Intermolecular Bond Lengths: Extrapolation to the Basis Set Limit on Uncorrected and BSSE-Corrected Potential Energy Hypersurfaces

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ABSTRACT: Geometry optimizations were carried out for the (HF)2, (H2O)2, and HF–H2O intermolecular complexes using the MP2/aug-cc-pVXZ [X = 2, 3, 4, and 5] theoretical models on both the uncorrected and counterpoise (CP) corrected potential energy hypersurfaces (PES). Our results and the available literature data clearly show that extrapolation of intermolecular distances to the complete basis set (CBS) limit is satisfactory on PESs corrected for BSSE. On the other hand, one should avoid such extrapolations using data obtained from uncorrected PESs. Also, fixing intramolecular parameters at their experimental values could cause difficulties during the extrapolation. As the available literature data and our results clearly show, the MP2/aug-cc-pVXZ [X = 2, 3, 4] data series of intermolecular distances obtained from the CP-corrected surfaces can be safely used for the purpose of CBS extrapolations. © 2000 John Wiley & Sons, Inc. J Comput Chem 22: 196–207, 2001

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Introduction

During the last few years significant progress has been made in the *ab initio* computation of intermolecular interactions. The rapid development in quantum chemical procedures (including methods, basis sets, and computer codes) and the expansion of computational resources changed the prospects of theoreticians working in the field of noncovalent chemistry. From the relevant developments we mention only those of importance for the present study. First, Dunning and coworkers have developed a family of correlation consistent [(aug)-cc-pV*(C)V] basis sets systematically extend the atomic radial and angular spaces as a function of the cardinal number X. Therefore, results obtained with them seem to provide an excellent opportunity to extrapolate energies as well as properties to the complete basis set (CBS) limit. In most systems studied, extrapolation of the total energy and many properties has been achieved with simple functional forms. The exponential form

$$A_X = A_{CBS} + ae^{-kX},$$  

(1)

and polynomials of the form

$$A_X = A_{CBS} + \sum_{k=3}^{k_{\text{max}}} a_k (X + \beta_k)^{-k},$$  

(2)

where $k_{\text{max}}$ is small, or similar polynomial forms with noninteger exponents have been extensively employed to estimate the CBS limit. The form of eq. (1), for example, suggests that properties obtained with three basis sets (e.g., X = D, T, and Q) gives an opportunity to extrapolate to the CBS limit of the investigated property. Second, much experience has been accumulated on how the correlation-consistent basis sets can be employed to estimate the CBS limit. In most systems studied, extrapolation of the total energy and many properties has been achieved with simple functional forms. The exponential form

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One of the principal difficulties during computation of intermolecular properties is due to the basis set superposition error (BSSE). Briefly, because one cannot use complete basis sets in practical computations, the description of “internal” monomer properties depends on the quality and location of the basis functions of the partner molecule(s). Due to BSSE, the calculated interaction energies become too large, and the predicted potential energy hypersurfaces are distorted.

The conventional way to correct for BSSE a posteriori is based on the Boys–Bernardi [counterpoise (CP)] scheme. On the other hand, the Chemical Hamiltonian Approach (CHA) introduced by Mayer eliminates the nonphysical terms of the Hamiltonian that are due to BSSE a priori, and therefore, represents an ultimate solution to the problem of BSSE. The CHA is available up to the MP2 level of theory, however, the actual implementation is not suitable for calculations as large as those presented here. For an excellent recent review on CHA, see ref. 21.

Using the CP scheme one has to recalculate the monomers in the basis of the whole supermolecule for every geometrical arrangement. For example, in the case of two interacting monomers A and B, the uncorrected interaction energy ($\Delta E$) can be calculated as

$$\Delta E = E_{AB}(AB) - E_A(A) - E_B(B),$$  

(3)

where $E_{AB}(AB)$ is the total energy of the complex, and $E_A(A)$ and $E_B(B)$ are the total energies of the monomers. [In the following, we will use subscripts to denote the molecular species in the energy expressions, while the letters in parentheses refer to the (composite) basis used in the calculation. For example, $E_A(A)$ is the energy of monomer A calculated by using its own basis set.] The CP-corrected interaction energy can be defined as

$$\Delta E_{\text{CP}} = E_{AB}(AB) - E_A(A) - E_B(B).$$  

(4)

Using eqs. (3) and (4), one can define the BSSE content of the interaction energy as

$$\delta_{\text{BSSE}} = \Delta E - \Delta E_{\text{CP}} = E_A(AB) - E_A(A) + E_B(AB) - E_B(B).$$  

(5)

Using eq. (5), one can define the CP-corrected potential energy surface (PES) of a dimer as

$$E_{\text{CP}} = E_{AB}(AB) - \delta_{\text{BSSE}} = E_{AB}(AB) + E_A(A) - E_A(AB) + E_B(B) - E_B(AB).$$  

(6)

