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The nature of metallophilic interactions in closed-shell d^8-d^8 metal complexes†

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We have quantum chemically analyzed the closed-shell d^8-d^8 metallophilic interaction in dimers of square planar $[M(CO)_2X_2]$ complexes ($M = Ni, Pd, Pt$; $X = Cl, Br, I$) using dispersion-corrected density functional theory at ZORA-BLYP-D3(BJ)/TZ2P level of theory. Our purpose is to reveal the nature of the $[X_2(CO)_2M] \cdots [M(CO)_2X_2]$ bonding mechanism by analyzing trends upon variations in M and X . Our analyses reveal that the formation of the $[M(CO)_2X_2]_2$ dimers is favored by an increasingly stabilizing electrostatic interaction when the M increases in size and by more stabilizing dispersion interactions promoted by the larger X . In addition, there is an overlooked covalent component stemming from metal-metal and ligand-ligand donor-acceptor interactions. Thus, at variance with the currently accepted picture, the d^8-d^8 metallophilicity is attractive, and the formation of $[M(CO)_2X_2]_2$ dimers is not a purely dispersion-driven phenomenon.

1. Introduction

Metallophilicity is the manifestation of a net attractive intra- or intermolecular interaction between metal centers (M), yielding metal-metal-bonded complexes.¹ This metal-metal interaction causes the spontaneous association, *i.e.*, self-assembly, of metal complexes forming aggregates with luminescence properties² and medicinal applications.³ Earlier work by Hoffman *et al.* showed that the closed-shell metallophilic interactions between linear d^{10} metal complexes are covalent due to the donor-acceptor interactions between the filled nd_{z^2} -type HOMO on one M and the $(n+1)s$ -type LUMO on the other M .⁴ However, Brands *et al.* showed that the donor-acceptor interactions cannot overcome the destabilizing steric Pauli repulsion between the two metal centers, and the metallophilic interactions are only attractive due to the additional stabilizing electrostatic interactions.⁵ In general, the closed-shell $d^{10}-d^{10}$ metallophilic interactions have a similar mechanistic picture to other well-known intermolecular interactions, such as hydrogen bonds.⁶

Nonetheless, alternative models have been used to explain metallophilicity. For example, Pykkö and coworkers concluded that the $d^{10}-d^{10} Au^+ \cdots Au^+$ attraction in staggered dimers of linear $AuPH_3Cl$ complexes is a dispersion-driven phenomenon, whereas the ligand-ligand repulsion hampers the formation of parallel dimers.⁷ This has been further supported by Chen *et al.* who found that $\pi-\pi$ interactions between face-to-face, eclipsed ligands are repulsive.⁸ On the other hand, Che *et al.* attributed the formation of d^8-d^8 and $d^{10}-d^{10}$ metal-complexes dimers to stabilizing ligand-ligand dispersion interactions.⁹

In the present study, we have quantum chemically investigated the bonding mechanism of eclipsed closed-shell d^8-d^8 metal complexes, using $[M(CO)_2X_2]_2$ dimers ($M = Ni, Pd, Pt$; $X = Cl, Br, I$) as model systems (Scheme 1). As will become clear in the following sections, our bonding analyses reveal that the stability of the $[M(CO)_2X_2]_2$ dimers stems from significantly attractive electrostatic interactions, on top of dispersion interactions promoted by the ligands. This closed-shell d^8-d^8 metal-metal interaction is, therefore, similar to that between d^{10}

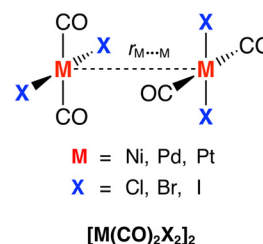
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Scheme 1 The studied $[M(CO)_2X_2]_2$ dimers.



metal centers, that is, it is also partially covalent in nature and, thus, not a pure dispersion-driven phenomenon.

2. Computational methods

2.1. Computational details

All calculations were carried out using the Amsterdam Modeling Suite (AMS) 2020.101 program.¹⁰ All stationary points and energies were obtained using relativistic, dispersion-corrected density functional theory at ZORA-BLYP-D3(BJ)/TZ2P. This approach has been benchmarked in previous work for bond energies of metal-metal complexes using CCSD(T)/CBS with mean absolute error of 0.7 kcal mol⁻¹.⁵ This approach comprises the BLYP level of the generalized gradient approximation (GGA); exchange functional developed by Becke (B), and the GGA correlation functional developed by Lee, Yang, and Parr (LYP)¹¹ (see Tables S4 and S5 in the ESI† for the Cartesian coordinates). The DFT-D3(BJ) method developed by Grimme and coworkers,¹² which contains the damping function proposed by Becke and Johnson,¹³ is used to describe non-local dispersion interactions. Scalar relativistic effects are accounted for using the zeroth-order regular approximation (ZORA).¹⁴ Molecular orbitals (MO) were expanded in a large, uncontracted set of Slater-type orbitals (STOs) containing diffuse functions: TZ2P.¹⁵ The basis set is of triple- ζ quality augmented with polarization functions, *i.e.*, one 3d and one 4f set on C, O, Cl; one 4d and one 4f set on Br; one 5d and one 4f set on I; one 4f set on Ni; one 5p and one 4f set on Pd; one 6p and one 5f set on Pt. All electrons were included in the variational process, *i.e.*, no frozen core approximation was applied. The accuracies of both the fitting scheme and the integration grid (Becke grid) were set to 'EXCELLENT'. The formation of *trans*-[M(CO)₂X₂] monomers and *trans*-[M(CO)₂X₂]₂ dimers (M = Ni, Pd, Pt; X = Cl, Br, I) is, in most cases, more favorable than the formation of *cis*-[M(CO)₂X₂] monomers and *cis*-[M(CO)₂X₂]₂ dimers (see Table S3 and Fig. S4 for bond energies and geometries, ESI†). The [Ni(CO)₂Br₂], [Ni(CO)₂I₂], and [Pd(CO)₂I₂] monomers, and their respective dimers, are the only exceptions in which the *cis* forms are more favorable (Table S3, ESI†). Thus, in this study, we always refer to the *trans*-[M(CO)₂X₂]₂ dimers. We have found here that the global-minimum structures of the [M(CO)₂X₂]₂ dimers are in the staggered conformation by performing a fully relaxed rotation around the M...M bond, in which the X-M...M-X dihedral angle is varied from 0° to 90° (see Fig. S5 and S6 in the ESI†). All optimized structures were confirmed to be true minima through vibrational analysis (no imaginary frequencies).

