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# The performance of CCSD(T) for the calculation of dipole moments in diatomics

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This work analyzes the accuracy of the coupled cluster with single, double, and perturbative triple excitation [CCSD(T)] method for predicting dipole moments. In particular, we benchmark CCSD(T) predictions for the equilibrium bond length, vibrational frequency, and dipole moment versus accurate experimental data. As a result, we find that CCSD(T) leads to accurate dipole moments. However, in some cases, it disagrees with the experimental values, and the disagreement can not be satisfactorily explained via relativistic or multi-reference effects. Therefore, our results indicate that benchmark studies for energy and geometry properties do not accurately describe other electron density magnitudes.

#### 1 Introduction

Coupled cluster with single, double, and perturbative triple excitations [CCSD(T)] is one of the most popular methods in electronic structure theory. Indeed, it serves as a benchmark reference in developing other electronic structure theory methods, such as density functional theory (DFT). It is sizeconsistent, and as a member of the coupled cluster family, it is systematically improvable. When utilized in combination with specific corrections, it is considered to approach subchemical accuracy in properties such as bond energies<sup>1</sup> at the complete basis set (CBS) limit.<sup>2</sup> DFT benchmarking studies most commonly focus on energetic properties. 1,3-17 Nevertheless, recently there has been increased interest in understanding the performance of other aspects of the wavefunction such as the electric dipole moment. 18-23

Most of the literature about benchmarking dipole moments is typically based on CCSD(T) performance, focusing on molecules consisting of light main-group elements. Nevertheless, molecules involving elements from the third row and heavier (Z > 18), especially transition-metal compounds, play an essential role in modern applications like catalysis and material synthesis due to their electronic and magnetic properties. 24-26 Many of these applications rely on an accurate description of the energetic properties and electron densities, so CCSD(T) is becoming popular among the different quantum chemistry methods. On the other hand, CCSD(T) generally relies on single

Slater-determinant Hartree-Fock references. As a result, its performance can sometimes be questionable in systems with multi-reference nature. 15,27 Another concern is the accuracy of the approximations that can be applied in CCSD(T) calculations. For example, frozen-core approximation is a popular choice when calculating systems with heavy elements. However, the computational cost of the CCSD(T) core-core and corevalence correlations quickly becomes impractical with an increasing number of electrons in the systems with effective core potentials (ECPs) applied.

Therefore, since the CCSD(T) method is central to modern quantum chemistry and for benchmarking other computational chemistry methods, it is necessary to compare its performance with that of other methods. In particular, it is possible to benchmark the performance of CCSD(T) against available experimental data on spectroscopic constants or molecular properties, such as the dipole moment. Fortunately, reliable experimental data regarding spectroscopic and molecular properties are increasing. In this regard, although small, diatomic molecules can be extremely effective benchmarking model systems as they show a wide variety of different bonding and spin configurations, they are expected to reflect some trends of polyatomic systems.7 Indeed, experimental diatomic test sets have become an attractive choice in various benchmarks for DFT and wavefunction methods for properties such as equilibrium geometries and bond energies. 1,3-17 However, investigations focusing on the performance of CCSD(T) against experimental dipole moments are still minimal for dipole moments.9,28

This work extensively studies the accuracy and limitations of the performance of CCSD(T) regarding dipole moments of diatomics. First, our study compares the performance of CCSD(T) methods with different basis sets against experimentally measured dipole moments collected recently in ref. 29.

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Table 1 Molecules in the dataset classified by classes of their constituent elements

Molecules
Alf $(X^1\Sigma^+)$ , Gaf $(X^1\Sigma^+)$ , InCl $(X^1\Sigma^+)$ , InF $(X^1\Sigma^+)$
GeTe $(X^1\Sigma^+)$ , GeO $(X^1\Sigma^+)$ , GeS $(X^1\Sigma^+)$ , PbO $(X^1\Sigma^+)$ , PbS $(X^1\Sigma^+)$ , SiO $(X^1\Sigma^+)$ ,
SiS $(X^1\Sigma^+)$ SnO $(X^1\Sigma^+)$ , SnS $(X^1\Sigma^+)$
BrO $(X^2\Pi_{3/2})$ , CF $(X^2\Pi)$ , IBr $(X^1\Sigma^+)$
CN $(X^2\Sigma^+)$ , CO $(X^1\Sigma^+)$ , CS $(X^1\Sigma^+)$ , CSe $(X^1\Sigma^+)$ , NO $(X^2\Pi_{1/2})$ , PN $(X^1\Sigma^+)$ , PO $(X^2\Pi)$ , SO $(X^3\Sigma^-)$
AgBr $(X^1\Sigma^+)$ , AgF $(X^1\Sigma^+)$ , AgI $(X^1\Sigma^+)$ , CuF $(X^1\Sigma^+)$ , YF $(X^1\Sigma^+)$
CON $(X^{\Sigma^+})$ , CO $(X^{1}\Sigma^+)$ , CS $(X^{1}\Sigma^+)$ , CSe $(X^{1}\Sigma^+)$ , NO $(X^{2}\Pi_{1/2})$ , PN $(X^{1}\Sigma^+)$ , PO $(X^{2}\Pi)$ , SO $(X^{3}\Sigma^-)$ AgBr $(X^{1}\Sigma^+)$ , AgF $(X^{1}\Sigma^+)$ , AgI $(X^{1}\Sigma^+)$ , CuF $(X^{1}\Sigma^+)$ , YF $(X^{1}\Sigma^+)$ HfO $(X^{1}\Sigma^+)$ , ScO $(X^{2}\Sigma^+)$ , ZrO $(X^{1}\Sigma^+)$

Second, and in line with the previous point, the equilibrium bond length and harmonic frequency are tested against the experimental data. The dataset consists of 32 diatomic molecules involving both main-group and transition metal elements with divergent bond natures. The dipole moments are obtained from well-controlled experimental measurements such as microwave spectroscopy, with reported uncertainties typically below 0.05 D.