According to eq. (6), one has to calculate five different total energies at every geometrical arrangement of the system to determine a CP-corrected PES. Of course, eq. (6) can be generalized to the case of...
an arbitrary number of subsystems, but the number of energy calculations necessary to determine the PES increases with the number of monomers enormously.\textsuperscript{23} There has been a continuous debate on the CP method in the literature. One of the main arguments against the CP scheme is that the composite basis of the complex \(AB\) is not available for monomer \(A\), as the Pauli exclusion principle precludes the use of the occupied orbitals of monomer \(B\).\textsuperscript{24} In practice, this means that the CP scheme overcompensates the BSSE. This effect was studied on a small analytical model by Mayer and Túri.\textsuperscript{25} They found that the CP scheme takes the leading BSSE terms properly into account. However, there are two additional terms, which are responsible for the propensity of the method to overcompensate the BSSE. The magnitude of these minor terms depends on the size and quality of the basis and, in accordance with the accumulated numerical experience,\textsuperscript{26–28} decreases as the basis set is enlarged. It has generally been accepted that one almost always has to correct for the BSSE, and that adequate basis sets have to be used to avoid the overcompensating nature of the CP method.

This status quo seems to have been challenged by new results obtained from calculations using the aug-cc-pV\(X\)Z basis sets of Dunning. As most of the pioneering investigations\textsuperscript{5, 13–15} have shown, it is rather important to correct for the BSSE if the standard cc-pV\(X\)Z basis sets are used for the calculations. In these cases, the CP-corrected properties are nearly always closer to the available experimental data than the uncorrected ones. However, the situation is substantially different in the case of the aug-cc-pV\(X\)Z basis sets. The general trend is that the uncorrected aug-cc-pV\(X\)Z results lie close to the corresponding experimental values, and the convergence of the CP-corrected aug-cc-pV\(X\)Z results to the CBS limit is slow. Based on these findings many of the authors of the pioneering studies\textsuperscript{14, 15} criticized the CP method and questioned the applicability of the CP scheme in conjunction with the aug-cc-pV\(X\)Z basis sets.

Dunning and coworkers have recently published a review\textsuperscript{4} about their activity devoted to the exploration of the limits of the CP scheme. These authors have shown that, in many cases, the convergence behavior of various molecular properties is significantly improved if the calculations are corrected for the BSSE. According to Dunning and coworkers,\textsuperscript{4} for many investigated properties the smooth convergence behavior of the results obtained with aug-cc-pV\(X\)Z basis sets is a pure illusion, as it is due to a fortuitous cancellation of the BSSE and the basis set incompleteness error (BSIE). Correction for the BSSE destroys the balance of the two errors; consequently, the CP-corrected data lie farther away from experiment than the uncorrected ones. However, the use of CP-corrected data is much safer for CBS limit extrapolation: the corrected data suffer only from the BSIE, which can be taken into account by a suitable extrapolation to the CBS limit. Anomalies related to the extrapolation to the CBS limit were most pronounced for weakly bound intermolecular complexes\textsuperscript{4} (van der Waals and hydrogen-bonded systems), although similar behavior was observed even for certain strongly bound systems.\textsuperscript{4}

Dunning and coworkers have investigated various molecular properties like interaction energies and equilibrium distances of di- and triatomic molecules. However, their work on real many-dimensional potential energy hypersurfaces (PEs) was limited to (HF)\(_2\), partly due to the lack of an efficient automated procedure for BSSE-free geometry optimizations. [The optimized geometries were computed from numerical (CP-corrected or uncorrected) gradients.] Those studies, which expressed scepticism\textsuperscript{14, 15} about the usefulness of the CP scheme when applied in connection with the aug-cc-pV\(X\)Z basis sets, were devoted to the characterization of PESs of hydrogen-bonded systems like the HF dimer\textsuperscript{15} and the water dimer.\textsuperscript{14, 15}

To resolve this apparent controversy, we decided to carry out geometry optimizations on prototypical hydrogen-bonded systems. Our goal is to investigate the behavior of the CP method in conjunction with the aug-cc-pV\(X\)Z basis sets for equilibrium geometric parameters. Therefore, we optimized the geometry of (HF)\(_2\) and (H\(2\)O)\(_2\) employing analytical gradients at both the uncorrected and BSSE-corrected MP2/aug-cc-pV\(X\)Z [\(X = D(2), T(3), Q(4)\), for (HF)\(_2\) also 5] levels of theory. The geometrical data obtained provide evidence in support of Dunning’s opinion described above. From the comparison of our data with the literature results available on (H\(2\)O)\(_2\), we evaluate the reliability of various approximations used in geometry optimizations (e.g., freezing a subset of the parameters), and show the importance of carrying out full geometry optimizations when extrapolation of geometrical parameters to the CBS limit is desired. Finally, we present optimized geometries and equilibrium interaction energies of the HF–H\(2\)O complex obtained on both uncorrected and BSSE-corrected MP2/aug-cc-pV\(X\)Z (\(X = D, T, Q\)) PESs.
**Computational Details**

All ab initio energy and gradient calculations utilized the Gaussian94 program system. For the CP geometry optimizations we used a program system described in ref. 30 with some small modifications in both the scripts and the various Gaussian links. The main idea behind the CP optimization is that various derivatives of eq. (6) can easily be calculated. For example, the gradient on the CP-corrected surface can be calculated as

\[
\frac{\delta E^{BB}}{\delta x} = \frac{\delta E_{AB}(AB)}{\delta x} + \frac{\delta E_{A}(A)}{\delta x} - \frac{\delta E_{A}(AB)}{\delta x} + \frac{\delta E_{B}(B)}{\delta x} - \frac{\delta E_{B}(AB)}{\delta x},
\]

where \( x \) is a geometry parameter. Our present implementation enables us to determine optimized geometries on the corrected PES in three to four optimization steps provided the initial geometry is the corresponding uncorrected optimized one. For rapid convergence it was important to use reliable Hessians in conjunction with simple Z-matrix-type coordinates during the GDIIS optimization. The Hessian was usually calculated on the uncorrected PES. In all correlated-level calculations the 1s core orbitals of O and F have been kept frozen.

