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Accuracy of DLPNO-CCSD(T) method for noncovalent bond dissociation enthalpies from coinage metal cation complexes.

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KEYWORDS: DLPNO-CCSD(T), Transition Metals, Dissociation Enthalpies, Binding Energies, Relativistic Effects, M06.

ABSTRACT. The performance of the domain based local pair-natural orbital coupled-cluster (DLPNO-CCSD(T)) method has been tested to reproduce the experimental gas phase ligand dissociation enthalpy in a series of Cu^+ , Ag^+ and Au^+ complexes. For 33 Cu^+ - non-covalent ligand dissociation enthalpies all-electron calculations with the same method result in MUE below 2.2 kcal/mol, although a MSE of 1.4 kcal/mol indicates systematic underestimation of the experimental values. Inclusion of scalar relativistic effects for Cu either via effective core potential (ECP) or Douglass-Kroll-Hess Hamiltonian, reduces the MUE below 1.7 kcal/mol and the MSE to -1.0 kcal/mol. For 24 Ag⁺ - non-covalent ligand dissociation enthalpies the DLPNO-CCSD(T) method results in a mean unsigned error (MUE) below 2.1 kcal/mol and vanishing mean signed error (MSE). For 15 Au⁺ - non-covalent ligand dissociation enthalpies the DLPNO-

CCSD(T) methods provides larger MUE and MSE, equal to 3.2 and 1.7 kcal/mol, which might be related to poor precision of the experimental measurements. Overall, for the combined dataset of 72 coinage metal ion complexes DLPNO-CCSD(T) results in a MUE below 2.2 kcal/mol and an almost vanishing MSE. As for a comparison with computationally cheaper density functional theory (DFT) methods, the routinely used M06 functional results in MUE and MSE equal to 3.6 and -1.7 kca/mol. Results converge already at CC-PVTZ quality basis set, making highly accurate DLPNO-CCSD(T) estimates to be affordable for routine calculations (single-point) on large transition metal complexes of > 100 atoms.

1. Introduction

Computational chemistry is routinely applied nowadays to support and integrate experimental studies in transition-metal catalysis.¹⁻⁷ The successful standalone experimental-free theoretical predictions in this field are far less common, however.⁸ While some failures in theoretical predictions are originated from the complexity of the systems themselves and can be ameliorated by proper inclusion of the effects deriving from incomplete sampling of the conformational space and/or solvation,^{1,9} the other failures are related to the accuracy of electronic structure methods. In general, scalar/vector relativistic effects, basis set completeness and multireference character of some systems should be properly addressed regardless on the electronic structure method used.¹⁰⁻¹¹ When it comes specifically to density functional theory (DFT) methods, which is the only affordable computational protocol to study systems of "realistic-size", one has to remember that the performance of the underlying exchange-correlation (XC) functionals is not uniform, and provides low to high accuracy predictions depending on the chemical system under study.¹² The careful "calibration" against highly accurate experimental measurements or wave

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function theory (WFT) calculations helps to improve upon the situation as well as latest developments in functional design.^{9,13-27} However, the question to what extent the chemical system under study is different from the ones used for calibration of a given functional and, more important, the amount of highly accurate measurements is limited.

On the other hand, so-called "ab initio" WFT methods²⁸ are rigorous and allow to systematically achieve chemical accuracy without any calibration against experimental data. In particular, the coupled cluster approximation with inclusion of doubles and triples excitations (CCSD(T)) is recognized as the golden standard to describe electronic structure, as long as no strong static correlation exists in the studied system. The drawback is that CCSD(T) is applicable only to small systems, since its cost scales as N⁷ with N being the basis set size of the system.

This has conveyed many efforts to reduce the computational cost of CCSD(T) by, for example, taking advantage of density fitting (or resolution of identity) approximations and new algorithms for two electron integrals transformation,²⁹⁻³² and by using localized molecular orbitals³³⁻³⁶ to optimize the selection of the most relevant excitations.³⁷⁻⁴⁰ Thanks to these technical advancements, CCSD(T) is now affordable for systems up to 100 atoms, including transition metals. Although different implementations of linear scaling CCSD(T) are quite effective, we found the DLPNO-CCSD(T)³⁴⁻³⁵ implementation the easiest to use for end users. DLPNO stands for domain based local pair natural orbital, since instead of canonical delocalized orbitals, pair natural orbitals⁴¹ are used and then localized,³⁴⁻³⁵ so that they can be classified into domains for proper sorting and selection of the most important excitations accounting for electronic correlation.

DLPNO-CCSD(T) accuracy has been assessed on either full CCSD(T) or experimental data for some systems of interest including non-convalent interactions,³² enzymatic reactions,⁴² organic reactions⁴²⁻⁴³ transition metals promoted reactions,⁴⁴⁻⁴⁷ conformational issues in transition metal structures,⁴⁴ and even extended to solid oxide crystals⁴⁸ and a small protein.³⁵ However, in the majority of the cases the DLPNO-CCSD(T) method was validated against experimental data on transition metals reactions in solvent, ⁴⁴⁻⁴⁷ and consequently the accuracy of the combined DLPNO-CCSD(T)/particular solvation model was assessed rather than DLPNO itself.

To test the performance of the DLPNO-CCSD(T) method from a different perspective, in the current work we present a systematic test study of the gas phase non-covalent ligand dissociation enthalpies in 72 complexes of Cu^+ , Ag^+ and Au^+ related to catalysis. To assess the performance of the DLPNO-CCSD(T) method itself the following steps were taken. First, to eliminate the effect of the solvent, only gas phase measurements were used to build a test set. Second, to discard the effects from the low-lying frequency modes, important for the molecular entropy, only performance in enthalpies was analyzed. Third, to exclude any influence of the scalar relativistic effects, only complexes giving closed shell reactants and products were selected. Finally, to avoid systems with multireference character, T1 diagnostic values were thoroughly monitored.

2. Computational Details

The ORCA⁴⁹ suite of programs was employed for all the calculations performed in the present work.

2.1 Geometry Optimizations. The geometry optimization was performed using the pure GGA PBE⁵⁰⁻⁵¹ functional as implemented in ORCA. The Grimme's D3(BJ)⁵² dispersion correction was activated via the "D3BJ" option to arrive at the PBE-D3(BJ) functional, because some of the complexes from the benchmark set are not small (see Figure 1) and the inclusion of dispersion interactions during the optimization might be essential.⁵³⁻⁵⁵ Default values were adopted for the self-consistent-field (SCF) and geometry optimization convergence criteria. Numerical integration of the exchange-correlation (XC) terms was performed using tighter-than-default "Grid 7" option (Lebedev770 and IntAcc=5.67 and no FinalGrid) to eliminate potential numerical noise. Geometries were characterized as true energy minima by the eigenvalues of the analytically calculated Hessian matrix. Translational, rotational, and vibrational partition functions for thermal corrections to give total enthalpies were computed within the ideal-gas, rigid-rotor, and harmonic oscillator approximations following standard procedures.

