Theoretical study of H$_2$O–HF and H$_2$O–HCl: Comparison with experiment

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Theoretical study of $\text{H}_2\text{O}–\text{HF}$ and $\text{H}_2\text{O}–\text{HCl}$: Comparison with experiment

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(Received 14 June 1984; accepted 16 July 1984)

The $\text{H}$ bonds in $\text{H}_2\text{O}–\text{HF}$ and $\text{H}_2\text{O}–\text{HCl}$ are studied and compared using $\textit{ab initio}$ molecular orbital methods and the results compared to experimental data. Basis sets used are: (i) triple valence 6-311G** and (ii) double $\xi$** with two sets of polarization functions. Electron correlation, included via second- and third-order Møller–Plesset perturbation theory, is found to have profound effects on both systems, particularly $\text{H}_2\text{O}–\text{HCl}$. Both $\text{H}$ bonds are strengthened substantially with a concomitant reduction in length. $\text{H}$-bond energies and geometries calculated at correlated levels are in excellent accord with available experimental information. In both systems, all levels of theory indicate the equilibrium geometry contains a pyramidal arrangement about the oxygen atom. However, the difference in energy between this structure and a $C_{2v}$ planar arrangement is found to be small enough that consideration of probability amplitudes in the ground vibrational level leads to nearly equal likelihood of observing either geometry. Agreement between experimental vibrational frequencies in $\text{H}_2\text{O}–\text{HF}$ and those calculated at correlated levels and involving quadratic, cubic, and quartic force constants is quite good. An explanation is offered for the increase in $\text{HX}$ bond length which occurs at SCF and correlated levels upon $\text{H}$-bond formation based upon nearly linear relationships between this length and subunit dipole moment and polarizability on the other. The dispersion energy is found to be a very sensitive, almost exactly linear function of the increase of $\text{H}–\text{X}$ bond length. This energy contributes substantially to the weakening of the $\text{HX}$ bond upon complexation.

INTRODUCTION

$\text{H}$ bonds of medium strength have long been a center of attention of spectroscopists and theoreticians alike. The relative simplicity of the equilibria in the gas phase was one factor in making the $\text{O}–\text{HX}$ bond in, e.g., ether–$\text{HX}$ and $\text{H}_2\text{O}–\text{HF}$, among the first complexes studied by gas-phase vibrational spectroscopy. It was in these systems that the band broadening and fine structure of the $\nu_v(\text{HX})$ stretching frequency were first observed. With refinement of experimental technique has come an accelerated interest in $\text{H}$-bonded systems. In 1975, Thomas was able to identify all of the low frequencies associated with $\text{H}$-bond deformations in $\text{H}_2\text{O}–\text{HF}$ and arrived at an estimate of the potential well depth. More recently, a series of papers has been published dealing with microwave studies of the same complex. By measurement of the properties of $\text{H}_2\text{O}–\text{HF}$ in vibrationally excited states, it has been possible to reconstruct the molecular geometry with unprecedented accuracy. Additionally, examination of the complex in a solid Ar matrix by Fourier-transform IR spectroscopy enabled Andrews and Johnson to provide evidence for an inversion motion. Matrix isolation techniques have also been used by Ault and Pimentel to identify the $\text{H}_2\text{O}–\text{HCl}$ complex. More recent pulsed-nozzle Fourier-transform microwave work by Legon and Willoughby led to detection of this complex in the gas phase as well. The latter authors concluded from their measurements that the vibrationally averaged structure of $\text{H}_2\text{O}–\text{HCl}$ is of $C_{2v}$ symmetry with a planar arrangement about the $\text{O}$ atom.