2.2. Bond analyses

Insight into the bonding mechanism is obtained by analyzing the potential energy curves for the formation of the [M(CO)₂X₂]₂ dimers (M = Ni, Pd, Pt; X = Cl, Br, I). The analyses are done by varying the M...M bond distance within the range of 2.5 Å to 4.5 Å, starting from the equilibrium geometry of the complex ($r_{M...M,eq}$) while keeping all other geometrical parameters frozen.

The interaction between [X₂(CO)₂M] and [M(CO)₂X₂] is analyzed using the activation strain model,¹⁶ which is a

fragment-based approach to understanding the energy profile associated with a chemical process in terms of the original reactants. Thus, the total bond energy ΔE is decomposed along the M...M bond distance $r_{M...M}$ (or just at one point along $r_{M...M}$) into the strain energy $\Delta E_{strain}(r_{M...M})$, which is associated with the geometrical deformation of the individual reactants as the process takes place, plus the actual interaction energy $\Delta E_{int}(r_{M...M})$ between the deformed reactants [eqn (1)].

$$\Delta E(r_{M...M}) = \Delta E_{strain}(r_{M...M}) + \Delta E_{int}(r_{M...M}) \quad (1)$$

In the equilibrium geometry, that is, for $r_{M...M} = r_{M...M,eq}$, this yields an expression for the bond energy $\Delta E(r_{M...M,eq}) = \Delta E = \Delta E_{strain} + \Delta E_{int}$. The interaction energy $\Delta E_{int}(r_{M...M})$ between the deformed reactants is further analyzed in the conceptual framework provided by the quantitative Kohn-Sham MO model.¹⁷ To this end, it is decomposed into physically meaningful terms [eqn (2)] using a quantitative energy decomposition analysis (EDA) as implemented in ADF.¹⁷

$$\begin{aligned} \Delta E_{int}(r_{M...M}) = & \Delta V_{elstat}(r_{M...M}) + \Delta E_{Pauli}(r_{M...M}) \\ & + \Delta E_{oi}(r_{M...M}) + \Delta E_{disp}(r_{M...M}) \end{aligned} \quad (2)$$

The usually attractive term ΔV_{elstat} corresponds to the classical Coulomb interaction between the unperturbed charge distributions of the deformed reactants and has four components: (i) the electron-electron repulsion between the electron densities of [X₂(CO)₂M] and [M(CO)₂X₂]; (ii) the nuclei-electron attraction between the nuclei of [X₂(CO)₂M] and the electron density of [M(CO)₂X₂]; (iii) the electron-nuclei attraction between the electron density of [X₂(CO)₂M] and the nuclei of [M(CO)₂X₂]; and (iv) the nuclei-nuclei repulsion between the nuclei of [X₂(CO)₂M] and [M(CO)₂X₂].¹⁷

The Pauli repulsion energy (ΔE_{Pauli}) comprises the destabilizing interactions between the fully occupied orbitals on either fragment and arises from the antisymmetrization of the Hartree wavefunction due to the Pauli principle. The orbital-interaction energy (ΔE_{oi}) accounts for charge transfer, that is, the interaction between occupied orbitals of one fragment with unoccupied orbitals of the other fragment, including the interactions of the highest occupied and lowest unoccupied MOs (HOMO-LUMO), and polarization, that is, empty-occupied orbital mixing on one fragment, due to the presence of another fragment. The dispersion energy ΔE_{disp} accounts for the dispersion corrections as introduced by Grimme *et al.*¹² To facilitate the analyses, the ASM and EDA were performed using the PyFrag 2019 program.¹⁸

3. Results and discussion

3.1. Bond strength and structure

Equilibrium geometries, dimerization energies (ΔE), and the M...M bond distances ($r_{M...M}$) of the stable C_s symmetrical [M(CO)₂X₂]₂ dimers (M = Ni, Pd, Pt; X = Cl, Br, I) calculated at the ZORA-BLYP-D3(BJ)/TZ2P level of theory are shown in Fig. 1. Upon the formation of the studied dimers, we have identified two main features: (i) the stability of the dimers significantly increases, and the M...M bond distance slightly shortens for



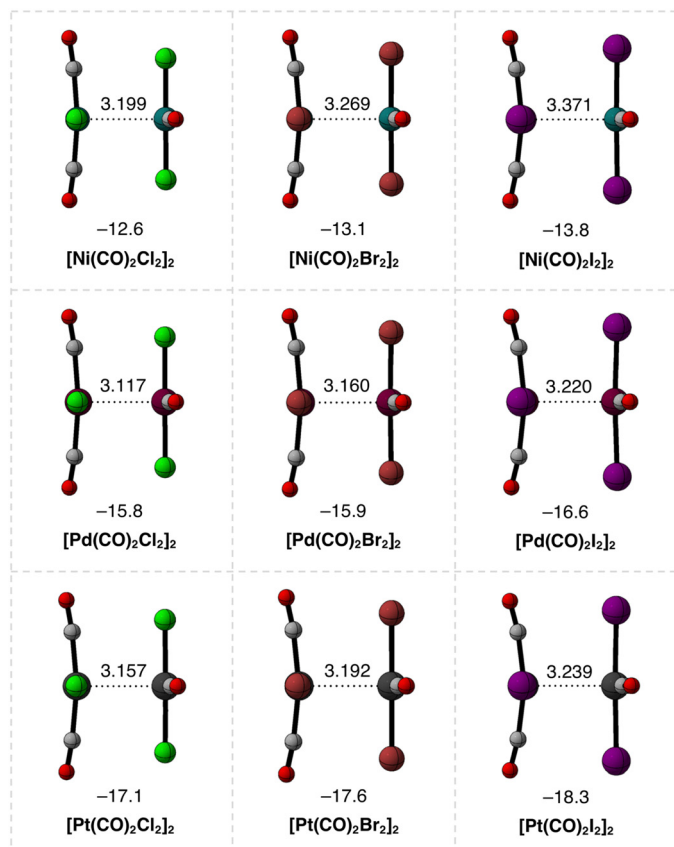


Fig. 1 Equilibrium geometries (in Å) and electronic bond energies (in kcal mol⁻¹) of the [M(CO)₂X₂]₂ dimers (M = Ni, Pd, Pt; X = Cl, Br, I). Computed at ZORA-BLYP-D3(BJ)/TZ2P.

heavier M; and (ii) the dimers are only slightly stronger but longer for heavier X ligands. For example, along the series from [Ni(CO)₂Br₂]₂ to [Pt(CO)₂Br₂]₂, ΔE becomes more stabilizing from -13.1 kcal mol⁻¹ to -17.6 kcal mol⁻¹ and $r_{M...M}$ decreases from 3.269 Å to 3.192 Å (Fig. 1). Along the series from [Pd(CO)₂Cl₂]₂ to [Pd(CO)₂I₂]₂, ΔE varies only from -15.8 kcal mol⁻¹ to -16.6 kcal mol⁻¹ and $r_{M...M}$ increases from 3.117 Å to 3.220 Å (Fig. 1).