**Results and Discussion**

**HF Dimer**

The HF dimer (see Fig. 1) is one of the most important prototypes of hydrogen-bonded systems. Therefore, it has been the subject of detailed experimental (cf., ref. 32 and references therein) and theoretical (cf., ref. 13 and references therein) studies. The structure of (HF)\textsubscript{2} was studied at the MP2/aug-cc-pVXZ (X = D, T, and Q) levels by Peterson and Dunning (henceforth, PD). In that article the geometry of the complex was fully optimized at the MP2/aug-cc-pVDZ and MP2/aug-cc-pVTZ levels on both the uncorrected and CP-corrected potential energy hypersurfaces. (PD used numerical gradients for the optimization.) With the aug-cc-pVQZ basis set PD carried out full geometry optimization on the uncorrected PES, while they optimized only the intermolecular F–F distance (\( r_{FF} \), Fig. 1) on the CP-corrected PES, keeping the other parameters fixed at their estimated MP2/aug-cc-pVQZ values. (For further details, see ref. 13.) The most important conclusions drawn by PD are as follows: (a) the uncorrected MP2/aug-cc-pVXZ interaction energies and geometries are closer to experiment than the corresponding corrected ones; (b) convergence of the CP-corrected binding energies and geometries is more regular than that of the uncorrected parameters; and (c) the CBS limit of the CP-corrected \( r_{FF} \) distance, 2.737 Å, is numerically indistinguishable from the uncorrected aug-cc-pVQZ optimized value. Overall, PD stressed the importance of the CP correction for the investigated properties.

As a first step in our investigation, we reoptimized the geometry of (HF)\textsubscript{2} at the levels for which PD published data. We were able to reproduce their results with only slight deviations by using our automated CP optimization algorithm (see Table I for the results). For the CP-corrected MP2/aug-cc-pVQZ case PD carried out partial optimizations by fixing most of the internal parameters. Our full optimization at the same level confirmed the adequacy of the approximations applied by PD, the fully optimized parameters (\( \alpha = 6.7^\circ, \beta = 111.7^\circ, r_{FF} = 2.753 \) Å) are close to the corresponding values (\( \alpha = 6.4^\circ, \beta = 111.15^\circ, r_{FF} = 2.753 \) Å) of PD.

As mentioned above, the uncorrected MP2/aug-cc-pVQZ \( r_{FF} \) distance seems to be converged, and agrees with the CBS limit, 2.737 Å, obtained on the CP-corrected surface by PD. (One has to note here that the CBS limit of \( r_{FF} \) determined by our geometry data, 2.741 Å, differs slightly from the corresponding value of PD, 2.737 Å.) At the same time, the actual value of the \( r_{FF} \) distance (2.753 Å) on the corrected PES is far from both the MP2/aug-cc-pVQZ and the corrected CBS limit values. This situation provides an opportunity to further investigate the importance of CP correction in estimating the CBS limit of geometrical parameters, like the \( r_{FF} \) distance in (HF)\textsubscript{2}. It seems to be worth investigat-
TABLE I.
Geometrical Parameters (Angles/Deg and Lengths/Å) and Total Energies ($E_{\text{tot}}/E_{\text{t}}$) of the HF Dimer Calculated at the Uncorrected and CP-Corrected MP2/aug-cc-pVXZ ($X = 2, 3, 4, 5$) Levels of Theory.

| $X$  | $r_{FF}$  | $\alpha$  | $\beta$  | $E_{\text{tot}}$ |
|------|-----------|-----------|----------|------------------|
| D    | 2.753$^a$ | 6.5 (6.6$^a$) | 110.2 (110.1$^a$) | $-200.519075^a$ |
| T    | 2.746$^a$ | 6.4$^a$   | 111.1$^a$ | $-200.689285^a$  |
| Q    | 2.736 (2.737$^a$) | 6.4$^a$ | 111.5 (111.6$^a$) | $-200.746906^a$  |
| 5    | 2.739     | 6.3       | 112.0     | $-200.768417$     |

| $N$  | $r_{FF}$  | $\alpha$  | $\beta$  | $E_{\text{tot}}$ |
|------|-----------|-----------|----------|------------------|
| D    | 2.812 (2.813$^b$) | 7.0$^a$   | 111.3$^a$ | $-200.518060$    |
| T    | 2.770 (2.772$^b$) | 6.7$^a$   | 111.7 (111.8$^b$) | $-200.688525$  |
| Q    | 2.759$^a$ | 6.7       | 111.7     | $-200.746495$    |
| 5    | 2.749     | 6.7       | 111.7     | $-200.768158$    |

$^a$ Ref. 6.
$^b$ Partial optimization. For details, see ref. 6. For the definition of the variables, see Figure 1. In those cases, when two values are presented for a variable, our optimized value does not agree with that of ref. 6 given in parentheses.