Complexes of the type $\text{H}_2\text{O}–\text{HX} (X = \text{F, Cl})$ have provided fertile ground for theoretical studies as well. For example, several explanations have been offered for the band profile of the $\nu_v(\text{HX})$ stretching frequency. While one hypothesis draws connections with predissociation, recent calculations by Bouteiller and Guissani have demonstrated the importance of anharmonic coupling between the low frequency $\nu(\text{O}–\text{F})$ and high frequency $\nu(\text{HX})$ stretching modes. SCF calculations were used by Lister and Palmieri to assign the total set of harmonic frequencies in $\text{H}_2\text{O}–\text{HF}$. Although these assignments were quite useful, their use of a relatively small basis set resulted in an incorrect prediction of the geometry of the complex. Due to inclusion of polarization functions, more recent calculations by Bouteiller, Allavena, and Leclercq were able to successfully predict a $C_5$ equilibrium geometry for the complex containing a pyramidal arrangement about the oxygen atom. These same investigators made an attempt to incorporate the effects of electron correlation upon the force constants within the complex but due to a somewhat limited list of configurations, found significant discrepancies between theoretical and experimental frequencies. Moreover, no attempt was made to study the influence of electron correlation upon the equilibrium structure of this complex or its contributions to the interaction energy. With regard to $\text{H}_2\text{O}–\text{HCl}$, Alagona et al. have provided some structural data with a medium-quality basis set at the SCF level; there is no information available concerning the role of electron correlation in this system.

Nevertheless, recent studies have underscored the importance of electron correlation to a true picture of $\text{H}$ bond-
For example, the variation–perturbation treatment of the water dimer by Jeziorski and van Hemert demonstrated that the contribution of dispersion to the total H-bond energy is over 1/3. In their study of the interaction between HF and CO, Benzol and Dykstra found that it is not possible to properly describe the relative stabilities of CO–HF and OC–HF without explicit account of electron correlation. Recent work in this laboratory has shown that dispersion plays an even more important role when second-row atoms such as Cl are involved.

For these reasons, a major goal of the present paper is an examination of the H$_2$O–HF and H$_2$O–HCl systems which incorporates electron correlation into the procedure. The structure and properties of these complexes are studied with special emphasis placed on the effects of correlation on the interactions and shape of potential energy surfaces. As such, we perform for the first time a detailed comparison of the proton-donor properties of HF and HCl at the correlated level of theory.

While calculations that include electron correlation are generally quite time consuming and demanding of computer resources, the recent implementation of Möller–Plesset perturbation theory into molecular orbital programs by Pople et al. represents a significant step forward. It has been recently demonstrated that second-order MP theory, applied within the supermolecule framework, can yield asymptotically the dispersion energy in a well-defined manner at the uncorrelated Hartree–Fock level. Our previous results indicate great usefulness of the MP2 method in the treatment of selected H bonds. However, this procedure has not been sufficiently tested at this point to assure that truncation of the perturbation expansion after the second-order term can produce data which are comparable with experiment. A second aim of this work is therefore a stringent test of the MP2 method; recently obtained high-quality experimental results for H$_2$O–HF and H$_2$O–HCl provide an exceptional opportunity for careful comparison with theoretical calculations.

**METHODS**

Electron correlation was included via Möller–Plesset perturbation theory to second (MP2) and third (MP3) orders, as implemented in the GAUSSIAN-80 package of computer codes. Previous experience has shown that MP3 treatment of H-bonded systems with the 6-311G* basis set can closely reproduce the first term, C$_r$R$^{-9}$, of the multiple-expanded dispersion energy based on the nonempirical Unsöld approximation and we accordingly perform some of our calculations with this basis set, denoted herein as A. On the other hand, a more accurate treatment of dispersion at the MP2 level requires at least two sets of polarization functions. Our doubly polarized basis set B was constructed as follows. The double-zeta [432/21] basis set, used previously by Jeziorski and van Hemert and by us, was applied to the H$_2$O molecule; the d-orbital exponents used were 0.4 and 1.5. For HF, the 6-31G basis set was augmented by two sets of d functions on F (exponents 0.25 and 1.0) and two sets of p functions on H (0.15 and 0.75). The latter exponents were chosen according to the suggestion of van Duijneveldt who obtained good dipole polarizabilities using them. The B basis set for Cl consists of the 6-6-31G set, supplemented by d functions with exponents 0.25 and 0.75, taken from van Duijneveldt.

