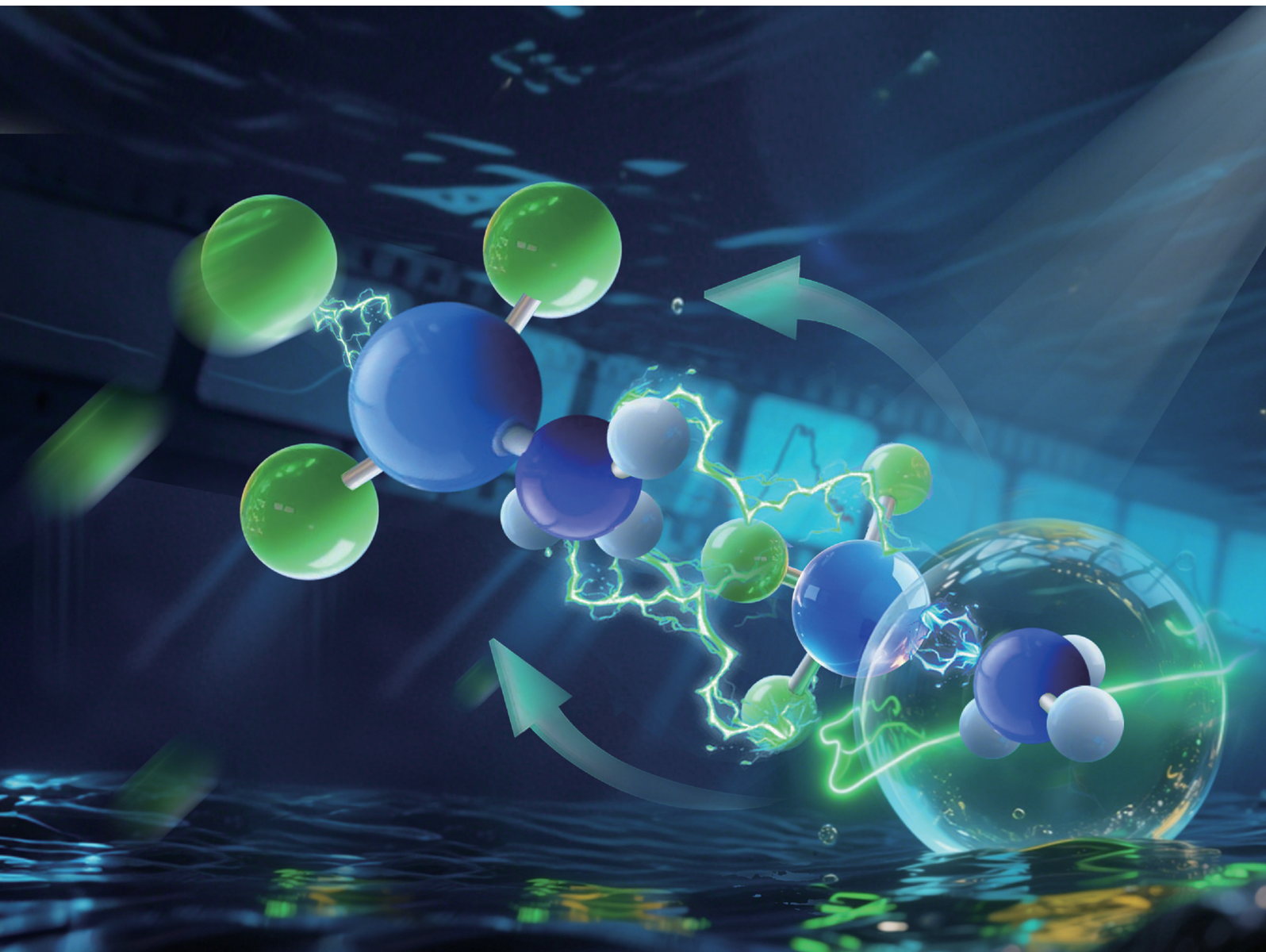


PCCP

Physical Chemistry Chemical Physics

rsc.li/pccp

25
YEARS
ANNIVERSARY



ISSN 1463-9076

PAPER

Xue-Bin Wang, Zhenrong Sun, Yan Yang *et al.*
Exploring the *trans* effect of the NH_3 ligand in platinum
halide complexes $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($\text{X} = \text{Cl}, \text{Br}, \text{I}$) using cryogenic
photoelectron spectroscopy and quantum chemical
calculations