The all-electron DEF2-TZVP⁵⁶ basis sets of the Karlsruhe group was used on all the elements apart from Ag and Au, along with corresponding auxiliary basis functions⁵⁷ needed to fit Coulomb potential to speed up the DFT calculations. Quasi-relativistic effective core potentials (ECP) of the Stuttgart type⁵⁸ were used to describe 28 inner electrons of Ag and 60 inner electrons of Au in combination with the corresponding DEF2-TZVP basis set.

Despite the complexes studied in the present work could be optimized with more rigorous WFT methods or more advanced hybrid meta-GGA DFT functionals, we found pure GGA DFT and density fitting algorithms more useful in light of its exclusive role in optimization of realistic-size TM complexes (>100 atoms). Indeed, the DLPNO-CCSD(T) method is aimed at molecules of this size, and its performance in conjunction with pure GGA for location of stationary points on the potential energy surface is especially interesting for routine applications.

If several conformations could be possible for some complexes, the structure of the most stable one was either taken from the literature or was found by geometry optimization with PBE functional of many manually-generated conformations.



Figure 1. The molecular structures of the largest Ag⁺ (above) and Cu⁺ (below) complexes which were investigated in the current work. Color coding: N(blue), C (turquoise), O (red), H (gray), Ag (gray), Cu (gray).

2.2 Single-Point Energy Evaluations.

The DLPNO-CCSD(T)³⁴⁻³⁵ method was applied for all single-point energy evaluations in the current work. The default "NormalPNO" DLPNO settings (TCutPairs = 10^{-4} , TCutPNO = 3.33 10^{-7} , TCutMKN = 10^{-3}) were used as recommended for most computational applications in terms of cost/efficiency ratio.⁴⁰

2.2.1 Reactions Involving Ag⁺ and Au⁺ Complexes.

Fully-relativistic ECP of the Stuttgart type⁵⁹ were used to describe 28 inner electrons of Ag and 60 inner electrons of Au in combination with the corresponding correlation consistent basis sets "CC-PVTZ-PP" and "CC-PVQZ-PP" of Peterson et al.⁶⁰ All other elements were described with all-electron correlation consistent basis sets of the CC-PVNZ⁶¹⁻⁶³ family (N=3 and 4). The correlation fitting basis sets (CC-PVNZ/C) developed by Hättig et al.⁶⁴ necessary for the resolution of identity approximation (RI) as a part of DLPNO scheme were used. Non relativistic Hamiltonian was used for all the calculations involving silver complexes. This combination of the basis sets will be further referred as "CC-PVNZ (ECP)".

2.2.2 Reactions Involving Cu⁺ Complexes.

Depending on the degree of inclusion of the scalar relativistic effects few strategies have been employed to describe ligand dissociation in the copper complexes. All strategies involve correlation fitting basis sets (CC-PVNZ/C) developed by Hättig et al.⁶⁴ for RI approximation.

2.2.2.1 All-electron non-relativistic calculations.

The non-relativistic Hamiltonian was used in conjunctions with all electron correlation consistent basis sets of CC-PVNZ (N = 3 and 4) family.⁶¹⁻⁶³ The copper atom was described with all

electron basis set of Peterson et al.⁶⁵ This combination of the basis sets will be termed "CC-PVNZ" from thereafter.

2.2.2.2 Effective core potential calculations.

The non-relativistic Hamiltonian was used in conjunction with all electron correlation consistent basis sets of the CC-PVNZ (N = 3 and 4) family⁶¹⁻⁶³ on all the elements but copper. Fully-relativistic ECPs of the Stuttgart type⁵⁹ were used to describe 10 inner electrons of Cu in combination with corresponding correlation consistent basis sets "CC-PVTZ-PP" and "CC-PVQZ-PP" of Peterson et al.⁶⁰ This combination of the ECP and the basis set corresponds to "CC-PVNZ (ECP)" strategy used earlier for solver complexes.

2.2.2.3 Scalar Relativistic Calculations.

The scalar relativistic Douglas-Kroll-Hess (DKH)⁶⁶ Hamiltonian was applied as implemented in the ORCA suite of programs. All electron correlation consistent basis sets re-contracted to be used in conjunction with DKH Hamiltonian were used.⁶⁷This combination of the correlation consistent basis and DKH Hamiltonian will be termed "CC-PVNZ (DKH)".

2.2.3 Complete Basis Set Extrapolation.

To eliminate the effects from basis set incompleteness, the extrapolation schemes for HF and correlation energies of individual species suggested by Helgaker et al.⁶⁸⁻⁷⁰ for two adjacent CC-PVNZ level basis sets were employed:

$$E_{HF}^{X} = E_{HF}^{\infty} + \alpha e^{-1.63X} (1)$$

 $E_{corl}^{X}=E_{corl}^{\infty}+\beta e^{-3}\left(2\right)$

 Where X = 3 and 4 for CC-PVTZ and CC-PVQZ basis sets, correspondingly; $E_{HF}^{\infty}/E_{corl}^{\infty}$ HF and correlation energies at CBS limit; α/β are parameters to be obtained from a system of the two equations. The total bond dissociation enthalpy at CBS limit for each molecule AB was evaluated via following equation:

$$\Delta H^{0}_{DLPNO-CCSD(T)} = E^{\infty}_{HF}(A) + E^{\infty}_{corl}(A) + H^{PBE-D3}_{corr}(A) + E^{\infty}_{HF}(B) + E^{\infty}_{corl}(B) + H^{PBE-D3}_{corr}(B) - (E^{\infty}_{HF}(AB) + E^{\infty}_{corl}(AB) + H^{PBE-D3}_{corr}(AB))$$
(3)

where H_{corr}^{PBE-D3} is the correction to the electronic energy to arrive to the enthalpy, see Section 2.1 for the details.

As for an indicative comparison with state of the art DFT functionals, we also evaluated M06 dissociation enthalpies on this study with equivalent basis sets. The formula we used for M06 CBS extrapolation being:

$$\Delta H_{M06}^{0} = E_{M06}^{\infty}(A) + H_{corr}^{PBE-D3}(A) + E_{M06}^{\infty}(B) + H_{corr}^{PBE-D3}(B) - (E_{M06}^{\infty}(AB) + H_{corr}^{PBE-D3}(AB))$$
(4)

where E_{M06}^{∞} has been extrapolated via formula (1).