### 2 Methodology

#### Computational approach

Our computations are performed at the core-correlated CCSD(T) level of theory using the CFOUR package<sup>30</sup> by means of unrestricted Hartree-Fock (UHF) orbitals as a reference. For elements with Z > 36, we use effective core potentials. The basis sets employed in this work include the augmented Dunning's weighted core-valence basis set aug-cc-pwCVT/QZ(-PP) to account for core-correlations<sup>2,31-46</sup> and a computationally much cheaper basis set, the segmented def2-QZVPP series basis sets developed by Ahlrichs et al. 47-49 In the case of closedshell molecules with def2-QZVPP basis, the results are provided via the Molpro package. 50,51

In this paper, we report the equilibrium dipole moment  $\mu_e$ and the zero-point vibrational corrected dipole moment  $\mu_0$ . To calculate  $\mu_0$ , for each molecule, we compute the potential energy curve in a point-wise manner using a grid of points ranging between  $0.4 \times R_e(\text{Exp})$  and  $3 \times R_e(\text{Exp})$ .† The spin states of molecules are determined as the spin state whose potential energy curve has the lowest energies compared with other spin states. The obtained spin states are consistent with experiments. Next, the equilibrium bond length  $R_e$  and harmonic vibrational frequency  $\omega_e$  are obtained by fitting the potential energy curve. The electric dipole moments  $\mu$  are calculated from analytic gradients at every single-point geometry, leading to the dipole moment at the equilibrium bond length  $\mu_e$  and the zero-point vibrational corrected dipole moment  $\mu_0$ . The latter includes vibrational average corrections, using the discrete variable representation (DVR) method for the vibrational wavefunctions,‡ and the overlap obtained by numerical integration. In this work, only the magnitude (not the direction) of the dipole moments is discussed.

For Dunning basis sets, the CBS limits are predicted using the standard two-point extrapolation scheme<sup>52-54</sup> as

Predicted CBS
$$(n_1/n_2)^{\text{corr}} = \frac{n_1^3 d_1 - n_2^3 d_2}{n_1^3 - n_2^3},$$
 (1)

where  $d_i$  denotes the correlation contribution of molecular property ( $\omega_e$  or  $\mu$ ) evaluated at a given basis set, specified by  $n_i$ . In particular,  $n_1$  and  $n_2$  are equal to 3 and 4 for the aug-ccpwCVTZ and aug-cc-pwCVQZ basis sets, respectively. In contrast, it is unnecessary to extrapolate for  $R_e$  since predictions at the quadruple- $\zeta$  level are already very close to convergence.

For elements calculated with the aug-cc-pwCVT/QZ-PP basis set, the relativistic effects have already been considered in the effective core potentials, showing that these have a minimal effect.<sup>55</sup> Nevertheless, a scalar relativistic effect correction has been obtained by the second-order Douglas-Kroll-Hess approximation (DK) implemented in Molpro for selected molecules when (aug-)cc-p(w)CVT/QZ-DK(3) basis sets<sup>56</sup> available.

#### 2.2 Dataset

Experimentally, the equilibrium bond length, harmonic frequency, and dipole moment have been measured in more than 100 diatomic molecules.<sup>29</sup> We selected 32 molecules to explore in depth with high-level quantum chemistry that was representative of the diatomics' chemical diversity; see Table 1 for molecules considered. We divide them into six classes to aid interpretation, with the number of molecules of each class shown in Fig. 1, following the same graphical representative, we will use in this paper to show results. The dataset includes various main-group metal and non-metal compounds showing

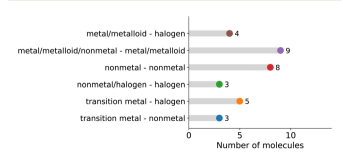


Fig. 1 Number of molecules in the present dataset classified by classes of their constituent elements.

<sup>†</sup> In particular, we use the points -0.4x, -0.35x, -0.3x, -0.24x, -0.18x, -0.12x,  $-0.08x, \ -0.04x, \ -0.02x, \ 0, \ 0.02x, \ 0.04x, \ 0.08x, \ 0.12x, \ 0.2x, \ 0.28x, \ 0.36x, \ 0.45x, \ 0.08x, \ 0.12x, \ 0.2x, \ 0.28x, \ 0.36x, \ 0.45x, \ 0.2x, \ 0.2x,$ 0.5x, 0.6x, 0.8x, 1.0x, 1.5x, 2.0x, 3.0x, where  $x = R_e(\text{Exp})$ .

<sup>±</sup> We use 200 DVR points to guarantee convergence on the vibrational energies

PbO 0.3 Uncertainty (D)
0
0
1 InC

Dipole moment (D) Fig. 2 Uncertainties of experimental dipole moments of the molecules included in the dataset

6

3

covalent and ionic bonds. In particular, the dataset contains 8 transition metal compounds.

The accuracy of the experimentally measured electric dipole moments depends on determining the magnitude of the applied electric field and its homogeneity. In constructing the current dataset, focus on molecules with dipole moments determined by high-resolution spectroscopic measurements, especially microwave spectroscopy and molecular beam electric resonance, excluding molecules with considerable uncertainty in dipole moments such as BF and RbI. As shown in Fig. 2, in this dataset, typical error bars of 0.1-5% are found, which translates into errors less than 0.05 D for most of the molecules under consideration. PbO, AgF, InCl, PbS, SnS and SnO are labeled in the figure since they show uncertainties over 0.1 D.

#### 3 Results and discussions

The performance of CCSD(T) on the equilibrium bond length,  $R_{\rm e}$ , harmonic vibrational frequency,  $\omega_{\rm e}$ , of diatomic molecules has been exhaustively investigated in the literature by comparing with experiments. Not only the accuracy of basis set families, 31,33,41 but also that of the conjunct relativistic pseudopotentials has been discussed and compared with fullelectron relativistic treatments. 36-40,42,43,57 Recently, it has been found that by using non-HF orbitals, the accuracy of CCSD(T)  $\omega_{\rm e}$  can be further improved for diatomic molecules consisting of row 2 and row 3 elements. 17 For transition metal diatomic molecules, the accuracy of CCSD(T) on  $R_e$  and  $\omega_e$ , as well as the influence of relativistic effect and multi-reference character have been investigated based on a dataset of 60 molecules.<sup>55</sup> Nevertheless, we first report the calculated  $R_e$  and  $\omega_e$  for the dataset employed in this work. Then, we present and discuss the theoretical predictions of dipole moments compared to their experimental values for the molecules in the dataset.

The performance of computations versus experimental data is tested via:

• Residuals defined as the difference between the experimental value of a molecular property and its computed value,  $x_i(\text{Exp.}) - x_i(\text{CCSD(T)})$ , where  $x_i$  is the experimental or calculated properties for molecule i.

· Root mean squared error (RMSE) defined as

$$RMSE = \sqrt{\frac{\sum_{i}^{N} (x_i(Exp) - x_i(CCSD(T))^2)}{N}},$$
 (2)

where N stands for the number of molecules in the dataset.