 Cite this: *Phys. Chem. Chem. Phys.*, 2025, 27, 12657

Exploring the *trans* effect of the NH₃ ligand in platinum halide complexes Pt(NH₃)ClX₂[−] (X = Cl, Br, I) using cryogenic photoelectron spectroscopy and quantum chemical calculations†

 Qixu Zhao,[‡] Jian Zhang,[‡] Xueying Li,^a Peng Tang,^a Fan Yang,^a Junyang Ma,^a Zhubin Hu,^a Haitao Sun,[‡] Xue-Bin Wang,[‡] Zhenrong Sun^{*ad} and Yan Yang[‡]

Cryogenic anion photoelectron spectroscopy, combined with quantum chemical calculations, was employed to investigate PtClX₂[−], Pt(NH₃)ClX₂[−] (X = Cl, Br, I), and their isomers. Photoelectron spectra recorded at 193 nm, supported by B3LYP-D3(BJ)/aug-cc-pVTZ(-pp) calculations, provided adiabatic (ADEs) and vertical detachment energies (VDEs) with excellent agreement between experimental and theoretical results. Coordination of the NH₃ ligand to PtClX₂[−] reduced the electron binding energy, and substantially elongated *trans* Pt–halogen bonds. Further computational analyses, including natural population analysis (NPA), frontier molecular orbital (FMO) studies, and dissociation energy calculations, all revealed significant changes in charge distributions and stability of *trans* halogen ligands. The results demonstrated that NH₃ coordination notably elevated FMO orbital energies, with the extent of this elevation correlating strongly with the *trans* halogen's orbital contributions. These findings provide new insights into ligand-induced electronic and structural transformation in platinum halide systems and establish a theoretical foundation for understanding the underlying molecular mechanisms that dictate the activities of platinum-based anticancer drugs.

 Received 1st March 2025,
 Accepted 1st May 2025

DOI: 10.1039/d5cp00807g

rsc.li/pccp

Introduction

The concept of the '*trans* effect' was first introduced by Chernetayev in 1926 during his study of Pt(II) square planar complexes, where he observed that coordinated groups in the *trans*

position influence each other within a complex. Today, the *trans* effect is a well-established phenomenon in ligand substitution reactions of square planar and octahedral transition metal complexes.^{1–3} It is defined as the influence of a coordinated group on the substitution rate of the ligand *trans* to itself, affecting the bond length and stability of the *trans* ligand.³ This phenomenon has found wide applications in fields such as medicinal chemistry and catalysis. For instance, the *trans* effect plays a pivotal role in the mechanisms of platinum-based anticancer drugs (e.g., cisplatin and oxaliplatin)^{4–6} and catalytic processes, such as electrocatalytic CO₂ reduction using Ru-based catalysts.^{7,8} In the case of platinum-based anticancer drugs, where the *trans* effect governs ligand substitution during interaction with DNA bases, the *cis*-configuration ensures that the opposite ligands of leaving groups are nitrogen-containing groups, such as NH₃. The NH₃ ligand exerts the *trans* effect, facilitates ligand dissociation and enhances the drug's activity.

A substantial body of research has been conducted on platinum halide complexes with various types of ligands, such as the Zeise's salt⁹ anions [PtX₃(C₂H₄)][−] (X = Cl, Br, I), among others. Utilizing high-resolution anion photoelectron spectroscopy combined with quantum chemical calculations, Hou *et al.*^{10,11} found that both the electron binding energy and the

^a State Key Laboratory of Precision Spectroscopy, and School of Physics and Electron Sciences, East China Normal University, Shanghai 200241, China.

