

View Article Online **PAPER**



Cite this: Phys. Chem. Chem. Phys. 2023, 25, 15371

Received 2nd March 2023, Accepted 9th May 2023

DOI: 10.1039/d3cp00978e

rsc.li/pccp

On the nature of inter-anion coinage bonds†

Jiayao Li,^a Qiuyan Feng,^a Changwei Wang*^a and Yirong Mo (1)*^b

To explore the binding energy profiles and elucidate the bonding nature in counter-intuitive anion---anion coinage bonds (CiBs), thirty-one complexes were constructed, and the inter-anion CiBs were studied theoretically. The metastability was evidenced by the characteristic potential wells in six cases, demonstrating that anions $[Au(CN)_4]^-$, $[Ag(CN)_2]^-$ and $[AuO]^-$ are appropriate building blocks for CiBs. The kinetic stability was further supported by ab initio molecular dynamics (AIMD) simulations and the analyses based on the local vibrational mode and quantum theory of atoms in molecules (QTAIM) methods. The anion---anion CiBs in the dimers of $[AuCl_a]^-$ and $[Au(CN)_a]^-$ previously observed in condensed phases were confirmed to be thoroughly repulsive under vacuum, but turned attractive in the crystal environment which was simulated using the solvation model based on density (SMD). However, the intrinsic strength of the inter-anion bonding is barely variated by the environment, as it is the combination of the inter-anion interaction and the environment effect that stabilizes the anion pairs. The block-localized wavefunction (BLW) method and its corresponding energy decomposition (BLW-ED) approach were further employed aiming at a chemically meaningful explanation for these counterintuitive phenomena. By inspecting the profiles of energy components, we identified the vital distinction between inter-anion CiBs and conventional non-covalent interactions lying in the electrostatic interaction, which variates nonmonotonically in the inter-anion complexes. The electrostatic interaction also dominates the depth of potential wells, which is commonly used to evaluate the kinetic stability, while Pauli exchange repulsion is the most repulsive factor preventing the formation of anion adducts. The importance of the Pauli exchange repulsion was further highlighted by comparing cases with and without metastability, in which the absence of a potential well is solely caused by the enhancement of the Pauli exchange repulsion.

1 Introduction

The spectrum of the noncovalent interactions has been remarkably expanded with newly recognized types, among which elements of both the p and d blocks on the periodic table can be directly involved as electrophilic sites. 1-11 Particularly, coinage bonds (CiBs, or regium bonds), 12,13 which refer to the close-shell interaction between the electron-deficient region on a coinage atom and the nucleophilic center of a Lewis base and have been determined and applied experimentally, 14-17 attract attention for their unusual nature. Unconventional cases or findings challenge and eventually enrich our understanding of noncovalent interactions even for the classical hydrogen bond. 18-20 A prominent example is the counterintuitive

inter-anion hydrogen bond, 18,21-26 which has already been utilized as a novel architectural linker in experimental studies. 23,27-32 The inter-anion hydrogen bond challenges the classical electrostatic model and the force field description of hydrogen bonds, given the incredible Coulombic repulsion between anions. 33,34 Similarly, inter-anion halogen bonds were also confirmed experimentally, 20,35-37 after pioneering theoretical predictions. 20 So far, the inter-anion/cation hydrogen bond, halogen bonds and many others have been well explored, 18,23,24,27,30-32,35,38-67 forming a diverse family of counterintuitive bonding. The inter-anion CiB term was first coined by Daolio et al.,14 who studied the crystal structures containing tetrachloridoaurates and subsequently characterized the Au···Cl short contact between [AuCl₄] anions by employing the quantum theory of atom in molecule (QTAIM)^{68,69} and natural bond orbital (NBO) theories.70-75 Notably, coinage metal complexes exhibit different oxidation states for central metals and various geometries, and may involve undiscovered inter-anion CiBs. Elucidating the nature of chemical bonds involving transition metals is also challenging due to their specific electronic structures.76-84

The history of CiBs may date back to the year 2000, when the Gerry lab studied the microwave spectra of Rg···M-X

^a Key Laboratory for Macromolecular Science of Shaanxi Province, School of Chemistry & Chemical Engineering, Shaanxi Normal University, Xi'an 710119, P. R. China. E-mail: snnu.changweiwang@snnu.edu.cn

^b Department of Nanoscience, Joint School of Nanoscience & Nanoengineering, University of North Carolina at Greensboro, Greensboro, NC 27401, USA. E-mail: y mo3@uncg.edu

[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d3cp00978e

complexes, where Rg, M and X stand for rare gas, group 11 and halogen atoms respectively.85-90 Complexes formed between M-X and a series of Lewis bases have been determined in experimental studies ever since. 91-95 Moreover, the Au···Cl bonding was spotted in the cytosine-derived gold(III) complex, according to the X-ray characterization and NBO analysis.⁹⁶ Various theoretical studies were carried out aiming at elucidating the bonding nature of CiBs, and the electrostatic model based on the σ -hole concept was usually adopted. ^{12,13,97–99} Before the remarkable work of Daolio et al., 14 other earlier experimental evidence for inter-anion CiBs can be found in the studies of the Leznoff group, who built coordination polymers using the [Au(CN)₄] units, and analyzed the Au···N interaction systematically in a subsequent work. 101

Theoretical investigations of inter-anion CiBs can follow the example of inter-anion hydrogen bonds, which have been intensively studied. Inter-anion hydrogen bonds can be analyzed from the perspectives of energy, force constant and electron density. The energy indicator was first introduced by Mata et al., who explored the energy profile along the interanion distance and found that there is a local minimum followed by a transition state (TS) impending the anion adduct from dissociation.²² Hence, the energy difference between the local minimum and TS (well-depth) serves as an energy measurement for the metastability. Alternatively, the local bond stretching force constant, 102-105 which is an "in situ" descriptor for the strength of a specified chemical bond, 106-112 has been utilized to describe the inter-anion hydrogen bonds. 113 Criteria for the inter-anion phenomena have also been developed based on the QTAIM approach, 68,69 which has been broadly applied to from the interpretations of novel chemical bonding¹¹⁴ to the developments of OM-based force fields for large-scale molecular simulations. 115,116 But the limitations of the QTAIM theory can be exemplified by the controversies over the $H \cdot \cdot \cdot H$ interaction in phenanthrene ^{117,118} and a series of intramolecular interactions. 119-121 Weinhold and Klein argued that the electrostatic interaction cannot be responsible for the inter-anion hydrogen bond, considering the Coulombic repulsion between net charges. Instead they proposed the charge transfer explanation and highlighted the governing role of orbital interactions with their NBO method.^{70–75} But this charge transfer model was queried by Frenking and Caramori,²¹ who stressed that it is unreliable to evaluate the electrostatic interaction based on a fallacious model where each entire anion is oversimplified to a point charge. There are also concerns that the NBO method tends to overestimate orbital interactions due to the non-optimization of bond orbitals, $^{21,33,122-124}$ and suffers from the basis set superposition error (BSSE), 125 though it has often been employed to probe the conventional and inter-anion CiBs. 12-14,98 By contrary, the electrostatic explanation of the kinetic stability has been supported by energy decomposition analyses (EDAs) with the key role of the electrostatic interaction in well-depths confirmed. 20,113,126 Notably, Alkorta et al. found that interanion interactions exhibit similar electron density distributions which are characterized by a series of chemical descriptors,

to their conventional counterparts, which can be well understood with the classical electrostatic model. 24,37,127 Similarly, a series of inter-cation non-covalent interactions have also been studied with their origins analyzed. 128-131

