# **RSC Advances**



PAPER

View Article Online

View Journal | View Issue



Cite this: RSC Adv., 2017, 7, 56054

# Theoretical investigation on donor—acceptor interaction between a carbonyl compound and an *N,N'*-dioxide—Sc(III) complex†

Zhishan Su, D Weiying He, Junming Wang, Yini Zuo and Changwei Hu D\*

Herein, metal-ligand bonding features in a chelation N,N'-dioxide—Sc(III) complex have been addressed using the DFT method at the M06/6-31+G\*\* Level. The donor–acceptor interaction between the carbonyl substrate and Sc(IIII)-based catalyst is analyzed in detail by the activation strain model (ASM), energy decomposition analysis (EDA), and natural orbital for chemical valence (NOCV) calculations. The orbital interaction is the major contributor to N,N'-dioxide— $[Sc(OTf)]^{2+}$  bonding, whereas the electrostatic interaction plays a more important role than orbital interaction in the activation of a carbonyl compound in hexacoordinate N,N'-dioxide—Sc(IIII) complexes. The substituents in the amide group of the N,N'-dioxide ligand (L) affect the electrostatic energy as well as the orbital energy between the  $CH_2O$  and Sc(IIII)-based catalyst by adjusting the Lewis acidity of the metal centre. The complex with *ortho*-diisopropylphenyl groups in the ligand exhibits a higher reactivity towards  $CH_2O$ . Compared to OiPr, the counter ion OTf in the Sc(IIII)-complex enhances the Lewis acidity of the metal centre and facilitates the activation of  $CH_2O$  by promoting electron density flow from  $CH_2O$  to the metal fragment. The high catalytic performance of the N,N'-dioxide—Sc(IIII) complex towards PhCHO and chalcone is attributed to their good nucleophilicity that results in a more stabilizing electrostatic and orbital interaction between the N,N'-dioxide— $[Sc(OTf)]^{2+}$  complex and carbonyl substrate.

Received 9th November 2017 Accepted 27th November 2017

DOI: 10.1039/c7ra12258f

rsc.li/rsc-advances

#### Introduction

The  $C_2$ -symmetric N,N'-dioxide (L) developed by Feng's group is a class of conformationally flexible ligands, which can be synthesized from cheap optically pure amino acids.1 The experimental investigations indicate that these ligands can act as neutral tetradentate ligands to bind a wide variety of metal ions; this results in the formation of effective and versatile chiral metal-complex catalysts for asymmetric catalysis. 1a,b After coordination at the metal centre, the chelating ligand is able to impose a relatively rigid environment, tuning the electron richness of metal centre and, in turn, the reactivity of the coordination complex.<sup>2</sup> In the catalytic processes involving N,N'dioxide-metal complexes, the carbonyl compound is one of the most popular substrates since it can coordinate towards the metal centre in a monodentate or bidentate fashion (if possible) via oxygen atoms to form reacting species. When only one coordinating O atom is available in a carbonyl compound,

Key Laboratory of Green Chemistry and Technology, Ministry of Education, College of Chemistry, Sichuan University, Chengdu, Sichuan, 610064, P. R. China. E-mail: changweihu@scu.edu.cn

 $\dagger$  Electronic supplementary information (ESI) available: Cartesian coordinates of all stationary points and energies; selected geometric and electronic parameters as well as results of for ETS-NOCV analysis Sc(III)-complexes. See DOI: 10.1039/c7ra12258f

another ligand, solvent or counter anion can occupy the remaining coordination site to construct an octahedral intermediate for subsequent conversion. The experimental observations and theoretical investigations indicate that the activity and selectivity of a chiral  $N_iN^i$ -dioxide–metal catalyst towards a carbonyl substrate depend closely on the inherent properties (such as Lewis acidity, ionic radius or coordination sphere properties) of the central metal ion as well as ligand structures (amide substituent or the amino acid backbone).<sup>3</sup>

The classical Dewar-Chatt-Duncanson model is generally employed to understand the metal-ligand interaction in organometallic chemistry.4,5 For transition metal-based systems, bonding is attributed to a synergic process of ligand → metal electron donation and metal → ligand back-donation.6 Many useful schemes, including charge decomposition analysis (CDA),7-10 atoms in molecule (AIM),11 block localized wave function energy decomposition (BLW-ED),12 interactionenergy partition analysis,13 lever electronic parameters,14 and molecular electrostatic potential (MEP),15 have been developed to describe this donor-acceptor interaction in metallic complexes. In addition, the results obtained from spectroscopic measurement, such as infrared stretching frequencies (e.g. Tolman electronic parameter (TEP))7,14a-c and NMR chemical shifts,16 can be used as indicators to evaluate the electronic donation of the ligand, or monitor structural as well as energetic changes in complexes with different ligands. Clot and coworkers calculated the vibrational frequency of CO in [Ni(CO)<sub>3</sub>L] complexes (L = P-donor ligands) by the B3PW91 method. The better the donor ability of phosphine (R<sub>3</sub>P) in the [Ni(CO)<sub>3</sub>R<sub>3</sub>P] complex, the lower the  $\nu(CO)$  vibration becomes as a result of back-donation from metal d orbitals into the low-lying antibonding  $\pi_{CO}^*$  orbital. The activation strain model (ASM)18-20 or distortion/interaction model21-23 combined with energy decomposition analysis (EDA)24 provides a quantitative tool to understand the physical factors controlling the bonding energy or activation barrier in reactions. Recently, as a useful approach, a combination of the extended transition state (ETS)13,25 scheme with the natural orbitals for the chemical valence (NOCV)26 method (namely ETS-NOCV scheme) is introduced to metallic complexes as well as organic molecules, offering a compact picture of chemical bond formation and its different components, as well as donor-acceptor properties in the molecular complex.<sup>2,27-29</sup> In the combined ETS-NOCV scheme, the orbital interaction energy ( $\Delta E_{\rm orb}$ ) between reacting fragments is expressed in terms of NOCV eigenvalues  $(\nu_k)$  as

$$\Delta E_{
m orb} = \sum_k \Delta E_{
m orb}{}^k = \sum_{k=1}^{M/2} 
u_k \Big[ -F_{-k,-k}^{
m TS} + F_{k,k}^{
m TS} \Big]$$

Where  $F_{-k,-k}^{TS}$  and  $F_{k,k}^{TS}$  are the diagonal Kohn-Sham matrix elements defined over NOCV with respect to the transition state (TS) density (at the midpoint between density of molecule and the sum of fragment densities). Each complementary NOCV pair represents one of the charge deformations  $\Delta \rho_k$ . The abovementioned components  $\Delta E_{\rm orb}^{\ \ k}$  provide energetic estimation of  $\Delta \rho_k$  that may be related to the importance of a particular electron flow channel for the bonding between considered molecular fragments.27b,c