Next, we performed our activation strain analysis¹⁶ (ASA) to decompose the bond energies (ΔE) into the strain (ΔE_{strain}) and

interaction (ΔE_{int}) energies (see Computational methods), and the results are shown in Table 1. The trends in the stability of the [M(CO)₂X₂]₂ dimers along M = Ni, Pd, Pt and X = Cl, Br, I are dominated by the stronger ΔE_{int} , whereas the weaker ΔE_{strain} varies only slightly along the same series. For example, along the series from [Ni(CO)₂Br₂]₂ to [Pt(CO)₂Br₂]₂, ΔE_{int} becomes more stabilizing from -14.0 kcal mol⁻¹ to -18.9 kcal mol⁻¹ and ΔE_{strain} becomes slightly more destabilizing from 0.9 kcal mol⁻¹ to 1.3 kcal mol⁻¹ (Table 1). Along the series from [Pd(CO)₂Cl₂]₂ to [Pd(CO)₂I₂]₂, ΔE_{int} becomes more stabilizing from -16.6 kcal mol⁻¹ to -17.9 kcal mol⁻¹ and ΔE_{strain} becomes slightly more destabilizing from 0.8 kcal mol⁻¹ to 1.3 kcal mol⁻¹ (Table 1). Note that the impact on ΔE_{int} and, thus, on the stability of the [M(CO)₂X₂]₂ dimers when varying M is larger than when varying X. As will become clear in the next sections, this is because of the difference in nature between the metal-metal interactions and those involving the ligands. This conclusion emerges from understanding the physical nature behind the observed trends in ΔE_{int} when the metal center M varies along Ni, Pd, and Pt and the ligands X vary along Cl, Br, and I.

3.2. Variation of M

To understand the origin of the increased stabilization of the [M(CO)₂X₂]₂ dimers upon varying M along Ni, Pd, and Pt, we further decomposed the ΔE_{int} into physically meaningful

Table 1 Activation strain and energy decomposition analyses (in kcal mol⁻¹) for the formation of the [M(CO)₂X₂]₂ dimers (M = Ni, Pd, Pt; X = Cl, Br, I) in their equilibrium geometries (in Å). Computed at ZORA-BLYP-D3(BJ)/TZ2P

M	X	$r_{M...M}$	ΔE	ΔE_{strain}	ΔE_{int}	ΔV_{elstat}	ΔE_{Pauli}	ΔE_{oi}	ΔE_{disp}
Ni	Cl	3.199	-12.6	0.6	-13.2	-12.3	20.3	-7.6	-13.6
	Br	3.269	-13.1	0.9	-14.0	-14.8	21.3	-8.0	-15.4
	I	3.371	-13.8	0.9	-14.7	-11.6	22.7	-7.7	-18.1
Pd	Cl	3.117	-15.8	0.8	-16.6	-24.1	34.8	-11.4	-15.8
	Br	3.160	-15.9	1.3	-17.2	-23.1	35.8	-12.0	-17.9
	I	3.220	-16.6	1.3	-17.9	-22.3	37.1	-12.0	-20.8
Pt	Cl	3.157	-17.1	0.9	-18.0	-28.0	39.5	-12.9	-16.6
	Br	3.192	-17.6	1.3	-18.9	-27.3	41.0	-13.8	-18.9
	I	3.239	-18.3	1.8	-20.1	-26.8	43.2	-14.2	-22.1



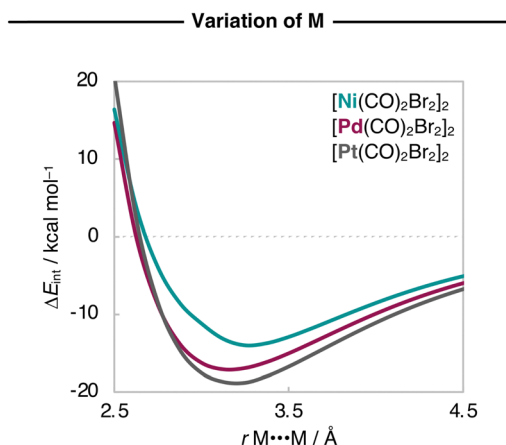


Fig. 2 Interaction energies (in kcal mol^{−1}) as a function of the M...M distance (in Å) starting from the equilibrium geometries of the representative [M(CO)₂Br₂]₂ dimers (M = Ni, Pd, Pt), while the geometry of the monomers is kept frozen. Computed at ZORA-BLYP-D3(BJ)/TZ2P.

terms, namely the electrostatic interactions (ΔV_{elstat}), steric Pauli repulsion (ΔE_{Pauli}), orbital interactions (ΔE_{oi}), and dispersion energy (ΔE_{disp}) using our energy decomposition analysis¹⁷ (EDA; see Computational methods), and the results are shown in Table 1. Our analyses reveal that, for the smaller Ni, we find that ΔE_{disp} is the largest attractive term in $[\text{X}_2(\text{CO})_2\text{M}] \cdots [\text{M}(\text{CO})_2\text{X}_2]$. However, as M increases in size, ΔV_{elstat} and ΔE_{oi} become significantly more stabilizing whereas ΔE_{disp} varies much less, which causes ΔV_{elstat} to dominate and become the most attractive term for Pd and Pt. For example, from $[\text{Ni}(\text{CO})_2\text{Br}_2]_2$ to $[\text{Pt}(\text{CO})_2\text{Br}_2]_2$, ΔE_{disp} varies only from -15.4 kcal mol^{−1} to -18.9 kcal mol^{−1}, whereas ΔV_{elstat} and ΔE_{oi} become significantly more stabilizing from -14.8 kcal mol^{−1} to -27.3 kcal mol^{−1} and from -8.0 kcal mol^{−1} to -13.8 kcal mol^{−1}, respectively (Table 1).