The geometry of (HF)$_2$ at the MP2/aug-cc-pV5Z level for the following reasons. If convergence of $r_{FF}$ was really manifested at the MP2/aug-cc-pVQZ level, the MP2/aug-cc-pV5Z value would coincide with the MP2/aug-cc-pVQZ value. On the other hand, any other MP2/aug-cc-pV5Z $r_{FF}$ value would question the extrapolation to the CBS limit using the uncorrected data. As it turns out, both the uncorrected ($\alpha = 6.3^\circ$, $\beta = 112.0^\circ$, $r_{FF} = 2.739$ Å) and CP-corrected ($\alpha = 6.7^\circ$, $\beta = 111.7^\circ$, $r_{FF} = 2.749$ Å) geometric parameters lie close to the corresponding MP2/aug-cc-pVQZ values. It is noted, that the difference between the corrected and uncorrected $r_{FF}$ values is substantial, 0.010 Å, at the MP2/aug-cc-pV5Z level of theory. The series of MP2/aug-cc-pVXZ ($X = D, T, Q, 5$) $r_{FF}$ data represent a minimum curve with the minimum between $X$ equal to Q and 5 (see Fig. 2). On the other hand, the CP-corrected MP2/aug-cc-pVXZ ($X = D, T, Q, 5$) $r_{FF}$ distances follow a monotonic curve.

One could extrapolate to the CBS limit of the $r_{FF}$ distance in three ways using the available data obtained on the corrected PES; for example, one could use the $\{2, 3, 4\}$, $\{3, 4, 5\}$, and $\{2, 3, 4, 5\}$ series for determining extrapolated geometric parameters according to eq. (1). Obviously, more reasonable values are expected for the latter choices. The CBS limit values of the $r_{FF}$ distance of (HF)$_2$ for the above listed series are reasonably close to each other at 2.741, 2.748, and 2.745 Å, respectively. (For the two to five fitting we used our own CP-corrected data, which slightly differ from those of PD.) One has to note that the MP2/aug-cc-pVQZ level, which represents the lowest level of theory employed, provides a much poorer approximation than that obtained at the MP2/aug-cc-pVTZ level. This effect can be seen, for example, in the huge BSSE content of the geometry. After correction for BSSE, the $r_{FF}$ distance is lengthened by 0.059, 0.024, 0.017 Å, and 0.010 Å at the MP2/aug-cc-pVDZ, MP2/aug-cc-pVTZ, MP2/aug-cc-pVQZ, and MP2/aug-cc-pV5Z levels of theory.

FIGURE 2. Comparison of the plain and CP-corrected $r_{FF}$ distances for (HF)$_2$. 

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respectively. Inferiority of the MP2/aug-cc-pVDZ geometry parameters, compared to the larger basis set results, is basically responsible for the differences between the CBS limit values obtained from the three fittings.

Nevertheless, as Chuang and Truhlar explored, one is tempted to use the inexpensive DZ and TZ data to obtain extrapolated geometric parameters corresponding to the basis set limit. However, instead of using the scheme of Chuang and Truhlar, we employed eq. (2) in the form of

\[ A_X = A_{CBS} + aX^{-3} \]  

(8)
to obtain estimates of the extrapolated \(r_{FF}\). Our noteworthy result is that the extrapolated \(r_{FF}\) distances, 2.743 and 2.752 Å in the uncorrected and BSSE-corrected cases, respectively, are very close to the corresponding MP2/aug-cc-pV5Z numbers. This is especially notable for the CP-corrected \(r_{FF}\) distance, in which case the MP2/aug-cc-pVTZ value deviates from the MP2/aug-cc-pV5Z value by 0.021 Å, while the extrapolated distance deviates only by 0.003 Å.

In summary, the smooth convergence behavior of the geometric parameters obtained directly with the aug-cc-pVXZ basis sets is due to a fortuitous cancellation of BSSE and BSIE. The simple exponential or polynomial functions employed to extrapolate to the CBS limit cannot work reliably if the points do not follow a monotonic curve, as is the case for the uncorrected MP2/aug-cc-pVXZ results. On the other hand, although the CP-corrected \(r_{FF}\) distances are usually farther away from the extrapolated values than the uncorrected distances (this is true for all but the MP2/aug-cc-pV5Z data), changes in the corrected geometric parameters are monotonic, and thus are in better accordance with the design philosophy of the aug-cc-pVXZ basis sets.

**WATER DIMER**

The water dimer (see Fig. 1) has a linear structure, which is known both from experiment and from *ab initio* calculations [ref. 15, and references therein]. Our results and the available MP2/aug-cc-pVXZ geometry data are summarized in Table II.