As for the basis set superposition error (BSSE), the standard counterpoise correction (CP)⁷¹ has not been applied in the present study for several reasons. First, with the two-point CBS extrapolation scheme⁶⁸⁻⁷⁰ based on CC-PVTZ/CC-PVQZ basis sets we approach the limit of a complete basis set at which *in theory* both the BSSE and basis set incompleteness should be virtually reduced to zero. Indeed, the mean signed error obtained for all 72 dissociation reaction turned out to be close to practically zero kcal/mol (see below) while in case of strong pollution by BSSE large and positive MSE would take place. Moreover, considering the complete basis set as a reference, several studies show that the extrapolation scheme is either as⁷² or more⁷³ accurate than the counterpoise correction for the evaluation of interaction energies and reaction barriers. Other references⁷³⁻⁷⁵ also demonstrate a better accuracy of the extrapolation scheme over the counterpoise method for the evaluation of an experimental property. Finally, recent studies on CCSD(T) evaluation of metal-ligand bond dissociations do not include counterpoise corrections.^{46,76,47}

2.3. The Benchmark Set.

The DLPNO-CCSD(T) method in its current implementation can only be applied to closed shell systems. This puts an important constraint on the potential TM ion reactions to be included in the benchmark set: both reactants and products have to be singlets. Since a dominant part of the TM ions have unfilled d-shell, there is a high chance that open shell states of the complexes formed by these metals might be more stable comparing to closed shell states. To verify it manually for every TM complex is a daunting task, and even worse, the results can also be dependent on the method chosen. The 11 group metals such as Cu, Ag and Au are known to have one 4s electron on a top of the filled d-shell which leads to the closed shell d¹⁰ configurations of the ions. Since the most stable closed-shell d¹⁰ configuration is unlikely to be changed by closed-shell non-covalent ligands in the TM complexes, we decided to focus only on Cu⁺, Ag⁺ and Au⁺ complexes in the current work. In total for the current study we selected 33, 24 and 15 non-covalent gas phase binding enthalpies/ZPE-energies for Cu⁺, Ag⁺ and Au⁺ complexes, respectively, for which reliable experimental data are available.

It should be noted that the dominant part of the experimental reaction energies are given as enthalpies at 298.15 K, and other are zero-point corrected energies (ZPE). To keep the study

consistent, all ZPE values were converted to enthalpies via following procedure. First, ZPE and enthalpic corrections were calculated at the PBE-D3(BJ)/DEF2-TZVP protocol, see above. Then, the subtraction of the ZPE correction from the experimental ZPE energies was followed by addition of the enthalpic correction to arrive at reaction enthalpies that were then compared to their experimental counterparts. If more than one experimental value was available for the reaction, the average was taken as the reference for comparisons with our calculations. To ensure that reactants and products are both singlets, the SP energy evaluations with multiplicities 3 (triplets) and 5 (quintet) at geometry optimization level were done on the optimized structures. In all cases the energies of open shell states were higher in energy. The selected dissociation reactions can be considered as a reference dataset for further benchmarking in the future. The 33, 24 and 15 non-covalent ligand dissociation reactions for Cu⁺, Ag⁺ and Au⁺ thus compose datasets CUNCDE33, AGNCDE24, and AUNCDE15, all together combined to the G11NCDE72 dataset. Optimized coordinates of all the systems composing these datasets are included in the SI.

 Table 1. Experimental dissociation enthalpies/energies used for benchmarking of DLPNO

 CCSD(T) calculations.

N⁰	Reaction	Expt. $\Delta H^{\circ}(298.15)$ or D_0 (kcal/mol) ^a	Avg. Δ H°(298.15) (kcal/mol) ^{b,c}
1	$(Ag•3H_3N)^+ = (Ag•2H_3N)^+ + H_3N$	14.6±0.1 ⁷⁷	14.6±0.1
2	$(Ag•3H_2O)^* = (Ag•2H_2O)^* + H_2O$	15.0±0.1 ⁷⁷	15.0±0.1
3	$(Ag•4H_3N)^+ = (Ag•3H_3N)^+ + H_3N$	13.0±0.1 ⁷⁷	13.0±0.1
4	$(Ag•4H_2O)^* = (Ag•3H_2O)^* + H_2O$	14.9±0.2 ⁷⁷	14.9±0.2
5	$(Ag \cdot 2H_3N)^+ = (Ag \cdot H_3N)^+ + H_3N$	36.9±0.8 ⁷⁷	36.9±0.8
6	$(Ag•2H_2O)^* = (Ag•H_2O)^* + H_2O$	25.4±0.3 ⁷⁷	25.4±0.3
7	$(Ag \cdot 3C_5H_5N)^{+} = (Ag \cdot 2C_5H_5N)^{+} + C_5H_5N$	16.7±0.2 ⁷⁷	16.7±0.2