For a consistent comparison, we extend our analysis to the entire reaction coordinate, as a function of the M...M bond distance ($r_{\text{M}\cdots\text{M}}$). Since ΔE_{strain} is small and the ΔE is dominated by ΔE_{int} , the analyses were done while the geometries of the fragments were kept frozen to that of the equilibrium geometries of the $[\text{M}(\text{CO})_2\text{X}_2]_2$ dimers. The resulting interaction

energy curves $\Delta E_{\text{int}}(r_{\text{M}\cdots\text{M}})$ for the representative $[\text{Ni}(\text{CO})_2\text{Br}_2]_2$, $[\text{Pd}(\text{CO})_2\text{Br}_2]_2$, and $[\text{Pt}(\text{CO})_2\text{Br}_2]_2$ series are graphically shown in Fig. 2. Herein, the $\Delta E_{\text{int}}(r_{\text{M}\cdots\text{M}})$ curves also become more stabilizing and the energy minimum is shifted towards shorter $r_{\text{M}\cdots\text{M}}$ as M varies along Ni, Pd, and Pt.

Next, we analyze the EDA terms as a function of $r_{\text{M}\cdots\text{M}}$ and the resulting diagrams for the representative $[\text{Ni}(\text{CO})_2\text{Br}_2]_2$, $[\text{Pd}(\text{CO})_2\text{Br}_2]_2$, and $[\text{Pt}(\text{CO})_2\text{Br}_2]_2$ series are graphically shown in Fig. 3. Our findings show that the increased stability of the $[\text{M}(\text{CO})_2\text{Br}_2]_2$ dimers as M varies along Ni, Pd, Pt is due to a greater electrostatic attraction between the monomers, as the electrostatic interaction curves $\Delta V_{\text{elstat}}(r_{\text{M}\cdots\text{M}})$ become significantly more stabilizing along the same series (Fig. 3). This is because the $[\text{X}_2(\text{CO})_2\text{M}] \cdots [\text{M}(\text{CO})_2\text{X}_2]$ electrostatic attraction is largely affected by the charge distributions around the metal centers within the monomers. In essence, there is a more effective electron–nuclei overlap as the electron density (ρ) of $[\text{M}(\text{CO})_2\text{Br}_2]_2$ becomes more diffuse, and the nucleus of the metal center increases in size (*i.e.*, has more protons) along the same series. For example, at the same point of the reaction coordinate, *e.g.*, $r_{\text{M}\cdots\text{M}} = 3.5$ Å, the negatively charged ρ of one fragment is more diffuse around the metal center and extends further towards the nucleus of the metal center of the other fragment, which becomes more positively charged as M varies along Ni, Pd, and Pt (see Fig. 4). As the two monomers approach each other, this attractive electron–nuclei overlap more quickly increases for heavier M and, consequently, the slope of the descending $\Delta V_{\text{elstat}}(r_{\text{M}\cdots\text{M}})$ curves increases along Ni, Pd, and Pt, shifting the equilibrium geometries of the $[\text{M}(\text{CO})_2\text{Br}_2]_2$ dimers to a shorter $r_{\text{M}\cdots\text{M}}$ as M varies along the same series.

Together with $\Delta V_{\text{elstat}}(r_{\text{M}\cdots\text{M}})$, the orbital interactions $\Delta E_{\text{oi}}(r_{\text{M}\cdots\text{M}})$ and the dispersion energy curves $\Delta E_{\text{disp}}(r_{\text{M}\cdots\text{M}})$ follow the same trend as $\Delta E_{\text{int}}(r_{\text{M}\cdots\text{M}})$ and become more stabilizing along M = Ni, Pd, Pt (Fig. 3). This effect is more pronounced in $\Delta E_{\text{oi}}(r_{\text{M}\cdots\text{M}})$ which arises, in part, from the donation of charge from the metal d_{z²}-type HOMO of one $\text{M}(\text{CO})_2\text{X}_2$ monomer into the metal s-type LUMO of the other monomer and will be explained in detail later (see Scheme 2). The $\Delta E_{\text{oi}}(r_{\text{M}\cdots\text{M}})$ curves are weakly attractive at longer $r_{\text{M}\cdots\text{M}}$ and, as soon as the d_{z²}-HOMO and s-LUMO start to overlap at

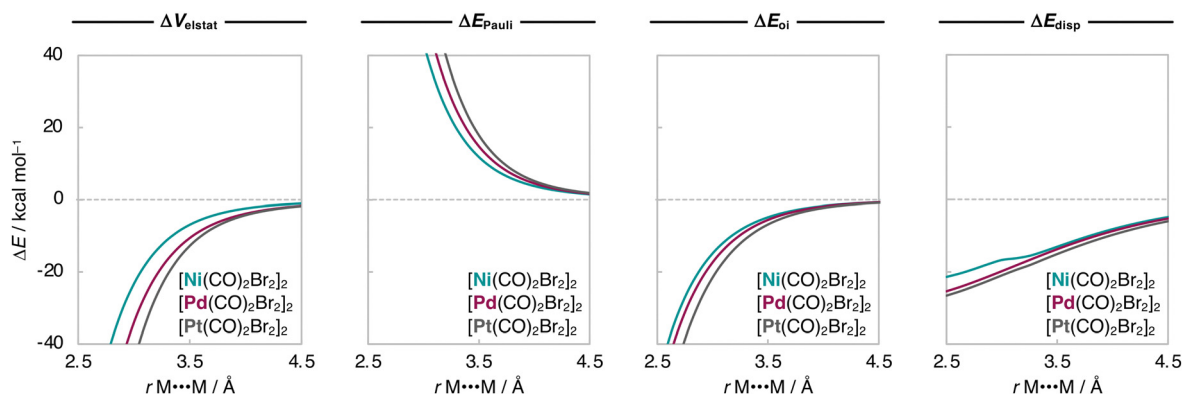


Fig. 3 Energy decomposition analysis (in kcal mol^{−1}) as a function of the M...M distance (in Å) for the representative $[\text{M}(\text{CO})_2\text{Br}_2]_2$ dimers (M = Ni, Pd, Pt), while the geometry of the monomers is kept frozen. Computed at ZORA-BLYP-D3(BJ)/TZ2P.



