Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2025, 16, 11619

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 22nd April 2025 Accepted 27th May 2025

DOI: 10.1039/d5sc02924d

rsc li/chemical-science

Privileged metal cluster complexes†

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Clusters are a prominent subject of interest in modern chemistry research, bridging atoms and materials or catalysts. Metal coordination and metal—metal bonding are crucial in determining the chemical structures and properties of metal clusters; however, formulating a universal principle for assessing their electronic activity remains challenging. Utilizing self-developed mass spectrometry, this study examines the gasphase reactions of rhodium and platinum clusters with common ligand molecules in forming metal complexes. We find that the Rh_n^{\pm} (n=1–35) and Pt_n^{\pm} (n=3–20) clusters readily react with CO and NO to form highly-selective products of cluster complexes. This illustrates the size-dependent saturable effect of sequential coordination, which is rooted in cluster stability alongside concurrent electron delocalization and local bonding. We introduce a new electronic rule, termed the electronic " $A\bar{a}$ " rule, to understand the adaptive balance of electron delocalization and averaged local bonding in stabilizing metal clusters, whether they are coordinated with ligands or not.

Introduction

Material innovations like metal complexes, metal-organic frameworks, and ligand-protected metal clusters illustrate how bonding principles shape the development of materials for a wide range of applications. The integration of chemical bonding theory with predictive modelling enables the rational design and creation of high-performance materials, advancing frontiers in nanotechnology, catalysis, chemo sensing, and beyond. The octet rule, or 8-electron rule, elucidates how atoms interact to attain a stable electron configuration by gaining, losing, or sharing electrons via chemical bonds, including multicentre-bonds and orbital hybridization. Similarly, the effective atomic number (EAN) rule, also known as the 18electron rule, offers a crucial framework for understanding the stability of transition metal (TM) complexes.1-4 Expanding on these fundamental bonding principles, additional rules such as Lauher's rule and 9N-L rule,5,6 as well as Wade's rule derived from polyhedral skeletal electron pair theory,7,8 have been introduced to provide a rational basis for understanding the

On the other hand, early studies have firmly established the jellium model to explain the electronic shell structures15-19 in metal clusters and superatoms.20-24 On this basis, Häkkinen and colleagues25 presented a perspective on coinage metal nanoclusters by emphasizing the delocalized valence electrons (n^*) of a superatomic metal core, defined as $n^* = N - \alpha - \beta$, where N is the number of valence electrons from metal atoms, α denotes the number of electrons captured by electrophilic ligands, and β indicates the total charge carried by the cluster. This perspective elucidates the fundamental principles of electronic delocalization and the superatomic nature of metal clusters; nevertheless, it overlooks the role of local bonding.26 A more comprehensive electron-counting rule is needed, one that integrates both electron delocalization and local bonding, to better predict structures and deepen the understanding of cluster stability.27 However, the preparation of pure metal clusters and their selective conversion into cluster complexes remains challenging. The challenges stem from the stringent

stability of polynuclear TM complexes. These electronic rules successfully explained the coordination and stability of bi-, tri-, and tetra-nuclear TM complexes by assuming full occupancy of their cluster valence molecular orbitals (CVMOs). For instance, $\mathrm{Co_4(CO)_{12}}$ features a T_{d} -symmetric $\mathrm{Co_4}$ core stabilized by twelve carbonyl ligands, yielding a total of 60 valence electrons allocated across 30 orbitals. However, challenges remain when applied to high-nuclearity complexes or core–shell metal clusters. A simple example is the trigonal bipyramidal complex $\mathrm{[Ni_5(CO)_{12}]^{2-}}$ which contains 76 valence electrons but only 36 valence orbitals. This discrepancy becomes particularly evident when the metallic core exhibits multi-centre bonding or when ligands adopt bidentate or bridging coordination, which are common in ligand-protected metal clusters. 14

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[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d5sc02924d

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conditions required to form metal-metal bonds, as well as the high surface energy of subnano metals, which makes them susceptible to collision-induced dissociation and unwanted contamination.

In this work, we employed a custom-designed mass spectrometer to investigate the reactions of Rh_n^\pm (n=1–35) and Pt_n^\pm (n=3–20) clusters with CO and NO. Our findings show that both rhodium and platinum clusters exhibit high reactivity with CO and NO, forming highly-selective carbonyl and nitrosyl complexes indicative of saturable coordination. To elucidate the structural stability and bonding characteristics of the resulting dominant complexes $\mathrm{M}_n(\mathrm{CO})_m^\pm$ and $\mathrm{M}_n(\mathrm{NO})_m^\pm$ ($\mathrm{M}=\mathrm{Rh}$, Pt), we performed quantum chemistry calculations. The results reveal size-dependent ligand coordination and electron delocalization on metal clusters. Based on this, we propose an new electron rule to elucidate the adaptive balance of delocalized electrons and averaged local bonding, referred to as the electronic " $A\bar{a}$ " rule. This rule highlights the self-adaptability of delocalized and localized electrons in stabilizing metal clusters of varying sizes.