Understanding the properties of the metal-ligand chemical bond is important for rationalizing and predicting the catalytic performance of N,N'-dioxide-metal complexes and developing new catalysts. To date, only few studies have been reported on the electronic properties of a chelating multidentate ligand as compared to those on mono or bidentate phosphines. 14b,30,31 The ranking of the donor ability of the chiral N,N'-dioxide ligand featuring four coordinating end groups still remains a challenge. In our previous calculations, 3a,b we studied the influence of the chiral backbone, linkage, and the substituent of aniline in the N,N'-dioxide-Mg(II) catalyst on its reactivity towards a dicarbonyl compound (isatin) and the corresponding chiral inductivity in asymmetric carbonyl-ene reaction. Herein, we changed the substituent of amide in the ligand in the N,N'dioxide-Sc(III) catalysts to investigate the catalytic behaviour of these catalysts towards three mono-carbonyl compounds

Scheme 1 Three L-proline-derived chiral N,N'-dioxide ligands (L1-L3).

(formaldehyde, benzaldehyde, and chalcone) via analysis of the donor-acceptor interaction (Scheme 1). Furthermore, the influence of the counterion from two scandium reagents (Sc(OTf)<sub>3</sub> and Sc(OiPr)<sub>3</sub>) on substrate-catalyst interaction (catalytic abilities) has been explored by the DFT method.

# Computational details

Geometry optimization and frequency calculations are performed via density functionary theory (DFT) using the Gaussian 09 program package.32 The M06 hybrid functional33 is adopted, in conjugation with the 6-31+G\*\* basis set.34 Natural bond orbital (NBO)35 analysis is employed to obtain further insight into the electronic properties of the system at the same theoretical level. The results of the activation strain model (ASM) on hexacoordinate chiral N,N'-dioxide-Sc(III) complexes are shown in Tables S1-S3,† in which the reaction energy  $\Delta E_r$  between metal-based fragment and ligand or carbonyl substrate in the complexes is decomposed into two chemical meaningful components: the distortion energy ( $\Delta E_{\text{strain}}$ ) and interaction energy ( $\Delta E_{int}$ ), by Gaussian 09. To understand the factors influencing the donor-acceptor interaction,  $\Delta E_{\text{int}}$  is further decomposed into electrostatic interaction ( $\Delta V_{\rm elstat}$ ), Pauli repulsion ( $\Delta E_{\text{Pauli}}$ ), and orbital interaction ( $\Delta E_{\text{oi}}$ ) (i.e.,  $\Delta E_{\text{int}} =$  $\Delta V_{\rm elstat} + \Delta E_{\rm Pauli} + \Delta E_{\rm oi}$ ) by energy decomposition analysis (EDA). EDA and ETS-NOCV calculations are performed by single-point calculation based on the optimized geometries obtained by Gaussian 09 using the Amsterdam density functional (ADF) program<sup>36</sup> at the M06/TZP level. We selected ScCl<sub>3</sub> and Sc(OTf)3 compounds as models to examine the multireference character of scandium compounds at the CCSD/6-31+G\*\*//M06/6-31+G\*\* level using the Gaussian 09 program. The corresponding T1 diagnostics are predicted to be 0.016 for ScCl<sub>3</sub> and 0.018 for Sc(OTf)<sub>3</sub>, which are smaller than the criteria 0.02, as suggested by Lee and Taylor. 37

#### Results and discussion

#### Donor-acceptor interaction between the ligand and [Sc(OTf)]<sup>2+</sup> species

Herein, we studied three Sc(III)-complexes (L-[Sc(OTf)]<sup>2+</sup>) formed by the coordination of N,N'-dioxide ligand (L1-L3) to [Sc(OTf)]<sup>2+</sup> in the gas phase. The corresponding complexes are denoted as L1-Sc(III)-OTf, L2-Sc(III)-OTf, and L3-Sc(III)-OTf for convenience. The coordination processes are exothermic by -275.8 to -290.8 kcal mol<sup>-1</sup>. As shown in Fig. S1,† the main structural parameters of these three complexes are similar, with the average distances between the four O-donors in the ligands to the Sc cation of 2.066-2.075 Å. NBO analysis indicates that the charge transfer (CT) from ligand to the [Sc(OTf)]<sup>2+</sup> moiety exhibits the following trend: L1-Sc(III)-OTf (0.942) > L2-Sc(III)-OTf (0.923) > L3-Sc(III)-OTf (0.913). As a result, the natural population analysis (NPA) of charge accumulated on the Sc(III) centre in L3-Sc(III)-OTf (1.629) is slightly larger than those of L1-Sc(III)-OTf (1.600) and L2-Sc(III)-OTf (1.622). As shown in Table 1, the bonding energy ( $\Delta E_{\rm BE}$ ) for L1-Sc(III)-OTf with an aliphatic cyclohexyl group is higher than those of the complexes

Table 1 Results of NBO and ETS-NOCV calculations for hexacoordinate complexes formed by ligand (L1–L3) with  $[Sc(OTf)]^{2+}$  fragments. The energies are in kcal mol<sup>-1</sup>

Complex	Charge <sup>a</sup>	$\mathrm{CT}^b$	$\Delta E_{ m Pauli}$	$\Delta V_{ m elstat}$	$\Delta E_{ m oi}$	$\Delta E_{ m orb}^{\sigma} \left(1 ight)$	$\Delta E_{ m orb}^{\sigma} \left( 2 \right)$	$\Delta E_{ m BE}{}^c$
L1–Sc(III)–OTf	1.600	0.942	165.0	-284.4	-283.8	-74.8	-97.1	-403.2
L2-Sc(III)-OTf	1.622	0.923	163.8	-272.1	-284.0	-73.4	-100.4	-392.3
L3-Sc(III)-OTf	1.629	0.913	165.3	-273.1	-292.7	-71.8	-96.0	-400.6

<sup>&</sup>lt;sup>a</sup> NPA charge accumulated on Sc(III) ion in the hexacoordinate complexes. <sup>b</sup> Charge transfer (CT) from the ligand (L1–L3) to the [Sc(OTf)]<sup>2+</sup> fragment. <sup>c</sup> Bonding energy (BE) between the ligand and the [Sc(OTf)]<sup>2+</sup> fragment obtained by ADF calculation.

containing aromatic substituent ligands (L2 and L3). These results indicate that there exists a strong interaction between L1 and [Sc(OTf)]<sup>2+</sup> fragment.