**RESULTS**

**Structure and vibrations of H$_2$O–HF**

The structural parameters of the H$_2$O–HX complexes are illustrated in Fig. 1 where R is defined as the distance between O and X atoms. β measures the deviation of the H atom from the O–X axis and α is the angle between this axis and the HOH bisector. Geometry optimizations of the H$_2$O–HF complex were carried out with basis set A and the results are presented in Table I. In these optimizations, the internal geometry of the H$_2$O molecule was held fixed in its experimentally determined structure; r(OH) = 0.957 Å and θ(HOH) = 104.5°. Gradient procedures were used to optimize the structure at the SCF level and stepwise procedures at the correlated levels.

We may see from the results at SCF as well as at correlated levels, that the optimized structure is of C$_s$ symmetry, with a pyramidal oxygen atom, in agreement with experimental findings as well as previous calculations with polarized basis sets. Slight nonlinearities of the H bond are noted in that the equilibrium values of β are somewhat greater than zero. The intermolecular separations R are smaller at correlated levels than the SCF value, with the MP2 distance slightly smaller than MP3. A similar pattern is noted for α, the pyramidalization angle of oxygen. Table I indicates that the MP3 procedure, in conjunction with basis set A, is capable of closely reproducing experimental data, listed in the last column of the table. The effects of correlation upon the SCF structure are as follows. The intermolecular distance is decreased, concomitant with an increase in the internal HF bond length and a trend toward a more perpendicular arrangement of the H$_2$O molecule; i.e., reduction of α. This tendency toward smaller α may be explained on elec-

![FIG. 1, C geometrical parameters for H$_2$O–HX (X = F, Cl). α measures the angle between the O–X axis and the OH$_2$ bisector.](image-url)
trostatic grounds as follows. The attraction between the dipole moments of the two molecules would favor large values of $\alpha$ which more closely align the moments. Correlation effects have been demonstrated previously to reduce the dipole moment of water and would thereby diminish the pull toward high $\alpha$. In this same regard, it is likely that the earlier predictions of a planar $O$ atom by unpolarized basis sets were due to their well known exaggeration of dipole moments.

The exact degree of pyramidalization of the oxygen atom is a product of a delicate balance between a number of opposing forces. Whereas dipole-dipole interactions favor the planar arrangement, a more perpendicular configuration arises from consideration of dipole-quadrupole terms. In addition to electrostatic effects, other forces such as exchange, polarization, charge transfer, and dispersion each have a different angular dependence. The net result is a very small energy difference between the planar and pyramidal geometries. The potential energy curve for the bending of the water molecule at the SCF level is shown in Fig. 2. This curve was computed by holding fixed the geometry of the complex at the values indicated in Table I and varying $\alpha$ between 105° and 255° ($\beta$ was set equal to 0° to assure symmetry of the curve). The symmetric potential contains two equivalent minima, corresponding to a pyramidal oxygen, separated by a configuration ($\alpha = 180^\circ$) containing a planar $O$ atom. Figures 3 and 4 illustrate the analogous potentials computed at the MP2 and MP3 levels (also using the SCF geometrical parameters and the A basis set). Superimposed on each potential energy curve is a series of vibrational levels obtained from the potential by the method of Somorjai and Hornig for evaluation of anharmonic frequencies in a double minimum potential.

FIG. 2. Total SCF/A energy of $H_2O-HF$ as a function of $\alpha$. Vibrational energy levels and associated wave functions are superposed.

where they are compared with experimental data. The first row contains the energy barrier to the inversion between the two symmetric minima. It is clear that, whereas the SCF barrier is quite small, the values obtained at correlated levels are several times larger and in much better accord with the experimental estimate. Also included in Table II are the energy differences between the ground vibrational level and the first two excited levels. The SCF treatment greatly overesti-

![FIG. 3. MP2/A potential for inversion of water in $H_2O-HF$.](image)

![FIG. 4. MP3/A energy of $H_2O-HF$ as a function of $\alpha$.](image)
bending of the water molecule is quite likely coupled to other
in isolation from the other geometrical parameters. The
important, perhaps, is the treatment of the bending potential

tions of basis set and finite perturbation expansion. More
ference.

TABLE II. Height of barrier and vibrational energy level differences* in
H2O-HF for the low-frequency in-plane bending mode corresponding to
water inversion.

