the marked delocalization of the shared proton.

Frequencies for the hydrogen stretch vibrations associated with the one-dimensional potential curves were calculated using Level 7.4,41 assuming that the vibration involved only motion of the H-bonded hydrogen atom. The effective mass of the oscillator was assumed to be 1 amu. The first three vibrational levels are superimposed on the potential curves in Figure 8. For Cl–C6H5NH2, the NH stretch transition is calculated to lie at 2845 cm⁻¹, comparing favorably with the experimental value (2920 cm⁻¹), and certainly better than the scaled harmonic prediction, which overestimates the experimental frequency by 150 cm⁻¹. For Cl–C6H5OH, the OH stretch frequency is calculated to lie at 2400 cm⁻¹, around 415 cm⁻¹ below the scaled harmonic frequency, and indeed below the series of bands apparent in the experimental spectrum between 2600 and 2800 cm⁻¹. This raises the possibility that the OH stretch band occurs below 2550 cm⁻¹, favoring the Cl–NH2...Ar, Alberti et al. found that the most favorable bond (D0 = 523 cm⁻¹; ref 45) is comparable in strength with Ar...aromatic bonds (for example, D0 = 314 ± 7 cm⁻¹ for C6H5•Ar; ref 46), the Ar atom is positioned so as to maximize the interaction with both the Cl– and the SBz subunit. For Cl–C6H5•Ar, Alberti et al. found that the most favorable location for the Ar atom is midway between the benzene ring and the Cl–, so that it is stabilized by interactions with both the anion and the Bz molecule. The situation is probably similar for the Cl–SBz complexes.

Figure 8. One-dimensional NH/OH stretch potential energy curves (calculated at the MP2/aug-cc-pVQZ level) for (a) C6H7NH3, (b) Cl–C6H7NH2, (c) C6H7OH, and (d) Cl–C6H5OH. In each case, the lowest three energy levels and associated wave functions are shown.

Figure 9. Correlation between (a) experimental vibrational redshift and (b) calculated bond elongation (at MP2/aug-cc-pVQZ level) and the difference in proton affinities (ΔPA) of Cl– and conjugate base Cl–. Filled triangles (▲) represent Cl–SBz complexes, whereas filled squares (■) represent complexes of Cl– and second-row hydride molecules.

The Cl– anion can be expected to be larger for the SBz molecules, because the PAs for C6H7CH2–, C6H7NH–, and C6H5O–(16.45, 15.89, and 15.16 eV) are lower than those of CH3–, NH2–, and OH– (18.13, 17.49, and 16.92 eV). This is reflected in the magnitudes of the vibrational red-shifts and the elongation of the XH bond (calculated at the MP2/aug-cc-pVQZ level), both of which are plotted against ΔPA in Figure 9. Note that for Cl–C6H5CH3, it is difficult to ascertain the magnitude of the red-shift because of the complexity of the spectrum and because both the aromatic and aliphatic CH stretch modes are affected by H-bonding. It is obvious from Figure 9 that the Cl–SBz complexes are associated with larger red-shifts and XH bond elongations than the complexes containing the corresponding second-row hydride molecule. For both sets of complexes, there is a near-linear correlation between ΔPA and the vibrational red-shift and the XH bond elongation.

Last, we consider the effect of the attached Ar atom. Although in each case the experimental spectrum does not provide direct information on the position of the Ar atom, presumably because the Ar...Cl– bond (D0 = 523 cm⁻¹; ref 45) is comparable in strength with Ar...aromatic bonds (for example, D0 = 314 ± 7 cm⁻¹ for C6H5•Ar; ref 46), the Ar atom is positioned so as to maximize the interaction with both the Cl– and the SBz subunit. For Cl–C6H5•Ar, Alberti et al. found that the most favorable location for the Ar atom is midway between the benzene ring and the Cl–, so that it is stabilized by interactions with both the anion and the Bz molecule. The situation is probably similar for the Cl–SBz complexes.

5. Summary

Infrared spectra have been recorded for Cl–C6H7CH2•Ar, Cl–C6H7NH2•Ar, and Cl–C6H5OH•Ar using vibrational predissociation spectroscopy. Accompanying ab initio calculations suggest that the lowest-energy conformers for Cl–C6H7NH2 and Cl–C6H5OH have the Cl– bound to a single hydrogen atom of the substituent group. In the case of Cl–C6H5CH3, the Cl– is hydrogen bonded to an ortho-hydrogen atom on the ring and to a hydrogen atom on the methyl group. Complexities in the infrared spectra are postulated to arise from Fermi resonances between the H-bonded XH stretch and overtone and

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(see Figures 5 and 7). One might expect that there should be similarities between the Cl–SBz complexes (Cl–C6H7CH2–, Cl–C6H7NH–, and Cl–C6H5O–(16.45, 15.89, and 15.16 eV) are lower than those of CH3–, NH2–, and OH– (18.13, 17.49, and 16.92 eV). This is reflected in the magnitudes of the vibrational red-shifts and the elongation of the XH bond (calculated at the MP2/aug-cc-pVQZ level), both of which are plotted against ΔPA in Figure 9. Note that for Cl–C6H5CH3, it is difficult to ascertain the magnitude of the red-shift because of the complexity of the spectrum and because both the aromatic and aliphatic CH stretch modes are affected by H-bonding. It is obvious from Figure 9 that the Cl–SBz complexes are associated with larger red-shifts and XH bond elongations than the complexes containing the corresponding second-row hydride molecule. For both sets of complexes, there is a near-linear correlation between ΔPA and the vibrational red-shift and the XH bond elongation.

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combination levels. There is a clear correlation between the proton affinity of the aromatic molecule’s conjugate base and both the vibrational red-shift and the elongation of the H-bonded XH group (Cl$^-$–C$_6$H$_5$CH$_3$ < Cl$^-$–C$_6$H$_5$NH$_2$ < Cl$^-$–C$_6$H$_5$OH).

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References and Notes


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