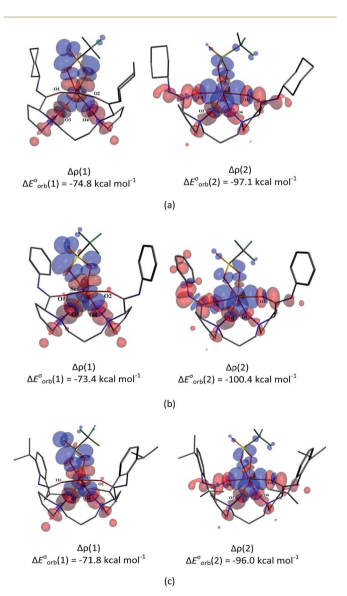


Fig. 1 Dominating contributions to the deformation density  $\Delta\rho$  associated with the coordination interaction between the ligand L1 and the  $[Sc(OTf)]^{2+}$  fragment in L1–Sc(III)–OTf (a), L2–Sc(III)–OTf (b), and L3–Sc(III)–OTf (c) complexes determined by the ETS-NOCV analysis. The contour value is  $|\Delta\rho|=0.001$  au. The blue/red contours correspond to accumulation/depletion of electron density.

EDA analysis suggests that the electrostatic and orbital interactions are the main contributors for ligand–[Sc(OTf)]<sup>2+</sup> interactions in these three complexes. For L1–Sc(III)–OTf, the electron-donating cyclohexyl group enhances the electrostatic interaction between L1 and [Sc(OTf)]<sup>2+</sup> fragments ( $\Delta V_{\rm elstat} = -284.4~{\rm kcal~mol^{-1}}$ ), contributing to a more stabilizing interaction energy. For L2–Sc(III)–OTf and L3–Sc(III)–OTf, the  $\Delta V_{\rm elstat}$  are comparable ( $-272.1~{\rm vs.}~273.1~{\rm kcal~mol^{-1}}$ ). Although introduction of *ortho-i*Pr groups into L3 slightly increases the destabilizing  $\Delta E_{\rm Pauli}$  (165.3 kcal mol<sup>-1</sup>) in L3–Sc(III)–OTf, the more stabilizing orbital interaction ( $\Delta E_{\rm oi} = 292.7~{\rm kcal~mol^{-1}}$ ) compensates for this unfavourable Pauli repulsion. Consequently, the  $\Delta E_{\rm BE}$  for L3–Sc(III)–OTf is 8.3 kcal mol<sup>-1</sup>, more stable than that for L2–Sc(III)–OTf.

To quantify the electron-transfer process between ligand and  $[Sc(OTf)]^{2+}$  moieties in these complexes, we visualized the deformation density  $(\Delta\rho)$ , and the corresponding orbital energies are shown in Fig. 1.  $\Delta\rho$  (1) represents the  $\sigma$ -donation from the N-oxide units of the ligand to the unoccupied  $d_{x_2-y_2}$  orbital of the Sc(III) centre, with  $\Delta E_{\rm orb}^{\sigma}$  (1) of -71.8 to -74.8 kcal mol $^{-1}$ . For  $\Delta\rho$  (2), the electronic density accumulation appears in the coordination region between four O-donors and metalfragments, thus indicating an electron flow from the ligand to the metal ion to strengthen O–Sc bonding. Interestingly, these effects also weaken the interaction between the counter OTf ion and metal centre, accompanying electronic density flow back from Sc(III) ion to OTf anion. This cooperation effect of coordination units in the chiral N,N'-dioxide-metal complex has also been observed in our previous study.<sup>3 $\alpha$ </sup>

# Effect of ligand in the Sc(m)-complex on the activation of $CH_2O$

The hexacoordinate complexes (L1–CH<sub>2</sub>O–OTf, L2–CH<sub>2</sub>O–OTf, and L3–CH<sub>2</sub>O–OTf) formed by coordinating formaldehyde (CH<sub>2</sub>O) to an L–[Sc(OTf)]<sup>2+</sup> complex were then investigated. Herein, the counter OTf anion occupies one coordinating site to stabilize the corresponding octahedral geometries (Fig. 2). Compared to free CH<sub>2</sub>O, the C=O bonds in these complexes are lengthened by 0.021–0.026 Å. Moreover, the corresponding Wiberg bond indices (WBI) decrease from 1.912 to 1.635–1.654. These results suggest that the C=O bonds of CH<sub>2</sub>O in Lm–CH<sub>2</sub>O–OTf (m=1–3) are significantly weakened. The NBO analysis indicates that the charge transfer (CT) occurs from CH<sub>2</sub>O to a catalyst accompanying the coordination process; this leads to electronic density re-distribution in the CH<sub>2</sub>O moiety.

Paper RSC Advances

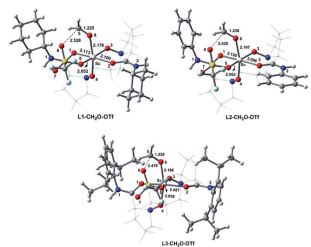


Fig. 2 Optimized geometries of the three hexacoordinate complexes L1–CH<sub>2</sub>O–OTf, L2–CH<sub>2</sub>O–OTf, and L3–CH<sub>2</sub>O–OTf. Selected bond lengths are in Å. The chiral backbone in ligands is shown in grey colour for clarity.

Consequently, the C=O bond becomes more polarized with an increase in dipole moment (from 2.533 Debye in free CH<sub>2</sub>O to 2.660–2.644 Debye). Accordingly, the C=O stretching vibration frequency ( $\nu_{\rm C=O}$ , cm<sup>-1</sup>) red-shifts from 1864 cm<sup>-1</sup> in free CH<sub>2</sub>O to 1747–1765 cm<sup>-1</sup> in the complexes. For L3–CH<sub>2</sub>O–OTf, the WBI of C=O bond is smaller than those of L1–CH<sub>2</sub>O–OTf and L2–CH<sub>2</sub>O–OTf. Moreover, the more negative charge accumulates on the O8 atom ( $-0.652~\nu s.~-0.621$  for L1–CH<sub>2</sub>O–OTf and -0.630 for L2–CH<sub>2</sub>O–OTf), which is consistent with the stronger polarity character of the C=O bond and lower  $\nu_{\rm C=O}$  (1765 cm<sup>-1</sup>) in the CH<sub>2</sub>O moiety. These results indicate that the catalyst with the L3 ligand exhibits slightly superior reactivity towards the CH<sub>2</sub>O substrate.