<table>
<thead>
<tr>
<th></th>
<th>SCF</th>
<th>MP2</th>
<th>MP3</th>
<th>Experiment</th>
</tr>
</thead>
<tbody>
<tr>
<td>E ¹</td>
<td>45</td>
<td>172</td>
<td>144</td>
<td>126</td>
</tr>
<tr>
<td>v = 1→ 0</td>
<td>134</td>
<td>85</td>
<td>96</td>
<td>64 ± 10</td>
</tr>
<tr>
<td>v = 2→ 0</td>
<td>257</td>
<td>251</td>
<td>255</td>
<td>267 ± 35</td>
</tr>
</tbody>
</table>

*All entries in cm⁻¹; calculated with basis set A.


dicates the spacing between the ground and first excited level;
MP values are again closer to the experimental information.
It is clear from the last row in Table II that the energy differ­
ence between the ground and second excited state is rather
insensitive to inclusion of electron correlation. The overall

A fundamentally important point concerns the corre­
respondence between the theoretical and experimental equilib­
rium geometries. By definition, the equilibrium structure refers to the absolute minimum in the potential energy surface of the molecule. Whereas it is possible in principle to locate this minimum by theoretical methods, the ground state vibrational motions may sometimes obscure the experimental elucidation of the minimum. In the SCF potential of Fig. 2, the ground vibrational level lies slightly higher in energy than the planar structure separating the two minima. The vibrational wave function, superposed on this energy level has its maximum amplitude for the planar structure with α = 180°, leading to a likelihood of observing this structure despite the fact that it is not the minimum of the potential. However, the flatness of the function indicates that the probability of observing pyramidal arrangements with α deviating from 180° by up to perhaps 30° is not much less than the planar geometry. The situation is somewhat different in the MP potentials depicted in Figs. 3 and 4 where the ground vibrational level lies below the barrier. Nonetheless, the wave functions are quite similar to the SCF function (despite the presence of a very shallow minimum at α = 180°) and the geometry is not strongly localized in the pyramidal configuration. In a dynamical sense, then, the SCF situation in Fig. 2 where the lowest vibrational level lies above the symmetric structure is virtually indistinguishable from the MP cases depicted in Figs. 3 and 4 where the opposite is true. We conclude that accommodation of a ground vibrational level below an energy barrier is not sufficient to guarantee a dyna­
ic distinction from a case where the level cannot be so placed; disparities arise only as the barrier height is further increased.

While the MP3 data compare fairly well with the experi­
mental information in Table II, there remains some differ­
ence. Part of this discrepancy is probably due to limita­
tions of basis set and finite perturbation expansion. More
important, perhaps, is the treatment of the bending potential in isolation from the other geometrical parameters. The bending of the water molecule is quite likely coupled to other

motions such as an intermolecular stretch or bending involving the H-bonding proton of HF. Treatment of the mode as a pure HOH bend was adopted here to match as closely as possible the one-dimensional model used by Kisiel, Legon, and Millen in their reconstruction of the potential.⁶

The preceding discussions have included a description of the anharmonicity of the bending of the water molecule relative to the HF subunit. The anomalous behavior of the band profile of the υ1 (HF) stretching frequency has also been explained on the basis of anharmonic effects, specifically as a coupling between the υ(O–F) and υ(FH) stretching modes.¹¹

In order to calculate anharmonic stretching frequencies, the energy was calculated for a two-dimensional grid of points in the vicinity of the equilibrium geometry. These energies E(r, R) were evaluated at both the SCF and MP2 levels in order to ascertain the effects of electron correlation upon the results. Except for r(HF) and R (O–F), all geometrical parameters were held fixed in the values optimized previously with basis set A. Table III contains the quadratic, cubic, and quartic force constants evaluated by numerical analysis of the computed potential energy surfaces. Comparison of the SCF and MP2 constants reveals the effects of electron correlation upon these constants. For the most part, these changes are qualitatively similar to the trends found previously by Bouteiller et al.¹⁴ There are, however, some large quantita­tive and even qualitative disparities in the results which have some bearing on the next point to be discussed concerning vibrational frequencies.