E-mail: yiyang@lps.ecnu.edu.cn

^b College of Chemistry & Chemical Engineering, Donghua University, Shanghai 201620, China

^c Physical Sciences Division, Pacific Northwest National Laboratory, Richland, WA 99352, USA. E-mail: xuebin.wang@pnnl.gov

^d Collaborative Innovation Center of Extreme Optics, Shanxi University, Taiyuan, Shanxi 030006, China. E-mail: zrsun@phy.ecnu.edu.cn

 † Electronic supplementary information (ESI) available: Initial geometries and optimized geometries of PtClX₂[−], Pt(NH₃)ClX₂[−] (X = Cl, Br, I) anions and the corresponding neutrals; lowest energy structures of PtClX₂, Pt(NH₃)ClX₂ (X = Cl, Br, I), and their isomers; the HOMO energy difference upon NH₃ coordination and orbital rearrangement between PtClX₂[−] and Pt(NH₃)ClX₂[−] (X = Br, I) anions; the LUMO plot of Pt(NH₃)Cl₃[−]; the ionic proportion of the chemical bonds Pt–Cl, Pt–Br, and Pt–I of PtClX₂[−], Pt(NH₃)ClX₂[−] (X = Cl, Br, I); the dissociation energies of different dissociation channels of PtClX₂[−], Pt(NH₃)ClX₂[−] and Pt(C₂H₄)ClX₂[−] (X = Cl, Br, I); and the structures and bonding energies of Pt(C₂H₄)ClX₂[−] (X = Cl, Br, I). See DOI: <https://doi.org/10.1039/d5cp00807g>

‡ These two authors contributed equally to this work.

interaction energy between the C_2H_4 ligand and the platinum halide decrease as the halide size increases. They also demonstrated how variations in ligand type influence the evolution of the electronic structure and molecular configuration in Zeise's family complexes. A wealth of research reporting on the physical and chemical properties of platinum halide complexes containing NH_3 ligands has also been well documented. For example, Zhang *et al.*^{12,13} focused on the vibrational properties of the NH_3 ligand in cisplatin. Warneke and Rohdenburg *et al.*^{14,15} reported the electron-induced decomposition of cisplatin, demonstrating the NH_3 ligand's role as a reducing agent to promote Cl dissociation. Pinter *et al.*¹⁶ studied the *trans* effect in ammine substitution reactions of *trans*- $TpPtCl_2NH_3$ ($T = NH_3, PH_3, CO, \text{ and } C_2H_4$) using energy decomposition and chemical valence analyses. Despite these efforts, the comprehensive impact of the NH_3 ligand's *trans* effect on the molecular configuration, electronic structure, and stability of halogenated platinum complexes has remained insufficiently explored to date. In addition, gas-phase photoelectron spectroscopy (PES) studies on halogenated platinum anions containing NH_3 ligands are notably lacking.

In this work, we employ cryogenic anion cluster photoelectron spectroscopy (CRACPES) and quantum chemical calculations to systematically investigate $PtClX_2^-$ ($X = Cl, Br, I$), $Pt(NH_3)ClX_2^-$ ($X = Cl, Br, I$), and their isomers. These anions provide an ideal model to explore the microscopic effects of the NH_3 ligand's *trans* effect in platinum halide anion complexes. The use of CRACPES enables precise measurements of electronic structures, stabilities, and energetics in the gas phase. The target anions, which cannot be prepared by electrospray ionization (ESI⁺) source directly, are produced through collision-induced dissociation (CID) in a two-stage ion funnel system.

Our results demonstrate that NH_3 , as an electron-donating ligand, upon binding to $PtClX_2^-$, significantly decreases the electron binding energy (eBE) and elongates the bond lengths between the halogen ligands and the platinum atom, particularly for the *trans* halogens. Furthermore, natural population analysis (NPA) and frontier molecular orbital (FMO) calculations reveal that the NH_3 ligand transports a substantial amount of its electron density along the NH_3 -Pt-*trans*-halogen axis and weakens the stability of the *trans* halogen ligands while elevating FMO energy levels. These findings provide critical insights into the fundamental role of the NH_3 ligand in platinum-based anticancer drugs and offer molecular-level guidance for the design of new metal-ligand anticancer therapeutics.