We present the results in two formats for each computational method investigated and molecular property under consideration. First, we present residual errors of all molecules, labeling molecules with significant errors. Second, we perform a statistical analysis of the errors by presenting the RMSE grouped by molecular class as designated in Table 1. In addition, when needed, we report the relative error  $r_{\rm E}$  as

$$r_{\rm E} = \frac{1}{N} \sum_{i}^{N} \frac{|x_i({\rm Exp.}) - x_i({\rm CCSD}({\rm T}))|}{x_i({\rm Exp.})}.$$
 (3)

Finally, we show results based on the aug-cc-pwCVQZ and def2-OZVPP basis sets. The def2-OVPP basis set has just over half the number of basis functions as auc-cc-pwCVQZ (actually very similar in size to aug-cc-pwCVTZ).

#### 3.1 Equilibrium bond length

When analyzing  $R_e$ , both aug-cc-pwCVQZ and def2-QZVPP basis sets give very accurate predictions, showing an RMSE  $\lesssim 0.008 \, \text{Å}$  $(r_{\rm E} \lesssim 0.2\%)$ , as shown in Fig. 3. In this figure, it is observed that non-metal diatomics, metal/metalloid/nonmetal-metal/

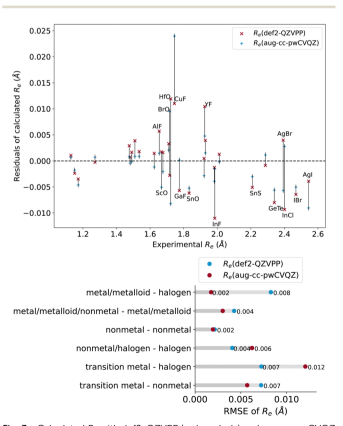


Fig. 3 Calculated R<sub>e</sub> with def2-QZVPP (red symbols) and aug-ccpwCVQZ (blue symbols) basis sets. Upper panel: residuals of the calculated  $R_{\rm e}$ Lower panel: RMSE of the computed  $R_{\rm e}$  for different classes of molecules.

metalloid molecules show the smallest RMSE  $\simeq 0.002$  Å  $(r_{\rm E} \simeq 0.1$ –0.2%), while some transition-metal-containing diatomics display more significant errors, with RMSE  $\simeq 0.01 \text{ Å}$  $(r_{\rm E} \simeq 0.5\%)$ . Furthermore, some outliers are noticeable, such as CuF, AgBr and AgI.

In the case of CuF, Aoto et al. have shown that the error of R<sub>e</sub> is reduced by including relativistic corrections, 55 and our results confirm these findings, as discussed in Section 3.3.3. In contrast, for some other transition-metal-containing molecules, for example, ScO or AgF, the scalar relativistic effect correction is negligible. Another source of discrepancy may be due to multi-reference effects; on that front, it has been shown that using multi-reference coupled-cluster theory for ScO and AgF leads to results similar to the single-reference results. 55 On the other hand, silver-halogen molecules have received little attention, except for AgF and AgCl, showing a negligible relativistic effect on the equilibrium bond length prediction. Therefore, in light of these results, we think that relativistic effects could not explain the significant error in the predictions of  $R_e$ for AgBr and AgI.

Our def2-QZPP calculations show systematically large  $R_e$ values further from the experiment than aug-cc-pwCVQZ calculations. For the aug-cc-pwCV basis set, increasing the number of basis functions usually improves the predictions of  $R_e$ towards experimental values. From the aug-cc-pwCVTZ basis set to the quadruple-ζ level, the RMSE for most classes of molecules can be significantly reduced to around 50%. Exceptions are the transition metal-containing molecules, for which the aug-cc-pwCVT/QZ basis set gives similar RMSEs. However, the reduction of basis functions in the def2-OZVPP basis sets does not necessarily worsen the  $R_e$  predictions. The RMSEs with the def2-QZVPP basis set are generally comparable with the aug-cc-pwCVQZ basis set for most molecules studied, except metal/metalloid halides. In particular, for transition metal halides, the def2-QZVPP basis set can be much closer to the experimental results than the aug-cc-pwCVQZ basis set.

#### 3.2 Vibrational harmonic frequency

The CCSD(T) performance for the vibrational harmonic frequency versus experimental data is shown in Fig. 4, where we can identify the absolute error of CCSD(T) calculations and its RMSE for each molecular class. The relative errors of CCSD(T) were compared to experimental  $r_{\rm E} \lesssim 2\%$  for most molecules. The RMSEs with the predicted CBS(aug-cc-pwCVT/QZ) and def2-QZVPP basis sets are 24.7 and 16.3 cm<sup>-1</sup>, respectively. We find that the calculations of  $\omega_e$  of diatomics involving maingroup metal elements are very accurate with both basis sets, leading to an RMSE  $\lesssim 10 \text{ cm}^{-1}$ . These findings align with those of Aoto et al., showing that the main-group nonmetal diatomics have large errors,55 and the most significant outliers include CN and NO. These are correlated to the poor UHF reference. It has been recently shown that these errors can be healed by using non-HF.<sup>17</sup>

For most molecules, the predictions of  $\omega_e$  are already close to experiments at the aug-cc-pwCVTZ level. The difference between the RMSEs from the aug-cc-pwCVTZ and aug-cc-

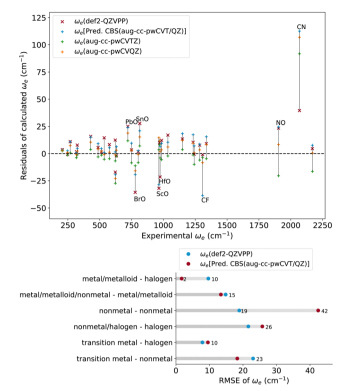


Fig. 4 Calculated  $\omega_{\rm e}$  with def2-QZVPP (red symbols) and augccpwCVQZ (blue symbols) basis sets. Upper panel: residuals of the calculated  $R_a$ . Lower panel: RMSE of the computed  $R_a$  for different classes of molecules.

pwCVQZ basis sets is only  $\sim 2.5 \text{ cm}^{-1}$ . Similar to  $R_e$ , the RMSE of the def2-QZVPP basis set is very close to that of the aug-cc-pwCVQZ basis set, sometimes even slightly better than that of the latter. However, for nonmetal diatomic molecules, the employment of the aug-cc-pwCV basis set significantly improves the results.