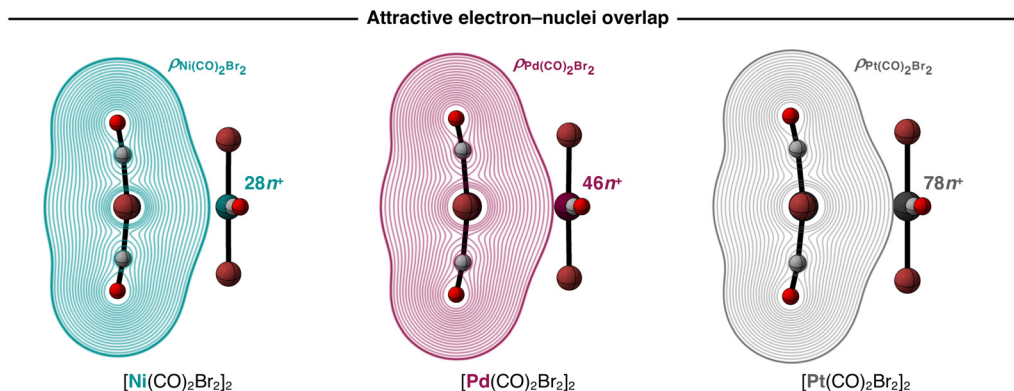


Fig. 4 Attractive overlap between the negatively charged electron densities and the positively charged nuclei (density contours from 0.0001 to 0.5000 a.u.) for the representative $[\text{M}(\text{CO})_2\text{Br}_2]_2$ dimers ($\text{M} = \text{Ni}, \text{Pd}, \text{Pt}$) at a consistent $\text{M} \cdots \text{M}$ bond distance ($r_{\text{M} \cdots \text{M}} = 3.5$ Å). Computed at ZORA-BLYP-D3(BJ)/TZ2P.

shorter $r_{\text{M} \cdots \text{M}}$, they become significantly stabilizing (Fig. 3). The $\Delta E_{\text{disp}}(r_{\text{M} \cdots \text{M}})$ curves, on the other hand, are relatively strong already at longer $r_{\text{M} \cdots \text{M}}$, but neither become much more stabilizing at shorter $r_{\text{M} \cdots \text{M}}$ nor significantly vary along $\text{M} = \text{Ni}, \text{Pd}, \text{Pt}$. In other words, the ΔE_{disp} term significantly contributes to the stability of the $[\text{M}(\text{CO})_2\text{X}_2]_2$ dimers but is almost insensitive to variations of the metal centers.

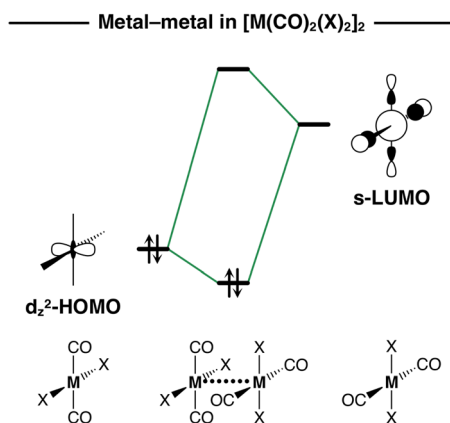
3.3. Variation of X

To understand why the ΔE_{int} , and thus the $[\text{M}(\text{CO})_2\text{X}_2]_2$ dimers, becomes slightly more stable upon varying X along Cl, Br, and I, we first analyze the EDA terms at the equilibrium geometries. Contrary to the variation along $\text{M} = \text{Ni}, \text{Pd}, \text{Pt}$, our analyses reveal that ΔE_{disp} becomes more stabilizing, whereas ΔV_{elstat} and ΔE_{oi} do not significantly change when X varies along Cl, Br, and I. For example, from $[\text{Pd}(\text{CO})_2\text{Cl}_2]_2$ to $[\text{Pd}(\text{CO})_2\text{I}_2]_2$, ΔE_{disp} becomes more stabilizing from -15.8 kcal mol $^{-1}$ to -20.8 kcal mol $^{-1}$, whereas ΔV_{elstat} and ΔE_{oi} vary from -24.1 kcal mol $^{-1}$ to -22.3 kcal mol $^{-1}$ and -11.4 kcal mol $^{-1}$ to -12.0 kcal mol $^{-1}$, respectively (Table 1).

Next, we analyze the ΔE_{int} and the EDA terms as a function of $r_{\text{M} \cdots \text{M}}$, and the resulting diagrams for the representative

$[\text{Pd}(\text{CO})_2\text{Cl}_2]_2$, $[\text{Pd}(\text{CO})_2\text{Br}_2]_2$, and $[\text{Pd}(\text{CO})_2\text{I}_2]_2$ series are graphically shown in Fig. 5 and 6. The $\Delta E_{\text{int}}(r_{\text{M} \cdots \text{M}})$ curves also become more stabilizing, and the energy minimum is shifted towards longer $r_{\text{M} \cdots \text{M}}$ as X varies along Cl, Br, and I. Our analyses show that the increased stability of the $[\text{Pd}(\text{CO})_2\text{X}_2]_2$ dimers as X varies along Cl, Br, and I is due to a greater attractive dispersion interaction promoted by the ligands as they increase in size along the same series. Note that the $\Delta E_{\text{disp}}(r_{\text{M} \cdots \text{M}})$ curves become significantly more stabilizing along $[\text{Pd}(\text{CO})_2\text{Cl}_2]_2$, $[\text{Pd}(\text{CO})_2\text{Br}_2]_2$, and $[\text{Pd}(\text{CO})_2\text{I}_2]_2$ (Fig. 6). Nevertheless, the equilibrium $r_{\text{M} \cdots \text{M}}$ slightly expands because the Pauli repulsion curves $\Delta E_{\text{Pauli}}(r_{\text{M} \cdots \text{M}})$ are steeper than the $\Delta E_{\text{disp}}(r_{\text{M} \cdots \text{M}})$ curves and push the equilibrium $r_{\text{M} \cdots \text{M}}$ to longer values as X varies along Cl, Br, and I (Fig. 6).

The $\Delta V_{\text{elstat}}(r_{\text{M} \cdots \text{M}})$ curves are almost insensitive to variations of X in the $[\text{Pd}(\text{CO})_2\text{X}_2]_2$ dimers and become only slightly more stabilizing along X = Cl, Br, I (Fig. 6). This is because, as aforementioned, the attractive electron-nuclei overlap occurs mainly at the $\text{M} \cdots \text{M}$ bond region and does not significantly change as the X ligands increase in size along X = Cl, Br, I.