Methods

CRACPES experiments

The experiments were conducted using the CRACPES system, which has been described in detail elsewhere.^{17,18} $KPt(NH_3)Cl_3$ (Shanghai Boer Chemical Reagents) was dissolved in a water/acetonitrile (CH_3CN) mixed solvent (1:4) to prepare a 2 mM sample solution. To generate $Pt(NH_3)ClX_2^-$ ($X = Br, I$), potassium halides (KBr and KI) were added to aliquots of this solution, facilitating ligand substitution reactions. $PtClX_2^-$

($X = Cl, Br, I$) anions were subsequently prepared *via* collision-induced dissociation (CID) of parent $Pt(NH_3)ClX_2^-$ ($X = Cl, Br, I$) anions. The target anions were cooled to 13 K in a two-stage linear ion trap before extraction into a TOF mass spectrometer for mass analysis. Each target anion was mass-selected, collimated, and momentum decelerated before being irradiated with 193 nm laser pulses (ExciStar™ XS 500, Coherent) in the interaction zone. Photodetached electrons were collected using a magnetic-bottle photoelectron spectrometer. The acquired original photoelectron TOF spectra were converted to electron kinetic energy (eKE) spectra and calibrated by the known spectra of I^- ,¹⁹ and MnO_4^- .²⁰ The final electron binding energy (eBE) spectra were obtained by subtracting the eKE spectra from the detachment photon energy, with an electron energy resolution ($\Delta E/E$) of $\sim 1.9\%$ (*i.e.*, ~ 30 meV for 1.6 eV KE electrons).

Computational methods

The structures of target anions and their corresponding neutral species were optimized using density functional theory (DFT) with the hybrid B3LYP-D3(BJ) exchange–correlation functional,²¹ which has been widely used in studying Pt-containing complexes with sufficient accuracy.^{22–24} The aug-cc-pVTZ-pp basis set with an effective core potential (ECP) was used for both Pt and I atoms to account for scalar relativistic effects,^{25,26} while the aug-cc-pVTZ basis set was used for H, N, Cl, and Br atoms.^{27,28} Various initial structures with different spin multiplicities for both anionic and neutral species were explored to identify global minima. Vibrational frequency calculations confirmed the absence of imaginary frequencies, ensuring that the optimized structures correspond to true minima. The adiabatic detachment energy (ADE) values were determined as the energy difference between the neutral and anionic species at their respective optimized geometries, including zero-point energy (ZPE) corrections. The vertical detachment energy (VDE) values were calculated as the energy difference between the neutral and anionic species, with the neutral geometry constrained to that of the optimized anion. Excited states of the neutral species were computed using time-dependent DFT (TDDFT)²⁹ at the CAM-B3LYP³⁰/aug-cc-pVTZ(-pp) level of theory. Spin-orbit coupling effects were not considered in these calculations, as these do not qualitatively impact experimental data interpretation.^{31–33} Since the three-coordinated platinum halide anions exhibit a pronounced *trans* effect upon NH_3 ligand coordination, we employed extended transition state-natural orbitals for chemical valence (ETS-NOCV),³⁴ natural population analysis (NPA), and ionic bonding contributions to investigate how *trans* effects influence charge redistribution and electronic interactions/transfer. Additionally, dissociation energy and binding energy calculations were performed to assess how the *trans* effect affects ligand stability at different coordination sites. For $Pt(NH_3)ClX_2^-$ and their isomers, ETS-NOCV analyses were performed using the Multiwfn 3.8 (dev) program.³⁵ In this approach, anions were divided into two fragments: the halogenated platinum anion and the NH_3 ligand, enabling a direct examination of how NH_3 impacts the electronic distribution of the halogenated platinum moiety. Additionally, NPA charge distributions and dissociation energies were calculated at the

B3LYP-D3(BJ)/aug-cc-pVTZ(-pp) level of theory. The binding energies were calculated with corrections for basis set superposition error (BSSE) using the counterpoise method, and zero-point energy (ZPE) corrections derived from vibrational frequency calculations at the B3LYP-D3(BJ)/aug-cc-pVTZ(-pp) level of theory. The ionic contributions of chemical bonds were evaluated using localized molecular orbital methods implemented in the Multiwfn 3.8 (dev) program, and the highest occupied molecular orbitals (HOMOs) and the ETS-NOCV results were visualized using the VMD³⁶ program. All quantum chemical calculations were performed using the Gaussian 16 Rev. C.02,³⁷ and GaussView 6 was used for structure visualization.