The first study devoted to the investigation of \((\text{H}_2\text{O})_2\) employing the aug-cc-pVXZ \((X = 2, 3, 4, 5)\) basis sets at the MP2 level was carried out by Feller, who determined the interaction energy of the complex at various correlated levels at a fixed geometry. The first geometry optimizations on the water dimer at the MP2(FC)/aug-cc-pVXZ \((X = D, T, Q)\) were carried out by Feller et al. on the uncorrected PES. Importance of fragment relaxation terms in the CP
scheme was investigated by Xantheas, who determined the fully relaxed geometry of \((H_2O)_2\) at the MP2/aug-cc-pVDZ and MP2/aug-cc-pVTZ levels on the uncorrected potential energy hypersurfaces. Furthermore, Xantheas carried out partial optimizations at the MP2/aug-cc-pVQZ and MP2/aug-cc-pV5Z levels varying the intermolecular \(r_{OO}\) distance (Fig. 1) and keeping other parameters fixed at their MP2/aug-cc-pVTZ value. Xantheas’ MP2/aug-cc-pV5Z value for the \(r_{OO}\) distance is 2.905 Å on the uncorrected and 2.913 Å on the CP-corrected PESs, respectively. In a recent article, Halkier et al. also investigated the structure of the water dimer employing the MP2/aug-cc-pV\(X\)Z models. These authors performed constrained optimizations, for example, they kept the water monomer parameters frozen at the corresponding experimental values. With this approximation, the geometry of \((H_2O)_2\) was determined at the MP2/aug-cc-pV\(X\)Z (\(X = 2, 3, 4\)) levels at both the uncorrected and the CP-corrected surfaces. (As a further approximation, the intermolecular valence angles were also frozen in the optimizations on the CP-corrected PES at their corresponding uncorrected values.) One should note here that, based on the analysis of their data, Halkier and coworkers argued against the computation of CP-corrected geometries at the MP2/aug-cc-pV\(X\)Z levels.

The results of our full geometry optimizations carried out on the uncorrected and CP-corrected PESs are shown in Table II. At the MP2/aug-cc-pVDZ and MP2/aug-cc-pVTZ levels we could reproduce Feller’s and Xantheas’ results with slight differences on both the uncorrected and corrected PESs. On the other hand, the \(r_{OO}\) distances obtained by Halkier et al. for the uncorrected surfaces differ considerably from the corresponding fully relaxed values. For example, the difference between the \(r_{OO}\) distances of Feller and Halkier is 0.016 Å at the MP2/aug-cc-pVTZ level. This discrepancy questions the validity of the approximations utilized by Halkier et al. Furthermore, Xantheas’ data determined at the MP2/aug-cc-pVQZ level from partial optimizations agree reasonably well with both Feller’s and our values obtained from full optimizations. Hereby, we can confirm the adequacy of the approximations employed by Xantheas in his partial optimizations on \((H_2O)_2\), for example, keeping all the intramolecular and the valence bending intermolecular parameters at their optimized MP2/aug-cc-pVTZ values in the subsequent aug-cc-pVQZ and aug-cc-pV5Z optimizations. Because of the reliability of the approximations employed by Xantheas and the cost of the MP2/aug-cc-pV5Z optimization we decided not to determine the geometry of \((H_2O)_2\) at the MP2/aug-cc-pV5Z level.

Concerning the extrapolation of the \(r_{OO}\) distance to the CBS limit, one can draw the following conclusions. The two series of \(r_{OO}\) distances, determined by Feller et al., Xantheas and Halkier et al. employing different approximations during the optimizations, follow a curve with a minimum. That is, Halkier’s aug-cc-pVTZ \(r_{OO}\) distance at 2.891 Å is shorter than the corresponding aug-cc-pVDZ (2.912 Å) and aug-cc-pVQZ (2.895 Å) values. This behavior is presumably due to the inadequacy of the constraints employed by Halkier et al. during the optimizations. In the data series of Feller et al. and Xantheas, the aug-cc-pVQZ \(r_{OO}\) distance (2.903 Å) is shorter than the corresponding aug-cc-pVDZ (2.916 Å), aug-cc-pVTZ (2.907 Å), and aug-cc-pV5Z (2.905 Å) values. This behavior of the aug-cc-pV\(X\)Z basis sets in the case of uncorrected calculations is very similar to what we observed in full geometry optimizations on \((HF)_2\). On the other hand, the CP-corrected \(r_{OO}\) distances converge more regularly, all three data series obtained by Xantheas, Halkier et al. and the present work ascend gradually when enlarging the cardinal number of the employed basis set (see Fig. 3). The extrapolated CBS limit of the \(r_{OO}\) distance obtained from Halkier’s data seems to be too short at 2.897 Å, indicating again the inadequacy of the constraints employed during the geometry optimizations. On the other hand, the extrapolated \(r_{OO}\) CBS limit values obtained from Xantheas’ partially optimized aug-cc-pV\(X\)Z \(X = 2, 3, 4, 5\) and our fully optimized aug-cc-pV\(X\)Z \(X = 2, 3, 4\) data using eq. (1) practically coincide at 2.910 Å. It is worth noting in this respect that the CBS limit values determined from.
the \(X = 2,3,4\) and \(X = 3,4,5\) series of Xantheas are also the same at 2.910 Å.