#### 3.3 Dipole moment

#### 3.3.1 Effect of methodology

3.3.1.1 Vibrational corrections on dipole moment predictions. Experimental dipole moments usually deviate from the theoretical ones at the equilibrium bond length due to the anharmonicity of the molecular interaction. Vibrational corrections have been shown to be important for accurate dipole moment calculations versus experimental values. 28,58-61 The importance of vibrational corrections depends on the anharmonicity of the underlying potential energy curve. In the case of dipole moment, as a first-order molecular property, it can be important to count for the deviation between the equilibrium bond length (given by the potential energy curve) and the most probable interatomic distance given by the ground state vibrational wave function. Therefore, apart from reporting the dipole moment at the equilibrium bond length,  $\mu_e$ , we calculate the vibrational average dipole moment  $\mu_0$  by numerically averaging the radial-dependent dipole moment with the vibrational ground state wavefunction of the molecule under consideration. In particular, we use a DVR approach to solve the single-

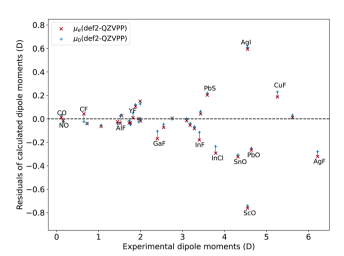


Fig. 5 Dipole moment errors calculated with def2-QZVPP basis set experimental results, with or without vibrational corrections

channel Schrödinger equation associated with the vibrational degrees of freedom over the obtained Born-Oppenheimer potential energy curve.

Fig. 5 and 6 show the residuals of CCSD(T) calculations on the dipole moments for the molecules of our dataset for the def2-QZVPP and CBS(aug-cc-pwCVT/QZ) basis set. As a result, we notice a trivial influence of the vibrational average on the dipole moment and consistent differences from the vibrational average with the two basis sets ( $\sim 0.01$  D, 2%). However, we find a few outliers: diatomics with light elements and short bond lengths such as CO ( $\sim$ 20%), NO ( $\sim$ 10%), CF ( $\sim$ 10%), AlF  $(\sim 4\%)$ , GaF  $(\sim 3\%)$ , and YF  $(\sim 2\%)$ . Indeed, for these outliers, the difference introduced by vibrational average correction correlates with their harmonic vibrational frequency. Finally, due to the slightly better performance of the vibrational average dipole moment, we will use  $\mu_0$  when referring to dipole moments from now on.

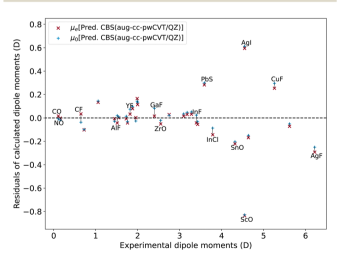


Fig. 6 Dipole moment errors calculated with aug-cc-pwCVQZ basis set experimental results, with or without vibrational versus corrections

3.3.1.2 Basis set family and size. A detailed study on the role of basis sets on the dipole moment is shown in Fig. 7 and 8, shown as the residuals of the calculated dipole moment with respect to the experiments. In particular, we investigate the performance of CCSD(T) dipole moments calculated with ccpwCV, aug-cc-pwCV, and def2-QZVPP basis sets. For most molecules with larger basis sets, the predicted magnitude of the dipole moment increases.

In the case of the cc-pwCV basis set, we observe a significant basis set size effect. For most molecules, increasing the size of the basis set from cc-pwCVTZ to cc-pwCVQZ reduces the underestimation of the dipole moment, improving the RMSE from 0.30 D to 0.24 D. In particular, when using the cc-pwCV basis set, molecules containing metal/metalloid elements are sensitive to the size of the basis set. For molecules with main-group metal/metalloid elements, increasing the size of the basis set from cc-pwCVTZ to cc-pwCVQZ halves the RMSE. In contrast, for the nonmetal-nonmetal and nonmetal/halogen-halogen molecules, the improvement of the dipole moment from the triple- $\zeta$  to quadruple- $\zeta$  level is less than 0.02 D. Therefore, it is

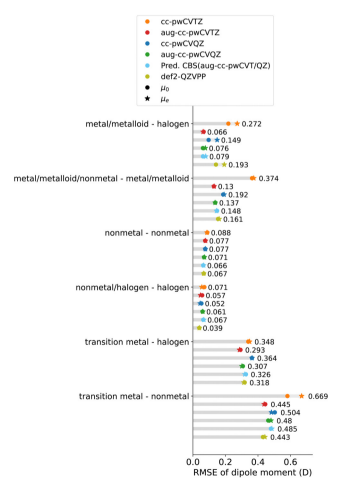


Fig. 7 RMSE of the CCSD(T) dipole moment calculated with cc-pwCVT/ QZ, aug-cc-pwCVT/QZ and def2-QZVPP basis sets. The dipole moments at the equilibrium bond length  $\mu_{\rm e}$  and the zero-point vibrational corrected dipole moment  $\mu_0$  are denoted as circles and stars, respectively

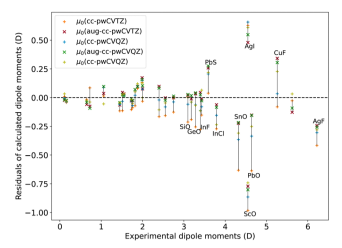


Fig. 8 The errors of dipole moment calculated with cc-pwCVT/QZ and aug-cc-pwCVT/QZ basis sets.

sufficient to use the cc-pwCVTZ basis set for molecules with nonmetal elements.

With the aug-cc-pwCV basis set, it is noticed that dipole moments are almost converged at the aug-cc-pwCVTZ level. As a result, the benefits of using the larger aug-cc-pwCVQZ basis set are  $\leq 0.01$  D, while in some cases, especially for diatomics with metal atoms, the aug-cc-pwCVTZ predictions are slightly more accurate than those of the aug-cc-pwCVQZ level. The role of augmented functions will be discussed in Section 3.3.1.3.

Furthermore, and surprisingly enough, for most molecules, using the def2-QZVPP basis set leads to a similar accuracy to that of the aug-cc-pwCVQZ basis set, even though the latter is computationally much more expensive. For some molecules, the def2-QZVPP basis set yields a dipole moment closer to the experimental one than the CBS(aug-cc-pwCVT/QZ).