Results and discussion

Photoelectron spectra of target anions

Fig. 1 presents the cryogenic anion photoelectron spectra of PtClX_2^- ($X = \text{Cl}, \text{Br}, \text{I}$) and $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Cl}, \text{Br}, \text{I}$) recorded

under 193 nm laser excitation. The observed spectral features, labelled as X and A–G, correspond to the transitions from the anionic ground state to the neutral ground and excited states upon electron detachment. The VDE is defined as the maximum of the X state in each spectrum, while the ADE is determined by the intersection of the tangent to the rising edge of the X state with the eBE axis, corrected for the instrumental resolution. Based on the experimental data, the ADEs values for PtClX_2^- and $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Cl}, \text{Br}, \text{I}$) were determined to be 4.26, 4.23, 4.08, 3.85, 3.79, and 3.96 eV, respectively, while the corresponding VDE values were found to be 4.46, 4.37, 4.23, 4.06, 4.01, and 4.09 eV. Fig. 1 reveals that the coordination of an NH_3 ligand to the three-coordinate platinum halide anions results in a notable decrease in the electron binding energy, shifting the overall spectra toward lower eBE values. Additionally, while the spectral features of PtCl_3^- and $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$ remain largely unchanged, the spectra of $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Br}, \text{I}$) exhibit noticeable changes compared to those of PtClX_2^- ($X = \text{Br}, \text{I}$) in the context of spectral broadening and splitting. Notably, in the