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8	$(Ag \cdot 4C_5H_5N)^{+} = (Ag \cdot 3C_5H_5N)^{+} + C_5H_5N$	17.9±0.2 ⁷⁷	17.9±0.2
9	$(Ag \cdot 2C_2H_4)^+ = (Ag \cdot C_2H_4)^+ + C_2H_4$	32.4±1.5; ⁷⁸ 30.1±1.3(30.1±1.3 (ZPE)) ⁷⁹	31.3(+2.7 / -2.4)
10	$(Ag•3CO)^{+} = (Ag•2CO)^{+} + CO$	13.5±1.8(13.1±1.8 (ZPE)) ⁸⁰	13.5±1.8
11	$(Ag•4CO)^{+} = (Ag•3CO)^{+} + CO$	10.6±0.9(10.8±0.9 (ZPE)) ⁸⁰	10.6±0.9
12	$(Ag•2CO)^{+} = (Ag•CO)^{+} + CO$	26.5±0.9(26.1±0.9 (ZPE)) ⁸⁰	26.5±0.9
13	$(Ag^{+} 2C_{6}H_{6})^{+} = (Ag C_{6}H_{6})^{+} + C_{6}H_{6}$	40.5±4.5(39.9±4.5 (ZPE)) ⁸¹	40.5±4.5
14	$(Ag \cdot 2C_2H_3N)^+ = (Ag \cdot C_2H_3N)^+ + C_2H_3N$	34.7±1.4 ⁸²	34.7±1.4
15	$(Ag \cdot C_2 H_4)^+ = Ag^+ + C_2 H_4$	33.7±3; ⁷⁸ (ZPE)) ⁷⁹ 32.9±3.0(32.2±3	33.3(+3.4 / -3.7)
16	$(Ag \bullet H_2O)^* = Ag^* + H_2O$	32.5±2.5 (31.6±2.5 (ZPE)); ⁸³ 33.3±2.2 ⁷⁷	32.9(+2.6 / -2.9)
17	$(Ag \cdot C_6 H_6)^+ = Ag^+ + C_6 H_6$	38.0±1.7(37.3±1.7 (ZPE)); ⁸⁴ 40.6±4.5 (39.9±4.5 (ZPE)) ⁸¹	39.3(+6.0 / -3.2)
18	$(Ag \cdot CO)^+ = Ag^+ + CO$	21.8±1.2(21.2±1.2 (ZPE)) ⁸⁰	21.8±1.2
19	$(Ag \cdot C_5 H_{10})^+ = Ag^+ + C_5 H_{10}$	38.2±4.5 (37.8±4.5 (ZPE)) ⁸¹	38.2±4.5
20	$(Ag \cdot C_3 H_6 O)^+ = Ag^+ + C_3 H_6 O$	38.0±4.5 (38.2±4.5 (ZPE)) ⁸¹	38.0±4.5
21	$(Ag \cdot C_5 H_8)^+ = Ag^+ + C_5 H_8$	39.5±4.5 (39.2±4.5 (ZPE)) ⁸¹	39.5±4.5
22	$\left(Ag \cdot C_2 H_3 N\right)^+ = Ag^+ + C_2 H_3 N$	39.4±1.4 ⁸²	39.4±1.4
23	$(Ag•3(C_2H_4))^+ = (Ag•2(C_2H_4))^+ + C_2H_4$	13.9±0.8(13.6±0.8 (ZPE)) ⁷⁹	13.9±0.8
24	$(Ag \cdot 4(C_2H_4))^+ = (Ag \cdot 3(C_2H_4))^+ + C_2H_4$	7.4±0.8(6.5±0.8 (ZPE)) ⁷⁹	7.4±0.8
25	$(Cu•2H_2O)^* = (Cu•H_2O)^* + H_2O$	39.6±3.0(39±3.0 (ZPE)); ⁸⁵ 40.7±1.6 ⁸⁶	40.1(+2.5 / -3.5)
26	$(Cu \cdot 2C_3H_6O)^+ = (Cu \cdot C_3H_6O)^+ + C_3H_6O$	50.2±1.6 ⁸⁷	50.2±1.6
27	$(Cu•3H_2O)^+ = (Cu•2H_2O)^+ + H_2O$	17.7±3.0(17.0±3.0 (ZPE)); ⁸⁵ 13.7±1.8; ⁸⁶ 16.4±0.2 ⁷⁷	15.9(+4.8 / -4.0)
28	$(Cu•4H_2O)^+ = (Cu•3H_2O)^+ + H_2O$	16.1±3.0(15.0±3.0 (ZPE)); ⁸⁵ 12.8±1.0; ⁸⁶ 16.7±0.2 ⁷⁷	15.2(+3.9 / -3.4)
29	$(Cu•3H_3N)^+ = (Cu•2H_3N)^+ + H_3N$	11.0±1.5; ⁸⁸ 17.0±3.1; ⁸⁹ 14.1±0.2 ⁷⁷	14.0(+6.1 / -4.5)
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30	$(Cu•4H_3N)^+ = (Cu•3H_3N)^+ + H_3N$	10.8±1.5; ⁸⁸ 12.9±3.1; ⁸⁹ 12.8±0.2 ⁷⁷	12.2(+3.8 / -2.9)
31	$(Cu•2C_6H_6)^+ = (Cu•C_6H_6)^+ + C_6H_6$	37.2±2.8(37.1±2.8 (ZPE)) ⁹⁰	37.2±2.8
32	$(Cu*3CO)^{+} = (Cu*2CO)^{+} + CO$	17.6±1.0(18.0±1.0 (ZPE)) ⁸⁰	17.6±1.0
33	$(Cu•4CO)^{+} = (Cu•3CO)^{+} + CO$	13.0±0.7(12.7±0.7 (ZPE)) ⁸⁰	13.0±0.7
34	$(Cu•2CO)^{+} = (Cu•CO)^{+} + CO$	41.6±0.7(41.0±0.7 (ZPE)) ⁸⁰	41.6±0.7
35	$(Cu•2H_3N)^+ = (Cu•H_3N)^+ + H_3N$	59.3±2.4; ⁸⁸ 52.1±3.1; ⁹¹ 53.8±3.1 ⁷⁷	55.1(+6.6 / -6.1)
36	$(Cu•3C_2H_6O)^+ = (Cu•2C_2H_6O)^+ + C_2H_6O$	13.1±1.0(13.1±1.0 (ZPE)) ⁹²	13.1 ±0.1
37	$(Cu•4C_2H_6O)^+ = (Cu•3C_2H_6O)^+ + C_2H_6O$	11.2±2.4(10.8±2.4 (ZPE)) ⁹²	11.2±2.4
38	$(Cu•2C_2H_4)^+ = (Cu•C_2H_4)^+ + C_2H_4$	41.6±3.0(41.5±3.0 (ZPE)) ⁹³	41.6±3.0
39	$(Cu•2C_2H_6O)^+ = (Cu•C_2H_6O)^+ + C_2H_6O$	46.5±1.9(46.1±1.9 (ZPE)) ⁹²	46.5±1.9
40	$(Cu•3C_2H_3N)^+ = (Cu•2C_2H_3N)^+ + C_2H_3N$	20.1±0.6 ⁹⁴	20.1±0.6
41	$(Cu•4C_2H_3N)^+ = (Cu•3C_2H_3N)^+ + C_2H_3N$	15.1±0.6 ⁹⁴	15.1±0.6
42	$(Cu \cdot 2C_2H_3N)^+ = (Cu \cdot C_2H_3N)^+ + C_2H_3N$	56.9±2.2 ⁹⁴	56.9±2.2
43	$(Cu•3C_{3}H_{6}O)^{+} = (Cu•2C_{3}H_{6}O)^{+} + C_{3}H_{6}O$	15.7±0.8 ⁹⁴	15.7±0.8
44	$(Cu•4C_{3}H_{6}O)^{+} = (Cu•3C_{3}H_{6}O)^{+} + C_{3}H_{6}O$	14.7±1.4 ⁹⁴	14.7±1.4
45	$(Cu•2C_4H_5N)^+ = (Cu•C_4H_5N)^+ + C_4H_5N$	44.2(44.0 (ZPE)) ⁹⁵	44.2
46	$(Cu \cdot H_2O)^+ = Cu^+ + H_2O$	35.9±3.0(35±3.0 (ZPE)); ⁸⁵ 38.4±1.8; ⁸⁶	37.2 (+3.0 / -5.2)
47	$(Cu \cdot C_3 H_6 O)^+ = Cu^+ + C_3 H_6 O$	48.1±1.1 ⁸⁷	48.1±1.1
48	$(Cu \cdot C_2H_4)^+ = Cu^+ + C_2H_4$	42.9±3.3(42.0±3.3 (ZPE)) ⁹³	42.9±3.3
49	$(Cu \cdot C_6 H_6)^+ = Cu^+ + C_6 H_6$	52.3±2.3(52.1±2.3 (ZPE)) ⁹⁰	52.3±2.3
50	$(Cu \cdot CO)^+ = Cu^+ + CO$	36.2±1.7(35.5±1.7 (ZPE)) ⁸⁰	36.2±1.7
51	$(Cu \cdot C_2 H_3 N)^+ = Cu^+ + C_2 H_3 N$	57.4±0.9 ⁹⁴	57.4±0.9
52	$\left(Cu_{\bullet}H_{3}N\right)^{+}=Cu^{+}+H_{3}N$	56 6+3 6 ^{.88} 51 6+3 1 ^{.89}	53.9(+6.3 / -5.4)