Obviously, theoretical methods with the capacity of quantify inter-anion physical factors, especially, the electrostatic and charge transfer interactions, are crucial for exploring the nature of inter-anion CiBs and resolving controversies, Many EDA methods have been developed and applied so far. 132-142 However, Foroutan-Nejad et al. pointed out that EDA approaches are path dependent. 143,144 Hence, it is necessary to examine the roles of key physical factors in the chemical bonding from multiple perspectives. For instance, the electrostatic interaction, which is homologous in most EDA approaches, 134,139,141,142,145-147 can be simulated using atomic multipole expansions as in force fields. 148-151 For the electron transfer interaction, ab initio valence bond (VB) theory 152,153 is preferable since it uses localized Lewis states to define a molecular wavefunction. As the simplest variant of VB theory, the block-localized wavefunction (BLW) method¹⁵⁴⁻¹⁵⁶ retains the characteristics of VB theory by using localized (fragmental) orbitals but simplifies the wavefunction with a single determinant for computational efficiency. Its energy decomposition (BLW-ED) approach 157,158 has also been developed and applied to inter-anion hydrogen, halogen and chalcogen bonds. 20,54,113

In this work, we intend to probe the nature of counterintuitive interactions in inter-anion CiBs theoretically. Representative coinage metal-containing anions with different geometries and oxidation states for central metals were selected 14,15,159-162 and their interactions with a series of simple anions were explored, with the goal of identifying new members of the inter-anion CiBs. In addition, the inter-anion CiBs found in crystal structures 14,100,101 were inspected in vacuum and condensed phase by employing the solvation model based on density (SMD). 163 Subsequently, the metastability was further examined using the QTAIM method, 68,69 the local vibrational mode theory, 104,105,164-166 and the ab initio molecular dynamics (AIMD) simulations. At last, key factors for the kinetic stability were elucidated by employing the BLW-ED approach. 157,158

2 Methods and computational details

2.1 BLW-ED approach

In the BLW-ED approach, the binding energy is defined as the overall variation in energy upon the formation of a complex from isolated and optimal monomers, and can be decomposed into several physically meaningful components, as shown in eqn (1). The deformation energy (ΔE_{def}) stands for the energy cost to distort monomer structures from their free and optimal states to the geometries in the complex. Bringing the deformed monomers from infinity to their positions in the complex without disturbing their respective electron densities leads to the frozen energy $(\Delta E_{\rm F})$, as defined in eqn (2), where the initial block-localized state (Ψ^{BLW0} in eqn (3)) is constructed using the wavefunction of distorted monomers (Φ_A^0 and Φ_B^0), and \hat{A} means

the antisymmetric operator. The frozen energy can be further separated into the electrostatic energy and Pauli exchange repulsion at the Hartree-Fock (HF) level (eqn (2)).

$$\Delta E_{\rm b} = \Delta E_{\rm def} + \Delta E_{\rm ele} + \Delta E_{\rm Pauli} + \Delta E_{\rm pol} + \Delta E_{\rm CT} + \Delta E_{\rm ec} \qquad (1)$$

$$\Delta E_{\rm F} = E(\Psi^{\rm BLW0}) - E(\Phi_{\rm A}^0) - E(\Phi_{\rm B}^0) = \Delta E_{\rm ele} + \Delta E_{\rm Pauli}$$
 (2)

$$\Psi^{\text{BLW0}} = \hat{A}(\Phi_{A}^{0}\Phi_{B}^{0}) \tag{3}$$

The electron densities of monomers can be adjusted within each of them to respond to the electric field generated by others. This process corresponds to the self-consistent optimization of the electron-localized state (Ψ^{BLW} in eqn (4)), and the subsequent stability gained is defined as the polarization energy (ΔE_{pol} in eqn (5)). The complex can be further stabilized by extending the movement of electrons to the entire complex, and this delocalization leads to the final HF wavefunction. The corresponding energy lowering is defined as the charge transfer energy, supplemented by the basis set superposition error (BSSE) (eqn (6)). Finally, the electron correlation contribution is defined as the difference in electron correlation energy between the complex and sum of deformed monomers (eqn (7)). In this work, the domain based local pair natural orbital coupled cluster method with single- and double-excitations (DLPNO-CCSD)¹⁶⁷ was employed for the calculation of electron correlations. Moreover, the BBSE is also divided into the HF and electron correlation parts, and assigned to the charge transfer and electron correlation components correspondingly (eqn (6) and (7)). Notably, the electron correlation term can be further decomposed based on the local energy decomposition (LED) analysis 168 within the DLPNO-CCSD(T) framework. However, the overall electron correlation is the least important contributor to the metastability among all energy terms as will be demonstrated, and therefore no further decomposition is conducted in this work.

$$\Psi^{\rm BLW} = \hat{A}(\Phi_{\rm A}\Phi_{\rm B}) \tag{4}$$

$$\Delta E_{\text{pol}} = E(\Psi^{\text{BLW}}) - E(\Psi^{\text{BLWO}})$$
 (5)

$$\Delta E_{\rm CT} = E(\Psi^{\rm HF}) - E(\Psi^{\rm BLW}) + BSSE^{\rm HF}$$
 (6)

$$\Delta E_{\rm ec} = E_{\rm corr}^{\rm AB} - E_{\rm corr}^{\rm A} - E_{\rm corr}^{\rm B} + {\rm BSSE}^{\rm corr}$$
 (7)

To study the binding energy in the crystal state, the SMD method is adopted to simulate the crystal environment. The total energy of the system (E^{SMD} in eqn (8)) is composed of the internal energy of the focused complex (E^{I}) and the energy of complex-surrounding interaction (V), which is dominated by the electrostatic forces. Therefore, the total binding energy in the crystal environment ($\Delta E_{\rm b}^{\rm SMD}$ in eqn (9)) can be simply divided into the intrinsic component ($\Delta E_{\rm b}^{\rm I}$), which stands for the difference in internal energy between the complex and optimal monomers, and the complex-surrounding component (ΔV) , which corresponds to the variation in the energy of the complex-surrounding interaction from the isolated and optimized monomers to the formation of the complex.

$$\Delta E^{\text{SMD}} = E^{\text{I}} + V \tag{8}$$

$$\Delta E_{\rm b}^{\rm SMD} = \Delta E_{\rm b}^{\rm I} + \Delta V \tag{9}$$

2.2 Computational details

Thirty-one anion ··· anion complexes were constructed and are shown in Scheme 1. The def2-TZVP basis set was selected and the small-core relativistic pseudopotential (def2-TZVP-PP) was utilized for gold and silver. 169,170 For complexes 1-6, the energy profiles along the inter-anion distance were evaluated by first performing constrained geometry optimizations, in which only the bond distance (R in Scheme 1) was fixed at a series of values with all the rest of the geometrical parameters optimized. A series of theoretical methods were tested, including B97-D3,¹⁷¹ PBE0-D3,¹⁷² M06-2X-D3,¹⁷³ B3LYP-D3,¹⁷⁴ CAM-B3LYP-D3,¹⁷⁵ ωB97x-D2¹⁷⁶ and SCS-MP2.¹⁷⁷ Afterwards, single point calculations were carried out at each point by employing the DLPNO-CCSD method, 167 with the resolution of the identity approximation 178 and frozen core technique adopted (RI-DLPNO-CCSD) for computational efficiency. Complex 4 is a crucial case because [Au(CN)₄]⁻ and I⁻ become thoroughly repulsive according to the RI-DLPNO-CCSD energy profile, while shallow potential wells $(-3.5 \text{ to } -5.2 \text{ kJ mol}^{-1})$ were observed based on all other methods except for ωB97x-D2 and SCS-MP2 (Fig. S1v-a and b, ESI†). In general, well-depths derived from ωB97x-D2 and SCS-MP2 are nearly the same and both larger than the

Au(III)	F ⁻	Cl ⁻	Br ⁻	ľ	CN-	OCI-
[Au(CN)	₄] ⁻ 1	2	3	4	5	6
[AuCl ₄]	· 7	8	9	10	11	12
[AuBr ₄]	· 13	14	15	16	17	18
[Aul ₄]	19	20	21	22	23	24

Au/Ag(I)	[Au(CN) ₂] ⁻	[Au	0] [.]	[Ag(CN) ₂]	
F ⁻	25	26	5	27	
	Y=CN-	CI ⁻	Br ⁻	I-	
[Au(III)Y ₄] ⁻ Dimer	28	29	30	31	

=CN⁻, Cl⁻, Br⁻ and I Y=CN⁻, Cl⁻, Br⁻ and I M=Au and Ag, Y=CN

Scheme 1 Complexes 1-31 studied in this work

RI-DLPNO-CCSD results (Fig. S1, ESI†). To examine the reliability of our above computations, we performed single point calculations for complex 1 with the perturbative triples (T) correction incorporated (i.e., RI-DLPNO-CCSD(T)), and also re-optimized the geometries using enlarged basis sets. It turned out that increased theoretical levels barely influence the welldepths (within 0.4 kJ mol⁻¹). Therefore, the ωB97x-D2 method was eventually selected for geometry optimizations and vibrational frequency calculations, and the RI-DLPNO-CCSD method was employed for the subsequent BLW-ED analysis of the binding energy profiles, aiming at a rigorous examination of the metastability.