Similarly to the case of \((\text{HF})_2\), use of eq. (8) for the extrapolation of aug-cc-pVDZ and aug-cc-pVTZ data gives very satisfactory results for the CP-corrected \(r_{\text{fo}}\) distances. While the difference between the aug-cc-pVTZ and aug-cc-pVQZ \(r_{\text{fo}}\) distances is 0.020 Å the extrapolated value differs by only 0.005 Å.

As a summary of the above considerations, we strongly advocate to perform full geometry optimizations in those studies whose aim is the extrapolation of geometrical parameters to the CBS limit. The reliability of the CBS limit of the \(r_{\text{fo}}\) distance determined from the fully relaxed CP-corrected aug-cc-pVXZ \(X = 2,3,4\) potential energy hypersurfaces is appealing compared to the aug-cc-pVXZ \(X = 2,3,4,5\) data obtained by Xantheas.35

HF–H2O COMPLEX

Determination of the structure of the HF–H2O complex proved to be a challenge for experimentalists and theoreticians alike. Theoretical studies36,37 showed that one can find stationary points on the HF–H2O PES with both \(C_s\) and \(C_{2v}\) symmetries. (In the case of the \(C_{2v}\) structure the \(\alpha\) and \(\beta\) angles (Fig. 1) are 0 and 180 degrees, respectively.) The \(C_{2v}\) species is favored by certain electrostatic arguments.38 Namely, the \(C_{2v}\) arrangement of the complex allows the dipole moment of H2O to be aligned with the dipole of HF. On the other hand, the interaction of the negative charges of the respective quadrupole tensor element of H2O and the positive end of the HF dipole clearly favors a structure where the plane of H2O is perpendicular to the direction of the HF dipole. The structure of HF–H2O can be considered as a compromise between these electrostatic and other (less important) nonelectrostatic effects. Indeed, although the \(C_s\) structure is more stable than the \(C_{2v}\) one, the energy difference between them is only 0.1 and 0.5 kcal mol\(^{-1}\) using a small basis set at the SCF and MP2 levels,36 respectively. The experimental39 estimate of the barrier is 0.4 kcal mol\(^{-1}\). Experimental determination of the angular features of the HF–H2O complex is difficult.30 The energy barrier between the \(C_s\) and \(C_{2v}\) structures is small, presumably close to the energy of the first vibrational level. In the case when the first vibrational level occurs above the top of the barrier, the experiment would predict a planar structure despite the existence of a double-well potential. If the vibrational level is slightly below the top of the barrier, the experiment can predict only a poorly defined structure.

The most recent theoretical investigation devoted to the structure of HF–H2O, including geometry optimizations, was carried out by Novoa et al.37 at the MP2/6-311++G(2d,2p) level of theory resulting in an energy difference of 0.45 kcal mol\(^{-1}\) between the \(C_s\) and \(C_{2v}\) structures. They have also calculated the equilibrium \(r_{\text{fo}}\) distance (2.663 Å) using a constrained optimization keeping the monomer parameters fixed. Finally, their interaction energies computed at the uncorrected and CP-corrected MP2/6-311++G(2d,2p) levels are −9.20 and −7.87 kcal mol\(^{-1}\), respectively.

Because of the available experimental and theoretical information, the HF–H2O complex is an ideal subject to test the CBS limit values of various properties determined from calculations performed at the MP2/aug-cc-pVXZ \(X = 2,3,4\) levels. To explore the quality of theoretical results obtained on the uncorrected and CP-corrected surfaces, we decided to calculate the equilibrium dissociation energy, the energy difference between the \(C_s\) and \(C_{2v}\) species, and the equilibrium geometry of HF–H2O. The experimental data were determined by Legon and coworkers39–41 using microwave rotational spectroscopy. The calculated total energies, energy barriers, and geometrical parameters obtained at the MP2/aug-cc-pVXZ \(X = 2,3,4\) levels are given in Table III. The CBS value for the total energy of H2O is −76.363558 \(E_h\), obtained from the total energies (−76.260910, −76.328992, and −76.351919 \(E_h\)) calculated at the MP2/aug-cc-pVDZ, MP2/aug-cc-pVTZ, and MP2/aug-cc-pVQZ levels, respectively.

The experimental41 equilibrium interaction energy \(\langle E_{\text{int}}\rangle\) of HF–H2O is −10.2 kcal mol\(^{-1}\), determined by Legon and coworkers from absolute intensities of rotational transitions. As mentioned before, the best theoretical values37 available are −9.20 and −7.87 kcal mol\(^{-1}\) obtained from uncorrected and CP-corrected calculations using a medium-size basis set and the MP2 method. The uncorrected aug-cc-pVXZ \(X = 2,3,4\) interaction energies shown in Table IV are appealingly close to each other. The CBS limit value of \(E_{\text{int}}\) obtained using the extrapolated total energies of the monomers and the complex, differs only slightly from these values. The interaction energies obtained at the CP-corrected PESs considerably differ from the corresponding uncorrected energies. However, this difference gradually decreases, considering the series of basis sets, going from the aug-cc-pVDZ set to the aug-cc-pVQZ one. The CP-corrected CBS
TABLE III.
Geometrical Parameters (Angles/Deg and Lengths/Å) and Total Energies ($E_{\text{tot}}/E_h$) of HF–H$_2$O Calculated at the Uncorrected and CP-Corrected MP2/aug-cc-pVXZ ($X = 2, 3, 4$) Levels of Theory.