Experimental and computational methods

The instrument leading to the findings in this study was described in detail elsewhere. ²⁸ In brief, a homemade reflection time-of-flight mass spectrometer (Re-TOFMS) is combined with a laser vaporization (LaVa) source together with a compact tube reactor ($\Phi=6$ mm, L=60 mm). Rh_n⁺ clusters were produced by laser ablation of a finely polished clean Rh disk (99.95% purity, $\Phi=16$ mm). A pulsed 532 nm laser (Nd:YAG) with an average energy of 15–35 mJ per pulse at a repetition rate of 10 Hz was

used for the experiments. The pulsed laser was well focused onto the surface of the Rh target which was installed on a motor-driven sample holder capable of both translational and rotational movement. A tube nozzle ($\Phi=1.35~{\rm mm}$, $L=35~{\rm mm}$) was installed at the outlet of the waiting room where the generated clusters were cooled by supersonic expansion under He buffer gas (99.999%) with a background pressure of 1.0 MPa. Both the buffer gas and reactant gas (0.1 MPa) were controlled by a pulse valve (10 Hz) respectively with the on-time at 150–250 µs (Series 9, General Valve).

All the structures were fully relaxed using density functional theory (DFT) calculations at the PBE/def2tzvpp level of theory *via* the Gaussian 16 package.²⁹⁻³² All energy calculations were corrected by zero-point vibrations, and frequency analyses were conducted to ensure that the structures at energy minima. Chemical bonding analyses were performed using the AdNDP method at the same level of theory, which enables an electron pair to delocalize over *n* atoms of the cluster. Molecular orbitals, nucleus-independent chemical shifts (NICS)³³ and DOS were calculated by using the Multiwfn program.³⁴ The PBE0 functional at the def2tzvpp level of theory was used to simulate photoelectron spectra.³⁵ Molecular visualization was accomplished using the VMD software,³⁶ and colouring of the structures was conducted using the VESTA software.³⁷

Results and discussion

Fig. 1 presents the mass spectra of Pt_n^+ and Pt_n^- (n=3-20) clusters in the absence and presence of varying doses of NO reactants. These Pt_n^{\pm} clusters were well controlled to exhibit a comparable size distribution centred at n=11-13, and they

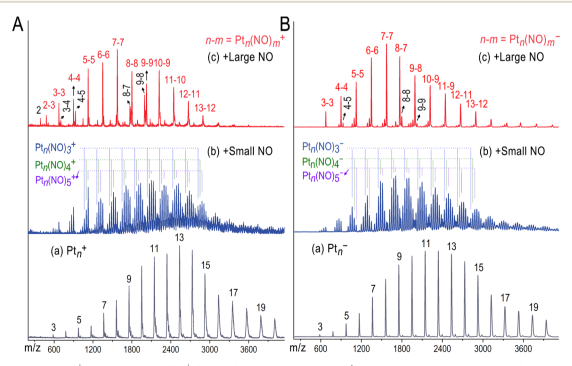


Fig. 1 NO reacting with Pt_n^{\pm} to produce $Pt_n(NO)_m^{\pm}$. (A and B) Mass spectra of the Pt_n^{+} and Pt_n^{-} clusters in the absence (a) and presence (b/c) of NO reactants (5% in He) with different doses controlled by a pulsed valve.

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exhibited comparable NO coordination reactions in the presence of small dose of NO reactants. However, when introducing sufficient NO reactants into the flow tube reactor, all the nascent Ptn + clusters react to form a highly selective coordination product for each. Specifically, the cationic Pt_n^+ clusters react to yield $Pt_3(NO)_3^+$, $Pt_4(NO)_4^+$, $Pt_5(NO)_5^+$, $Pt_6(NO)_6^+$, $Pt_{7}(NO)_{7}^{\ +}, \quad Pt_{8}(NO)_{8}^{\ +}, \quad Pt_{9}(NO)_{9}^{\ +}, \quad Pt_{10}(NO)_{9}^{\ +}, \quad Pt_{11}(NO)_{10}^{\ +},$ $Pt_{12}(NO)_{11}^+$, $Pt_{13}(NO)_{12}^+$, etc. These products remain consistent even when the size distribution of the nascent clusters changes (Fig. S1-S4, ESI†), with a few exceptions for Pt₄⁺ and Pt_{8.9}⁺ which form $Pt_4(NO)_{4,5}^+$, $Pt_8(NO)_{7,8}^+$ and $Pt_9(NO)_{8,9}^+$ respectively, likely due to the critical points of the transition from planar to threedimensional structure, and to the core-shell structure of the metallic cores. Similarly, the anionic Pt_n clusters demonstrate comparable NO reactivity, exclusively generating $Pt_n(NO)_n^-$ ($n \le n$ 7) with a strict 1:1 metal-to-ligand ratio. This stoichiometric correspondence where each Pt atom coordinates one NO molecule mirrors the monolayer protection phenomenon observed in wet-chemically synthesized nanoclusters. This 1:1 stoichiometric ratio in these $Pt_n(NO)_n$ clusters is reminiscent of the face-centered cubic (fcc) packing arrangement seen in ABtype ionic crystals for certain simple metal complexes, as well as NO adsorption on each Pt atom site of a Pt surface (Fig. S17, ESI†). Also, it bears similarity to the monolayer-protected metal clusters. Notably, the larger Pt_n^- clusters $(n \ge 8)$ adsorb one fewer NO molecule to form $Pt_n(NO)_{n-1}$, which could be associated with the formation of core-shell structures.