Starting from the optimized structures of the three complexes, we performed the relaxed potential energy surface (PES) scan by increasing the distance between CH<sub>2</sub>O and Sc(III) centre  $(R_{O8\cdots Sc}, \text{ Å})$  by 0.05 Å at the M06/6-31+G\*\* level. The variation of abovementioned electronic descriptors are shown in Tables S1-S3.† As shown in Fig. 3a and b,  $\nu_{C=O}$  and the corresponding WBI for the C5=O8 bond exhibit an increasing tendency with  $R_{O8\cdots Sc}$ . The corresponding correlation equations are shown in Table 2, with  $R^2$  of 0.991-0.997 and 0.999. Although the WBI of the C=O bond of the CH<sub>2</sub>O moiety in L1- $CH_2O$ -OTf and L2- $CH_2O$ -OTf is comparable (1.654 vs. 1.640), the C=O bond in L2-CH2O-OTf becomes remarkably less polarized upon lengthening the O8···Sc distance. As a result, a steeper variation of  $\nu_{C=O}$  is observed for L2-CH<sub>2</sub>O-OTf (Fig. 3a). Different from  $\nu_{C=O}$ , the WBI of O8···Sc interaction decreases linearly as the distance between CH2O and catalyst increases (Fig. 3b). Good linear correlations are observed for the three complexes, with an  $R^2$  of 0.995–0.997 (Table 3).

Activation strain model (ASM) analysis indicates that the interaction energy term  $\Delta E_{\rm int}$  makes a major contributions to the CH<sub>2</sub>O-L-[Sc(OTf)]<sup>2+</sup> interaction in the three abovementioned complexes as compared to the deformation energy

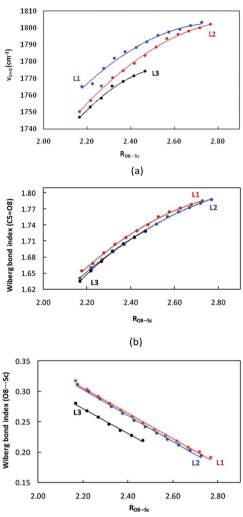


Fig. 3 Variation of the C=O stretching vibration frequency ( $\nu_{C=O}$ , cm $^{-1}$ ) (a), the corresponding Wiberg bond index (WBI) of the C=O bond (b), and WBI of the O8···Sc interaction (c) in L1–CH<sub>2</sub>O–OTf, L2–CH<sub>2</sub>O–OTf, and L3–CH<sub>2</sub>O–OTf complexes with the increasing distance between O8 and Sc(III) ( $R_{O8\cdots Sc}$ , Å).

(c)

 $\Delta E_{
m strain}$ . Furthermore, the deformation energies of the metalfragment  $(\Delta E_{\text{strain}(CAT)})$  are significantly more destabilizing than that of the  $CH_2O$  fragment ( $\Delta E_{\text{strain}(CH,O)}$ ). Similarly, there also exist good correlations between  $R_{O8\cdots Sc}$  and  $\Delta E_{int}$ , with the coefficient of 0.997-0.999 (Fig. 4). Compared to that in L1- $CH_2O$ -OTf and L2- $CH_2O$ -OTf,  $\Delta E_{int}$  in L3- $CH_2O$ -OTf is clearly more stabilizing at any given point along the O8...Sc distance; this indicates the stronger donor-acceptor interaction between CH<sub>2</sub>O and L3-[Sc(OTf)]<sup>2+</sup> moiety, as well as Lewis acidity of Sc(III) ion in L3-CH2O-OTf. These results are in good agreement with a larger electron-deficient character for the Sc(III) centre in L3-CH<sub>2</sub>O-OTf (NPA charge accumulated on Sc(III) cation is 1.647). Although the *ortho-i*Pr groups in ligand L3 increase the Pauli repulsion between CH<sub>2</sub>O and Sc-fragment ( $\Delta E_{Pauli}$  = 40.1 kcal mol<sup>-1</sup>), the more stabilizing energies (electrostatic interaction  $\Delta V_{\rm elstat}$  and orbital interaction  $\Delta E_{\rm oi}$ ) compensates for this destabilizing steric repulsion; this leads to a more

Table 2 Correlations between  $\nu_{C=O}$  (cm<sup>-1</sup>), Wiberg bond index (WBI) of C=O bond in the CH<sub>2</sub>O moiety, the O8···Sc interaction as well as  $\Delta E_{int}$  (kcal mol<sup>-1</sup>) with O8···Sc distance ( $R_{O8\cdots Sc}$  Å) for L1-CH<sub>2</sub>O-OTf, L2-CH<sub>2</sub>O-OTf, and L3-CH<sub>2</sub>O-OTf complexes

Ligand	$N^a$	Equations	$R^2$
L1	12	$\nu_{\text{C}=\text{O}} = -96.9 \times (R_{\text{O8}\text{Sc}})^2 + 544.9 \times (R_{\text{O8}\text{Sc}}) + 1037.4$	0.991
		$WBI_{C5=O8} = -0.227 \times (R_{O8\cdots Sc})^2 + 1.357 \times (R_{O8\cdots Sc}) - 0.224$	0.999
		$WBI_{O8\cdots Sc} = -0.214 \times (R_{O8\cdots Sc}) + 0.774$	0.997
		$\Delta E_{\rm int} = 11.6 \times (R_{\rm O8\cdots Sc})^2 - 34.4 \times (R_{\rm O8\cdots Sc}) - 19.6$	0.997
L2	13	$\nu_{\text{C}=\text{O}} = -91.5 \times (R_{\text{O8}\cdots\text{Sc}})^2 + 537.3 \times (R_{\text{O8}\cdots\text{Sc}}) + 1015.0$	0.997
		$WBI_{C5=O8} = -0.170 \times (R_{O8\cdots Sc})^2 + 1.085 \times (R_{O8\cdots Sc}) + 0.091$	0.999
		$WBI_{O8\cdots Sc} = -0.212 \times (R_{O8\cdots Sc}) + 0.770$	0.997
		$\Delta E_{ m int} = 9.2 \times \left( R_{ m O8\cdots Sc} \right)^2 - 21.6 \times \left( R_{ m O8\cdots Sc} \right) - 39.4$	0.998
L3	7	$\nu_{\text{C}==\text{O}} = -161.7 \times (R_{\text{O8}\cdots\text{Sc}})^2 + 841.1 \times (R_{\text{O8}\cdots\text{Sc}}) + 683.4$	0.995
		$WBI_{C5=O8} = -0.355 \times (R_{O8\cdots Sc})^2 + 1.958 \times (R_{O8\cdots Sc}) - 0.942$	0.999
		$WBI_{O8\cdots Sc} = -0.204 \times (R_{O8\cdots Sc}) + 0.720$	0.995
		$\Delta E_{\rm int} = 21.7 \times (R_{\rm O8\cdots Sc})^2 - 80.0 \times (R_{\rm O8\cdots Sc}) + 27.5$	0.999

<sup>&</sup>lt;sup>a</sup> Number of data points employed in the correlation.