The procedure described in Ref. 28 for extracting the vibrational frequencies from all force constants up through fourth order was applied to the data in Table III and the

TABLE III. Quadratic, cubic, and quartic force constants for H2O-HF [in mdyn/Å²], calculated with basis set A.

<table>
<thead>
<tr>
<th></th>
<th>SCF</th>
<th>MP2</th>
</tr>
</thead>
<tbody>
<tr>
<td>F</td>
<td>9.967</td>
<td>8.497</td>
</tr>
<tr>
<td>r²</td>
<td>0.123</td>
<td>1.128</td>
</tr>
<tr>
<td>rR</td>
<td>0.228</td>
<td>0.335</td>
</tr>
<tr>
<td>r²</td>
<td>-78.15</td>
<td>-68.96</td>
</tr>
<tr>
<td>rR</td>
<td>2.433</td>
<td>2.264</td>
</tr>
<tr>
<td>r²</td>
<td>0.317</td>
<td>0.995</td>
</tr>
<tr>
<td>R³</td>
<td>-1.013</td>
<td>-1.789</td>
</tr>
<tr>
<td>r²</td>
<td>4.835</td>
<td>4.885</td>
</tr>
<tr>
<td>r²</td>
<td>25.14</td>
<td>24.486</td>
</tr>
<tr>
<td>r²</td>
<td>-24.94</td>
<td>-10.01</td>
</tr>
<tr>
<td>R³</td>
<td>-1.783</td>
<td>-9.354</td>
</tr>
<tr>
<td>r²</td>
<td>6.600</td>
<td>8.194</td>
</tr>
</tbody>
</table>

TABLE IV. Stretching frequencies in H2O-HF (in cm⁻¹).

<table>
<thead>
<tr>
<th></th>
<th>SCF</th>
<th>MP2</th>
<th>SCF</th>
<th>SCF</th>
<th>SCF</th>
<th>SCF</th>
<th>Expt</th>
</tr>
</thead>
<tbody>
<tr>
<td>v(FH)</td>
<td>4013</td>
<td>3764</td>
<td>3824</td>
<td>4051</td>
<td>4031</td>
<td>3946</td>
<td>3608</td>
</tr>
<tr>
<td>v(O–F)</td>
<td>204</td>
<td>236</td>
<td>198</td>
<td>222</td>
<td>242</td>
<td>242</td>
<td>198</td>
</tr>
</tbody>
</table>

*Reference 4.
Reference 29.
TABLE V. Calculated geometries and interaction energies.

<table>
<thead>
<tr>
<th></th>
<th>H₂O-HF</th>
<th>H₂O-HCl</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>SCF</td>
<td>B</td>
</tr>
<tr>
<td></td>
<td>MP2</td>
<td></td>
</tr>
<tr>
<td>R (O=O), Å</td>
<td>2.70</td>
<td>2.65*</td>
</tr>
<tr>
<td>r, Å</td>
<td>0.907</td>
<td>0.921*</td>
</tr>
<tr>
<td>∆r, Å</td>
<td>0.011</td>
<td>0.012*</td>
</tr>
<tr>
<td>∆E\text{SCF}, kcal/mol</td>
<td>-9.38</td>
<td>-9.19</td>
</tr>
<tr>
<td>∆E\text{MP2}, kcal/mol</td>
<td>-10.51</td>
<td>-11.03b</td>
</tr>
</tbody>
</table>

*Geometry optimized with MP3.

b MP3 value is -10.49.

results are presented in Table IV. It may be seen that the application of MP2 to the A basis set improves the agreement of \( \nu(HF) \) with the experimental value markedly. In fact, the MP2/A result is the best calculated value to date, much superior to the previous estimates with the 6-31G* basis set. In the case of the lower frequency vibration \( \nu(O=O) \), all theoretical data lead to overestimates of the experimental data, probably associated with the exaggerations of the strength of the H bond. To test the effects of further extension of the basis set calculations, were carried out with basis set C which contains the same atomic orbitals as does B for the H₂O subunit but the description of HF is improved by use of a [742/31] basis set suggested previously by Lischka.¹⁰

As may be seen in the appropriate column of Table IV, the larger basis set decreases both stretching frequencies. Using the comparison between MP2 and SCF/A results as a guide, it is expected that inclusion of correlation to the SCF/C data would lead to excellent reproduction of experimental frequencies.