3.3.1.3 Influence of diffuse functions. It has been shown that for dipole moments, diffuse functions augmented to the basis set play an important role in hybrid and double-hybrid density functionals, and wave function-based methods can be applied to get closer dipole moment predictions to CCSD(T). 22,62 To further explore if diffuse functions help CCSD(T) predictions to approach experiments, the cc-pwCVT/QZ basis sets have been used to calculate the dipole moments and compared to the results from augmented basis sets, shown in Fig. 8. As discussed in Section 3.3.1.2, from the cc-pwCVTZ to the quadruple- $\zeta$  level, the prediction of the dipole moment's magnitude increases. Adding the augmentation functions further increases the dipole moment's magnitude, leading, in some cases, to its overestimation. Molecules containing metal elements are more sensitive to augmentation than nonmetal molecules because of the longer-range nature of the wave function. On average, the RMSE can be reduced by 0.10 D by augmenting the cc-pwCVTZ basis set, while at the quadruple-ζ level, the overall improvement from augmentation is only 0.03 D, much smaller than the triple- $\zeta$  level. Therefore, the improvement from including augmented functions is negligible at the quadruple-ζ level for dipole moments of most molecules.

Exceptions are metal/metalloid halides (metal/metalloidhalogen and transition metal-halogen molecules), for which the improvement of RMSE by employing diffuse functions at the quadruple- $\zeta$  level is  $\sim 0.07$  D. Therefore, for these molecules, it is suggested to use augmented basis sets.

3.3.2 Overall performance of CCSD(T) on the dipole moments. The performance of CCSD(T) predictions on the dipole moment  $\mu_0$  with the CBS(aug-cc-pwCVT/QZ) and def2-QZVPP basis sets is shown in Fig. 9. The overall performance of the Dunning's and def2-basis sets are very similar, showing an RMSE of 0.215 D and 0.209 D, respectively. Significant errors over 0.2 Debye are only seen for molecules with dipole moments greater than 3 Debye, though there is no correlation between dipole moment error and its absolute value.

In particular, diatomics containing main-group elements, especially non-metal elements, are well described with  $r_{\rm E} \lesssim$ 5%: RMSE ≤ 0.08 D for nonmetal-nonmetal molecules and RMSE  $\leq 0.5$  D for nonmetal halides. For molecules containing main-group metals/metalloids, the RMSE becomes larger ( $\lesssim 0.15$  D), but still close to the experimental uncertainty, except SnO and PbS.

However, in stark contrast to the main-group molecules, larger errors are observed in systems involving transition metals. In particular, with the predicted CBS(aug-cc-pwCVT/ QZ), we observe an RMSE of 0.32 D ( $r_E = 5.5\%$ ) and 0.51 D ( $r_E =$ 6.9%) for transition metal halides and other transition metalnon metal diatomics, respectively, much larger than the experimental uncertainty.

The source of the discrepancy between CCSD(T) and experiments can be various. As the predicted CBS is obtained from a relatively large basis set, we believe the errors are not associated with the size of the basis set. Another source is the possible multi-reference character of the molecules. However, in the current dataset, the transition-metal-containing molecules or their analogs are generally dominated by singe-reference characters.<sup>55,63</sup> Besides, we notice that the residuals of

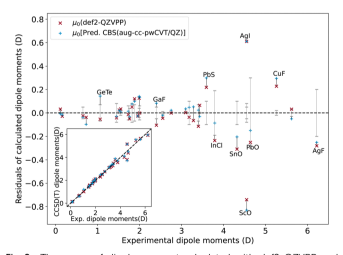


Fig. 9 The errors of dipole moment calculated with def2-QZVPP and aug-cc-pwCVQZ basis sets. The error bars of experimental measurements are shown in grav

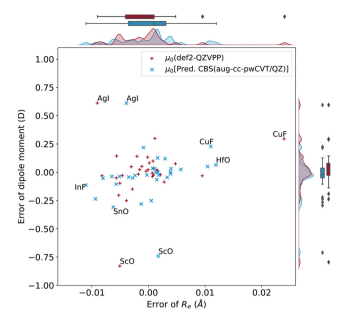


Fig. 10 Errors of dipole moments as a function of errors of  $R_e$  with def2-QZVPP and predicted CBS(aug-cc-pwCVT/QZ)

CCSD(T) predictions from experiments are not explicitly correlated with the experimental uncertainties. Although for some molecules, for example, PbO and InCl, the residual is closer to the experimental uncertainty, for other molecules, like PbS, the residual is much larger than the experimental uncertainty. In the following section, the possible sources of errors will be explored further in detail.

**3.3.3 Origin of the errors.** One possible source of error in calculated dipole moments is an inaccurate prediction of bond lengths (consider classically that dipole moment is charge separation multiplied by bond length). We explore this possibility in Fig. 10, but find no strong relationship between the errors in the equilibrium bond length and the errors in the dipole moment, which agrees with previous studies on the nature of the dipole moment of diatomic molecules.<sup>29</sup> For example, both aug-cc-pwCV and def2-basis sets overestimate the dipole moment of AgI and CuF and simultaneously underestimate the dipole moment of ScO, even if  $R_e$  has been accurately predicted with aug-cc-pwCVQZ for these molecules. In contrast, several molecules with precise dipole moments show more significant errors in the prediction of  $R_e$ , e.g., HfO,

IBr, InF. Similarly, the errors of dipole moment are not correlated with the errors of  $\omega_e$ . These results suggest that benchmark studies based only on energetic properties may fail to foresee other properties relative to electron density.

As noticed in Fig. 9, there are a few outliers that require some extra discussion. Naturally, one would expect that the error resides in the single-reference nature of CCSD(T) calculations. However, it has been previously shown that the molecules displaying significant errors on dipole moments show a dominant single-reference nature.55

Another possible source of errors is the non-relativistic treatment. We perform CCSD(T) calculations, including the scalar relativistic correction, to investigate the role of the relativistic effect. In particular, we focus on 9 molecules showing significant errors in the dipole moment. The dipole moments  $\mu_e$  calculated at experimental geometry, with or without relativistic correction, are summarized in Table 2. We notice that the inclusion of the relativistic effect slightly decreases the magnitude of the predicted dipole moment. The difference of  $\mu_e$  introduced by the relativistic effect is commonly tiny, 0.01-0.07 D (0.3-1.5%), except CuF, for which the difference is 0.14 D (2.7%). For the molecules whose dipole moments are overestimated by non-relativistic CCSD(T) treatments (e.g., PbS, AgI, CuF), the relativistic dipole moment becomes closer to the experiment. In contrast, for other molecules, the underestimation of the dipole moment is further aggravated by the relativistic correction. Overall, including the relativistic correction, the RMSE of the 9 molecules can be slightly improved from 0.252 D to 0.235 D.

To further investigate the origin of the errors, in the following paragraphs, we analyze individual molecules with the most significant errors, including ScO, CuF and AgI.