We have also analysed the reactions of Rh_n^{\pm} clusters with NO and CO gases to form complexes. Significantly, in the presence of sufficient CO reactant, the Rh_n⁺ clusters exhibit remarkable reaction selectivity to form a predominant cluster complex for each, seen as Rh₃(CO)₉⁺, Rh₄(CO)₁₂⁺, Rh₅(CO)₁₄⁺, Rh₆(CO)₁₆⁺, $Rh_7(CO)_{17}^+$, $Rh_8(CO)_{18}^+$, $Rh_9(CO)_{19}^+$, $Rh_{10}(CO)_{18}^+$, $Rh_{11}(CO)_{19}^+$, $Rh_{12}(CO)_{20}^{+}$, and $Rh_{13}(CO)_{22}^{+}$ (Fig. S5-S7, ESI†). Further increase of CO doses did not form complexes of a larger coordination number, indicating saturation effect of the CO coordination on the Rh_n^+ clusters. Although the observed $Rh(CO)_4^+$, Rh₃(CO)₉⁺, Rh₄(CO)₁₂⁺, and Rh₅(CO)₁₄⁺ species match previously reported carbonyl complexes,38-45 the larger clusters $Rh_n(CO)_m^+$ $(n \ge 6)$ —including $Rh_6(CO)_{16}^+$, $Rh_7(CO)_{17}^+$, $Rh_8(CO)_{18}^+$, $Rh_9(CO)_{19}^+$, and $Rh_{10}(CO)_{18}^+$ —offer new insights into electron delocalization and multicentre bonding in polynuclear complexes. Similarly, the Rh_n^+ clusters in reacting with NO also form highly-selective metal cluster complexes Rh_n(- $NO)_m^+$ (Fig. S8 and S9, ESI†). It is important to note that in these cluster complexes, not all metal atoms achieve 18-electron coordination. Instead, the entire system reflects a dynamic coordination equilibrium, which is accompanied by electron delocalization.

Using density functional theory (DFT) calculations, we determined the lowest energy structures of the $Pt_n(NO)_m^{\pm}$ clusters. For small $\operatorname{Pt}_n^{\pm}$ clusters (n < 8), favourable coordination of all metal atoms is achieved, allowing partial electron delocalization. We focused particularly on the Pt₈(NO)₇ cluster within a critical size. To validate its structure, we conducted photoelectron spectroscopy (PES) experiments using both 266 nm and 355 nm laser ionization, as shown in Fig. 2a. The simulated PES spectrum of the lowest energy structure (isomer a) aligns well with the experimental data, verifying the coreshell structure of $Pt_8(NO)_7$. In contrast, the other low-lying isomers (isomers b and e) exhibit significantly different vertical ionization energy values (X') and PES profiles. This supports the prediction that larger Pt_n^{\pm} clusters $(n \ge 8)$ adsorb one less NO molecule to form $Pt_n(NO)_{n-1}^{\pm}$ products simply due to their core-shell structures.

Fig. 2b illustrates the chemical bonding analysis for using adaptive natural density partitioning (AdNDP).46 The AdNDP analysis shows that each Pt atom (5d⁹6s¹) in the cluster possesses five d-type lone pairs (d¹⁰), indicating that the unpaired π^* electron of the NO molecule is coupled with the 5d electrons of the Pt atom within symmetrymatching regulation. This behaviour is consistent across other $Pt_n(NO)_m^{\pm}$ cluster complexes, although the extent of delocalization depends on the number of Pt atoms and NO molecules. Notably, the $Pt_8(NO)_7$ cluster exhibit two types of 8c-2e bonds, displaying distinct superatomic characteristics of S and P orbitals. Furthermore, a multi-centre bonding (22c-2e) pattern

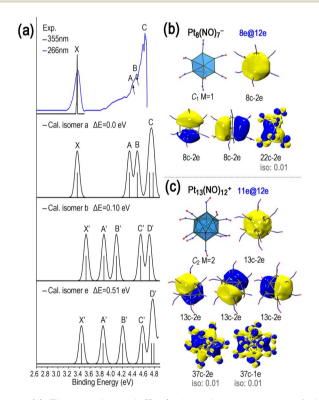


Fig. 2 (a) The experimental (Exp.) photoelectron spectra of the Pt₈(NO)₇ clusters, obtained using 355 nm (black curve) and 266 nm (blue curve) laser ionization, in comparison with the calculated (Cal.) spectra of low-lying isomers. The peaks marked with X, A, B and C are assigned to the electronic transitions involved in the transformation process of $Pt_8(NO)_7$ anion to the electron-detached $Pt_8(NO)_7$ neutral. The geometry and AdNDP bonding analysis of the (b) $Pt_8(NO)_7$ and (c) Pt₁₃(NO)₁₂ clusters respectively. Also labelled are symmetries, spin multiplicities (M), and the electron counting (xe@ne) by considering both delocalized electrons and average number of locally bonding electrons for surface metal atoms. The polyhedral patterns of the metallic cores are shown to guide eye. The isovalues for the patterns of AdNDP analyses are set to 0.02 unless otherwise stated.

is observed, indicating weak global electron delocalization across the entire ${\rm Pt_8(NO)_7}^-$ cluster. Similarly, we conducted AdNDP analysis for the ${\rm Pt_{13}(NO)_{12}}^\pm$ cluster, whose metallic core (${\rm Pt_{13}}$) has been previously shown to adopt a core–shell structure. 47,48 Notably, the ${\rm Pt_{13}(NO)_{12}}^+$ cluster also exhibit two types of 13c–2e bonds (Fig. 2c), reflecting superatomic S and P orbitals, along with weak global delocalization through a 37c–2e bond spanning the entire cluster.