| $X$ | Plain | CP-Corrected |
|-----|-------|--------------|
|     | $E_{\text{tot}}$ | $r_{FO}$ | $\alpha$ | $\beta$ | $E_{\text{tot}}$ | $r_{FO}$ | $\alpha$ | $\beta$ |
|     |       |     |       |       |       |     |       |       |
| Cs  |       |     |       |       |       |     |       |       |
| 2   | $-176.531107$ | 2.658 | 129.5 | 1.4  | $-176.529305$ | 2.701 | 130.6 | 1.5  |
| 3   | $-176.684156$ | 2.643 | 130.7 | 1.4  | $-176.683198$ | 2.662 | 132.0 | 1.5  |
| 4   | $-176.735847$ | 2.640 | 132.4 | 1.4  | $-176.735257$ | 2.654 | 132.6 | 1.4  |
| CBS | $-176.762208$ | 2.639 | —     | —    | $-176.761869$ | 2.652 | —     | —    |
|     |       |     |       |       |       |     |       |       |
| C$_{2v}$ |       |     |       |       |       |     |       |       |
| 2   | $-176.530314$ | 2.666 | 0.0   | 180.0 | $-176.528674$ | 2.710 | 0.0   | 180.0 |
| 3   | $-176.683503$ | 2.655 | 0.0   | 180.0 | $-176.682605$ | 2.670 | 0.0   | 180.0 |
| 4   | $-176.735318$ | 2.648 | 0.0   | 180.0 | $-176.734751$ | 2.661 | 0.0   | 180.0 |
| CBS | $-176.761803$ | 2.636 | —     | —    | $-176.761465$ | 2.658 | —     | —    |

The calculated energy barriers ($\Delta E$) belonging to the transition from the Cs to the C$_{2v}$ species of HF–H$_2$O are shown in Table IV. $\Delta E$ gradually decreases, going from the smallest to the larger basis sets. It is worth noting that the $\Delta E$ values obtained at the plain and CP-corrected surfaces are very close to each other at the MP2/aug-cc-pVQZ level of theory. The CBS limit values of the same parameter obtained at the plain and the CP-corrected PESs numerically coincide at 0.25 kcal mol$^{-1}$. The CBS limit of $\Delta E$ is considerably smaller than the corresponding equilibrium experimental\textsuperscript{40} data (0.36 kcal mol$^{-1}$), again showing the importance of inclusion of higher order correlation terms. It is worth noting that the good agreement with experiment obtained in previous theoretical studies\textsuperscript{36, 37} is clearly due to cancellation of errors, for example, the quality of the basis sets employed in those investigations was unsatisfactory to consistently deal with the problems of BSSE and BSIE.

TABLE IV.
Equilibrium Interaction Energies ($E_{\text{int}}$ in kcal/mol) and Energy Splitting ($\Delta E$ in kcal mol$^{-1}$) between the Cs and C$_{2v}$ Species of HF–H$_2$O Calculated at the Uncorrected and CP-Corrected MP2/aug-cc-pVXZ ($X = 2, 3, 4$) Levels of Theory.

| $X$ | $E_{\text{int}}$ | $\Delta E$ |
|-----|-----------------|------------|
|     | Plain | CP-Corrected | Plain | CP-Corrected |
|     |       |             |       |             |
| 2   | $-9.0$ | $-7.9$ | 0.50  | 0.40  |
| 3   | $-9.0$ | $-8.4$ | 0.41  | 0.37  |
| 4   | $-8.9$ | $-8.5$ | 0.33  | 0.32  |
| CBS\textsuperscript{a} | $-8.8$ | $-8.6$ | 0.25  | 0.25  |

\textsuperscript{a}The CBS limit values were calculated using the respective CBS limit total energies shown in Table III.
On the other hand, the CBS limit of $r_{FO}$ is 2.652 Å on the CP-corrected PES, representing a difference of 0.013 Å between the two extrapolated values. Legon and coworkers have determined the experimental value of $r_{FO}$ at 2.662 Å from the microwave rotational spectrum of HF–H₂O. Direct comparison of the experimental $r_0$ and theoretical $r_e$ values is not fully valid, but it is clear that the CP-corrected CBS limit $r_{FO}$ distance lies closer to experiment than the corresponding uncorrected value (see Fig. 4). However, one has to note here that the vibrational effects on hydrogen bond lengths can be substantial resulting in a case when the uncorrected CBS limit value is closer to the unmeasured experimental $r_e$ value. Another interesting anomaly concerns the change of the actual value of $r_{FO}$ going from the Cₘ to the C₂ᵥ species. One expects that the intermolecular bond length increases when climbing the transition state (C₂ᵥ species) region. All the data listed in Table III supports this statement, but the CBS limit of $r_{FO}$ obtained at the plain PESs contradicts it. In our opinion, this strange behavior is again due to the unreliability of the CBS limit values of geometry parameters obtained on the uncorrected PES.