In the case of CuF, the dipole moment is overestimated by 0.28 D at the CCSD(T)/CBS(aug-cc-pwCVT/QZ) level compared to the experimental value of 5.26(2) D, taken from the recent supersonic molecular beam high-resolution optical Stark spectrum.73 Furthermore, we observe a small divergence of 0.02 Å between experimental R<sub>e</sub> and CCSD(T)/aug-cc-pwCVQZ prediction. At experimental  $R_{\rm e}$ , the predicted dipole moment  $\mu_{\rm e}$  is 5.420 D with CCSD(T)/aug-cc-pwCVQZ. Interestingly, when applying DK relativistic corrections to CCSD(T) with the corresponding relativistic-contracted Douglas-Kroll basis sets (Cu:augcc-pwCVQZ-DK; F:aug-cc-pwCVQZ-DK), the experimental Re is

Table 2 Dipole moment at experimental equilibrium bond length  $\mu_{ev}$  calculated with non-relativistic or scalar relativistic CCSD(T)

		Non-rela	ativistic	Scalar re	elativistic
Molecule	$\mu_0$ (Exp.) (D)	μ <sub>e</sub> (D)	Basis set	μ <sub>e</sub> (D)	Basis set
AgF	6.22(20)	5.991	Ag:cc-pwCVTZ-PP; F:cc-pVQZ	5.956	Ag:cc-pwCVTZ-DK; F:cc-pVQZ-DK
AgBr	5.62(3)	5.789	Ag:cc-pwCVTZ-PP; Br:cc-pVQZ	5.716	Ag:cc-pwCVTZ-DK; Br:cc-pVQZ-DK
AgI	4.55(5)	5.139	Ag:cc-pwCVTZ-PP; I:cc-pwCVTZ-PP	5.087	Ag:cc-pwCVTZ-DK; I:cc-pwCVTZ-DK3
CuF	5.26(2)	5.420	Cu:aug-cc-pwCVQZ; F:aug-cc-pwCVQZ	5.278	Cu:aug-cc-pwCVQZ-DK; F:aug-cc-pwCVQZ-DK
InCl	3.79(19)	3.629	In:aug-cc-pwCVQZ-PP; Cl:aug-cc-pwCVQZ	3.616	In:aug-cc-pwCVQZ-DK3; Cl:aug-cc-pwCVQZ-DK
PbO	4.64(30)	4.479	Pb:aug-cc-pwCVQZ-PP; O:aug-cc-pCVQZ	4.460	Pb:aug-cc-pwCVQZ-DK3; O:aug-cc-pCVQZ-DK
PbS	3.59(10)	3.726	Pb:aug-cc-pwCVQZ-PP; S:aug-cc-pCVTZ	3.669	Pb:aug-cc-pwCVQZ-DK3; S:aug-cc-pCVTZ-DK
SnO	4.32(10)	4.106	Sn:aug-cc-pwCVQZ-PP; O:aug-cc-pCVQZ	4.074	Sn:aug-cc-pwCVQZ-DK3; O:aug-cc-pCVQZ-DK
SnS	3.18(10)	3.190	Sn:aug-cc-pwCVQZ-PP; S:aug-cc-pCVQZ	3.147	Sn:aug-cc-pwCVQZ-DK3; S:aug-cc-pCVQZ-DK

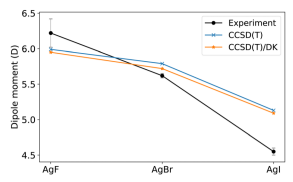


Fig. 11 Dipole moments of AgF, AgBr, and AgI. The theoretical predictions are calculated at experimental  $R_{\rm e}$  with CCSD(T)/(Ag:cc-pwCVTZ-PP; F/Br: cc-pVQZ; I:cc-pwCVTZ-PP), and CCSD(T)/(Ag:cc-pwCVTZ-DK; F/Br:cc-pVQZ-DK; I:cc-pwCVTZ-DK) with scalar DK relativistic corrections. The experimental uncertainties are shown in gray.

perfectly reproduced with an error of 0.001 Å, and the dipole moment obtained is 5.28 D, which is consistent with previous reports<sup>87</sup> and perfectly reproduces experimental values. This finding aligns with the previous studies on CuH, AgH, and AuH, <sup>88</sup> where small changes in the predicted  $R_{\rm e}$  by the inclusion of relativistic effect have been observed to cause considerable change in  $\mu_{\rm e}(R_{\rm e})$ .

The analysis of the Stark effect on the rotational spectrum of AgI shows a dipole moment of 4.55(5) D.66 However, our predicted CBS(aug-cc-pwCVT/QZ) gives 5.15 D, including the vibrational average correction, which overestimates the experimental measurement. However, it is in close agreement with previous theoretical predictions.<sup>89</sup> In contrast, for AgF (isovalent analogues of AgI), CCSD(T) underestimates the dipole moment. Furthermore, silver-halogen molecules, experimentally, show a drastic decrease in the dipole moment as the halogen element becomes heavier, whereas, theoretically, the decrease is less steep, as shown in Fig. 11. The disagreement between CCSD(T) calculations and experimental results can not be healed by including a scalar relativistic effect. The difference of dipole moment obtained by relativistic CCSD(T)/(Ag:cc-pwCVTZ-DK; I:cc-pwCVTZ-DK) and non-relativistic CCSD(T)/(Ag:cc-pwCVTZ-PP; I:cc-pwCVTZ-PP) is only 0.05 D at the experimental  $R_e$ . Additionally, we notice that there is an earlier measurement from the same group, giving  $\mu_0$  = 5.10(15) D, 90 which agrees with our CCSD(T)/(aug-cc-pwCVT/QZ) predictions. Therefore, the disagreement calls for revision.

#### 4 Conclusions

In this work, we have benchmarked the performance of CCSD(T) for predicting dipole moments by comparing computational results based on large basis sets with accurate experimental results. In particular, we study 32 diatomic molecules with diverse bonding and elemental composition whose experimental dipole moments are available. Thus, we use experimental information to benchmark a computational approach directly. We also consider the accuracy of the equilibrium bond length and vibrational harmonic frequency and compare them to the accuracy of the dipole moment prediction.

We find that single-reference CCSD(T) calculations, using the def2- and aug-cc-pwCVX (where X = T and Q) family of basis set, satisfactorily describe the dipole moment of most of the molecules in the dataset, with errors usually less than 0.15 D, especially for molecules containing only main-group elements. However, this accuracy is contingent on selecting the most appropriate basis sets depending on the molecule under consideration. For instance, we have shown that for the cc-pwCV basis set at the triple- $\zeta$  level, diffuse functions play an essential role in molecules containing metal elements. However, augmentation with diffuse functions slightly improves the dipole moment predictions at the quadruple-ζ level where the basis set is already large enough. Similarly, the basis set incompleteness error is evident at the triple-ζ level when employing the cc-pwCV basis set, while the augmented counterpart gives almost converged results. Surprisingly, the def2-QZVPP basis set shows good comparable performance compared to the much larger aug-ccpwCVOZ basis set. Therefore, the def2-OZVPP basis set should be the choice to predict dipole moments in larger systems.