Table 3 Results of ASM, EDA, and ETS-NOCV calculations for hexacoordinate complexes. The energies are in kcal  $mol^{-1}$ 

					$\Delta E_{ m strain}$								
Ligand	Substituent	Counterion	Carbonyl substrate	$\Delta E_{ m int}^{a}$	CAT	Carbonyl substrate	Sum	$\Delta E_{ m r}$	$\Delta E_{ m Pauli}$	$\Delta V_{ m elstat}$	$\Delta E_{ m oi}$	$\Delta E_{ m orb} \left( 1  ight)$	$\Delta E_{ m orb}$ (2)
L1	Cyclohexyl	OTf	CH <sub>2</sub> O	-39.0(-37.0)	14.4	0.7	15.0	-24.0	37.1	-43.6	-30.5	-31.5	-15.6
L2	Ph	OTf	CH <sub>2</sub> O	-42.4(-40.8)	15.0	0.8	15.8	-26.6	39.6	-46.6	-33.8	-34.5	-16.8
L3	2,6- <i>i</i> Pr	OTf	$CH_2O$	-43.6(-41.1)	19.1	0.8	20.0	-23.6	40.1	-47.3	-33.9	-34.1	-18.2
L3	2,6- <i>i</i> Pr	O <i>i</i> Pr	$CH_2O$	-32.6(-30.9)	2.3	0.3	2.6	-30.0	22.3	-34.3	-18.9	-25.7	-3.3
L3	2,6- <i>i</i> Pr	OTf	PhCHO	-56.4(-53.1)	20.0	2.0	22.0	-34.4	42.7	-56.8	-39.0	-34.4	-21.6
L3	2,6- <i>i</i> Pr	OTf	Chalcone	-71.7(-67.0)	21.7	4.6	26.3	-45.4	47.7	-65.7	-49.0	-33.5	-27.0

<sup>&</sup>lt;sup>a</sup> Interaction energies obtained by ADF are shown in bracket.

stabilizing  $\Delta E_{\rm int}$  (-41.1 kcal mol<sup>-1</sup>). Compared to the case of L3–Sc(III)–OTf, the orbital interaction  $\Delta E_{\rm oi}$  between the CH<sub>2</sub>O-fragment and Sc(III)-fragment is smaller than electrostatic energy  $\Delta V_{\rm elstat}$  by 13.4 kcal mol<sup>-1</sup> (-33.9 vs. -47.3 kcal mol<sup>-1</sup>) in L3–CH<sub>2</sub>O–OTf. Therefore, the electrostatic interaction is the major driving force to form L3–CH<sub>2</sub>O–OTf by interaction between CH<sub>2</sub>O and L3–[Sc(OTf)]<sup>2+</sup> moiety.

For L1, the electron-donating cyclohexyl groups in the amide units enhance the electronic density transfer from the ligand to

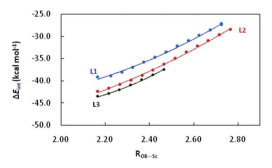


Fig. 4 Variation of the interaction energy  $\Delta E_{\rm int}$  (kcal mol<sup>-1</sup>) between CH<sub>2</sub>O and L-[Sc(OTf)]<sup>2+</sup> moiety with the increasing distance between O8 and Sc(III) ( $R_{\rm O8\cdots Sc}$ , Å) in L1-CH<sub>2</sub>O-OTf, L2-CH<sub>2</sub>O-OTf, and L3-CH<sub>2</sub>O-OTf complexes.

the Sc(III) center in the L1-CH<sub>2</sub>O-OTf complex; this decreases the Lewis acidity of metal ion ( $\Delta E_{\rm int} = -37.0 \text{ kcal mol}^{-1}$ ). As expected, both  $\Delta V_{\rm elstat}$  and  $\Delta E_{\rm oi}$  are less stabilizing than those in L3-CH<sub>2</sub>O-OTf by 3.7 and 3.4 kcal mol<sup>-1</sup>, respectively. With respect to L2, the phenyl group on the right brachial amide tends to be placed in the same plane to the carbonyl of amines in L2-CH<sub>2</sub>O-OTf. A good conjugate effect between the phenyl group and carbonyl of amines compensates for the electronicdeficient character of Sc(III) cation in L2-CH2O-OTf by strengthening the electron density transfer from ligand to metal, decreasing the stabilizing attracting term ( $\Delta V_{\rm elstat}$  and  $\Delta E_{oi}$ ) between CH<sub>2</sub>O-fragment and Sc(III)-fragment by 1.5 kcal mol<sup>-1</sup>. As a result, the  $\Delta E_{\rm int}$  of L2-CH<sub>2</sub>O-OTf is less that of L3-CH<sub>2</sub>O-OTf (-40.8)stabilizing than  $-41.1 \text{ kcal mol}^{-1}$ ).

We visualized the deformation density associated with the orbital interaction between  $CH_2O$  and the Sc(III)-based fragments in the three complexes. The two dominating components of deformation density,  $\Delta\rho$  (1) and  $\Delta\rho$  (2), with the corresponding energy values are presented in Fig. 5 and S3.† As shown in Fig. 5, the accumulation of electron density appears in the region between the O8 atom and the scandium centre in L3– $CH_2O$ -OTf; this indicates electrons flowing from  $\sigma$ -donor O lone electron pair of  $CH_2O$  to unoccupied d orbital of Sc(III) ion

Paper RSC Advances

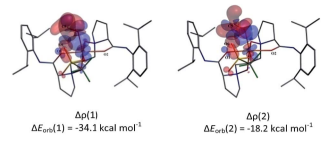


Fig. 5 Dominating contributions to the deformation density  $\Delta\rho$  describing the coordination interaction between CH<sub>2</sub>O and scandium-based fragments for L3–CH<sub>2</sub>O–OTf complex by ETS-NOCV analysis. The contour value is  $|\Delta\rho|=0.001$  au. The blue/red contours correspond to accumulation/depletion of electron density.

to form the Sc–O bond. Simultaneously, the depletion of electron density on  $\pi$  double C=O bond of CH<sub>2</sub>O suggests the weakening of  $\pi$  bonds in CH<sub>2</sub>O. With respect to the second NOCV-based deformation density contribution  $\Delta\rho$  (2), it presents 36% of the interfragment orbital interaction, accompanying the weakening of C–H as well as C=O bonds in CH<sub>2</sub>O. In addition, the slight electronic density accumulation between the O6 atom of OTf anion and the C8 atom of CH<sub>2</sub>O stabilizes the hexacoordinate complexes, further favouring the activation of CH<sub>2</sub>O. Similar results are obtained for L1–CH<sub>2</sub>O–OTf and L2–CH<sub>2</sub>O–OTf, as shown in Fig. S3 in the ESI.†

As shown in Table S4 and Fig. S4 in the ESI,† CDA analysis indicates that the main molecular orbital involving the donoracceptor interaction and electronic density transfer between  $\mathrm{CH_2O}$  and  $\mathrm{L3-Sc}(\mathrm{III})$  in the  $\mathrm{L3-CH_2O-OTf}$  complex is the occupied HOMO-146 orbital. It is constructed from the HOFO-5 of CH<sub>2</sub>O fragment and LUFO-2, LUFO-3, and LUFO-6 orbitals of the L3-Sc(III) fragment. The HOFO-5 is the hybridized orbital of  $\sigma(CH_2)$  and s atomic orbital of  $CH_2O$ . The total contribution of LUFO orbitals in the L3-Sc(III) fragment is smaller than that of the HOFO-5 of CH<sub>2</sub>O moiety by 53.2% (66.4% vs. 13.2%). Interestingly, the contribution of the unoccupied orbital in the Sc(m)-based fragment decreases in the order of L3 > L2 > L1, which is in good agreement with electron density transfer from CH2O to catalysts. Thus, the combination of stronger electrostatic interactions and the more stabilizing orbital interaction contributes to the superior reactivity of L3-[Sc(OTf)]<sup>2+</sup> towards  $CH_2O$ .