		53.5±3.177	
53	$(Cu \cdot C_5 H_5 N)^+ = Cu^+ + C_5 H_5 N$	58.8±2.5 ⁹⁶	58.8±2.5
54	$(Cu \cdot C_4 H_4 N_2)^+ = Cu^+ + C_4 H_4 N_2$	60.1±2.3 ⁹⁷	60.1±2.3
55	$(Cu \cdot C_5 H_5 N_5)^+ = Cu^+ + C_5 H_5 N_5$	71.0±2.6(70.3±2.6 (ZPE)) ⁹⁸	71.0±2.6
56	$(Cu \cdot C_2 H_6 O)^+ = Cu^+ + C_2 H_6 O$	44.5±2.9(44.2±2.9 (ZPE)) ⁹²	44.5±2.9
57	$(Cu \cdot C_4 H_5 N)^+ = Cu^+ + C_4 H_5 N$	59.5±2.0(59.0±2.0 (ZPE)) ⁹⁵	59.5±2.0
58	(Au•styrene) ⁺ = Au ⁺ + styrene	44.0±0.9(42.7±0.9 (ZPE)) ⁹⁹	44.0±0.9
59	(Au•phenylacetylene) ⁺ = Au ⁺ + phenylacetylene	44.3±0.9(43.1±0.9 (ZPE)) 99	44.3±0.9
60	$(Au \cdot C_6 H_6)^+ = Au^+ + C_6 H_6$	39.3±0.7(38.7±0.7 (ZPE)) 99	39.3±0.7
61	$(Au \cdot 1-pentene)^+ = Au^+ + 1-pentene$	42.8±1.2(42.0±1.2 (ZPE)) 99	42.8±1.2
62	$(Au \cdot 1-pentyne)^+ = Au^+ + 1-pentyne$	44.0±1.6(43.4±1.6 (ZPE)) 99	44.0±1.6
63	$(Au \cdot 2\text{-pentyne})^+ = Au^+ + 2\text{-pentyne}$	44.2±0.9(44.0±0.9 (ZPE)) 99	44.2±0.9
64	$(Au \cdot cyclooctane)^{+} = Au^{+} + cyclooctane$	44.2±0.5(43.6±0.5 (ZPE)) 99	44.2±0.5
65	$(Au \cdot 1,5$ -cyclooctadiene) ⁺ = Au^+ + 1,5-cyclooctadiene	44.6±0.7(44.5±0.7 (ZPE)) 99	44.6±0.7
66	$(Au \cdot 1, 3$ -cyclooctadiene) ⁺ = Au^+ + 1,5-cyclooctadiene	42.8±0.7(42.2±0.7 (ZPE)) 99	42.8±0.7
67	$(Au \cdot CO)^+ = Au^+ + CO$	48.7±3.5(48.0±3.5 (ZPE)) ¹⁰⁰	48.7±3.5
68	$(Au \cdot H_2O)^+ = Au^+ + H_2O$	41.2±2.3(40.1±2.3 (ZPE)) ¹⁰¹	41.2±2.3
69	$(Au \cdot 2H_2O)^+ = (Au \cdot H_2O)^+ + H_2O$	45.7±3.5(45.0±3.5 (ZPE)) ¹⁰¹	45.7±3.5
70	$(Au•3H_2O)^+ = (Au•2H_2O)^+ + H_2O$	23.7±4.6(23.1±4.6 (ZPE)) ¹⁰¹	23.7±4.6
71	$(Au \cdot 4H_2O)^+ = (Au \cdot 3H_2O)^+ + H_2O$	21.8±4.6(20.8±4.6 (ZPE)) ¹⁰¹	21.8±4.6
72	$(Au \cdot C_6 F_5 H)^+ = Au^+ + C_6 F_5 H$	31.2±4.0(31.0±4.0 (ZPE)) ¹⁰²	31.2±4.0

^a Denoted as (ZPE). ^b Final experimental (average) Δ H°(298.15) used to compare with theoretical values. ^c To account for uncertainties if more than one experimental value is available, we considered the reference higher uncertainty as the difference between (1) the highest experimental enthalpy plus its corresponding positive uncertainty and (2) the average of the experimental values. As well, we considered the reference lower uncertainty as the difference between (1) the lowest experimental uncertainty plus its corresponding negative uncertainty plus its corresponding negative uncertainty plus its corresponding to the experimental value.

3. Results and Discussion

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First, we discuss the DLPNO-CCSD(T) performance on the ligand dissociation enthalpies of Ag^+ complexes. Then, we proceed to the reactions involving Cu^+ ions. A keen attention is payed to importance of scalar relativistic effects. After that, we discuss the results obtained for Au^+ complexes. Finally, we comment on the overall performance obtained for all three coinage metal ion reactions bot for DLPNO-CCSD(T) and M06. Practical recommendations shall be given.

3.1 Ag⁺ Bond Dissociation Enthalpies, the AGNCDE24 Dataset.

The DLPNO-CCSD(T) bond dissociation enthalpies obtained with "CC-PVTZ (ECP)", "CC-PVQZ (ECP)" and CBS (ECP) basis sets are presented in Chart 1 along with the corresponding experimental values. In general, the accuracy in MUE below 3 kcal/mol has been achieved in reproducing the dissociation enthalpies for practically all the Ag⁺ complexes. Few dissociation enthalpies, however, are somewhat poorly reproduced; in particular large errors of 5.4, 7.0, 7.1, 4.1 and 4.5 kcal/mol were obtained for reactions 5, 13, 14, 16 and 22, respectively. To have a better understanding of an inconsistency between calculated numbers and their experimental counterparts, additional measurements of Deng and Kebarle¹⁰³ providing the total dissociation enthalpy of the two non-covalent ligands (AgL₂⁺ = Ag⁺ + L₂) were analyzed.

In short, Deng and Kebarle estimated the sum of the enthalpies of reactions 13 and 17 (dissociation of the two benzene molecules from $(Ag \cdot 2C_6H_6)^+$) to be 68.2±3.0 kcal/mol. The DLPNO-CCSD(T)/CBS (ECP) calculations predict this value to be 71.4 kcal/mol which is only slightly above the experimental values, and almost within the experimental uncertainty. Since the enthalpy of the reaction 17 is predicted quite well by our estimates, we believe that the dissociation enthalpy of the reaction 13 might be slightly overestimated in the experiments of Ho et al.⁸¹ Also, the enthalpy of reactions 14 and 22 (acetonitrile dissociation from $(Ag \cdot 2C_2H_3N)^+$

and $(Ag \cdot 2C_2H_3N)^+$) are both overestimated comparing to the experimental measurements of Shoeib et al.⁸² This results in overestimation of the sum of the enthalpies of reaction 14 and 22 by 11.6 kcal/mol relative to experimental value of Shoeib et al.⁸² of 74.1 On the other hand, Deng and Kebarle estimate the sum of the enthalpies of reaction 14 and 22 to be 84.8±3.0 kcal/mol, which is perfect agreement with our calculations.