H-bond energies

Although the MP3 treatment with the singly polarized A basis set has been seen above to adequately reproduce experimental information concerning the equilibrium geometries and vibrational frequencies, this approach is somewhat less satisfactory for study of the energy of interaction between the subsystems. Table V contains geometries optimized with each basis set at both SCF and correlated levels. The fourth row lists the interaction energies computed at the SCF level for each geometry; analogous MP2 energies are contained in the last row.

Since the SCF and MP geometries are significantly different, there is some ambiguity in assigning a value to the increase in H-bond energy associated with correlation. Let us consider, for example, basis set A calculations of the H₂O--HF system. As may be seen in the first column of Table V, MP2 treatment of the geometry optimized at the SCF level raises the interaction energy from -9.38 kcal/mol to -10.51, an increase of -1.13. However, the SCF geometry does not correspond to the bottom of the MP potential and consequently the above procedure does not indicate the full magnitude of the correlation effect. Optimization of the geometry at the MP level further increases the interaction energy to -11.03 kcal/mol.

From a computational point of view, the full effect of correlation is obtained by comparison of complete correlation treatment [\( \Delta E_{\text{MP}} \) / SCF geometry] with the SCF results [\( \Delta E_{\text{SCF}} \) / SCF geometry]. On the other hand, for the purpose of studying radial and angular dependence of the dispersion component, \( \Delta E_{\text{MP}} \) and \( \Delta E_{\text{SCF}} \) must refer to the same geometry. For this reason, the dispersion contribution to the H-bond energy will be referred to below as the difference between \( \Delta E_{\text{MP}} \) and \( \Delta E_{\text{SCF}} \), both evaluated with the MP geometry. For the systems being examined here, the MP2 interaction energies computed with the SCF and MP geometries differ by about 0.5 kcal/mol. Thus, one would underestimate the complexation energies by this amount if MP2 were simply applied to a geometry optimized at the SCF level. Coupling this fact with the difference in SCF interaction energies between the two geometries would lead to a more severe underestimate of the contribution of dispersion to the stability of each complex. For example, using SCF geometries for H₂O--HF with basis set B, correlation contributes 1.39 kcal/mol (9.15 - 7.76), or 15% to the total interaction energy, whereas the corresponding contribution with MP2 geometries is 2.58 kcal/mol or 27%.

We expect our MP2/B estimate of the interaction energy in H₂O--HF to be fairly reliable. In addition to its good representation of electrostatic attraction (see Table VI for calculated values of subsystem dipole moments) and adequate framework for correlation and induction, the superposition error is rather small. Evaluation of this error by the counterpoise procedure yields a value of 0.7 kcal/mol at the SCF level. Subtraction of this quantity from the MP2/B interaction energy in Table V provides our best theoretical estimate of -8.9 kcal/mol. Cancellation is expected between small additional corrections that might be added. For example, the absence of orbitals, needed for good representation of quadrupole polarizabilities [and \( R^{-10} \) term of dispersion energy] will probably lead to a slight understi-

TABLE VI. SCF and experimental dipole moments (\( \mu \)).

<table>
<thead>
<tr>
<th></th>
<th>A</th>
<th>B</th>
<th>Experiment</th>
</tr>
</thead>
<tbody>
<tr>
<td>H₂O</td>
<td>2.17</td>
<td>2.06</td>
<td>1.85*</td>
</tr>
<tr>
<td>HF</td>
<td>2.00</td>
<td>1.86</td>
<td>1.83*</td>
</tr>
<tr>
<td>HCl</td>
<td>...</td>
<td>1.18</td>
<td>1.09*</td>
</tr>
</tbody>
</table>

* Reference 31.

References 32.