The geometric structures and AdNDP analyses of the small $Pt_n(NO)_m^{\pm}$ clusters are presented in Fig. 3a and b (more details in Fig. S10–S39, ESI†). Among them, the lowest energy structures of $Pt_3(NO)_3^{\pm}$, $Pt_4(NO)_4^{\pm}$, $Pt_5(NO)_5^{\pm}$, $Pt_6(NO)_6^{\pm}$, and $Pt_7(NO)_7^{\pm}$ exhibit either planar or polyhedral geometry for their metallic cores, with each surface platinum atom bonded to a single terminal nitrosyl ligand. Interestingly, these $Pt_n(NO)_m^{\pm}$ clusters (n=3–7) all demonstrate electron localization and delocalization, showing diverse multi-centre bonds such as 3c–2e, 4c–2e, 5c–2e, *etc.*, as well as weak global delocalization involving all-atom multicentre bonds. The delocalized electrons not only form multicentre bonds but also contribute to superatomic features, resulting in balanced coordination and

enhanced stability. What is more interesting is that each Pt atom on the surface of these $Pt_n(NO)_m^{\pm}$ clusters achieve a 12-electron coordination, which could be favoured for the six $5d \cdot 6s$ valence orbitals of Pt. This is reasonable as the 6p orbital is significantly higher in energy, with a substantial energy gap (>6 eV) between the 6 s and 6p orbital levels. The electron delocalization and localization in these clusters were further verified by analyses on density of states (DOS) and canonical molecular orbitals (Fig. S40–S48, ESI†).

For comparison, Fig. 3c illustrates the geometries and AdNDP analyses of the $Rh_n(NO)_m^+$ (n=1-6) clusters. These $Rh_n(NO)_m^+$ clusters exhibit a variety of isomers (Fig. S49–S53, ESI†). Only the most stable ones are selected for bonding analysis (Fig. S54–S60, ESI†). The NO molecule contains an unpaired electron in its antibonding π^* orbital, and this unpaired single electron can engage with the electrons of the Rh atom, facilitating a favourable d^{10} electron configuration. Among the $Rh_n(NO)_m^+$ clusters, the larger ones ($n \ge 6$) display pronounced electron delocalization, reflecting their superatomic nature; in comparison, the smaller clusters (n=1-5) exhibit π -electron delocalization,⁴⁹ with metal–metal bond

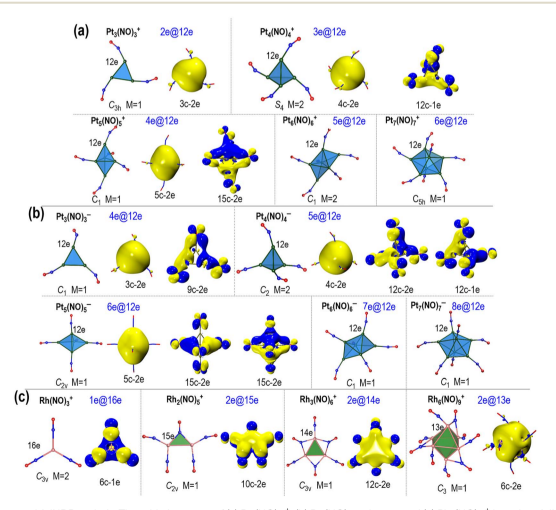


Fig. 3 Geometry and AdNDP analysis. The orbital patterns of (a) $Pt_n(NO)_m^+$, (b) $Pt_n(NO)_m^-$ clusters and (c) $Pt_n(NO)_m^+$ based on AdNDP analysis. Also labelled are symmetries, spin multiplicities (M), and the electron counting (xe@ne) according to the " $A\bar{a}$ " rule which takes into account the itinerant electrons and the average number of localized electrons at surface atoms. The polyhedral patterns of the metallic cores are shown to guide eye. The isovalues for the patterns of AdNDP analyses are set to 0.02.

lengths extended due to ligand interactions. Specifically, $\mathrm{Rh(NO)_3}^+$ features a 6c–1e multi-centre bond associated with π delocalization, and there are 16 electrons (16e) contributed by the Rh atom and three NO ligands. The delocalized orbitals in these $\mathrm{Rh}_n(\mathrm{NO})_m^+$ primarily consist of antibonding π^* orbitals of NO, resulting in full occupation of the five d-type lone pairs.