The enthalpy of reaction 16 (dissociation of H₂O molecule from $(Ag \cdot H_2O)^+$) is underestimated by our DLPNO-CCSD(T) calculations, and amounts to 28.7 kcal/mol at CBS (ECP) extrapolation while the experimental estimates are 32.5 ± 2.5^{83} and $33.3\pm2.2.^{77}$ However, we predict the sum of the enthalpy of reactions 16 and 6 (dissociation of the two H₂O molecules from $(Ag \cdot 2H_2O)^+$) to be 55.6 kcal/mol, which matches perfectly the estimate of Deng and Kebarle equal to 56.5 ± 3.0 . Since the enthalpy of reaction 6 is predicted quite close to the measurements of Holland et al.⁷⁷ we believe that the enthalpy reported by Holland et al.⁷⁷ and Aribi et al.⁸³ for reaction 16 might be slightly overestimated.

The enthalpy of reaction 5 (dissociation of NH₃ from $(Ag \cdot 2NH_3)^+$) is calculated to be 42.3 kcal/mol, which is somewhat larger than 36.9 kcal/mol documented by Holland et al.⁷⁷ As for the binding of NH₃ in $(Ag \cdot NH_3)^+$, our calculations predict corresponding dissociation enthalpy of 43.4 kcal/mol. Unfortunately, no experimental value is available on the dissociation of NH₃ from $(Ag \cdot NH_3)^+$ for a comparison. Nevertheless, our theoretical estimation of dissociation of the two NH₃ molecules from $(Ag \cdot 2NH_3)^+$ complex, 85.7 kcal/mol, is in perfect agreement with the experimental estimates of Deng and Kebarle (85.6±3.0).¹⁰³ Since pronounced overestimation of the enthalpy of reaction 5 has also been noticed in other DFT¹⁰⁴ and CCSD(T) calculations,¹⁰⁵ perhaps re-examination of the data of Holland et al. would be of interest.

Thus we believe that in addition to the approximations used in the theoretical calculations, documented disagreements can also be attributed partly to the inconsistencies between different experimental measurements.



Chart 1. Experimental and DLPNO-CCSD(T)/CC-PVNZ (ECP) dissociation enthalpies obtained for transition metal ion – non covalent ligand bonds in 24 Ag^+ complex.

In Chart 2 the overall errors obtained for the Ag⁺ dataset are presented. With CC-PVTZ (ECP) basis set the MUE/MSE are equal to 2.1 and 0.6 kcal/mol, respectively. Upon the increase of the

basis set to CC-PVQZ (ECP) quality the MUE remains the same and MSE gets equal to 0.3 demonstrating only a slight tendency to underestimation. Finally, at CBS limit the MUE gets equal 2.1 kcal/mol and MSE is 0.0 kcal/mol, indicating that the cases where enthalpies are overestimated are exactly compensated by the cases where the enthalpies are underestimated. Bearing in mind that average uncertainty associated with experimental enthalpies is in general near 1-2 kcal/mol we consider the achieved accuracy as acceptable. Another encouraging observation is that in terms of MUEs the results are converged already at CC-PVTZ (ECP) basis set making the calculations affordable for large TM complexes of hundreds of atoms.



Chart 2. DLPNO-CCSD(T)/CC-PVNZ (ECP) mean unsigned (MUE) and mean signed (MSE) errors with respect to experimental values obtained for 24 dissociation enthalpies involving Ag⁺ complexes.

3.2 Cu⁺ Binding Enthalpies, the CUNCDE33 Dataset.

3.2.1 Non Relativistic All Electron Calculations.

The DLPNO-CCSD(T) enthalpies calculated for dissociation of non-covalent ligands from Cu⁺ complexes are given in Chart 3. Comparing to the performance obtained for Ag^+ complexes, a clear tendency to underestimation can be identified. This is especially pronounced if MUE and MSE are plotted for all 33 Cu⁺ reactions, see Chart 4. Thus, with the CC-PVTZ basis set the MUE/MSE errors are equal to 2.7 and 2.0 kcal/mol which is considerably larger comparing to what was observed for Ag^+ complexes. Upon increase in the basis set to CC-PVQZ the MUE and MSE get improved only by 0.4 and 0.2 kcal/mol, respectively. Finally, at CBS extrapolation the MUE and MSE are equal to 2.1 and 1.4 kcal/mol, correspondingly. While MUE gets equal to what was obtained for Ag^+ complexes in the CBS limit, MSE does not vanish and remains equal to 1.4 kcal/mol, demonstrating that the dissociation enthalpies are systematically underestimated.

Since DLPNO-CCSD(T) estimates have been shown to be within 1 kcal/mol of CCSD(T) results,⁴⁰ truncation in the localization scheme is unlikely to be responsible for the underestimation as well as the basis sets which are of acceptable quality for the calculations at this level of theory. It is especially interesting that for seemingly trivial reaction 50 (CO dissociation from CuCO⁺) our documented enthalpy at the CBS limit is 30.9 kcal/mol, which is 5.3 kcal/mol than experimentally measured quantity. In principle, scalar relativistic effects, which are proportional to $Z^{4.34}$ with Z being the atomic number, according to the Wood-Boring calculations¹⁰⁶ have been shown to be non-negligible for Cu, as documented in the literature.^{44,76,107-110} Indeed, Matito et al.¹¹¹ estimated the CO bond dissociation energy in (CuCO)⁺ to be equal 37.3 kcal/mol at CCSD(T) Dirac – Coulomb Hamiltonian level. On the other hand, Frenking et al.¹¹² using relativistic Stuttgart-type ECP on Cu atom estimated this energy to be 31.2 kcal/mol, which is more in line with our non-relativistic estimates. However,

e.g. in the popular def2 family of the basis sets of Ahlrichs and co-workers⁵⁶ widely used for routine DFT and WFT calculations nowadays, relativistic ECPs start to be used only from Rb implying that the relativistic effects up to Kr could be neglected. Therefore, we believe that it is worth to explore the influence of scalar relativistic effects on Cu^+ bonding energies.

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Chart 3. Experimental and DLPNO-CCSD(T)/CC-PVNZ dissociation enthalpies obtained for transition metal ion – non covalent ligand bonds in 33 Cu^+ complex.



Chart 4. DLPNO-CCSD(T)/CC-PVNZ mean unsigned (MUE) and mean signed (MSE) errors with respect to experimental values obtained for 33 dissociation enthalpies involving Cu⁺ complexes.

To further explore the influence of the scalar relativistic effects on the dissociation enthalpies of Cu^+ complexes we carried out DLPNO-CCSD(T) calculations with Stuttgart ECP on copper as well as all-electron relativistic calculations with the DKH Hamiltonian.