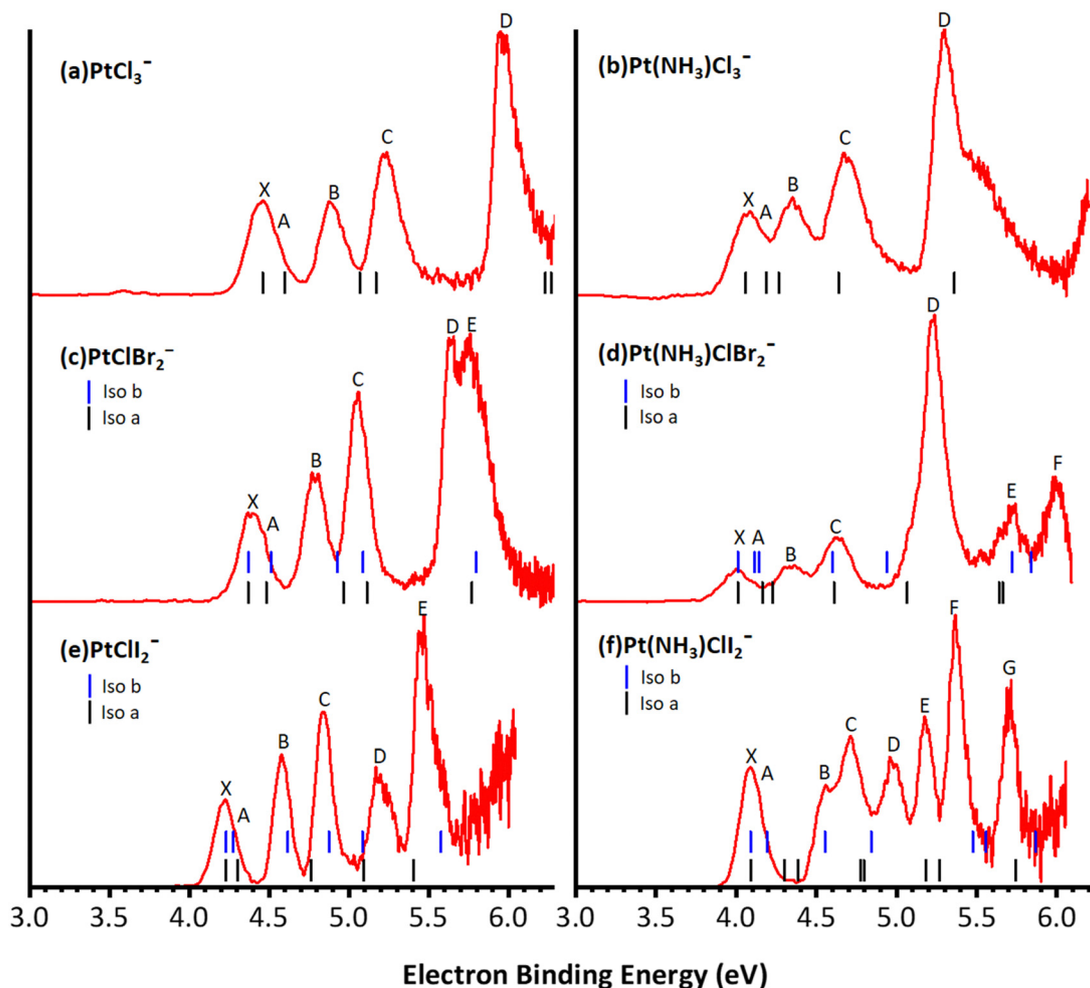


Fig. 1 The 193 nm cryogenic photoelectron spectra of (a) PtCl_3^- , (b) $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$, (c) PtClBr_2^- , (d) $\text{Pt}(\text{NH}_3)\text{ClBr}_2^-$, (e) PtCl_2^- , and (f) $\text{Pt}(\text{NH}_3)\text{Cl}_2^-$. Vertical bars indicate the energies of the ground and excited states of neutral species, calculated using TDDFT at the CAM-B3LYP/aug-cc-pVTZ(-pp) level of theory. In (c)–(f), black and blue bars represent the transition energies derived from different isomers. The calculated first VDE is shifted to align with the experimental X peak in each spectrum.

spectrum of PtClBr_2^- , the D and E peaks appear to originate from the splitting of a single spectral feature, arising presumably from the copresence of two isomers with distinct halogen coordination geometries in the experiments. This hypothesis will be further examined through theoretical calculations in subsequent sections.

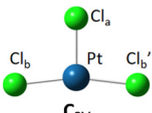
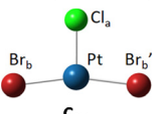
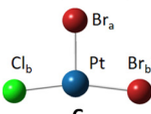
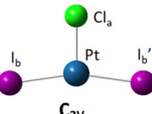
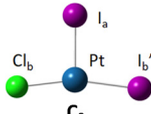
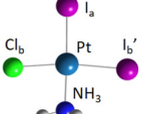
Optimized structures, calculated ADEs and VDEs

The DFT optimized, lowest-energy structures of PtClX_2^- and $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Cl}, \text{Br}, \text{I}$), are summarized in Table 1 (see Fig. S1–S3, ESI† for the details about how to construct the initial structures with different spin multiplicities, and the corresponding optimized neutral molecules). The most stable structures in Table 1 all feature a singlet electronic state with an overall similar platinum trihalide framework, in which one halide at the apex (X_a) is distinct from the other two at the base ($X_b, X_{b'}$) that forms a large obtuse $X_b\text{--Pt--}X_{b'}$ angle ($> 160^\circ$). PtCl_3^- has two identical Pt--Cl_b bonds of 2.310 Å, and a slightly shorter Pt--Cl_a bond at 2.247 Å. Upon coordination of NH_3 to PtCl_3^- from the base, *trans*- to the apical Cl, all Pt--Cl bonds elongate, with the elongation of the apical Cl nearly twice bigger than the base Cl (0.071 vs. 0.042 Å), while the $\text{Cl}_b\text{--Pt--Cl}_{b'}$ angle increases from 166° to 172° . For PtClBr_2^- , two low-lying isomers are identified

with the one featuring Cl at the apex (iso a) being slightly less stable by 0.07 eV compared to the other with the Br on the apical position (iso b). Attaching one NH_3 molecule from the base, *trans*-positioned to the apical halide in PtClBr_2^- results in two corresponding isomers of $\text{Pt}(\text{NH}_3)\text{ClBr}_2^-$, in which the coordination of NH_3 induces similar geometric changes for the PtClBr_2^- moiety as described in the PtCl_3^- case above, *i.e.*, substantially more elongation for the $\text{Pt--}X_a$ bond than $\text{Pt--}X_b$ with increased $X_b\text{--Pt--}X_{b'}$ bond angles. Notably, binding NH_3 to PtClBr_2^- reverses the relative stability of the two isomers of PtClBr_2^- with the iso a of $\text{Pt}(\text{NH}_3)\text{ClBr}_2^-$ featuring a *trans*-Cl becoming more stable. Similarly, two isomers of PtClI_2^- are located with iso a featuring an apical Cl lying 0.2 eV higher in energy than iso b. Attaching one NH_3 molecule from the base of PtClI_2^- largely eliminates this relative energy difference and again reverses the energy order with iso a of the ammonium complex become the most stable one, but only by 0.05 eV (Table 1). The same geometric changes brought by NH_3 are observed in $\text{Pt}(\text{NH}_3)\text{ClI}_2^-$ as well.