AIMD simulations in vacuum at 298 K were carried out at the ωB97x-D2/def2-TZVP(-PP) theoretical level for the anion adducts which exhibit metastability in vacuum. Local minima were chosen as the initial structures and the initial atomic velocities were obtained based on the Boltzmann-Maxwell distribution randomly. The Berendsen thermostat was utilized and a step size of 1 fs was selected. Simulations with 5000 steps were accomplished for each case with the quantitative statics performed using results after the equilibration (i.e., 1500–5000 fs in each simulation).

Geometrical optimizations, vibrational frequency calculations, BLW-ED and GKS-EDA at the HF level were performed using the in-house version of GAMESS (US). 179 Orca 180,181 was used for the electron correlation energies and AIMD simulations. In addition, local bond stretching force constants of inter-anion CiBs were derived from the local vibrational mode theory. 104,105,164-166 QTAIM analysis was performed using the Multiwfn package. 182 QM-based atomic charges and multipoles were obtained by using the natural population analysis (NPA) and the GDMA¹⁸³ program respectively. For these NPA and GDMA calculations, the ωB97x functional and the 6-311G basis set were selected except that for Au and Ag the Lanl2DZ pseudopotentials were adopted. Afterwards, the electrostatic energy was reevaluated along the inter-anion distance using the atomic multipole expansion scheme in the AMOEBA polarizable force field 184-187 incorporated in the Tinker software. 188

3 Results and discussions

Binding energy profiles for complex 1 ($[Au(CN)_4]^- \cdots F^-$) and 4 $([Au(CN)_4]^- \cdots I^-)$ are plotted in Fig. 1. The metastability of $[Au(CN)_4]^- \cdots F^-$ was evidenced by the local minimum, which can trap the anion adduct. On the contrary, the binding energy between [Au(CN)₄]⁻ and I⁻ monotonously gets repulsive as the inter-anion distance shrinks, suggesting a repulsive nature of the inter-anion interaction. All cases were similarly examined (Fig. S2, ESI†) and the characteristic potential wells were found in complexes 1, 2, 5, 6, 26 and 27 (colored in blue in Scheme 1). Among all tetracoordinated coinage metal complexes, only $[Au(CN)_4]^-$ can form inter-anion CiBs. This is due to the fact that cyanide is the strongest electron-withdrawing group tested, and can considerably reduce the electron density of Au. Furthermore, a considerable barrier height (-42.5 kJ mol⁻¹)

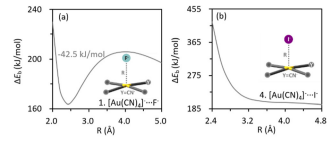


Fig. 1 Profiles of binding energies along the inter-anion distances for complexes 1 (a) and 4 (b) computed at the RI-DLPNO-CCSD//ωB97x-D2/ def2-TZVP theoretical level

was found in $[Au(CN)_4]^- \cdots F^-$, in which F^- has the smallest radius among anions and therefore, the closest contact with the coinage center. Metastability was absent in [Au(CN)₂]-···F-(complex 25), while observed in $[Ag(CN)_2]^- \cdots F^-$ (complex 27), providing us comparative examples to clarify the roles of coinage metals to the metastability. Inter-anion CiB was also observed in $[AuO]^- \cdots F^-$ (complex 26), which exhibits a linear structure as the conventional CiBs illuminated previously in a theoretical study.¹¹ Besides, substitution reactions were observed in complexes 7, 13, 19, 23 and 24. For the example of complex 7, one Cl in [AuCl₄] is switched with F after reaching the (Coulombic) barrier height, and the infinitely separated Cl⁻ and [AuF(Cl)₃]⁻ anions are eventually observed (Fig. S2g, ESI†).

Intriguingly, the dimer of [AuCl₄]⁻ turns more destabilizing as the inter-anion distance decreases, suggesting that the experimentally observed inter-anion interaction cannot hold in vacuum. This repulsive nature was also confirmed by reevaluating the binding energy profile at higher theoretical levels in vacuum (Fig. S2a and c, ESI†). Obviously, the packing effect in the crystal structure plays an essential role in stabilizing the anion ... anion pairs. The effect of the crystal environment on the binding energy profile was subsequently simulated using the SMD method, with a series of dielectric constants tested (Fig. 2). Indeed, the characteristic potential well appeared when the environmental effect was incorporated, and even negative binding energies were observed. Importantly, the intrinsic binding energy in the [AuCl₄] dimer is barely variated by the SMD method (Fig. 2b), while the solute-solvent term, which stands for the interaction between anion pair and the crystal environment in this work, turns more stabilizing drastically at short ranges (Fig. 2c) and gives rise to the local minimum. In other words, it is the interaction between the anion pair and crystal environment that stabilizes the system in condensed phases. Similarly, the key role of the surrounding to the inter-anion CiBs between [Au(CN)₄] anions was also confirmed (Fig. 2d-f). The intrinsic binding energies of aforementioned dimers were not further decomposed because of the absence of metastability. Besides, the absence of metastability in vacuum was observed in all AuY_4^- (Y = CN⁻, Cl⁻, Br⁻ and I⁻) dimers (Fig. S2ab-ae, ESI†). In the following part, we performed theoretical analyses only for cases exhibiting inter-anion CiBs in vacuum.

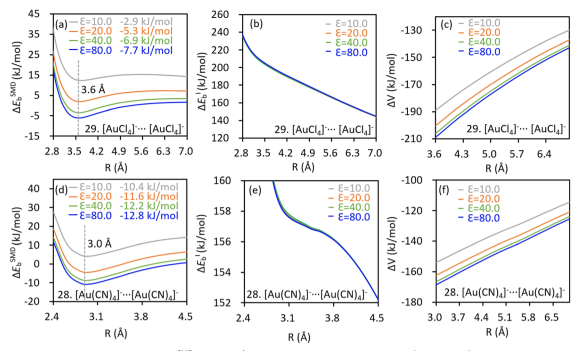


Fig. 2 Profiles of total binding energy (a and d, ΔE_b^{SMD} in kJ mol⁻¹), intrinsic binding energy (b and e, ΔE_b^I in kJ mol⁻¹) and the solute–solvent interaction term (c and f, ΔV in kJ mol⁻¹) along the inter-anion distance for complexes [AuCl₄]⁻...[AuCl₄]⁻ and [Au(CN)₄]⁻...[Au(CN)₄]⁻, obtained from single point calculations at the ωB97X-D2/def2-TZVP theoretical level with a series of dielectric constants selected.