Non-relativistic predictions are generally accurate enough for most molecules. Although, for some molecules like CuF, scalar relativistic corrections may have an essential role in the dipole moment calculations. For some molecules, *e.g.*, ScO, AgI, we observe significant divergence with experiments that can not satisfactorily be explained by the multi-reference or relativistic effect. We recommend for these unusual molecules a careful consideration of both experimental and theoretical (*e.g.*, multi-reference coupled-cluster) results to ensure a better understanding of dipole moment in these systems. Furthermore, an extension of the current dataset, *e.g.*, alkali metal molecules, would be desirable for future benchmarks.

Finally, we have shown that errors in the prediction of the dipole moment do not correlate with the accuracy of the equilibrium bond length, showing that errors in the dipole moment must be explained by errors in the electron distribution, not just differing bond lengths and reinforcing the fact that different properties are predicted with different accuracies with computational approximations. Therefore, benchmark studies on methods regarding energetic and geometric properties do not guarantee strong performance for other properties.

#### Conflicts of interest

There are no conflicts to declare.

# **Appendices**

# Appendix: Experimental *versus* calculated equilibrium bond length $R_{\rm e}$ , harmonic vibrational frequency $\omega_{\rm e}$ and electric dipole moment $\mu$

Table 3 lists the experimental electric dipole moments employed in this work, including the pertinent references.

Table 3 The experimental and calculated dipole moments in Debye

1	l																														
μ <sub>0</sub> (def2- QZVPP)	5.652 5.940	5.160	1.540	1.709	0.625	1.407	0.144	1.932	2.116	5.488	2.295	3.217	1.999	1.006	3.496	0.688	3.555	3.285	0.128	2.750	2.001	4.389	3.809	1.584	3.809	3.100	1.716	4.011	3.143	1.871	2.505
μ <sub>e</sub> (def2- QZVPP)	5.631 5.898	5.144	1.478	1.727	0.694	1.426	0.123	1.960	2.140	5.446	2.230	3.198	1.981	0.995	3.473	0.685	3.497	3.219	0.143	2.757	1 984	4.373	3.792	1.578	3.788	3.082	1.695	3.995	3.125	1.830	2.478
$\mu_0$ [CBS(aug-cc-pwCVT/QZ)]	5.569	5.161	1.535	1.728	0.613	1.421	0.104	1.932	2.131	5.555	2.482	3.334	2.133	1.204	3.400	0.628	3.703	3.424	0.139	2.773	1 979	4.490	3.889	1.559	3.721	3.133	1.744	4.116	3.227	1.895	2.530
μ <sub>e</sub> [CBS(aug-cc- pwCVT/QZ)]	5.548 5.927	5.145	1.471	1.717	0.685	1.444	0.129	1.961	2.155	5.513	2.416	3.314	2.114	1.192	3.376	0.624	3.646	3.358	0.156	2.780	1 959	4.471	3.872	1.552	3.713	3.114	1.723	4.098	3.210	1.853	2.500
$\mu_0$ (aug-cc-pwCVQZ)	5.530 5.964	5.098	1.540	1.737	0.611	1.407	0.094	1.943	2.147	5.566	2.483	3.323	2.103	1.157	3.405	0.634	3.707	3.425	0.134	2.764	1 976	4.486	3.866	1.566	3.748	3.125	1.720	4.103	3.197	1.890	2.530
μ <sub>e</sub> (aug-cc- pwCVQZ)	5.509 5.922	5.082	1.476	1.729	0.680	1.431	0.119	1.972	2.171	5.524	2.417	3.303	2.084	1.146	3.381	0.630	3.650	3.358	0.149	2,771	1 958	4.467	3.849	1.559	3.721	3.106	1.699	4.085	3.180	1.848	2.502
$\mu_0$ (aug-cc-pwCVTZ)	5.493	5.030	1.560	1.734	0.591	1.387	0.090	1.954	2.163	5.603	2.498	3.315	2.076	1.096	3.420	0.651	3.728	3.445	0.132	2.751	1 978	4.489	3.849	1.581	3.779	3.118	1.703	4.097	3.170	1.890	2.549
$\mu_{\rm e}$ (aug-cc-pwCVTZ)	5.473 5.935	5.015	1.496	1.747	0.662	1.413	0.114	1.983	2.186	5.561	2.431	3.295	2.057	1.086	3.396	0.647	3.672	3.378	0.144	2.758	1 962	4.470	3.833	1.574	3.793	3.099	1.683	4.079	3.153	1.848	2.521
$\mu_0$ (cc-pwCVQZ)	5.916	5.208	1.484	1.693	0.619	1.390	0.104		2.142	5.294	2.381	3.220	2.074	1.151	3.352		3.637	3.288	0.121	2.715	1 983	4.305	3.796	1.568	3.685	3.041	1.701	3.955	3.142	1.840	2.471
$\mu_{\rm e}$ (cc-pwCVQZ)	5.877	5.191	1.423	1.710	0.687	1.415	0.127		2.165	5.256	2.318	3.202	2.056	1.140	3.330		3.581	3.227	0.135	2.722	1 966	4.289	3.780	1.564	3.727	3.025	1.681	3.939	3.125	1.800	2.445
$\mu_0$ (cc-pwcvtz)	5.594 5.803	5.178	1.401	1.673	0.635	1.335	0.121		2.123	5.180	2.235	3.028	1.969	1.073	3.280	0.811	3.518	3.124	0.115	2.626	1 959	4.004	3.629	1.566	3.567	2.886	1.626	3.688	2.990	1.749	2.395
$\mu_{\rm e}$ (cc-pwCVTZ)	5.574 5.767	5.161	1.343	1.690	0.700	1.362	0.143		2.146	5.144					3.258	0.687	3.465	3.066	$^{0}$ 0.126	2.634			3.614	1.566	3.419	5 2.871	1.608	3.675	2.974	1.711	
e Exp.		$(4.55(5)^{99})$	$(1.515(4)^{6})$	$X^2\Pi_{3/}$ 1.76(4) <sup>68</sup>	I 0.65(5) <sup>68</sup>						$2^{+} 2.45(5)^{74}$		$^{+}$ 2.00(6) <sup>76</sup>	$^{+}$ 1.06(7) <sup>77</sup>	$3.431(5)^{78}$		$3.79(19)^{79}$	$3.40(7)^{74}$	$I_{1/} 0.1595(15)^{80}$		1 1 88(7)82				$^{+}$ 4.55(8) <sup>84</sup>		$^{+}$ 1.73(6) <sup>85</sup>	$4.32(10)^{76}$	$(3.18(10))^{76}$	$^{+}$ 1.82(8) <sup>86</sup>	
Molecule State Exp.	$\begin{array}{ccc} AgBr & X^1\Sigma^+ \\ AgF & X^1\Sigma^+ \end{array}$			BrO X²∏	$\frac{2}{X^2}\Pi$			CS $X^1\Sigma^+$											NO $X^2\Pi_{1/}$		PO X <sup>2</sup> TI		PbS $X^1\Sigma^+$							$X^1\Sigma^+$	