3.2.2 Relativistic ECP Calculations.

The DLPNO-CCSD(T) results obtained with Stuttgart relativistic ECP on copper atom combined with the corresponding "CC-PVNZ-PP" basis set and Dunning "CC-PVNZ" basis sets on other elements are presented in Figure 5. The general tendency to underestimation of the dissociation enthalpies documented previously for Cu^+ complexes has disappeared. This is clearly from

inspection of Figure 6, in which the MUE and MSE are given for all 33 Cu⁺ reactions. Already at "CC-PVTZ (ECP)" level the MUE and MSE errors are 1.7 kcal/mol and -0.2 kcal/mol.



Chart 5. Experimental and DLPNO-CCSD(T)/CC-PVNZ (ECP) dissociation enthalpies obtained for transition metal ion – non covalent ligand bonds in 33 Cu^+ complex.

Comparing to all-electron "CC-PVTZ" results the MUE decreases by almost 1 kcal/mol while MSE is reduced by more than 1.5 kcal/mol, and is essentially vanishing. Upon the increase in the basis set to quadruple- ζ quality only minor (~0.1 kcal/mol) improvement in the MUE was achieved, while the MSE was increased to -0.5 kcal/mol, underlying the tendency to overestimate the enthalpies. Finally, at the CBS level the MUE is 1.7 kcal/mol and the MSE is - 1.0 kcal/mol.

It should also be mentioned that for all-electron CBS dissociation enthalpies MUEs larger than 4.0 kcal/mol were found for 5 reactions (34, 38, 50, 54, 57), whereas using ECP on copper results in a MUE larger than 4 kcal/mol only for reaction 53. For this reaction the DLPNO-CCSD(T)/CBS(ECP) calculations predict the enthalpy change to be 7.8 kcal/mol larger than the experimental estimates of Rodgers et al.⁹⁶ However, we believe that such large disagreement cannot be ascribed only to the approximations associated with our calculations. Indeed, alternative photodissociaton threshold measurements¹¹³ predict enthalpy of reaction 53 to be ~65.1 (enthalpy corrected) which is only 1.5 lower than our theoretical estimate. Usually, photodissociation measured enthalpies are less reliable than collision induced dissociation measured enthalpies, and that is why only latter are included in Table 1 to calibrate the DLPNO-CCSD(T) performance. Probably, the experimental collision induced dissociation enthalpy value for reaction 53 should be re-examined.

Importantly, the problematic reaction 50 for which larger errors were documented with allelectron non-relativistic basis sets is now perfectly described with the MUE/MSE in CBS limit

-0.5

-1

(ECP)

equal to 0.5 kcal/mol. The dissociation enthalpy of reaction 50 is 35.7 kcal/mol, which essentially matches the experimental value of 36.2±1.7 kcal/mol by Meyer and co-authors.⁸⁰ While this result is in close agreement with the relativistic calculations of Matito et al.,¹¹¹ it differs from the estimate of Frenking et al.¹¹² by 4.5 kcal/mol. This difference can be explained by better correlation consistent basis sets and CBS extrapolation used in the present work. It should also be mentioned that the difference between relativistic (ECP) and non-relativistic all electron enthalpies is especially impressive for reaction 50 since it amounts to 4.8 kcal/mol. ■ MUE ■ MSE 1.5 Error (kcal/mol) 0.5

CC-PVTZ CC-PVQZ CBS (ECP)

(ECP)



3.2.3 Relativistic All Electron Douglass-Kroll-Hess Calculations

To ensure that the difference between ECP and all-electron results are due to relativistic effects we performed DLPNO-CCSD(T) DKH calculations with "CC-PVTZ-DKH" basis sets which were re-contracted⁶⁷ to be used for the DKH calculations. It must be noted that both the number and the values of the exponents are identical in the CC-PVNZ-DKH to those in the CC-PVNZ basis sets. The only difference between CC-PVNZ and CC-PVNZ-DKH is the contraction coefficients, which means that the basis sets of CC-PVNZ and CC-PVNZ-DKH are exactly of the same quality.

The absolute reaction enthalpies with DKH calculations turned out to be very similar to those obtained using ECP on copper and are given in the SI (Charts S1 and S2). The MUE/MSEs obtained for all 33 Cu⁺ reactions are similar to the MUE/MSE obtained with ECP. The average unsigned difference between dissociation energies obtained with ECP and all-electron DKH results are 0.2, 0.1 and 0.2 kcal/mol for "CC-PVTZ", "CC-PVQZ" and CBS basis sets. This confirms that the difference documented between ECP and all-electron non-relativistic results is almost entirely due to relativistic effects. Moreover, the relativistic contribution to the reaction energies is not uniform for all the reactions. The difference between (Δ E(DLPNO-CCSD(T)/CBS(DKH)) - Δ E(DLPNO-CCSD(T)/CBS)) strongly depends on the reactions, it varies from -1.7 kcal/mol (reaction 43) to 5.1 kcal/mol (reaction 57). This underlines that the relativistic effects cannot be ignored at least for Cu complexes if highly accurate absolute or relative dissociation reaction energies are of importance.

3.3 Au⁺ Bond Dissociation Enthalpies, the AUNCDE15 Dataset.

The DLPNO-CCSD(T) bond dissociation enthalpies obtained with "CC-PVTZ (ECP)", "CC-PVQZ (ECP)" and CBS (ECP) basis sets for 15 Au⁺ complexes are presented in Chart 7 along

with the corresponding experimental values. Comparing to the results obtained for Ag^+ and Cu^+ complexes larger discrepancies between our theoretical estimates and the experimental dissociation enthalpies can be noticed. Indeed, the overall MUE and MSE errors in CBS limit obtained for all 15 enthalpies are 3.2 and 1.7 kcal/mol, respectively.



Chart 7. Experimental and DLPNO-CCSD(T)/CC-PVNZ (ECP) dissociation enthalpies obtained for transition metal ion – non covalent ligand bonds in 15 Au^+ complex.

However, the error between our theoretical estimates and the experimental values is not equally spread for 15 reactions. The smallest errors have been obtained for reactions 58 - 66. Thus, MUE/MSE calculated only for those reactions are 1.6 and 0.6 kcal/mol, which is comparable to what was obtained for Cu⁺ and Ag⁺ complexes. Interestingly, the experimental uncertainties for the reactions 58 - 66 are in the range of 0.5 - 1.6 kcal/mol (average expt. uncertainty for these 9