The kinetic stabilities of complex $[Au(CN)_4]^- \cdots F^-$ were also evidenced by the AIMD simulations, in which dissociation of the anion adduct was not observed. As shown in Fig. 3a, the bond distance deviates from the average value (\bar{R}) insignificantly. AIMD results of the other complexes exhibiting the inter-anion CiBs are shown in Fig. S3 (ESI†). Notably, the maximum/minimum deviations of the bond distance correlate with the localized bond force constants, which stand for the curvatures of binding energy profiles around the local minima, in an appropriate linear manner (Fig. 3b). In other words, a steeper curve leads to less significant deviations of bond lengths in the AIMD simulation. For example, complex 26 ([AuO] -···F) exhibits the greatest bond force constant (0.982 mDyn \mathring{A}^{-1}) and therefore, the least deviation in bond length (Fig. S3e, ESI†). Besides, the average bond distance in the AIMD simulations is close to the optimal value (Table 1). The interanion CiBs were further supported by the QTAIM theory,

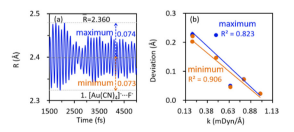


Fig. 3 (a) Evolution of the Au···F⁻ distance over time (after the 1500st fs) in the AIMD simulation of complex 1 and (b) correlation between the maximum/minimum deviations of the bond distances from the average values and localized bond force constants

Table 1 Bond distances (R, in Å), average bond distances (\bar{R} , in Å) from the AIMD simulations, electron density (ρ , in a.u.) and the Laplace ($\nabla^2 \rho$, in a.u.) values at the BCPs of complexes with metastability

No.	Complexes	R	$ar{R}$	ho	$\nabla^2 \rho$
1	$[Au(CN)_4]^-\cdots F^-$	2.360	2.360	0.052	0.226
2	[Au(CN)₄] -···Cl	2.980	3.024	0.026	0.075
5	$[Au(CN)_4]^-\cdots CN^-$	2.620	2.656	0.042	0.112
6	$[Au(CN)_4]^-\cdots OCl^-$	2.535	2.561	0.040	0.142
26	$[AuO]^{-}\cdots F^{-}$	2.157	2.159	0.072	0.354
27	$[Ag(CN)_2]^-\cdots F^-$	2.284	2.274	0.050	0.254

because a bond critical point (BCP) with appropriate electron density and positive Laplace values were observed in each case with metastability, suggesting close-shell interactions.

Roles of individual energy components in the metastability can be revealed intuitively by examining their corresponding profiles along the inter-anion distances (Fig. 4). In each complex, the polarization, charge transfer and electron correlation interactions all become more attractive as anions approach each other, suggesting a preference for the local minimum over the transition states (TS), which has a stretched inter-anion distance. Conversely, both the Pauli exchange repulsion and deformation energy grow more destabilizing as the inter-anion distance shrinks, contributing negatively to the well depths. The repulsive electrostatic interaction increases as the length of inter-anion CiB shrinks at long ranges, but turns to decrease drastically after reaching the Coulombic repulsion barrier, indicating a positive contribution to the well depth. The decreasing trend of electrostatic interaction can be rationalized by the nucleus-electron attraction, which can overturn the

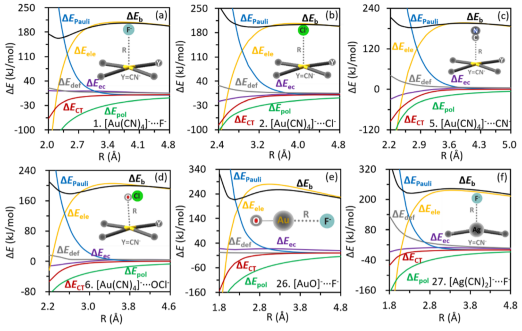


Fig. 4 Variations of energy components along with the inter-anion distances in (a) $[Au(CN)_4]^-\cdots F^-$; (b) $[Au(CN)_4]^-\cdots Cl^-$; (c) $[Au(CN)_4]^-\cdots CN^-$; (d) $[Au(CN)_4]^-\cdots OCl^-$; (e) $[AuO]^-\cdots F^-$; (f) $[Ag(CN)_2]^-\cdots F^-$

electrostatic repulsion among electrons and among nuclei at short ranges. Actually, the nonmonotonic profile of electrostatic interaction was also observed in other inter-anion interactions, 54,113 and can be regarded as the key distinction between inter-anion and conventional interactions, in which all energy terms variate monotonically along the bond length. The electrostatic energy profiles of complexes 1, 26 and 27 were reevaluated using the QM-based atomic multipoles, and the vital nonmonotonicity was reproduced (Fig. S4, ESI†). Hence, the electrostatic interaction in inter-anion CiBs can be simulated and explained based on the classical distributed atomic multipole expansion.

The origin of potential wells can be clarified by examining the differences in energy components between local minima and the corresponding TSs as listed in Table 2. Negative values mean the stabilities (lowering in energy) gained from a TS to a local minimum, following the convention in the BLW-ED scheme. All the well depths ($\Delta\Delta E_{\rm b}$) are dominated by the electrostatic interaction, with the polarization interaction acting as the second most important factor. Both the charge transfer and electron correlation contribute positively to the kinetic stability, while Pauli exchange repulsion and deformation tend to shallow

the potential wells. The governing role of electrostatic interactions is in line with its decreasing trend at short ranges as shown in Fig. 4. The positive role of polarization can be rationalized by the considerable polarizability of the coinage donor (Table S1, ESI†). All the aforementioned contributions of physical factors were also observed in our previous studies of inter-anion hydrogen, halogen and chalcogen interactions. 20,54,113 The electrons are shifted from F or OCl to the metal complexes according to the natural population analysis (NPA in Table S2, ESI†), exhibiting a similar pattern as the conventional non-covalent interactions (from Lewis base to acid). Besides, we redrew the energy profiles by using the def2-TZVPD basis set and found that the metastability and the lengths of inter-anion CiBs (Table S3, ESI†) are barely variated.

The potential well vanishes when Ag(I) in $[Ag(CN)_2]^- \cdots F^-$ (complex 27) is replaced by Au(1) (complex 25). We analyzed the role of the central metal by comparing individual energy terms in complexes 27 and 25 at a series of fixed inter-anion distances, ranging from the local minimum of $[Ag(CN)_2]^- \cdots F^-$ to the TS. Fig. 5a illustrates the correlations of important energy components. The electrostatic interactions in both cases are pretty close within the interval of distances selected, according to the linear correlation with a slope of 1.0. Interestingly, the

Table 2 The difference in energy components (in kJ mol⁻¹) between local minima and transition states (TSs)

No.	Complexes	$\Delta\Delta E_{ m def}$	$\Delta \Delta E_{ m ele}$	$\Delta \Delta E_{\rm Pauli}$	$\Delta\Delta E_{\rm pol}$	$\Delta\Delta E_{\rm CT}$	$\Delta\Delta E_{ m ec}$	$\Delta\Delta E_{ m b}$
1	$[Au(CN)_4]^-\cdots F^-$	5.0	-101.7	135.9	-69.1	-16.8	4.2	-42.5
2	$[Au(CN)_4]^- \cdots Cl^-$	2.8	-20.4	41.1	-18.0	-6.9	-5.6	-7.1
5	$[Au(CN)_4]^-\cdots CN^-$	7.2	-80.9	124.3	-29.9	-23.0	-10.9	-13.3
6	$[Au(CN)_4]^- \cdots OCl^-$	3.6	-53.4	89.1	-33.8	-14.2	-9.6	-18.3
26	[AuO]-···F-	0.7	-121.9	198.1	-77.2	-36.9	-3.0	-40.2
27	$[Ag(CN)_2]^-\cdots F^-$	26.3	-77.0	82.1	-40.6	-6.6	-2.0	-17.7

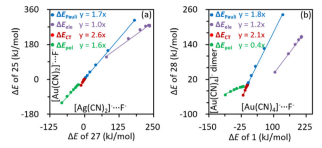


Fig. 5 Correlations of energy components calculated at a series of fixed inter-anion distances between complexes with and without kinetic stability: (a) $[Au(CN)_2]^- \cdots F^- vs$. $[Ag(CN)_2]^- \cdots F^-$ and (b) $[Au(CN)_4]^-$ dimer Vs [Au(CN)₄]⁻···F⁻