Reference 33.
mate. On the other hand, the reverse trend may be expected from truncation of the MP expansion at second order. Although no calculations have been performed to date which incorporate the full fourth order of perturbation theory for H bonds, polarizability calculations suggest that considerable cancellation occurs between the third and full fourth orders.\(^{34}\)

The enthalpy of formation of the H\(_2\)O–HF complex has been measured in the gas phase\(^4\) and is equal to \(-6.2 \pm 1\) kcal/mol at 298 K. Before comparison of our theoretical estimate of the electronic energy with this experimental quantity, it is first necessary to make adjustments for changes in translational, rotational, and vibrational energy, as well as addition of a \(\Delta PV\) term. All the corrections, exclusive of the vibrational term, are straightforward to calculate from standard thermodynamical formulas and amount to \(-2.4\) kcal/mol. Evaluation of the vibrational correction requires knowledge of all the frequencies in both the complex and the isolated subsystems. Much of this information is provided by Lister and Palmieri\(^{12}\) who provide experimental data where available and supplement the remaining frequencies with calculated quantities. Using the frequencies supplied by these authors, we arrive at a vibrational energy difference between the complex and subsystems of 4.4 kcal/mol at 298 K. Combining all the adjustments together with our computed \(\Delta E^*\) of \(-8.9\) kcal/mol leads to a value of \(-6.9\) kcal/mol for \(\Delta H^\circ\) which falls well within the range of experimental uncertainty.

**Comparison of H\(_2\)O–HCl with H\(_2\)O–HF**

The complex of H\(_2\)O with HCl was studied using the doubly polarized B basis set only. Like H\(_2\)O–HF, the optimized geometry of this complex belongs to the C\(_2v\) point group. The binding potential for the OH\(_2\) molecule at both the SCF and MP levels is illustrated in Fig. 5. As in the previous case of H\(_2\)O–HF, correlation substantially raises the barrier for this motion and shifts the equilibrium value of \(\alpha\) away from the planar configuration. The O–H–Cl bridge is slightly nonlinear with \(\beta\) equal to 2.8° at the SCF level and 0.9° with MP2. The optimized values of \(R\) (O–Cl) and \(r\) (HCl) are contained in Table V where is may be seen that the MP2/B H–bond length of 3.19 Å is rather close to the experimental estimate of 3.21 Å.\(^9\)

Although experimental data\(^9\) indicate a planar (C\(_2v\)) geometry for the H\(_2\)O–HCl complex, the large amplitude motion of the H\(_2\)O unit and associated vibrational averaging makes it difficult to distinguish between this geometry and the C\(_3\) pyramidal arrangement. As may be seen in Fig. 5, the calculated barrier for inversion between the two equivalent C\(_7\) geometries is 116 cm\(^{-1}\) at the MP2/B level. Comparison with the data for H\(_2\)O–HF in Figs. 2–4 makes it clear that as in the previous case, the ground vibrational energy will occur close to the top of the barrier and consequently the wave function will have little difference in probability amplitude between the planar and pyramidal geometries. The observation of a C\(_2v\) structure would therefore be consistent with a double well potential with low barrier as predicted by the calculations.

In their attempt to reconstruct the molecular geometry of H\(_2\)O–HCl from measured rotational constants, Legon and Willoughby\(^9\) suggested that the internal HOH angle increases by 4° upon complexation. We checked this hypothesis theoretically at the MP2 level. It was found that while some increase in this angle was observed, the magnitude of this increase was only 0.5°.

The energetic data in Table V indicate that the H–bond in H\(_2\)O–HCl is somewhat weaker than in H\(_2\)O–HF. Comparing the results within the framework of the B basis set, the O–H–F bond is stronger than O–H–Cl by 3.6 kcal/mol at the SCF level and 3.0 at MP2. Another important distinction concerns the contribution of correlation to the stability of each bond. This contribution is somewhat higher in H\(_2\)O–HCl (2.4 vs 1.9 kcal/mol). If one is considering the relative contributions of correlation, this difference is even more striking. Whereas correlation amount to 20% of the total interaction energy in H\(_2\)O–HF, this term increases to 37% in H\(_2\)O–HCl. Using the previous definition of dispersion as the difference between \(\Delta E^{\text{MP2}}\) and \(\Delta E^{\text{SCF}}\) with MP geometry, the percentage contribution of dispersion is 27% in H\(_2\)O–HF and 42% in H\(_2\)O–HCl. Another indication of the relative importance of correlation in the two systems comes from a comparison of equilibrium H-bond lengths. The distance between oxygen and fluorine is diminished by 0.06 Å by correlation while the shortening of the O–Cl separation is three times that amount.