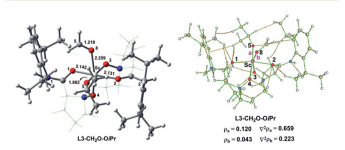


Fig. 6 Optimized geometry and AIM analysis of hexacoordinate L3– $\mathrm{CH_2O-O}$ iPr. Selected bond lengths are in Å. The chiral backbone in the ligand is shown in grey colour for clarity.

#### Effect of counterion on the activation of CH2O

To understand the effect of counterion in the catalyst on Lewis acidity of the metal ion as well as the activation of carbonyl, we replaced the OTf anion in L3-CH<sub>2</sub>O-OTf with iso-propoxy ion (OiPr). The corresponding optimized geometries of the hexacoordinate complexes L3-CH2O-OiPr are shown Fig. 6. Compared to those for L3-CH<sub>2</sub>O-OTf, the O8···Sc distances are longer by 0.094 Å for L3-CH<sub>2</sub>O-OiPr. Furthermore, the C=O bond length of CH<sub>2</sub>O follows the order L3-CH<sub>2</sub>O-OTf > L3-CH<sub>2</sub>O-OiPr. This result indicates that CH2O becomes less weakened in L3-CH<sub>2</sub>O-OiPr. The O5-Sc distance in L3-CH<sub>2</sub>O-OiPr is shorter significantly than those of L3-CH<sub>2</sub>O-OTf (1.862 Å vs. 2.058 Å); this indicates a stronger interaction between OiPr and Sc(III) centre. This result could be attributed to the stronger basicity of OiPr anion than that of OTf anion. Suffering large steric repulsion from OiPr, tetradentate N,N-dioxide ligand moves slightly far away from the metal centre. Accordingly, the average O-Sc distance in ligand in L3-CH<sub>2</sub>O-OiPr is as long as 2.131 Å. This effect also weakens the coordinate interaction of CH<sub>2</sub>O to Sc(III) ion; this leads to the less stabilizing  $\Delta V_{\rm elstat}$  (-34.3 kcal mol<sup>-1</sup>) and  $\Delta E_{\rm oi}$  $(-18.9 \text{ kcal mol}^{-1})$ . NBO analysis indicate that the energy level of LUMO orbitals in two complexes has the order of L3-CH<sub>2</sub>O-O*i*Pr > L3-CH<sub>2</sub>O-OTf. Accordingly, the electrophilic indices  $\omega$  of complexes are 9.4 eV for L3-Sc(III)-OTf and 8.1 eV for L3-Sc(III)-OiPr. Similar to the case of L3-CH<sub>2</sub>O-OTf, the  $\Delta V_{\rm elstat}$  is still a major contributor in L3-CH2O-OiPr, which is also confirmed by AIM analysis (Fig. 6). The positive Laplacian of electronic density  $\nabla^2 \rho$  at (3, -1) bonding critical points (a and b) is observed between CH<sub>2</sub>O or OiPr ion to metal centre, which is similar to that of L3-CH<sub>2</sub>O-OTf in Fig. S5.† Furthermore,  $\Delta E_{int}$  exhibits the following order L3-CH<sub>2</sub>O-OTf > L3-CH<sub>2</sub>O-OiPr, which is in good agreement with the abovementioned results of electrophilic indices. The inferior reactivity of the catalyst with Sc(OiPr)<sub>3</sub> as a precursor was also observed in the experiments.3d Therefore, the counter ion could adjust the electrophilicity of the Sc(III)-complex by affecting  $\Delta V_{\rm elstat}$  and  $\Delta E_{\rm oi}$ . The Lewis acidity of Sc(III) ion in L3-CH<sub>2</sub>O-OTf is stronger than that of L3-CH2O-OiPr, consequently, exhibiting high reactivity towards the nucleophilic CH<sub>2</sub>O substrate.

#### Variation of the carbonyl substrate

We used two more carbonyl substrates (PhCHO and chalcone) to further evaluate the reactivity of L3-[Sc(OTf)]<sup>2+</sup>. The

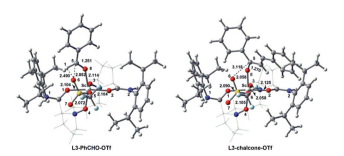


Fig. 7 Optimized geometries of the hexacoordinate complexes L3–PhCHO-OTf and L3-chalcone-OTf. Selected bond lengths are in Å. The chiral backbone in ligands is shown in grey colour for clarity.