Examining the aforementioned optimized structures evidences that the coordination of the NH_3 ligand to platinum trihalides leads to an overall elongation of the Pt--halogen bonds. Notably, the bond length of the $\text{Pt--}X_a$, positioned *trans* to NH_3 , increases more significantly, showing a clear *trans*

Table 1 Lowest-energy structures of PtClX_2^- and $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Cl}, \text{Br}, \text{I}$) anions optimized at the B3LYP-D3(BJ)/aug-cc-pVTZ-(pp) level of theory. The relative energy ΔE^a (in eV), bond length (in Å) and bond angles (in degree) are provided

Anions	PtClX_2^- (singlet)	$\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ (singlet)
		
Pt–Cl _a	2.247	2.318
Pt–Cl _b	2.310	2.352
Pt–Cl _{b'}	2.310	2.352
∠ Cl _a –Pt–Cl _b	97	94
∠ Cl _b –Pt–Cl _{b'}	166	172
	Iso a ($\Delta E = 0.07$)	Iso a ($\Delta E = 0$)
		
Pt–Cl _{a/b}	2.248	2.310
Pt–Br _{a/b}	2.439	2.377
Pt–Br _{b'}	2.439	2.434
∠ Br _{a/b} –Pt–Cl _{a/b}	98	97
∠ Br _{a/b} –Pt–Br _{b'}	165	97
	Iso a ($\Delta E = 0.2$)	Iso b ($\Delta E = 0$)
		
Pt–Cl _{a/b}	2.251	2.315
Pt–I _{a/b}	2.610	2.548
Pt–I _{b'}	2.610	2.595
∠ I _{a/b} –Pt–Cl _{a/b}	98	97
∠ I _{a/b} –Pt–I _{b'}	164	97
		Iso a ($\Delta E = 0$)
		
		Iso b ($\Delta E = 0.05$)
		
Pt–Cl _{a/b}		2.317
Pt–I _{a/b}		2.666
Pt–I _{b'}		2.668
∠ I _{a/b} –Pt–Cl _{a/b}		94
∠ I _{a/b} –Pt–I _{b'}		172
		Iso a ($\Delta E = 0$)
		
		Iso b ($\Delta E = 0.03$)
		
Pt–Cl _{a/b}		2.357
Pt–Br _{a/b}		2.451
Pt–Br _{b'}		2.480
∠ Br _{a/b} –Pt–Cl _{a/b}		94
∠ Br _{a/b} –Pt–Br _{b'}		95

^a ΔE represents the energy difference between the isomer structure and the lowest-energy structure.

Table 2 The experimental and calculated ADEs and VDEs of PtClX_2^- and $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Cl}, \text{Br}, \text{I}$) anions. The calculations were performed at the B3LYP-D3(BJ)/aug-cc-pVTZ(-pp) level of theory

Anions	ADE (eV)		VDE (eV)	
	Exp. ^a	Calc.	Exp. ^a	Calc.
PtCl_3^-	4.26	4.18	4.46	4.35
$\text{Pt}(\text{NH}_3)\text{Cl}_3^-$	3.85	3.83	4.06	4.00
PtClBr_2^- (iso a)	4.23	4.13	4.37	4.28
PtClBr_2^- (iso b)		4.12		4.26
$\text{Pt}(\text{NH}_3)\text{ClBr}_2^-$ (iso a)	3.79	3.86	4.01	4.00
$\text{Pt}(\text{NH}_3)\text{ClBr}_2^-$ (iso b)		3.84		3.98
PtClI_2^- (iso a)	4.08	3.79	4.23	4.12
PtClI_2^- (iso b)		3.99		4.10
$\text{Pt}(\text{NH}_3)\text{ClI}_2^-$ (iso a)	3.96	3.80	4.09	3.90
$\text{Pt}(\text{NH}_3)\text{ClI}_2^-$ (iso b)		3.77		3.86

^a The experimental uncertainty of ADEs and VDEs is 0.02 eV.