Comprehensive DFT calculations were also performed to analyse the structures and bonding characteristics of the Rh_n(-CO)_m⁺ (n = 1-7) clusters (Fig. S61–S71, ESI†). Because of the primary distinction between carbonyl and nitrosyl complexes in the bonding and antibonding orbitals of the CO and NO molecules (Fig. S72, ESI†), all Rh_n(CO)_m⁺ species exhibit four d-type lone pairs, which differs from that of the Rh_n(NO)_m⁺ clusters. The lowest energy structures of Rh(CO)₄⁺, Rh₂(CO)₇⁺, Rh₃(CO)₉⁺, Rh₄(CO)₁₂⁺, Rh₅(CO)₁₄⁺, Rh₆(CO)₁₆⁺, Rh₇(CO)₁₇⁺, and Rh₈(CO)₁₈⁺ reveal that carbonyl ligands bind to Rh atoms through two distinct modes: bridging and terminal. Previous studies have unveiled the stability of Rh(CO)₄⁺ as a stable 16e Rh-complex with a planar tetragonal configuration. ^{50,51} This also aligns with the aforementioned 16e-stability of Rh(NO)₃⁺.

For these polynuclear complexes, the cluster valence electrons are distributed across the entire cluster, enabling the formation of multicentre bonds. For example, $Rh_3(CO)_9^+$ features a 3c–2e bond, with each Rh atom attaining 14 electrons through covalent bonds and coordination interaction. Likewise, $Rh_2(CO)_7^+$ exhibits a 6c–1e bond derived from the d orbitals of Rh atoms and a π^* antibonding orbital of CO. The $Rh_4(CO)_{12}^+$ and $Rh_5(CO)_{14}^+$

species also demonstrate multi-centre bonding defined by orbital hybridization; while $Rh_6(CO)_{16}^+$ adopts an octahedral Rh_6 core, with multicentre bonds reflecting electron delocalization and aromaticity (Fig. S73 and S74, ESI†). We have also studied the anionic $Rh_n(CO)_m^-$ systems (Fig. S75–S81, ESI†), which find similar structures and AdNDP patterns pertaining to multicenterbonding and electron delocalization especially for $n \ge 6$. Additionally, the DOSs and orbital analyses align with the results of the AdNDP analysis (Figs. S82–S84, ESI†).

The formation of predominant carbonyl and nitrosyl complexes motivated us to further study the high selectivity of NO and CO coordination with Pt and Rh clusters. Among them, nitrosyl complexes exhibit significant electron delocalization due to the involvement of half-filled antibonding π^* orbitals along the molecular axis. In contrast, the π^* orbitals of CO remain vacant, facilitating backdonation from the metal to the CO ligands and promoting favourable local coordination.¹⁰ Considering that these cluster complexes exhibit delocalized electrons and distinct local coordination preferences, we propose the electronic " $A@\bar{a}$ " or " $A\bar{a}$ " rule. This rule integrates adaptive delocalized electrons (A) and localized electrons on average (\bar{a}) to assess the stability of metal clusters. In this " $A\bar{a}$ " rule, the total number of valence electrons (N_{CVE}) for a metal cluster complex is determined by the sum of delocalized and localized valence electrons, i.e.,

$$N_{\text{CVE}} = A + \bar{a} \cdot n' + c \tag{1}$$

Table 1 A list showing the different electronic counting rules of carbonyl and nitrosyl complexes^a

$Rh_n(CO)_m^+$	CVMO	9N-L	CVE	$Aar{a}$	$Rh_n(NO)_m^+$	CVMO	9N-L	CVE	$Aar{a}$
Rh(CO) ₄ ⁺	9	9	16e	16e	Rh(NO) ₃ ⁺	9	9	17e	1e@16e
$Rh_2(CO)_7^+$	17	17	31e	1e@15e	$Rh_2(NO)_5^+$	17	17	32e	2e@15e
$Rh_3(CO)_9^+$	24	24	44e	2e@14e	$Rh_3(NO)_6^+$	24	24	44e	2e@14e
$Rh_4(CO)_{12}^+$	30	30	59e	3e@14e	$Rh_4(NO)_8^+$	30	30	59e	3e@14e
$Rh_{5}(CO)_{14}^{+}$	36	36	72e	4e@14e	$Rh_5(NO)_9^+$	36	36	71e	3e@14e
Rh ₆ (CO) ₁₆ ⁺	43	42	85e	5e@14e	$Rh_6(NO)_9^+$	43	42	80e	2e@13e
$Pt_n(NO)_m^+$	CVMO	9 N- L	CVE	Aā	$\operatorname{Pt}_n(\operatorname{NO})_m^-$	CVMO	9N-L	CVE	Αā
Pt ₃ (NO) ₃ +	24	24	38e	2e@12e	$Pt_3(NO)_3^-$	24	24	40e	4e@12e
$Pt_4(NO)_4^+$	30	30	51e	3e@12e	$Pt_4(NO)_4^-$	30	30	53e	5e@12e
$Pt_5(NO)_5^+$	36	36	64e	4e@12e	$Pt_5(NO)_5^-$	36	36	66e	6e@12e
$Pt_6(NO)_6^+$	43	42	77e	5e@12e	$Pt_6(NO)_6^-$	43	42	79e	7e@12e
$Pt_7(NO)_7^+$	49	48	90e	6e@12e	$Pt_7(NO)_7^-$	49	48	92e	8e@12e
$Pt_8(NO)_8^+$	_	57	103e	8e@12e	$Pt_8(NO)_7^-$	_	57	102e	8e@12e
$Pt_{13}(NO)_{12}^{+}$	_	82	165e	11e@12e	$\mathrm{Pt}_{13}(\mathrm{NO})_{12}^{-}$	_	82	167e	13e@12e
$Ru_n(CO)_m^+$	CVMO	9N-L	CVE	Αā	$\operatorname{Co}_n(\operatorname{CO})_m^+$	CVMO	9N-L	CVE	$A\bar{a}$
Ru(CO) ₅ ⁺	9	9	17e	17e	Co(CO) ₅ ⁺	9	9	18e	18e
$Ru_2(CO)_9^+$	17	17	33e	2e@16e	$Co_2(CO)_8^+$	17	17	33e	1e@16e
$Ru_3(CO)_{12}^+$	24	24	47e	5e@14e	$Co_3(CO)_{10}^+$	24	24	46e	2e@15e
$Ru_4(CO)_{14}^{+}$	30	30	59e	7e@13e	$Co_4(CO)_{12}^+$	30	30	59e	3e@14e
$Ru_5(CO)_{16}^+$	36	36	71e	9e@12e	$Co_5(CO)_{14}^{+}$	36	36	72e	4e@14e
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^a CVE is the total number of cluster valence electrons; CVMO is the number of cluster valence molecular orbitals; 9N-L denotes the estimated quantity of cluster valence molecular orbitals, excluding the count of relative antibonding orbitals associated with the polyhedral skeleton edges of metal–metal bonds; " $A\bar{a}$ " values relate to the delocalized electrons (A) and averaged local electrons (\bar{a}) for surface metal atoms.