Scheme 2 Schematic molecular orbital diagram of the metal-metal donor-acceptor interactions in $[\text{M}(\text{CO})_2\text{X}_2]_2$ dimers. The DFT orbitals are shown in Fig. S3 of the ESI.†

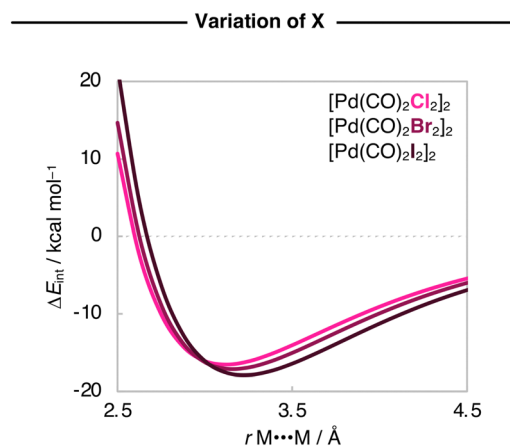


Fig. 5 Interaction energies (in kcal mol $^{-1}$) as a function of the $\text{M} \cdots \text{M}$ distance (in Å) starting from the equilibrium geometries of the representative $[\text{Pd}(\text{CO})_2\text{X}_2]_2$ dimers ($\text{X} = \text{Cl}, \text{Br}, \text{I}$), while the geometry of the monomers is kept frozen. Computed at ZORA-BLYP-D3(BJ)/TZ2P.



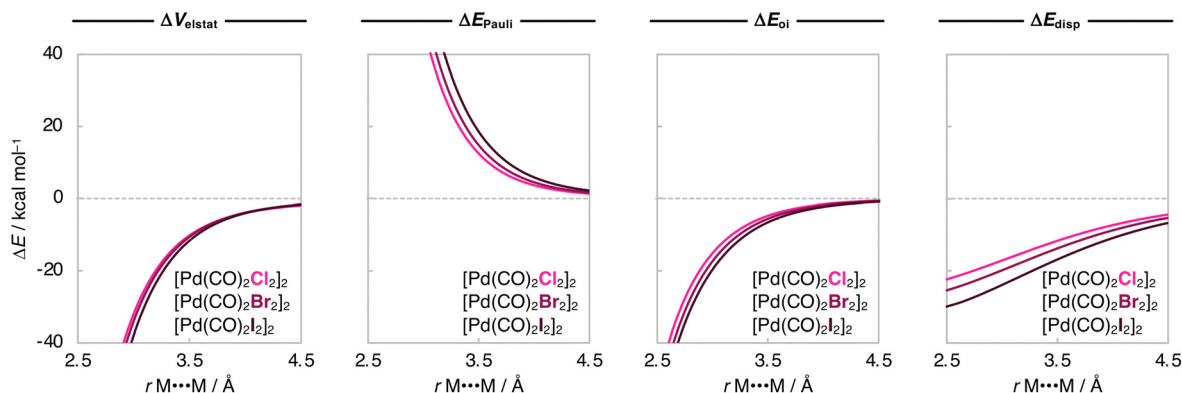


Fig. 6 Energy decomposition analysis (in kcal mol⁻¹) as a function of the M...M distance (in Å) for the representative [Pd(CO)₂X₂]₂ dimers (X = Cl, Br, I), while the geometry of the monomers is kept frozen. Computed at ZORA-BLYP-D3(BJ)/TZ2P.

Instead, the increased size of X causes the ρ of the monomers to extend further along the CO...X region, where there is not a pronounced electron-nuclei overlap (see Fig. S1, ESI[†]). Consequently, the electrostatic attraction between the monomers in the [M(CO)₂X₂]₂ only marginally increases when X varies along Cl, Br, and I.

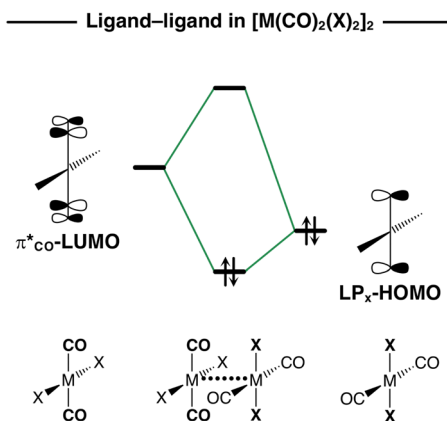
Our analyses along variations of M and X have revealed two important interactions for the bonding mechanism and, thus, stability of the [M(CO)₂X₂]₂ dimers. On the one hand, there are electrostatic interactions that increase in relevance as the metal becomes bigger. On the other hand, there are attractive dispersion interactions stemming from the ligands. Despite this difference in nature between the metal-metal and ligand-ligand interactions, they also share some similarities. Similar to the trend upon varying M along Ni, Pd, and Pt, the $\Delta E_{oi}(r_{M...M})$ curves also become more stabilizing when X varies along Cl, Br, and I in the [Pd(CO)₂X₂]₂ dimers (Fig. 6). This is due to the stabilizing ligand-ligand donor-acceptor interaction of the C=O π^* -type (π^*_{CO}) LUMO of one monomer with the X lone-pair-type (LP_X) HOMO of the other monomer (see Scheme 3). In the following section, we explain why the $\Delta E_{oi}(r_{M...M})$ curves become more stabilizing along M = Ni, Pd, Pt and X = Cl, Br, I in the [M(CO)₂X₂]₂ dimers.

3.4. Metal-metal versus ligand-ligand donor-acceptor interactions

In previous sections, we showed that the stability of the [M(CO)₂X₂]₂ dimers is mainly governed by the electrostatic attraction, which becomes more important for larger M and dispersive interactions promoted by the ligands. However, both metal-metal and ligand-ligand interactions have a covalent component stemming from donor-acceptor interactions that has been hitherto overlooked. Note that the ΔE_{oi} between the monomers in [M(CO)₂X₂]₂ dimers, which accounts for donor-acceptor interactions, becomes more stabilizing along M = Ni, Pd, Pt and X = Cl, Br, I (*vide supra*). Next, we explain the physical nature behind these trends.