Table 4 The experimental and calculated equilibrium bond length  $R_e$  (in Å) and harmonic vibrational frequency  $\omega_e$  (in cm<sup>-1</sup>). The experimental values are taken from ref. 91 and 92, or the same reference of the experimental dipole moment of the corresponding molecule

Molecule	State	R <sub>e</sub> (Exp.)	R <sub>e</sub> (aug-cc-pwCVTZ)	R <sub>e</sub> (aug-cc- pwCVQZ)	R <sub>e</sub> (def2- QZVPP)	$\omega_{\rm e}$ $\omega_{\rm e}$ (aug-cc-(Exp.) pwCVTZ)	$\omega_{\rm e}$ (aug-cc-pwCVQZ)	$\omega_{\rm e}$ [CBS(aug-cc-pwCVT/QZ)]	$\omega_{\rm e}$ (def2-QZVPP)
AgBr	$X^1\Sigma^{\scriptscriptstyle +}$	2.393	2.394	2.387	2.397	247.7 246.4	248.6	250.2	246.9
AgF	$X^1\Sigma^+$	1.983	1.982	1.979	1.982	513.5 512.0	515.7	518.1	515.0
AgI	$X^1\Sigma^+$	2.545	2.542	2.536	2.541	206.5 208.9	209.7	210.2	210.1
AlF		1.654	1.660	1.656	1.660	802.3 793.9	801.5	804.9	804.1
BrO	$X^2\Pi_{3/2}$	1.717	1.728	1.726	1.714	778.7 767.5	762.8	631.2	743.0
CF	$\overset{2}{X}{}^{2}\Pi$	1.272	1 277	1.273	1.272	1308.1 1304.5	1299.7	1269.4	1307.2
CN	$X^2\Sigma^+$	1.172	1.170	1.167	1.168	2068.6 2160.4	2175.5	2181.2	2108.1
CO	$X^1\Sigma^+$	1.128	1.132	1.129	1.129	2169.8 2153.4	2170.3	2177.5	2174.4
CS CS	$X^1\Sigma^+$		1.539	1.536	1.537	1285.1 1279.1	1288.5	1293.7	1292.9
CSe	$X^1\Sigma^+$	1.676	1.678	1.674	1.678	1035.4 1033.1	1041.4	1045.8	1052.4
CuF	$X^1\Sigma^+$	1.745	1.773	1.769	1.756	622.7 595.3	599.7	603.3	605.6
GaF	$X^1\Sigma^+$		1.778	1.775	1.769	622.2 615.4	621.0	623.7	634.7
GeO	$X^1\Sigma^+$	1.625	1.627	1.623	1.626	985.5 979.8	989.0	994.9	997.7
SeS	$X^1\Sigma^+$	2.012	2.019	2.012	2.013	575.8 571.1	577.1	581.3	584.2
БеТе	$X^1\Sigma^+$	2.340	2.343	2.335	2.332	323.9 322.7	326.1	328.7	331.7
IfO	$X^1\Sigma^+$	1.723	1.719	1.715	1.735	974.1 970.1	976.0	980.7	952.5
Br	$X^1\Sigma^+$	2.469	2.478	2.464	2.463	268.6 269.2	276.1	279.5	279.7
nCl	$X^1\Sigma^+$	2.401	2.413	2.404	2.392	317.4 313.6	316.4	318.6	319.2
nF	$X^1\Sigma^+$	1.985	1.988	1.984	1.974	535.4 530.1	534.5	536.3	549.9
10	$X^2\Pi_{1/2}$	1.151	1.152	1.149	1.148	1904.2 1883.7	1912.5	1928.7	1927.6
PN	$\overset{2}{X}{}^{1}\Sigma^{+}$	1.491	1 496	1.491	1.493	1337.2 1332.5	1345.6	1352.8	1346.4
PO	_		1.482	1.477	1.479	1233.3 1232.0	1242.9	1250.8	1243.7
bO	$X^1\Sigma^+$	1.922	1.924	1.919	1.922	721.0 732.8	739.9	745.1	746.1
bS	$X^1\Sigma^+$	2.287	2.297	2.288	2.286	429.4 433.2	439.9	444.6	445.3
O	$X^3\Sigma^-$	1.481		1.481	1.482	1149.2 1152.9	1161.3	1167.7	1162.7
cO	$X^2\Sigma^+$	1.668	1.674	1.663	1.670	965.0 968.6	979.6	936.3	932.9
iO	$X^1\Sigma^+$	1.510	1.515	1.511	1.514	1241.6 1231.6	1242.3	1248.6	1241.0
iS	$X^1\Sigma^{\scriptscriptstyle +}$	1.929	1.939	1.931	1.933	749.6 741.1	752.6	759.2	753.1
nO	$X^1\Sigma^{\scriptscriptstyle +}$	1.833	1.832	1.827	1.826	814.6 821.6	829.9	835.6	842.3
nS	$X^1\Sigma^{\scriptscriptstyle +}$	2.209	2.215	2.206	2.204	487.3 484.2	491.4	496.5	492.9
F.	$X^1\Sigma^{\scriptscriptstyle +}$	1.926	1.935	1.930	1.936	631.3 628.9	634.4	637.7	629.2
rO	$X_1\Sigma^+$	1.712	1.719	1.714	1.715	969.8 977.9	981.4	981.6	979.8

Similarly, it includes the calculated dipole moments employing different basis sets.

Table 4 lists the experimental and calculated equilibrium bond length  $R_{\rm e}$  and harmonic vibrational frequency  $\omega_{\rm e}$ , employing different basis sets.

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