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here A denotes the delocalized electrons, encompassing π delocalization and multicentre bonding electrons within superatomic states. The variable n' signifies the number of metal atoms involved in surface coordination, excluding the core metal atoms. Meanwhile; \bar{a} indicates the average number of localized valence electrons including lone pairs and bonding electrons, which is typically less than 18 for transition metals. The term c accounts for the charge and localized valence electrons of the core metal atoms. Since the values of N_{CVE} and n'can be determined from the cluster formula, we can predict the \bar{a} value if A is known, estimate A by subtracting the localized electrons, or determine n' once A and \bar{a} are established (to infer a core-shell structure if $n \neq n'$). Notably, both A and \bar{a} can be determined by the AdNDP analysis. For instance, Rh2(CO)7+ displays a 6c-1e delocalized bond, with each Rh atom possessing 15 electrons, yielding an electronic formula of 1e@15e. Similarly, Rh₃(CO)₉⁺ corresponds to 2e@14e; Rh₄(CO)₁₂⁺ to 3e@14e, $Rh_5(CO)_{14}^+$ to 4e@14e, and $Rh_6(CO)_{16}^+$ to 5e@14e, as detailed in Table 1. This electronic " $A\bar{a}$ " rule is also applicable to the $Rh_n(NO)_m^{\pm}$ and $Pt_n(NO)_m^{\pm}$ systems. Notably, all the observed prominent $Pt_n(NO)_m^+$ and $Pt_n(NO)_m^-$ products demonstrate a preference of the xe@12e configuration, highlighting the importance of the "Aā" rule in explaining cluster stability and optimal coordination. It is worth mentioning that the DFT calculations in this study were performed on smaller systems, but the "Aā" rule is applicable and advantageous for

According to the " $A\bar{a}$ " rule, we have examined the electronic configurations and structure stability of the $Co_n(CO)_m^+$ and $Ru_n(CO)_m^+$ clusters in previous investigations, ^{52–54} as well as a recently reported $Fe_6C(CO)_{16}$ cluster, ⁵⁵ all of which adhere to the xe@ne regulation (Fig. S85–S89, ESI†). Considering the number of ligands that act as monodentate (μ_1 , 2e or 3e donor, including the single-site, bridge-site and hollow-site modes) and bidentate (μ_2 , e.g., 4e donor) in a M_nX_m (M, metal; X, ligand) cluster to be represented by x and y respectively, the relationship for stable cluster complexes (e.g., CO) can be expressed as:

understanding larger metal clusters.

$$x + y = m;$$
 $\sum_{i=1}^{n} (2x_i + 4y_i) = n'(\bar{a} - v)$ (2)

here, ν denotes the valence electrons of the metal atom, while \bar{a} can be derived from eqn (1). This framework allows for the quantification of μ_1 -CO ligands, aiding in structure prediction. The electronic " $A\bar{a}$ " rule is also believed to be applicable for evaluating structural stability of metal cluster complexes involving phosphines, olefins, alkyls, and other types of ligands, as well as small molecules having both multicentre bonds and hybrid orbitals (e.g., O_3 within 3c–4e and sp^2 hybridization).