The donor-acceptor interactions in the [M(CO)₂X₂]₂ dimers comprise of two main components, namely, the metal-metal d_{z^2} -HOMO-s-LUMO and the ligand-ligand π^*_{CO} -LUMO- LP_X -HOMO interactions (see Schemes 2 and 3). The relevance of a donor-acceptor interaction can be estimated by the magnitude of its orbital stabilization, which is proportional to its HOMO-LUMO overlap squared (S^2) divided by its respective orbital energy gap ($\Delta\epsilon$) [see eqn (3)]. For a consistent comparison, we report these values for the metal-metal and ligand-ligand donor-acceptor interactions at $r_{M...M} = 3.5$ for all [M(CO)₂X₂]₂ dimers in Table 2.

$$\Delta E_{oi} \propto S^2/\Delta\epsilon \quad (3)$$



Scheme 3 Schematic molecular orbital diagram of the ligand-ligand donor-acceptor interactions in [M(CO)₂X₂]₂ dimers. The DFT orbitals are shown in Fig. S3 of the ESI[†].

The $\Delta E_{oi}(r_{M...M})$ curves for the [M(CO)₂X₂]₂ dimers become more stabilizing along M = Ni, Pd, Pt because the metal-metal donor-acceptor interactions between the d_{z^2} -type HOMO of one monomer and the s-type LUMO of the other monomer are strengthened along the same series. When the metal center increases in size along Ni, Pd, and Pt, the bond overlap S between the d_{z^2} -HOMO and the s-LUMO increases as both orbitals become more diffuse, resulting in larger orbital stabilization and, thus, stronger donor-acceptor interactions along the same series. For example, in the series of [Ni(CO)₂Br]₂, [Pd(CO)₂Br]₂, and [Pt(CO)₂Br]₂ at $r_{M...M} = 3.5$, S increases from 0.09 to 0.17 to 0.18, as both the d_{z^2} -HOMO and the s-LUMO further extend towards the other metal center (see Fig. 7a, but also the overlap densities in Fig. S2a, ESI[†]). Consequently, $S^2/\Delta\epsilon$ times 10³ increases from 1.0 to 3.6 to 4.0 along the same series (Table 2).



Table 2 Orbital interaction energies (in kcal mol^{−1}), orbital overlap, energy gap ($\Delta\epsilon$, in eV), and orbital stabilization for the metal–metal and ligand–ligand donor–acceptor interactions in the $[M(\text{CO})_2\text{X}_2]_2$ dimers ($M = \text{Ni}, \text{Pd}, \text{Pt}$; $X = \text{Cl}, \text{Br}, \text{I}$) at a consistent $M \cdots M$ bond distance ($r_{M \cdots M} = 3.5$ Å). Computed at ZORA-BLYP-D3(BJ)/TZ2P

M	X	ΔE_{oi}	Metal–metal				Ligand–ligand			
			S	$\Delta\epsilon^a$	$10^3 \times S^2/\Delta\epsilon$	% ^b	S	$\Delta\epsilon^a$	$10^3 \times S^2/\Delta\epsilon$	% ^b
Ni	Cl	−3.8	0.11	8.2	1.5	53	0.07	3.8	1.3	47
	Br	−4.9	0.09	7.8	1.0	36	0.08	3.5	1.8	64
	I	−5.8	0.08	7.3	0.9	25	0.09	3.1	2.6	75
Pd	Cl	−4.7	0.18	8.4	3.9	67	0.08	3.4	1.9	33
	Br	−5.6	0.17	8.1	3.6	58	0.09	3.1	2.6	42
	I	−6.5	0.16	7.7	3.3	47	0.10	2.7	3.7	53
Pt	Cl	−5.8	0.20	8.2	4.9	67	0.09	3.3	2.5	33
	Br	−6.8	0.18	8.1	4.0	55	0.10	3.0	3.3	45
	I	−8.0	0.17	7.8	3.7	44	0.11	2.6	4.7	56

^a See HOMO and LUMO energies in Tables S1 and S2 (ESI). ^b Contribution of the associated orbital stabilization to the total metal–metal + ligand–ligand orbital stabilization.

The ΔE_{oi} term in the $[M(\text{CO})_2\text{X}_2]_2$ dimers is not fully dominated by the metal–metal donor–acceptor interactions and is, in part, made by ligand–ligand donor–acceptor interactions. This is confirmed by the trends in the $\Delta E_{\text{oi}}(r_{M \cdots M})$ curves that also become more stabilizing along $X = \text{Cl}, \text{Br}, \text{I}$ (Fig. 6). The bond overlap S for the $\pi^*_{\text{CO}}\text{-LUMO-LP}_X\text{-HOMO}$ interaction is, in general, significantly smaller than that of the metal–metal

donor–acceptor interactions (Table 2). However, as the X ligands become larger and less electronegative along Cl, Br , and I , the $\text{LP}_X\text{-HOMO}$ extends further towards the $\pi^*_{\text{CO}}\text{-LUMO}$ and becomes higher in energy (see Fig. S2b and Table S2 for the orbital energies, ESI†). This results in a larger S and a smaller $\Delta\epsilon$ between the $\pi^*_{\text{CO}}\text{-LUMO}$ and the $\text{LP}_X\text{-HOMO}$. For example, in the series of $[\text{Pd}(\text{CO})_2\text{Cl}_2]_2$, $[\text{Pd}(\text{CO})_2\text{Br}_2]_2$, and $[\text{Pd}(\text{CO})_2\text{I}_2]_2$ at $r_{M \cdots M} = 3.5$, S increases from 0.08 to 0.09 to 0.10 and $\Delta\epsilon$ decreases from 3.4 eV to 3.1 eV to 2.7 eV. Consequently, $S^2/\Delta\epsilon$ times 10^3 increases from 1.9 to 2.6 to 3.7 along the same series (Table 2).

Since the ΔE_{oi} term is not made of one single component, our analyses show that there is an interplay between the metal–metal and ligand–ligand donor–acceptor interactions in the $[M(\text{CO})_2\text{X}_2]_2$ dimers along $M = \text{Ni}, \text{Pd}, \text{Pt}$ and along $X = \text{Cl}, \text{Br}, \text{I}$. Therefore, to understand the relative importance of the metal–metal and ligand–ligand interactions, we computed the magnitude of their orbital stabilization according to eqn (3) and estimated their relative contribution to the total orbital stabilization (metal–metal + ligand–ligand). The results are collected in Table 2.