[Au(CN)₂]⁻···F⁻ even exhibits more stabilizing polarization and charge transfer interactions than $[Ag(CN)_2]^- \cdots F^-$ (slopes greater than one). This is understandable because [Au(CN)₂] has a higher polarizability than [Ag(CN)₂] (Table S1, ESI†), and therefore, stronger polarization. Moreover, Au(I) has a larger size than Ag(1), leading to stronger inter-anion orbital overlap. Nevertheless, the enhanced orbital overlap also increases the destabilizing Pauli exchange repulsion, which eventually leads to the absence of metastability in $[Au(CN)_4]^- \cdots F^-$. We also compared the BLW-ED results of the [Au(CN)₄] dimer with the complex $[Au(CN)_4]^- \cdots F^-$ (Fig. 5b). In general, from the TS to the local minimum of $[Au(CN)_4]^- \cdots F^-$, the variation of the overall stabilizing factors is -187.6 kJ mol⁻¹, which is almost the same as the corresponding change in the $[Au(CN)_4]^-$ dimer (-183.3 kJ mol⁻¹). However, the enhancement in Pauli exchange repulsion is much greater in the [Au(CN)₄] dimer (240.5 vs. 135.9 kJ mol⁻¹) and eventually erases the metastability. Liner correlations between energy components of these two complexes were also observed (Fig. 5b). Comparable electrostatic interactions were found, while stronger charge transfer interaction was discovered in the [Au(CN)₄] dimer. It is reasonable because F is the weakest electron donor studied in this work. F- has more a concentrated electron density distribution than [Au(CN)₄]⁻, and therefore imposes a stronger electric field to [Au(CN)₄]⁻, leading to stronger polarization in $[Au(CN)_4]^- \cdots F^-$.

4. Conclusions

Binding energy profiles of thirty-one anion pairs involving CiBs were explored along the inter-anion distances, employing stateof-the-art theoretical methods. It turns out that the anions [Au(CN)₄]⁻, [Ag(CN)₂]⁻ and [AuO]⁻ are all possible candidates for the formation of inter-anion CiBs in vacuum. The kinetic stability was further supported by the AIMD simulations, the QTAIM theory and the local vibrational analyses. Two general characters can be derived from the cases with metastability. One is the coordination of CN⁻ on the coinage metal, and the other is the small size of the anion acting as the Lewis base. This is understandable because cyanide usually leads to a remarkable electron deficiency (relatively) of the metal, and

the anions with small radius may approach the central metal closely.

The inter-anion CiBs in $[AuCl_4]^- \cdots [AuCl_4]^ [Au(CN)_4]^- \cdots [Au(CN)_4]^-$ complexes identified in experimentally obtained crystal structures were proved to be absent in vacuum, but slightly attractive when the crystal environment is simulated with the SMD method. However, it should be emphasized that the intrinsic strength of the inter-anion interaction was barely variated by the environment, and the changes of the binding energy profiles originate from the solute-solvent term, which stands for the interaction between the anion pair and the crystal environment in this work.

Variations in individual energy components along the inter-anion distances were inspected and the distinction of inter-anion CiBs from conventional non-covalent interactions was found to be the electrostatic interaction, which variates nonmonotonically in anion ··· anion complexes. The distinctive nonmonotonicity of electrostatic profiles was captured by the classical atomic multipole expansion, offering us a concise model to rationalize and simulate the electrostatic interaction in these counterintuitive phenomena. The electrostatic interaction also governs the well-depths. In addition, the polarization, charge transfer and electron correlation all prefer the local minima over TSs, while the deformation energy, and especially the Pauli exchange repulsion become more destabilizing from TSs to local minima, playing negative roles in the well-depths. The significance of Pauli exchange repulsion in the metastability was highlighted by comparing the BLW-ED results of cases with and without metastability at a series of fixed inter-anion distances. In detail, from [Ag(CN)2]-...F to $[Au(CN)_2]^- \cdots F^-$, the increasing size of the central metal strengthens the destabilizing Pauli exchange repulsion, which eliminates the potential well. Similarly, the Pauli exchange repulsion is remarkably enhanced when F in [Au(CN)₄] ···· F is replaced by $[Au(CN)_4]^-$, leading to the absence of metastability.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

CW acknowledges the support from the Natural Science Foundation of China (No. 22073060). This work was performed in part at the Joint School of Nanoscience and Nanoengineering, a member of the National Nanotechnology Coordinated Infrastructure (NNCI), which is supported by the National Science Foundation (Grant ECCS-2025462).

Notes and references

1 A. Daolio, A. Pizzi, M. Calabrese, G. Terraneo, S. Bordignon, A. Frontera and G. Resnati, Angew. Chem., Int. Ed., 2021, 60, 20723-20727.

- 2 A. Bauza, I. Alkorta, J. Elguero, T. J. Mooibroek and A. Frontera, Angew. Chem., Int. Ed., 2020, 59, 17482-17487.
- 3 A. Bauza, T. J. Mooibroek and A. Frontera, Angew. Chem., Int. Ed., 2013, 52, 12317-12321.
- 4 J. Fanfrlik, A. Prada, Z. Padelkova, A. Pecina, J. Machacek, M. Lepsik, J. Holub, A. Ruzicka, D. Hnyk and P. Hobza, Angew. Chem., Int. Ed., 2014, 53, 10139-10142.
- 5 S. Zahn, R. Frank, E. Hey-Hawkins and B. Kirchner, Chem. --Eur. J., 2011, 17, 6034-6038.
- 6 A. Daolio, A. Pizzi, G. Terraneo, A. Frontera and G. Resnati, ChemPhysChem, 2021, 22, 2281-2285.
- 7 I. Alkorta, J. Elguero and A. Frontera, Crystals, 2020, 10, 180.
- 8 S. Mirdya, A. Frontera and S. Chattopadhyay, CrystEngComm, 2019, 21, 6859-6868.
- 9 L. Guan and Y. Mo, J. Phys. Chem. A, 2014, 118, 8911-8921.
- 10 A. C. Legon, Phys. Chem. Chem. Phys., 2010, 12, 7736-7747.
- 11 A. C. Legon and N. R. Walker, Phys. Chem. Chem. Phys., 2018, 20, 19332-19338.
- 12 A. Frontera and A. Bauza, Chem. Eur. J., 2018, 24, 7228-7234.
- 13 R. Wang, Z. Wang, X. Yu and Q. Li, ChemPhysChem, 2020, 21, 2426-2431.
- 14 A. Daolio, A. Pizzi, G. Terraneo, M. Ursini, A. Frontera and G. Resnati, Angew. Chem., Int. Ed., 2021, 60, 14385-14389.
- 15 A. Pizzi, M. Calabrese, A. Daolio, M. Ursini, A. Frontera and G. Resnati, CrystEngComm, 2022, 24, 3846-3851.
- 16 M. L. N. Pina, A. Frontera and A. Bauza, J. Phys. Chem. Lett., 2020, 11, 8259-8263.
- 17 J. Halldin Stenlid, A. J. Johansson and T. Brinck, Phys. Chem. Chem. Phys., 2018, 20, 2676-2692.
- 18 F. Weinhold and R. A. Klein, Angew. Chem., Int. Ed., 2014, 53, 11214-11217.
- 19 X. Zhang, H. Dai, H. Yan, W. Zou and D. Cremer, J. Am. Chem. Soc., 2016, 138, 4334-4337.
- 20 C. Wang, Y. Fu, L. Zhang, D. Danovich, S. Shaik and Y. Mo, J. Comput. Chem., 2018, 39, 481-487.
- 21 G. Frenking and G. F. Caramori, Angew. Chem., Int. Ed., 2015, 54, 2596-2599.
- 22 I. Mata, I. Alkorta, E. Molins and E. Espinosa, Chem-PhysChem, 2012, 13, 1421-1424.
- 23 M. O. Miranda, D. J. R. Duarte and I. Alkorta, Chem-PhysChem, 2020, 21, 1052-1059.
- 24 L. M. Azofra, J. Elguero and I. Alkorta, J. Phys. Chem. A, 2020, 124, 2207-2214.
- 25 I. Mata, E. Molins, I. Alkorta and E. Espinosa, J. Phys. Chem. A, 2015, 119, 183-194.
- 26 I. Iribarren, M. M. Montero-Campillo, I. Alkorta, J. Elguero and D. Quinonero, Phys. Chem. Chem. Phys., 2019, 21, 5796-5802.
- 27 E. M. Fatila, E. B. Twum, A. Sengupta, M. Pink, J. A. Karty, K. Raghavachari and A. H. Flood, Angew. Chem., Int. Ed., 2016, 55, 14057-14062.
- 28 J. S. McNally, X. P. Wang, C. Hoffmann and A. D. Wilson, Chem. Commun., 2017, 53, 10934-10937.
- 29 E. M. Fatila, E. B. Twum, J. A. Karty and A. H. Flood, Chem. - Eur. J., 2017, 23, 10652-10662.