Conclusions

To summarize, we report an investigation in exploring and understanding the privileged metal cluster complexes. We systematically studied the gas-phase reactivity of Rh_n^\pm (n=1–35) and Pt_n^\pm (n=3–20) clusters with CO and NO. Interestingly, these metal clusters exhibit remarkable selectivity in the NO- and CO-

coordination reactions when exposed to sufficient gas reactants, leading to the formation of saturable carbonyl and nitrosyl complexes. Typical examples include a series of $Pt_n(NO)_n^{\pm}$ (n=3-8) and $Pt_n(NO)_{n-1}^{\pm}$ (n=8-14) products, as well as several Rh cluster complexes such as $Rh_4(CO)_{12}^{+}$, $Rh_4(NO)_8^{+}$; $Rh_5(CO)_{14}^{+}$, $Rh_5(NO)_9^{+}$; $Rh_6(CO)_{16}^{+}$, $Rh_6(NO)_9^{+}$; $Rh_7(CO)_{17}^{+}$, and $Rh_7(NO)_{10}^{+}$. Notably, the NO complexes favour significant electron delocalization pertaining to the involvement of half-filled antibonding π^* orbitals along the NO molecule axis. In contrast, the π^* orbitals of CO are vacant, which facilitates metal-to-ligand backdonation and reinforces local coordination. We introduce an electronic " $A\bar{a}$ " rule to evaluate the structural stability of metal cluster complexes, clarifying adaptive balance of delocalized electrons and averaged local bonding.

Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

The manuscript was written through contributions of all authors.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We thank Prof. Gernot Frenking in Philipps-Universität Marburg, Prof. Jinlong Yang in University of Science and Technology of China, Prof. Mingfei Zhou and GuanJun Wang in Fudan University, for their helpful discussions. This work was supported by the National Natural Science Fundation of China (Grant No. 92261113 and 22272180), CAS Project for Young Scientists in Basic Research (Grant No. YSBR-050), CAS Key Research Program of Frontier Sciences (QYZDBSSW-SLH024). SNK was supported by a grant from US Department of Energy (DOE) under the award DE-SC0006420.

Notes and references

- 1 G. Frenking and N. Fröhlich, Chem. Rev., 2000, 100, 717-774.
- 2 L. Andrews and A. Citra, Chem. Rev., 2002, 102, 885-912.
- 3 X. Jiang, Z. Xiao, W. Zhong and X. Liu, *Coord. Chem. Rev.*, 2021, 429, 213634.
- 4 A. Fielicke, Chem. Soc. Rev., 2023, 52, 3778-3841.
- 5 J. W. Lauher, J. Am. Chem. Soc., 1978, 100, 5305-5315.
- 6 A. Tang and S. Li, Chin. Sci. Bull., 1982, 28, 25-27.
- 7 F. Lips and S. Dehnen, *Angew. Chem., Int. Ed.*, 2011, **50**, 955–959.
- 8 D. M. P. Mingos, 50th Anniversary of Electron Counting Paradigms for Polyhedral Molecules: Bonding in Clusters, *Intermetallics and Intermetalloids*, University of Oxford, Oxford, UK, 2021.

9 R. Bau, M. H. Drabnis, L. Garlaschelli, W. T. Klooster,

- Z. W. Xie, T. F. Koetzle and S. Martinengo, *Science*, 1997, 275, 1099–1102.
- 10 X. Wu, L. Zhao, J. Jin, S. Pan, W. Li, X. Jin, G. Wang, M. Zhou and G. Frenking, *Science*, 2018, **361**, 912–916.
- 11 E. Kapiloff and K. M. Ervin, *J. Phys. Chem. A*, 1997, **101**, 8460–8469.
- 12 E. K. Parks, K. P. Kerns and S. J. Riley, *J. Chem. Phys.*, 2000, **112**, 3384–3393.
- 13 Z. Lin, Chin. J. Struct. Chem., 2024, 43, 100254.

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- 14 G. Pacchioni and N. Roesch, *Inorg. Chem.*, 1990, 29, 2901–2908.
- I. Katakuse, T. Ichihara, Y. Fujita, T. Matsuo, T. Sakurai and H. Matsuda, *Int. J. Mass Spectrom. Ion Processes*, 1985, 67, 229–236.
- 16 W. Ekardt, Phys. Rev. Lett., 1984, 52, 1925-1928.
- 17 W. D. Knight, K. Clemenger, W. A. De Heer, W. A. Saunders, M. Y. Chou and M. L. Cohen, *Phys. Rev. Lett.*, 1984, **52**, 2141– 2143.
- 18 W. A. De Heer, Rev. Mod. Phys., 1993, 65, 611-676.
- 19 M. Brack, Rev. Mod. Phys., 1993, 65, 677-732.
- 20 X. Zhang, Y. Wang, H. Wang, A. Lim, G. Gantefoer, K. H. Bowen, J. U. Reveles and S. N. Khanna, *J. Am. Chem. Soc.*, 2013, 135, 4856–4861.
- 21 Z. Luo and A. W. Castleman Jr, Acc. Chem. Res., 2014, 47, 2931–2940.
- 22 A. C. Reber and S. N. Khanna, *Acc. Chem. Res.*, 2017, **50**, 255–263.
- 23 E. A. Doud, A. Voevodin, T. J. Hochuli, A. M. Champsaur, C. Nuckolls and X. Roy, *Nat. Rev. Mater.*, 2020, 5, 371–387.
- 24 P. Jena, H. Fang and Q. Sun, in *Superatoms: Principles, Synthesis and Applications*, 2021, pp. 15–51.
- 25 M. Walter, J. Akola, O. Lopez-Acevedo, P. D. Jadzinsky, G. Calero, C. J. Ackerson, R. L. Whetten, H. Groenbeck and H. Hakkinen, *Proc. Natl. Acad. Sci. U. S. A.*, 2008, 105, 9157–9162.
- 26 Z. Luo, A. C. Reber, M. Jia, W. H. Blades, S. N. Khanna and A. W. Castleman Jr, *Chem. Sci.*, 2016, 7, 3067–3074.
- 27 Z. Luo, A. W. Castleman Jr and S. N. Khanna, *Chem. Rev.*, 2016, **116**, 14456–14492.
- 28 H. Zhang, H. Wu, Y. Jia, L. Geng, Z. Luo, H. Fu and J. Yao, *Rev. Sci. Instrum.*, 2019, **90**, 073101.
- 29 J. Paier, R. Hirschl, M. Marsman and G. Kresse, *J. Chem. Phys.*, 2005, **122**, 234102.
- 30 W. Florian and A. Reinhart, *Phys. Chem. Chem. Phys.*, 2005, 7, 3297–3305.
- 31 F. Weigend, Phys. Chem. Chem. Phys., 2006, 8, 1057-1065.
- 32 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai,