It is well known that the length of the HX bond is increased upon formation of a H bond such as H\(_2\)O–HX. The magnitude of this bond stretch is provided in the third row of Table V as \(\Delta r\). For H\(_2\)O–HF this lengthening is 0.012 Å at the SCF level; the corresponding value for H\(_2\)O–HCl is 0.009 Å. Correlation increases this stretch to 0.017 Å in the former system and to 0.015 Å in the latter. Concomitant with these bond lengthenings is a reduction in the associated \(F_c\) force constant. At the SCF level, this constant is reduced by 1.5 mdyn/Å following complexation; the analogous value at the MP2 level is 2.0. Based on previous experience, the correlation-induced bond stretches are probably somewhat exaggerated. Future calculations including higher-order perturbation effects would be quite useful in order to determine the true magnitude of the effect of correlation upon the bond length.

A very interesting and useful relationship is observed if one plots the contribution of electron correlation to the total interaction energy against the \(r\) (HX) bond length in the hy-
FIG. 6. Dispersion energy $E^D$ shown as a function of $r(\text{HX})$. The lower scale is for $\text{H}_2\text{O}-\text{HF}$ and the upper for $\text{H}_2\text{O}-\text{HCl}$.

drogen halide. The data in Fig. 6 indicate a very nearly exact linear relation between these two quantities. That is, as the proton is shifted away from the halogen atom, the attractive dispersion between the two molecules is increased accordingly. The slope of the curve for $\text{H}_2\text{O}-\text{HF}$ is dramatically steeper than for $\text{H}_2\text{O}-\text{HCl}$: the values of $dE^D/dr$ are $-34.6$ and $-8.3$ kcal/mol Å, respectively. Treating the dispersion energy in terms of interactions between the polarizabilities of the individual molecules, the linear relationship implies that the polarizability of HX, like its dipole moment, increases linearly with bond length. This rather dramatic increase of dispersion stabilization occurs at the expense of the energy required to stretch the H-X bond.

Early theories of H bonding attempted to explain the observed lengthening of the HX bond upon complexation on the basis of charge transfer to a vacant antibonding orbital of the proton donor. The results described here indicate that the bond stretch may be ascribed instead to a number of other effects. The increased dipole moment arising from the bond stretch would be expected to magnify the stabilizing electrostatic forces. Attractive induction forces would also be increased as a result of the greater polarizability (and dipole moment) of the HX molecule. These stabilizing considerations explain the amount of $\Delta r$ accounted for at the SCF level and they basically agree with the earlier concepts. What is new, is our observation that dispersion forces play an important additional role in the process of lengthening of the HX bond.

We may summarize the differences between the H bonds in $\text{H}_2\text{O}-\text{HF}$ and $\text{H}_2\text{O}-\text{HCl}$ as follows. HF has a greater dipole moment than does HCl and H-bonds involving HF will hence contain a greater amount of electrostatic stabilization. The smaller contribution of dispersion to the H bond in $\text{H}_2\text{O}-\text{HF}$ is a result of the lower polarizability of the HF molecule. On the other hand, the high sensitivity of the polarizability of HF to the bond length allows the latter deficiency to be made up by small stretches of the bond upon complexation. Calculated $\Delta r$ for $\text{H}_2\text{O}-\text{HF}$ is slightly larger than that for $\text{H}_2\text{O}-\text{HCl}$, which agrees with the experimentally observed larger low-frequency shifts of $v_x(\text{HX})$ in H-bonded complexes involving HF.

ACKNOWLEDGMENTS

Financial support was provided by grants to S. S. from NIH (GM29391 and AM01059) and from the Research Corporation. Allocations of computer time from SIU are acknowledged.

26. This is only a qualitative picture; quantitative analysis would require consideration of possible lack of convergence of the multipole expansion of electrostatic energy due to penetration effects.