- 30 W. Zhao, A. H. Flood and N. G. White, Chem. Soc. Rev., 2020, 49, 7893-7906.
- 31 N. G. White, CrystEngComm, 2019, 21, 4855-4858.
- 32 T. S. C. MacDonald, B. L. Feringa, W. S. Price, S. J. Wezenberg and J. E. Beves, J. Am. Chem. Soc., 2020, 142, 20014-20020.
- 33 F. Weinhold, Angew. Chem., Int. Ed., 2003, 42, 4188-4194.
- 34 F. Weinhold, Molecules, 2022, 27, 377.
- 35 J. M. Holthoff, E. Engelage, R. Weiss and S. M. Huber, Angew. Chem., Int. Ed., 2020, 59, 11150-11157.
- 36 C. Loy, J. M. Holthoff, R. Weiss, S. M. Huber and S. V. Rosokha, Chem. Sci., 2021, 12, 8246-8251.
- 37 D. Quinonero, I. Alkorta and J. Elguero, Phys. Chem. Chem. Phys., 2016, 18, 27939-27950.
- 38 A. Strate, T. Niemann, D. Michalik and R. Ludwig, Angew. Chem., Int. Ed., 2017, 56, 496-500.
- 39 R. Barbas, R. Prohens, A. Bauza, A. Franconetti and A. Frontera, Chem. Commun., 2019, 55, 115-118.
- 40 D. A. Cullen, M. G. Gardiner and N. G. White, Chem. Commun., 2019, 55, 12020-12023.
- 41 R. Prohens, A. Portell, M. Font-Bardia, A. Bauza and A. Frontera, Chem. Commun., 2018, 54, 1841-1844.
- 42 F. Zapata, L. Gonzalez, A. Bastida, D. Bautista and A. Caballero, Chem. Commun., 2020, 56, 7084-7087.
- 43 G. Dos Passos Gomes, G. Xu, X. Zhu, L. M. Chamoreau, Y. Zhang, O. Bistri-Aslanoff, S. Roland, I. V. Alabugin and M. Sollogoub, Chem. - Eur. J., 2021, 27, 8127-8142.
- 44 J. M. Holthoff, R. Weiss, S. V. Rosokha and S. M. Huber, Chem. - Eur. J., 2021, 27, 16530-16542.
- 45 E. M. Fatila, M. Pink, E. B. Twum, J. A. Karty and A. H. Flood, Chem. Sci., 2018, 9, 2863-2872.
- 46 A. Knorr, P. Stange, K. Fumino, F. Weinhold and R. Ludwig, ChemPhysChem, 2016, 17, 458-462.
- 47 T. Niemann, P. Stange, A. Strate and R. Ludwig, ChemPhysChem, 2018, 19, 1691-1695.
- 48 D. Quinonero, I. Alkorta and J. Elguero, ChemPhysChem, 2020, 21, 1597-1607.
- 49 R. Wysokinski, W. Zierkiewicz, M. Michalczyk and S. Scheiner, ChemPhysChem, 2020, 21, 1119–1125.
- 50 R. Wysokinski, W. Zierkiewicz, M. Michalczyk and S. Scheiner, ChemPhysChem, 2021, 22, 818–821.
- 51 W. Zierkiewicz, R. Wysokinski, M. Michalczyk and S. Scheiner, ChemPhysChem, 2020, 21, 870–877.
- 52 F. Weinhold, Inorg. Chem., 2018, 57, 2035-2044.
- 53 W. Zhao, J. Tropp, B. Qiao, M. Pink, J. D. Azoulay and A. H. Flood, J. Am. Chem. Soc., 2020, 142, 2579-2591.
- 54 D. Fan, L. Chen, C. Wang, S. Yin and Y. Mo, J. Chem. Phys., 2021, 155, 234302.
- 55 S. Scheiner, J. Chem. Phys., 2020, 153, 140901.
- 56 S. Scheiner, R. Wysokinski, M. Michalczyk and W. Zierkiewicz, J. Phys. Chem. A, 2020, 124, 4998-5006.
- 57 J. Chen, K. Qian, K. Xiao, J. Luo, H. Li, T. Ma, U. Kortz, M. Tsige and T. Liu, Langmuir, 2020, 36, 10519-10527.
- 58 F. Khorrami and M. H. Kowsari, J. Phys. Chem. B, 2020, 124, 3770-3783.

59 A. Grabarz, M. Michalczyk, W. Zierkiewicz and S. Scheiner, Molecules, 2021, 26, 2116.

- 60 J. K. Philipp and R. Ludwig, Molecules, 2020, 25, 4972.
- 61 L. M. Azofra, J. Elguero and I. Alkorta, *Phys. Chem. Chem. Phys.*, 2020, 22, 11348–11353.
- 62 T. Niemann, P. Stange, A. Strate and R. Ludwig, *Phys. Chem. Chem. Phys.*, 2019, 21, 8215–8220.
- 63 R. Wysokinski, W. Zierkiewicz, M. Michalczyk and S. Scheiner, Phys. Chem. Chem. Phys., 2021, 23, 13853–13861.
- 64 A. Gogoi, D. Dutta, A. K. Verma, H. Nath, A. Frontera, A. K. Guha and M. K. Bhattacharyya, *Polyhedron*, 2019, **168**, 113–126.
- 65 A. Knorr and R. Ludwig, Sci. Rep., 2015, 5, 17505.
- 66 S. Scheiner, J. Comput. Chem., 2022, 43, 1814-1824.
- 67 L. Andreo, R. M. Gomila, E. Priola, A. Giordana, S. Pantaleone, E. Diana, G. Mahmoudi and A. Frontera, *Cryst. Growth Des.*, 2022, 22, 6539–6544.
- 68 I. Mata, E. Molins, I. Alkorta and E. Espinosa, *J. Chem. Phys.*, 2009, **130**, 044104.
- 69 R. F. W. Bader, Atoms in Molecules: A Quantum Theory, Oxford University Press, Oxford, UK, 1990.
- 70 F. Weinhold and C. R. Landis, Chem. Educ. Res. Pract., 2001, 2, 91-104.
- 71 A. E. Reed, L. A. Curtiss and F. Weinhold, *Chem. Rev.*, 1988, **88**, 899–926.
- 72 F. Weinhold and C. R. Landis, *Discovering Chemistry With Natural Bond Orbitals*, Wiley, Hoboken, NJ, 2012.
- 73 F. Weinhold, C. R. Landis and E. D. Glendening, *Int. Rev. Phys. Chem.*, 2016, 35, 399-440.
- 74 E. D. Glendening, C. R. Landis and F. Weinhold, *J. Comput. Chem.*, 2019, 40, 2234–2241.
- 75 E. D. Glendening, C. R. Landis and F. Weinhold, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2011, 2, 1–42.
- 76 G. Ohanessian and W. A. Goddard, Acc. Chem. Res., 1990, 23, 386–392.
- 77 M. Kaupp, Angew. Chem., Int. Ed., 2001, 40, 3534-3565.
- 78 S. Radenković, D. Danovich, S. Shaik, P. C. Hiberty and B. Braïda, *Comput. Theor. Chem.*, 2017, **1116**, 195–201.
- 79 J. Joy, D. Danovich, M. Kaupp and S. Shaik, *J. Am. Chem. Soc.*, 2020, **142**, 12277–12287.
- 80 J. B. Mann, T. L. Meek, E. T. Knight, J. F. Capitani and L. C. Allen, J. Am. Chem. Soc., 2000, 122, 5132–5137.
- 81 D. G. Liakos and F. Neese, *J. Chem. Theory Comput.*, 2011, 7, 1511–1523.
- 82 M. Kaupp, J. Comput. Chem., 2007, 28, 320-325.
- 83 J. M. Galbraith, A. Shurki and S. Shaik, *J. Phys. Chem. A*, 2000, **104**, 1262–1270.
- 84 S. Dordevic, S. Radenkovic, S. Shaik and B. Braida, *Molecules*, 2022, 27, 490.
- 85 C. J. Evans, A. Lesarri and M. C. L. Gerry, *J. Am. Chem. Soc.*, 2000, 122, 6100–6105.
- 86 J. M. Michaud and M. C. L. Gerry, J. Am. Chem. Soc., 2006, 128, 7613–7621.
- 87 J. M. Thomas, N. R. Walker, S. A. Cooke and M. C. L. Gerry, J. Am. Chem. Soc., 2004, 126, 1235–1246.
- 88 C. J. Evans and M. C. L. Gerry, *J. Chem. Phys.*, 2000, **112**, 9363–9674.