- T. Vreven, K. Throssell, J. A. Montgomery Jr., J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman and D. J. Fox, *Gaussian 16 Rev. B.01*, Gaussian, Inc., Wallingford, CT, 2016.
- 33 Z. F. Chen, C. S. Wannere, C. Corminboeuf, R. Puchta and P. V. Schleyer, *Chem. Rev.*, 2005, **105**, 3842–3888.
- 34 T. Lu and F. Chen, J. Comput. Chem., 2012, 33, 580-592.
- 35 C. Adamo and V. Barone, J. Chem. Phys., 1999, 110, 6158-6170.
- 36 W. Humphrey, A. Dalke and K. Schulten, *J. Mol. Graph.*, 1996, 14, 33–38.
- 37 K. Momma and F. Izumi, *J. Appl. Crystallogr.*, 2011, **44**, 1272–
- 38 C. Chi, J. Cui, Z. H. Li, X. Xing, G. Wang and M. Zhou, *Chem. Sci.*, 2012, 3, 1698–1706.
- 39 G. Wang, J. Cui, C. Chi, X. Zhou, Z. H. Li, X. Xing and M. Zhou, *Chem. Sci.*, 2012, 3, 3272–3279.
- 40 J. Cui, X. Zhou, G. Wang, C. Chi, Z. H. Li and M. Zhou, *J. Phys. Chem. A*, 2014, **118**, 2719–2727.
- 41 P. Ferrari, J. Vanbuel, N. M. Tam, M. T. Nguyen, S. Gewinner, W. Schollkopf, A. Fielicke and E. Janssens, *Chem.–Eur. J.*, 2017, 23, 4120–4127.
- 42 A. Fielicke, G. Von Helden, G. Meijer, B. Simard, S. Dénommée and D. M. Rayner, *J. Am. Chem. Soc.*, 2003, 125, 11184–11185.
- 43 A. Fielicke, G. Von Helden, G. Meijer, D. B. Pedersen, B. Simard and D. M. Rayner, *J. Am. Chem. Soc.*, 2005, 127, 8416–8423.
- 44 I. Swart, F. M. F. D. Groot, B. M. Weckhuysen, D. M. Rayner, G. Meijer and A. Fielicke, *J. Am. Chem. Soc.*, 2008, **130**, 2126– 2127.
- 45 Q. Xu, S. Inoue, Y. Souma and H. Nakatani, *J. Organomet. Chem.*, 2000, **606**, 147–150.
- 46 D. Y. Zubarev and A. I. Boldyrev, *Phys. Chem. Chem. Phys.*, 2008, **10**, 5207–5217.
- 47 N. Watari and S. Ohnishi, *J. Chem. Phys.*, 1997, **106**, 7531–7540.
- 48 T. Imaoka, H. Kitazawa, W.-J. Chun, S. Omura, K. Albrecht and K. Yamamoto, *J. Am. Chem. Soc.*, 2013, **135**, 13089–13095.
- 49 Y. Gao, X. Lei, R. Cheng, S. Lin and Z. Luo, *Phys. Chem. Chem. Phys.*, 2024, **26**, 28019–28024.
- 50 A. D. Brathwaite, H. L. Abbott-Lyon and M. A. Duncan, *J. Phys. Chem. A*, 2016, **120**, 7659–7670.
- 51 R. Crabtree, *The Organometallic Chemistry of the Transition Metals*, John Wiely & Sons Inc., Hoboken, New Jersey, 2001.
- 52 S. Fedrigo, T. L. Haslett and M. Moskovits, *Chem. Phys. Lett.*, 1999, **307**, 333–338.
- 53 K. P. Kerns, E. K. Parks and S. J. Riley, *J. Chem. Phys.*, 2000, **112**, 3394–3407.
- 54 D. Yubero Valdivielso, C. Kerpal, W. Schollkopf, G. Meijer and A. Fielicke, *Dalton Trans.*, 2023, **52**, 9929–9939.
- 55 C. R. Cobb, R. K. Ngo, E. J. Dick, V. M. Lynch and M. J. Rose, Chem. Sci., 2024, 15, 11455–11471.