reactions is 0.9 kcal/mol). On the other hand, for the reactions 67 - 72 significantly larger experimental uncertainties are documented, namely 2.3 - 4.6 kcal/mol (average expt. uncertainty for these 6 reactions is 3.8 kcal/mol). This indicates that larger errors obtained for reactions 67 -72 are not necessarily related to theoretical method itself but can be consequence of larger experimental uncertainties and less precise measurements. Comparison of our study with other theoretical studies supports this hypothesis. For reaction 67 (dissociation of CO from AuCO⁺) our CBS estimate is 42.3 kcal/mol which is 6.4 kcal/mol smaller comparing to measurements of Schwarz et al.¹⁰⁰ of 48.7±3.5. Gordon and co-workers¹¹⁴ have estimated the CO dissociation energy from AuCO⁺ to be $\Delta E_0 = 43.9$ kcal/mol at essentially CCSD(T)/aug-CC-PVTZ level (ECP on Au) which results in a final enthalpy $\Delta H^{\circ} = 44.7$ kcal/mol after our enthalpic correction is applied. Schwarz et al.¹¹⁵ estimated CO binding energy in AuCO⁺ to be 44.1 kcal/mol (ΔH° = 44.9 kcal/mol) at similar level of theory. Since both literature-based AuCO⁺ dissociation enthalpies are neither CP-corrected, nor CBS-extrapolated, it is reasonable to assume that due to BSSE these values should be reduced by 1-2 kcal/mol, which would result in perfect agreement with our estimation. In addition, for reaction 67 we also tested the influence of diffuse functions. Thus, our DLPNO-CCSD(T) aug-CC-PVTZ/aug-CC-PVQZ extrapolation results in the dissociation enthalpy of 42.5 kcal/mol which indicates that the diffuse functions, as expected for cationic systems, are not responsible for our deviations with the experimental measurements. Our CBS estimates for reactions 68, 69, 70 and 71 (dissociation of H₂O from Au(H₂O)⁺, Au(H₂O)⁺, $Au(H_2O)_3^+$, $Au(H_2O)_4^+$) are 35.9, 42.1, 17.2 and 16.5 kcal/mol, respectively. The experimental dissociation enthalpies of reactions 68 - 71 are systematically higher, namely 41.2, 45.7, 23.7 and 21.8 kcal/mol. On the other hand, our DLPNO-CCSD(T evaluations) agree reasonably well with CCSD(T)/aVDZ estimates of Lee at al.: 116 36.2, 41.7, 17.3 and 16.0 kcal/mol.



Chart 8. DLPNO-CCSD(T)/CC-PVNZ (ECP) mean unsigned (MUE) and mean signed (MSE) errors with respect to experimental values obtained for 24 dissociation enthalpies involving Ag⁺ complexes.

3.4 Overall Performance.

3.4.1 DLPNO-CCSD(T)

The overall performance of the DLPNO-CCSD(T) methods obtained for 72 transition metal ion $(Cu^+, Ag^+ \text{ and } Au^+)$ – non-covalent ligands enthalpies is depicted in Chart 9. For copper, the ECP results were selected. We remark that DKH all-electron calculations with CC-PVTZ(DKH) basis sets would lead to similar conclusions. Already at CC-PVTZ level the overall coinage metals MUE is 2.1 kcal/mol and the MSE is 0.4 kcal/mol. Upon increase of the basis set to CC-PVQZ quality the MUE does not change and the MSE becomes 0.2 kcal/mol, which is quite small. Finally, at CBS limit slightly higher MUE is obtained, namely 2.2 kcal/mol, while MSE gets

smaller, namely -0.1 kcal/mol. It has to be noticed that the accuracy close to 2.1 kcal/mol has been obtained, and, which is very important, the calculations are essentially converged at the triple- ζ quality basis set. That means that DLPNO-CCSD(T)/CC-PVTZ calculations are already very accurate, and high quality CCSD(T) estimates can be obtained for very large molecules, since very time-consuming CC-PVQZ calculations could be avoided.



Chart 9. DLPNO-CCSD(T)/CC-PVNZ (ECP) mean unsigned (MUE) and mean signed (MSE) errors with respect to experimental values obtained for all 72 dissociation enthalpies involving Cu⁺, Ag⁺ and Au⁺ complexes.

3.4.2. M06

Finally, in the light of the insightful discussion promoted by Xu et al. on whether practical CCSD(T) calculations agree better than DFT when compared to experimental data for dissociation energies of bonds to transition metals,⁷⁶ we decided to extend the current research with one representative DFT method, since the main scope of this work is not an extensive

benchmark of DFT methods. Due to its excellent work in describing of transition metal – noncovalent ligand bond breaking,^{1,117-118} we selected the M06 functional of Zhao et al.^{15,17} for the evaluation of the dissociation enthalpies of our dataset. The errors obtained for all 72 dissociation reactions involving coinage metal ions are given in Chart 10. First, the overall MUE obtained for M06 is 3.3 kcal/mol at CBS limit and the MSE is -1.7 kcal/mol. Considering that DFT methods are clearly faster than DLPNO-CCSD(T), the overall performance of the M06 functional is remarkable. Indeed, it results in a small tendency to overestimate the dissociation enthalpy relative to DLPNO-CCSD(T), resulting in negative MSEs. Finally, it has to be noticed that M06 dissociation enthalpies are essentially converged at the triple- ζ quality basis set. In this regard, our results can be regarded as another small support to consider DFT as a reference for transition metals chemistry,⁷⁶ although it is clear than more functionals and other transition metals have to be considered in the tests.¹¹⁹⁻¹²⁰



Chart 10. M06/CC-PVNZ (ECP) mean unsigned (MUE) and mean signed (MSE) errors with respect to experimental values obtained for all 72 dissociation enthalpies involving Cu^+ , Ag^+ and Au^+ complexes.

Conclusions

The recently developed DLPNO-CCSD(T) method was tested in reproducing 72 non-covalent ligand – transition metal ion $(Ag^+, Cu^+ and Au^+)$ gas phase dissociation enthalpies measured experimentally. The best protocol we used, namely DLPNO-CCSD(T)/CC-PVTZ(ECP) results in remarkable accuracy, with overall MUE 2.1 kcal/mol. All large deviations of our theoretical estimates from the experimentally documented values were explained. It was shown that the found deviations are not necessarily due to shortcomings of DLPNO-CCSD(T) scheme but could be attributed to quite large errors, documented or not, in the experimental enthalpies. Scalar relativistic effects turned out to be non-negligible for copper complexes, and their inclusion in the calculations either through relativistic ECP or through the Hamiltonian were proved to be fundamental, moreover, ECP and DKH based dissociation enthalpies turned out to be of the same quality for copper. Gratifying, results are converged already at the CC-PVTZ quality basis set, which allows to routinely obtain highly accurate DLPNO-CCSD(T) dissociation enthalpies for the transition metal complexes consisting of > 100 atoms. The results encourage to apply DLPNO-CCSD(T) for single-point energy evaluations in calculations related to transition metal catalysis. However, since there is increasing evidence that for some specific problem in TM chemistry⁷⁶ and non-covalent interactions¹²¹ the accuracy of CCSD(T) cannot be considered satisfactory, more tests involving other metal complex would be a good subject for the future study. Finally, overall good performance is provided by the M06 functional, with MUE and MSE of 3.3 kcal/mol and -1.7 kcal/mol.

ASSOCIATED CONTENT

Supporting Information. Cartesian coordinates (Å), M06, DLPNO-CCSD(T), with different basis sets, T1 diagnostic values, enthalpic and ZPE corrections, tabulated enthalpies and errors forming the basis of Charts 1 - 7. This material is available free of charge via the Internet at http://pubs.acs.org.

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