- 89 C. J. Evans and M. C. L. Gerrya, J. Chem. Phys., 2000, 112, 1321–1329.
- 90 C. J. Evans, D. S. Rubinoff and M. C. L. Gerry, *Phys. Chem. Chem. Phys.*, 2000, 2, 3943–3948.
- 91 S. G. Francis, S. L. Matthews, O. Poleshchuk, N. R. Walker and A. C. Legon, *Angew. Chem., Int. Ed.*, 2006, 45, 6341–6343.
- 92 S. J. Harris, A. C. Legon, N. R. Walker and D. E. Wheatley, Angew. Chem., Int. Ed., 2010, 49, 181–183.
- 93 C. Medcraft, E. Gougoula, D. M. Bittner, J. C. Mullaney, S. Blanco, D. P. Tew, N. R. Walker and A. C. Legon, *J. Chem. Phys.*, 2017, 147, 234308.
- 94 S. L. Stephens, W. Mizukami, D. P. Tew, N. R. Walker and A. C. Legon, *J. Chem. Phys.*, 2012, **136**, 064306.
- 95 J. C. Mullaney, S. L. Stephens, D. P. Zaleski, M. J. Sprawling, D. P. Tew, N. R. Walker and A. C. Legon, *J. Phys. Chem. A*, 2015, **119**, 9636–9643.
- 96 A. Terron, J. Buils, T. J. Mooibroek, M. Barcelo-Oliver, A. Garcia-Raso, J. J. Fiol and A. Frontera, *Chem. Commun.*, 2020, 56, 3524–3527.
- 97 J. H. Stenlid and T. Brinck, J. Am. Chem. Soc., 2017, 139, 11012–11015.
- 98 W. Zierkiewicz, M. Michalczyk and S. Scheiner, *Phys. Chem. Chem. Phys.*, 2018, **20**, 22498–22509.
- 99 C. Trujillo, G. Sánchez-Sanz, J. Elguero and I. Alkorta, *Struct. Chem.*, 2020, **31**, 1909–1918.
- 100 C. J. Shorrock, H. Jong, R. J. Batchelor and D. B. Leznoff, *Inorg. Chem.*, 2003, 42, 3917–3924.
- 101 A. R. Geisheimer, J. E. Wren, V. K. Michaelis, M. Kobayashi, K. Sakai, S. Kroeker and D. B. Leznoff, *Inorg. Chem.*, 2011, 50, 1265–1274.
- 102 D. Cremer and E. Kraka, Dalton Trans., 2017, 46, 8323-8338.
- 103 E. Kraka and D. Cremer, *Int. J. Quantum Chem.*, 2019, **119**, e25849.
- 104 Y. Tao, W. Zou, S. Nanayakkara and E. Kraka, J. Chem. Theory Comput., 2022, 18, 1821–1837.
- 105 D. Cremer, A. Wu, A. Larsson and E. Kraka, J. Mol. Model., 2000, 6, 396–412.
- 106 Y. Tao, W. Zou and E. Kraka, *Chem. Phys. Lett.*, 2017, **685**, 251–258.
- 107 V. Oliveira, E. Kraka and D. Cremer, *Inorg. Chem.*, 2017, **56**, 488–502.
- 108 W. Zou, X. Zhang, H. Dai, H. Yan, D. Cremer and E. Kraka, J. Organomet. Chem., 2018, 865, 114–127.
- 109 R. Kalescky, E. Kraka and D. Cremer, J. Phys. Chem. A, 2013, 117, 8981–8995.
- 110 V. Oliveira, D. Cremer and E. Kraka, *J. Phys. Chem. A*, 2017, 121, 6845–6862.
- 111 D. Setiawan, E. Kraka and D. Cremer, *J. Phys. Chem. A*, 2015, **119**, 1642–1656.
- 112 D. Sethio, V. Oliveira and E. Kraka, *Molecules*, 2018, 23, 2763.
- 113 L. Chen, Q. Feng, C. Wang, S. Yin and Y. Mo, *J. Phys. Chem. A*, 2021, **125**, 10428–10438.
- 114 S. Shaik, D. Danovich, W. Wu and P. C. Hiberty, *Nat. Chem.*, 2009, **1**, 443–449.

- 115 F. Jimenez-Gravalos and D. Suarez, J. Chem. Theory Comput., 2021, 17, 4981-4995.
- 116 T. Xu, W. Wang and S. Yin, J. Chem. Theory Comput., 2018, 14, 3728-3739.
- 117 S. Grimme, C. Muck-Lichtenfeld, G. Erker, G. Kehr, H. Wang, H. Beckers and H. Willner, Angew. Chem., Int. Ed., 2009, 48, 2592-2595.
- 118 Y. Mo, J. Phys. Chem. A, 2012, 116, 5240-5246.
- 119 E. Cerpa, A. Krapp, A. Vela and G. Merino, Chem. Eur. J., 2008, 14, 10232-10234.
- 120 L. J. Farrugia, C. Evans and M. Tegel, J. Phys. Chem. A, 2006, 110, 7952-7961.
- 121 P. Cassam-Chenaï and D. Jayatilaka, Theor. Chem. Acc., 2001, 105, 213-218.
- 122 F. M. Bickelhaupt and E. J. Baerends, Angew. Chem., 2003, 115, 4315-4320.
- 123 P. R. Schreiner, Angew. Chem., Int. Ed., 2002, 41, 3579-3582.
- 124 Y. Mo, W. Wu, L. Song, M. Lin, Q. Zhang and J. Gao, Angew. Chem., Int. Ed., 2004, 43, 1986-1990.
- 125 A. J. Stone, J. Phys. Chem. A, 2017, 121, 1531-1534.
- 126 P. R. Horn, Y. Mao and M. Head-Gordon, Phys. Chem. Chem. Phys., 2016, 18, 23067-23079.
- 127 I. Alkorta, I. Mata, E. Molins and E. Espinosa, Chem. Eur. J., 2016, 22, 9226-9234.
- 128 A. Mele, G. Romano, M. Giannone, E. Ragg, G. Fronza, G. Raos and V. Marcon, Angew. Chem., Int. Ed., 2006, 45, 1123-1126.
- 129 M. Holz and K. J. Patil, Ber. Bunsenges. Phys. Chem., 1991, 95, 107-113.
- 130 S. Grimme and J. P. Djukic, *Inorg. Chem.*, 2011, **50**, 2619–2628.
- 131 O. Shih, A. H. England, G. C. Dallinger, J. W. Smith, K. C. Duffey, R. C. Cohen, D. Prendergast and R. J. Saykally, J. Chem. Phys., 2013, 139, 035104.
- 132 K. Morokuma, Acc. Chem. Res., 1977, 10, 294-300.
- 133 R. Z. Khaliullin, A. T. Bell and M. Head-Gordon, Chem. -Eur. J., 2009, 15, 851-855.
- 134 B. Jeziorski, R. Moszynski and K. Szalewicz, Chem. Rev., 1994, 94, 1887-1930.
- 135 K. Kitaura and K. Morokuma, Int. J. Quantum Chem., 1976, **10**, 325-340.
- 136 R. Z. Khaliullin, A. T. Bell and M. Head-Gordon, J. Chem. Phys., 2008, 128, 184112.
- 137 P. Su and H. Li, J. Chem. Phys., 2009, 131, 014102.
- 138 P. Su, H. Liu and W. Wu, J. Chem. Phys., 2012, 137, 034111.
- 139 W. Chen and M. S. Gordon, J. Phys. Chem., 1996, 100, 14316-14328.
- 140 R. Z. Khaliullin, E. A. Cobar, R. C. Lochan, A. T. Bell and M. Head-Gordon, J. Phys. Chem. A, 2007, 111, 8753-8765.
- 141 P. Su, Z. Jiang, Z. Chen and W. Wu, J. Phys. Chem. A, 2014, **118**, 2531-2542.
- 142 P. Su, Z. Tang and W. Wu, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2020, 10, e1460.
- 143 D. M. Andrada and C. Foroutan-Nejad, Phys. Chem. Chem. Phys., 2020, 22, 22459-22464.
- 144 J. Poater, D. M. Andrada, M. Sola and C. Foroutan-Nejad, Phys. Chem. Chem. Phys., 2022, 24, 2344-2348.