**RSC Advances** Paper

optimized geometries for the hexacoordinate complexes L3-PhCHO-OTf and L3-chalcone-OTf are shown in Fig. 7. As shown in Table S5,† NBO analysis indicates that more negative charges accumulated on the O atom of free PhCHO or chalcone substrates because of electron delocalization of the adjacent phenyl ring (-0.561 and -0.597 vs. -0.533 in CH<sub>2</sub>O). Consequently, the nucleophilicity of PhCHO and chalcone is significantly enhanced to 2.02 and 2.81 eV, respectively, as compared to that of CH<sub>2</sub>O (1.73 eV). This effect favours the coordination interaction between PhCHO or chalcone and Sc(III) centre, with shorter O8···Sc distances of 2.114 Å for L3-PhCHO-OTf and 2.058 Å for L3-chalcone-OTf, respectively. EDA analysis indicates that the interaction energies  $\Delta E_{\rm int}$  between carbonyl substrates and scandium-based fragment are -53.1 kcal mol<sup>-1</sup> for L3-PhCHO-OTf and -67.0 kcal mol<sup>-1</sup> for L3-chalcone-OTf, which are more stabilizing than that of L3-CH<sub>2</sub>O-OTf (-41.1 kcal mol<sup>-1</sup>, Table S3†). Furthermore, the NOCV deformation density as well as the corresponding energy  $\Delta E_{\rm orb}^{\sigma} =$  $\Delta E_{\rm orb}$  (1) +  $\Delta E_{\rm orb}$  (2) for L3-PhCHO-OTf and L3-chalcone-OTf complexes are higher than that of L3-CH2O-OTf (56.0 and 60.5 kcal mol<sup>-1</sup> vs. 52.3 kcal mol<sup>-1</sup>). These results indicate that L3-[Sc(OTf)]<sup>2+</sup> exhibits higher reactivity for PhCHO and chalcone substrates as compared to that for CH2O. Contrary to PhCHO, the chalcone with a more significant  $\pi$ -conjugated structure tends to coordinate to L3-[Sc(OTf)]<sup>2+</sup>, with most stabilizing  $\Delta V_{\rm elstat}$  (-65.7  $kcal mol^{-1}$ ) and (-49.0 kcal mol<sup>-1</sup>) for carbonyl-Sc(III)-fragment interaction. The C=O bonds of PhCHO and chalcone in these complexes are lengthened by 0.039 and 0.044 Å, with  $\Delta \nu_{\rm C=O}$  of 149.3 and 200.5 cm<sup>-1</sup>, respectively, indicating that C=O bonds are activated significantly in L3-PhCH<sub>2</sub>O-OTf and L3-chalcone-OTf. These results are in good agreement with the excellent reactivity of chiral N,N'-dioxide-Sc(III) catalysts for the transformation of chalcone or PhCHO substrates in organometallic catalysis. 1a,b

#### Conclusions

Theoretical investigation on donor-acceptor interaction between a ligand and carbonyl compound with a metal ion in N,N'-dioxide-Sc(III) complex is performed using the DFT method. The effect of the substituent of the ligand and counter ion on Lewis acidity as well as the reactivity of the Sc(III)-complex towards carbonyl compound is studied. These calculations reveal the following conclusions:

- (1) The orbital attraction is the major contributor to the interaction between the chiral N,N'-dioxide ligand and Sc(III)based fragment in the L-[Sc(OTf)]<sup>2+</sup> complex. The ligand with the cyclohexyl group in amide exhibits a stronger coordination ability towards the Sc(III) centre as compared to the ligands with an aromatic ring.
- (2) The electrostatic interaction is the major driving force to form a hexacoordinate complex by interaction between CH2O and L3-[Sc(OTf)]<sup>2+</sup> moiety. Variation of amide moiety in substituents of the ligands adjusts electrostatic energy as well as orbital energy between CH<sub>2</sub>O and L-[Sc(OTf)]<sup>2+</sup> fragment in hexacoordinate complexes, affecting the activation of carbonyl compound. The combination of stronger  $\Delta V_{\rm elstat}$  and more

stabilizing  $\Delta E_{oi}$  contributes to the high catalytic performance of complex with a 2,6-diisopropylphenyl group towards CH<sub>2</sub>O.

- (3) The counter anion in the complex tunes the energy level of the LUMO orbital of N,N'-dioxide–Sc(III) complex; this affects the electrophilicity of the L-Sc(III)-complex towards the carbonyl substrate. The Lewis acidity of the metal ion in the complex could be adjusted by changing Pauli repulsion, electrostatic, as well as orbital interaction between the carbonyl compound and metal centre.
- (4) PhCHO and chalcone with high nucleophilicity exhibit excellent reactivity in the presence of L-[Sc(OTf)]<sup>2+</sup> complex; this may be related to more stabilizing electrostatic and orbital interaction. The activation process accompanies with more significantly electron density transfer from a carbonyl compound to a Sc-based fragment.

### Conflicts of interest

There are no conflicts of interest to declare.

# Acknowledgements

We thank the National Natural Science Foundation of China (No. 21290182, 21321061 and 21572141), 111 Project (B17030) and the Program for New Century Excellent Talents in the University of China (No. NCET-13-0390) for providing the financial support. We thank Prof. X. M. Feng and Prof. X. H. Liu for helpful discussions.

#### Notes and references

- 1 (a) X. H. Liu, L. L. Lin and X. M. Feng, Acc. Chem. Res., 2011, 44, 574; (b) X. H. Liu, L. L. Lin and X. M. Feng, Org. Chem. Front., 2014, 1, 298; (c) K. Shen, X. H. Liu, L. L. Lin and X. M. Feng, Chem. Sci., 2012, 3, 327; (d) X. H. Liu, H. F. Zheng, Y. Xia, L. L. Lin and X. M. Feng, Acc. Chem. Res., 2017, 50, 2621.
- 2 M. Fusè, I. Rimoldi, E. Cesarotti, S. Rampino and V. Barone, Phys. Chem. Chem. Phys., 2017, 19, 9028.
- 3 (a) J. M. Wang, Z. S. Su, N. Yang and C. W. Hu, J. Org. Chem., 2016, 81, 6444; (b) J. M. Wang, Y. N. Zuo, C. W. Hu and Z. S. Su, Catal. Sci. Technol., 2017, 7, 2183; (c) Y. N. Zuo, Z. S. Su, J. M. Wang and C. W. Hu, Catal. Today, 2017, 298, 130-137; (d) Q. Zhang, X. Xiao, L. L. Lin, X. H. Liu and X. M. Feng, Org. Biomol. Chem., 2011, 9, 5748; (e) Y. Y. Chu, X. H. Liu, W. Li, X. L. Hu, L. L. Lin and X. M. Feng, Chem. Sci., 2012, 3, 1996; (f) M. S. Xie, X. H. Liu, Y. Zhu, X. H. Zhao, Y. Xia, L. Lin and X. M. Feng, Chem.-Eur. J., 2011, 17, 13800.
- 4 M. J. S. Dewar, Bull. Soc. Chim. Fr., 1951, 18, C71-C79.
- 5 J. Chatt and L. A. Duncanson, J. Chem. Soc., 1953, 3, 2939.
- 6 M. Mitoraj and A. Michalak, Organometallics, 2007, 26, 6576.
- 7 G. Frenking and N. Fröhlich, Chem. Rev., 2000, 100, 717.
- 8 S. Dapprich and G. Frenking, J. Phys. Chem., 1995, 99, 9352.
- 9 A. W. Ehlers, S. Dapprich, S. F. Vyboishchikov and G. Frenking, Organometallics, 1996, 15, 105.