When the metal centers are large and the ligands are small and more electronegative, the donor–acceptor metal–metal interactions are strengthened and dominate over the weakened ligand–ligand interactions. For example, for $[\text{Pt}(\text{CO})_2\text{Cl}_2]_2$ at $r_{M \cdots M} = 3.5$ Å, the metal–metal interactions are up to 67% of the total orbital stabilization. On the other hand, the ligand–ligand

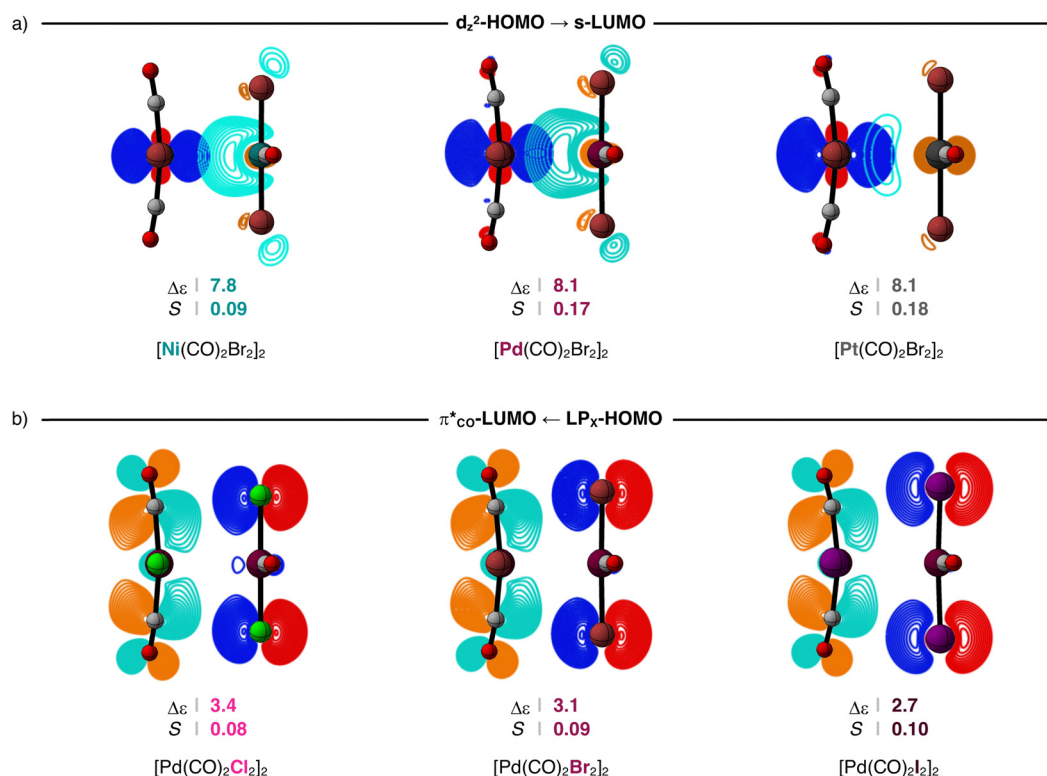


Fig. 7 Stabilizing HOMO–LUMO energy gap (in eV) and orbital overlap for (a) the metal–metal and (b) ligand–ligand donor–acceptor interactions for the representative $[M(\text{CO})_2\text{Br}_2]_2$ ($M = \text{Ni}, \text{Pd}, \text{Pt}$) and $[\text{Pd}(\text{CO})_2\text{X}_2]_2$ ($X = \text{Cl}, \text{Br}, \text{I}$) dimers at a consistent $M \cdots M$ bond distance ($r_{M \cdots M} = 3.5$ Å; contour plots from 0.9 to 0.04 a.u.). Computed at ZORA-BLYP-D3(BJ)/TZ2P.



donor–acceptor interactions only dominate when the metal center is small and the ligands are large and less electronegative, like in the $[\text{Ni}(\text{CO})_2\text{I}_2]_2$ dimer, in which the ligand–ligand donor–acceptor interactions are up to 75% of the total orbital stabilization at $r_{\text{M}\cdots\text{M}} = 3.5 \text{ \AA}$ (Table 2).

4. Conclusions

The stability of square planar $[\text{X}_2(\text{CO})_2\text{M}] \cdots [\text{M}(\text{CO})_2\text{X}_2]$ dimers increases as the metal centers M vary along Ni, Pd, and Pt, and the X ligands vary along Cl, Br, and I. Our quantum chemical bonding analyses show that the dimers are formed due to stabilizing electrostatic interactions on top of stabilizing dispersion interactions promoted by the ligands. Both interactions become more stabilizing as M and X increase in size. In addition, our analyses revealed an overlooked covalent component stemming from metal–metal and ligand–ligand donor–acceptor interactions. These findings emerge from our quantitative Kohn–Sham molecular orbital analyses using dispersion-corrected relativistic density functional theory.

The stability of the studied $[\text{M}(\text{CO})_2\text{X}_2]_2$ dimers increases as M varies along Ni, Pd, and Pt as the electron density of one monomer becomes more diffuse around M and more effectively interpenetrates towards the nucleus of M on the other monomer, resulting in a stronger electron–nuclei electrostatic attraction. Larger X ligands further increase the stability of $[\text{M}(\text{CO})_2\text{X}_2]_2$ due to a greater dispersion interaction as X increases in size along Cl, Br, and I. The dispersion component in $[\text{X}_2(\text{CO})_2\text{M}] \cdots [\text{M}(\text{CO})_2\text{X}_2]$, which is almost insensitive to variations in the metal center, only dominates when M is small, and X is large.

The overlooked covalent component in $[\text{X}_2(\text{CO})_2\text{M}] \cdots [\text{M}(\text{CO})_2\text{X}_2]$ comprises the metal–metal donor–acceptor interaction of the nd_{z^2} -type HOMOs of one monomer with $(n + 1)s$ -type LUMOs of the other monomer as well as the ligand–ligand donor–acceptor interaction of the $\text{C}=\text{O}$ π^* -type LUMO on one monomer with the X lone-pair-type HOMO on the other monomer. The metal–metal donor–acceptor interaction dominates when M is large, resulting in a larger stabilizing nd_{z^2} -HOMO– $(n + 1)s$ -LUMO overlap. On the other hand, the ligand–ligand donor–acceptor interactions dominate when M is small and X becomes less electronegative. In this situation, the metal–metal donor–acceptor interaction is weakened, and the ligand–ligand donor–acceptor interaction is favored due to a smaller π^*_{CO} -LUMO– LP_X -HOMO energy gap.

Data availability

The data that support the findings of this study are available in the ESI,[†] of this article.

Conflicts of interest

There are no conflicts to declare.

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