Concerning the extrapolated $r_{FO}$ distances obtained by using eq. (8), one finds tendencies similar to those observed in the cases of (HF)₂ and (H₂O)₂. The extrapolated C₂ᵥ $r_{FO}$ distances, 2.650 and 2.653 Å, in the uncorrected and CP-corrected cases, respectively, are very close to the corresponding CBS value (2.658 Å) determined by using eq. (1) on the CP-corrected PESs. The extrapolated Cₘ $r_{FO}$ distances, 2.637 and 2.646 Å determined on the plain and CP-corrected PESs lie close to the corresponding CBS limit values (2.639 and 2.652 Å) obtained by using eq. (1). It is worth noting here that the extrapolated [eq. (8)] C₂ᵥ $r_{FO}$ value is longer than the corresponding Cₘ value in both the plain and CP-corrected cases. This means that bond length extrapolation based on eq. (8) is more reliable in the case of the HF–H₂O complex than application of eq. (1) because the expected change of parameter $r_{FO}$ is at least qualitatively given back by the former technique.

**COMMENT ON THE TRENDS OF INTERMOLECULAR DISTANCES IN THE (HF)₂, (H₂O)₂, AND HF–H₂O SERIES**

One of the referees of the present article suggested to investigate trends observed for the intermolecular bond lengths in the (HF)₂, (H₂O)₂, and HF–H₂O series. The two most interesting observations are as follows: (1) the differences between the plain aug-cc-pVQZ and the corresponding (CP-corrected) CBS values of the intermolecular distances are 0.005–0.008, 0.007, and 0.012 Å for (HF)₂, (H₂O)₂, and HF–H₂O, respectively; (2) the differences between the CP-corrected aug-cc-pVQZ and (CP-corrected) CBS values of the intermolecular bond lengths are 0.008–0.012, 0.007, and 0.002 Å, gradually decreasing in the (HF)₂, (H₂O)₂, and HF–H₂O series. (It is to be noted that three different CBS values were obtained for (HF)₂, resulting in uncertainties in the above-mentioned differences between the aug-cc-pVQZ and CBS limit values.)

We begin with the explanation of tendency (2), because in this case differences between two BSSE-free quantities are evaluated resulting in a less complex situation than that of point (1). The interaction between the monomers of the complexes in the (HF)₂, (H₂O)₂, and HF–H₂O series is getting definitely stronger. This means that the importance of very weak interactions, like dispersion forces, is decreasing, while the strength of the electrostatic and charge transfer interactions is increasing for the series of complexes investigated. Of course, one expects more reliable results for HF–H₂O than for (HF)₂ by using the same basis set in the calculations. Equivalent with this statement is that the real physical part of the interaction is more satisfied (e.g., it is getting closer to the CBS limit) for HF–H₂O than for (HF)₂ at a given level of theory provided that a reasonable basis set is applied in the particular calculations.

The difference between the plain aug-cc-pVQZ and (CP-corrected) CBS limit bond distances depends clearly on the magnitude of the BSSE and BSIE. The magnitude of the BSSE also depends on
two main factors in our case. The first and more important is the choice of the basis set applied. Plotting the BSSE content of the interaction energy vs.
different basis sets results in a curve that exhibits a maximum. The flexibility of a small basis set is insufficient for the appearance of a significant BSSE-type correlation energy contribution. With the growing size of the basis set such contributions appear and the BSSE content of the interaction energy increases. Using even larger basis sets the BSSE content will again decrease. It is clear that the basis sets applied in the present study represent the postmaximum regions of such BSSE content curves, but the actual locations of the aug-cc-pVXZ \((X = 2, 3, 4,\) and \(5\)) points and the slopes of the curves are clearly different for the \((\text{HF})_2, (\text{H}_2\text{O})_2,\) and \(\text{HF}–\text{H}_2\text{O}\) cases. In this respect, similarly to the description of the physical part of the interaction, the strength of the interaction is very important. One is evidently important to note here that these observations further stress the importance of correction for the BSSE.

**Conclusions**

Geometry optimizations were carried out for the \((\text{HF})_2, (\text{H}_2\text{O})_2,\) and \(\text{HF}–\text{H}_2\text{O}\) intermolecular complexes using the MP2/aug-cc-pVXZ \((X = 2, 3, 4,\) and \(5)\) theoretical models on both the uncorrected and CP-corrected potential energy hypersurfaces. Our results and the available literature data clearly show that extrapolation of intermolecular distances to the CBS limit is satisfactory on PESs corrected for BSSE. Evaluation of two extrapolation schemes \([\text{eqs. (1) and (8)}]\) suggests that reliable extrapolated geometry parameters can be obtained even with a simple two-point formula based on aug-cc-pVDZ and aug-cc-pVTZ data. Application of the extrapolation to obtain the CBS limit of intermolecular geometry parameters seems to be worth it if various species are to be compared on the PES of the complex investigated. Also, fixing intramolecular parameters at their experimental values could cause difficulties during the extrapolation. As the available literature data and our results clearly show, the MP2/aug-cc-pVXZ \((X = 2, 3, 4)\) data series of intermolecular distances obtained from the CP-corrected surfaces can be safely used for the purpose of CBS extrapolations.

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