Paper

10 G. Frenking, K. Wichmann, N. Fröhlich, J. Grobe, W. Golla,

- D. Le Van, B. Krebs and M. Läge, Organometallics, 2002, 21,
- 11 V. Tognetti, L. Joubert, P. Cortona and C. Adamo, J. Phys. Chem. A, 2009, 113, 12322.
- 12 Y. Mo, J. Gao and S. D. Peyerimhoff, J. Chem. Phys., 2000, 112, 5530.
- 13 T. Ziegler and A. Rauk, Inorg. Chem., 1979, 18, 1755.
- 14 (a) C. A. Tolman, Chem. Rev., 1977, 77, 313; (b) N. Fey, Dalton Trans., 2010, 39, 296; (c) S. L. Mukerjee, S. P. Nolan, C. D. Hoff and R. L. Delavega, Inorg. Chem., 1988, 27, 81; (d) N. Fey, A. G. Orpen and J. N. Harvey, Coord. Chem. Rev., 2009, 253, 704.
- 15 (a) C. H. Suresh and N. Koga, *Inorg. Chem.*, 2002, **41**, 1573; (b) O. Kühl, Coord. Chem. Rev., 2005, 249, 693; (c) D. J. Nelson and S. P. Nolan, Chem. Soc. Rev., 2013, 42, 6723.
- 16 (a) F. Cheng, A. L. Hector, W. Levason, G. Reid, M. Webster and W. Zhang, *Inorg. Chem.*, 2010, 49, 752; (b) Q. Teng and H. V. Huynh, Inorg. Chem., 2014, 53, 10964; (c) H. V. Huynh, Y. Han, R. Jothibasu and J. A. Yang, Organometallics, 2009, 28, 5395; (d) K. Verlinden, H. Buhl, W. Frank and C. Ganter, Eur. J. Inorg. Chem., 2015, 2416.
- 17 L. Perrin, E. Clot, O. Eisenstein, J. Loch and R. H. Crabtree, Inorg. Chem., 2001, 40, 5806.
- 18 W. J. van Zeist, R. Visser and F. M. Bickelhaupt, Chem.-Eur. J., 2009, 15, 6112.
- 19 I. Fernandez and F. M. Bickelhaupt, Chem. Soc. Rev., 2014, 43, 4953.
- 20 I. Fernandez, Phys. Chem. Chem. Phys., 2014, 16, 7662.
- 21 D. H. Ess and K. N. Houk, J. Am. Chem. Soc., 2007, 129, 10646.
- 22 X. Hong, Y. Liang, A. K. Griffith, T. H. Lambert and K. N. Houk, Chem. Sci., 2014, 5, 471.
- 23 Y. F. Yang, Y. Liang, F. Liu and K. N. Houk, J. Am. Chem. Soc., 2016, 138, 1660.
- 24 M. von Hopffgarten and G. Frenking, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2012, 2, 43.
- 25 T. Ziegler and A. Rauk, Theor. Chim. Acta, 1977, 46, 1.
- 26 (a) A. Michalak, M. Mitoraj and T. Ziegler, J. Phys. Chem. A, 2008, 112, 1933; (b) A. Michalak, M. Mitoraj and T. Ziegler, J. Mol. Model., 2008, 14, 681.
- 27 (a) M. Mitoraj, M. Parafiniuk, M. Srebro, M. Handzlik, A. Buczek and A. Michalak, J. Mol. Model., 2011, 17, 2337; (b) M. Mitoraj, J. Phys. Chem. A, 2011, 115, 14708; (c) R. Kurczab, M. Mitoraj, A. Michalak and T. Ziegler, J. Phys. Chem. A, 2010, 114, 8581; (d) G. A. Ardizzoia, M. Bea, S. Brenna and B. Therrien, Eur. J. Inorg. Chem., 2016, 3829.
- 28 M. Hamdaoui, M. Ney, V. Sarda, L. Karmazin, C. Ailly, N. Ieffert, S. Dohm, A. Hansen, S. Grimme and J. P. Djukic, Organometallics, 2016, 35, 2207.
- 29 M. Mitoraj, A. Michalak and T. Ziegler, J. Chem. Theory Comput., 2009, 5, 962.
- 30 Y. Canac and C. Lepetit, Inorg. Chem., 2017, 56, 667.

- 31 C. Flener Lovitt, G. Frenk and G. S. Girolami, Organometallics, 2012, 31, 4122.
- 32 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Akatsuji, M. Aricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery, J. J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, T. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, Yazyev, A. J. Austin, R. Cammi, C. Pomelli, W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, O. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, Gaussian 09, revision D.01, Gaussian, Inc., Wallingford, CT, 2013.
- 33 Y. Zhao and D. G. Truhlar, Acc. Chem. Res., 2008, 41, 157.
- 34 (a) T. Clark, J. Chandrasekhar, G. W. Spitznagel and P. V. R. Schleyer, J. Comput. Chem., 1983, 4, 294; (b) V. A. Rassolov, J. A. Pople, M. A. Ratner and T. L. Windus, J. Chem. Phys., 1998, 109, 1223.
- 35 A. E. Reed, L. A. Curtiss and F. Weinhold, Chem. Rev., 1988, 88, 899.
- 36 E. J. Baerends, T. Ziegler, J. Autschbach, D. Bashford, A. Bérces, F. M. Bickelhaupt, C. Bo, P. M. Boerrigter, L. Cavallo, D. P. Chong, L. Deng, R. M. Dickson, D. E. Ellis, M. vanFaassen, L. Fan, T. H. Fischer, C. F. Guerra, M. Franchini, A. Ghysels, A. Giammona, S. J. A. van Gisbergen, A. W. Götz, J. A. Groeneveld, O. V. Gritsenko, M. Grüning, S. Gusarov, F. E. Harris, P. V. D. Hoek, C. R. Jacob, H. Jacobsen, L. Jensen, J. W. Kaminski, G. V. Kessel, F. Kootstra, A. Kovalenko, M. V. Krykunov, E. V. Lenthe, D. A. McCormack, A. Michalak, M. Mitoraj, S. M. Morton, J. Neugebauer, V. P. Nicu, L. Noodleman, Ρ. Osinga, S. Patchkovskii, M. Pavanello, P. H. T. Philipsen, D. Post, C. C. Pye, W. Ravenek, J. I. Rodríguez, P. Ros, P. R. T. Schipper, H. V. Schoot, G. Schreckenbach, J. S. Seldenthuis, M. Seth, J. G. Snijders, M. Solà, M. Swart, D. Swerhone, G. T. Velde, P. Vernooijs, Versluis, L. Visscher, O. Visser, F. Wang, T. A. Wesolowski, E. M. van Wezenbeek, G. Wiesenekker, S. K. Wolff, T. K. Woo and A. L. Yakovlev, ADF2016, Theoretical Chemistry, Vrije Universiteit, Amsterdam, The Netherlands, http://www.scm.com.
- 37 (a) W. Y. Jiang, N. J. DeYonker and A. K. Wilson, J. Chem. Theory Comput., 2012, 8, 460; (b) T. J. Lee and P. R. Taylor, Int. J. Quantum Chem., Quantum Chem. Symp., 1989, \$23, 199.