- 145 C. D. Sherrill, Acc. Chem. Res., 2013, 46, 1020-1028.
- 146 M. O. Sinnokrot and C. D. Sherrill, J. Am. Chem. Soc., 2004, 126, 7690-7697.
- 147 Y. Geng, T. Takatani, E. G. Hohenstein and C. D. Sherrill, J. Phys. Chem. A, 2010, 114, 3576-3582.
- 148 S. A. Bojarowski, P. Kumar, C. M. Wandtke, B. Dittrich and P. M. Dominiak, J. Chem. Theory Comput., 2018, 14, 6336-6345
- 149 C. Liu, J. P. Piquemal and P. Ren, J. Chem. Theory Comput., 2019, 15, 4122-4139.
- 150 Q. Wang, J. A. Rackers, C. He, R. Qi, C. Narth, L. Lagardere, N. Gresh, J. W. Ponder, J. P. Piquemal and P. Ren, J. Chem. Theory Comput., 2015, 11, 2609-2618.
- 151 A. J. Stone, J. Phys. Chem. A, 2011, 115, 7017-7027.
- 152 W. Wu, P. Su, S. Shaik and P. C. Hiberty, Chem. Rev., 2011, 111, 7557-7593.
- 153 Z. Chen and W. Wu, J. Chem. Phys., 2020, 153, 090902.
- 154 Y. Mo and S. D. Peyerimhoff, J. Chem. Phys., 1998, 109, 1687-1697.
- 155 Y. Mo, J. Chem. Phys., 2003, 119, 1300-1306.
- 156 Y. Mo, L. Song and Y. Lin, J. Phys. Chem. A, 2007, 111, 8291-8301.
- 157 Y. Mo, J. Gao and S. D. Peyerimhoff, J. Chem. Phys., 2000, 112, 5530-5538.
- 158 Y. Mo, P. Baob and J. Gao, Phys. Chem. Chem. Phys., 2011, 13, 6760-6775.
- 159 M. De Santis, S. Rampino, L. Storchi, L. Belpassi and F. Tarantelli, Inorg. Chem., 2019, 58, 11716-11729.
- 160 M. A. Rawashdeh-Omary, M. A. Omary and H. H. Patterson, J. Am. Chem. Soc., 2000, 122, 10371-10380.
- 161 R. Zhang, Y. Yu, T. C. Steimle and L. Cheng, J. Chem. Phys., 2017, 146, 064307.
- 162 T. Ichino, A. J. Gianola, D. H. Andrews and W. C. Lineberger, J. Phys. Chem. A, 2004, 108, 11307-11313.
- 163 A. V. Marenich, C. J. Cramer and D. G. Truhlar, J. Phys. Chem. B, 2009, 113, 6378-6396.
- 164 Y. Tao, W. Zou, D. Sethio, N. Verma, Y. Qiu, C. Tian, D. Cremer and E. Kraka, J. Chem. Theory Comput., 2019, 15, 1761-1776.
- 165 E. Kraka, W. Zou and Y. Tao, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2020, 10, e1480.
- 166 W. Zou, Y. Tao, M. Freindorf, D. Cremer and E. Kraka, Chem. Phys. Lett., 2020, 748, 137337.
- 167 Y. Minenkov, E. Chermak and L. Cavallo, J. Chem. Theory Comput., 2015, 11, 4664-4676.
- 168 C. Geng, S. Ye and F. Neese, Angew. Chem., Int. Ed., 2010, 49, 5717-5720.
- 169 F. Weigend and R. Ahlrichs, Phys. Chem. Chem. Phys., 2005, 7, 3297-3305.
- 170 F. Weigend, Phys. Chem. Chem. Phys., 2006, 8, 1057-1065.
- 171 S. Grimme, J. Comput. Chem., 2006, 27, 1787-1799.
- 172 C. Adamo and V. Barone, *J. Chem. Phys.*, 1999, **110**, 6158–6170.
- 173 Y. Zhao and D. G. Truhlar, Theor. Chem. Acc., 2007, 120, 215-241.
- 174 P. J. Stephens, F. J. Devlin, C. F. Chabalowski and M. J. Frisch, J. Phys. Chem., 1994, 98, 11623-11627.

- 175 T. Yanai, D. P. Tew and N. C. Handy, Chem. Phys. Lett., 2004, 393, 51-57.
- 176 J. D. Chai and M. Head-Gordon, Phys. Chem. Chem. Phys., 2008, 10, 6615-6620.
- 177 S. Grimme, J. Chem. Phys., 2003, 118, 9095-9102.
- 178 T. Shen, Z. Zhu, I. Y. Zhang and M. Scheffler, J. Chem. Theory Comput., 2019, 15, 4721-4734.
- 179 M. W. Schmidt, K. K. Baldridge, J. A. Boatz, S. T. Elbert, M. S. Gordon, J. H. Jensen, S. Koseki, N. Matsunaga, K. A. Nguyen, S. Su, T. L. Windus, M. Dupuis and J. A. Montgomery Jr, J. Comput. Chem., 1993, 11, 1347-1363.
- 180 F. Neese, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2011, 2, 73-78.
- 181 F. Neese, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2022, 12, e1606.

- 182 T. Lu and F. Chen, J. Comput. Chem., 2012, 33, 580-592.
- 183 A. J. Stone, J. Chem. Theory Comput., 2005, 1, 1128-1132.
- 184 P. Ren and J. W. Ponder, J. Phys. Chem. B, 2003, 107, 5933-5947.
- 185 J. W. Ponder, C. Wu, P. Ren, V. S. Pande, J. D. Chodera, M. J. Schnieders, I. Haque, D. L. Mobley, D. S. Lambrecht, J. Robert, A. DiStasio, M. Head-Gordon, G. N. I. Clark, M. E. Johnson and T. Head-Gordon, J. Phys. Chem. B, 2010, 114, 2549-2564.
- 186 P. Ren, C. Wu and J. W. Ponder, J. Chem. Theory Comput., 2011, 7, 3143-3161.
- 187 Y. Shi, Z. Xia, J. Zhang, R. Best, C. Wu, J. W. Ponder and P. Ren, J. Chem. Theory Comput., 2013, 9, 4046-4063.
- 188 J. A. Rackers, Z. Wang, C. Lu, M. L. Laury, L. Lagardere, M. J. Schnieders, J. P. Piquemal, P. Ren and J. W. Ponder, J. Chem. Theory Comput., 2018, 14, 5273-5289.