effect, whereas the Pt- X_b bonds adjacent to the NH_3 ligand exhibit only minor changes, much less influenced by this interaction. For the mixed ligand platinum trihalides PtClX_2^- ($X = \text{Br}, \text{I}$), iso a in which the sole Cl is at the apex is less favourable than iso b that features the apical Br or I. Binding NH_3 *trans* to the apex of PtClX_2^- preferentially stabilizes more on iso a, rendering iso a the most stable structure for $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$. Additionally, Fig. S2 (ESI[†]) presents the optimized structure of the PtCl_3^- under different spin multiplicities, showing that its D_{3h} -symmetric triplet-state structure is higher in energy by 0.29 eV, compared to its lower-symmetry singlet ground-state configuration. In addition to B3LYP-D3(BJ), we employed CAM-B3LYP and MP2 with the aug-cc-pVTZ(-pp) basis set to optimize the structures of PtClX_2^- and $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Cl}, \text{Br}, \text{I}$) and compute their corresponding energies. The optimized geometries from all three methods were in close agreement. However, B3LYP-D3(BJ) provided the best match to the experimental ADE and VDE values, and its results were therefore adopted. Detailed comparisons are provided in Fig. S4 and Table S1 (ESI[†]).

Table 2 compares the experimental and theoretical ADE and VDE values for PtClX_2^- and $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Cl}, \text{Br}, \text{I}$). Excluding the results of iso a of PtClI_2^- which is 0.2 eV less stable and unlikely contributed to the experiments, an excellent agreement between the experimental and theoretical results is found with the mean absolute errors (MAEs) of 0.096 eV for ADE and 0.106 eV for VDE. Additionally, the excited states of the neutral molecules were calculated using the anionic structures (Table 3) and compared with the corresponding spectra (see the vertical lines in Fig. 1). The simulated TDDFT stick spectra reasonably cover the observed spectral bands, but obvious deviations exist, presumably due to the lack of consideration of multi-reference nature and spin-orbit splitting effects in the calculations.

Trans effect displayed in charge distribution, MO interaction, and bonding characteristics

To further investigate the *trans* effect in platinum halide complexes upon NH_3 coordination, the NPA charge distributions of

Table 3 Experimentally measured and theoretically calculated VDEs and excitation energies of neutral excited states from TDDFT calculations, at the CAM-B3LYP/aug-cc-pVTZ(-pp) level. The VDE of the ground state X was calculated at the B3LYP-D3(BJ)/aug-cc-pVTZ(-pp) level

Anions	States	VDEs (eV)	
		Expt. ^c	Calc.
PtCl_3^-	X	4.46	4.35
	A	4.55	4.59
	B	4.87	5.06
	C	5.23	5.17
	D	5.98	6.23/6.27
$\text{Pt}(\text{NH}_3)\text{Cl}_3^-$	X	4.06	4.00
	A	4.16	4.19
	B	4.35	4.26
	C	4.67	4.64
	D	5.30	5.35
PtClBr_2^- (iso a, b)	X	4.37	4.28 ^a , 4.26 ^b
	A	4.46	4.48 ^a , 4.51 ^b
	B	4.77	4.97 ^a , 4.93 ^b
	C	5.05	5.11 ^a , 5.09 ^b
	D	5.63	5.80 ^b
	E	5.75	5.77 ^a
$\text{Pt}(\text{NH}_3)\text{ClBr}_2^-$ (Iso a, b)	X	4.01	4.00 ^a , 3.98 ^b
	A	4.10	4.16 ^a , 4.12 ^b /4.14 ^b
	B	4.34	4.23 ^a
	C	4.63	4.61 ^a , 4.60 ^b
	D	5.23	5.07 ¹ , 4.94 ^b
	E	5.72	5.64 ^a /5.66 ^a , 5.72 ^b
	F	5.98	5.84 ^b
PtClI_2^- (iso a, b)	X	4.23	4.12 ^a , 4.10 ^b
	A	4.29	4.30 ^a , 4.27 ^b
	B	4.58	4.76 ^a , 4.62 ^b
	C	4.84	4.87 ^b
	D	5.17	5.10 ^a , 5.09 ^b
	E	5.46	5.41 ^a , 5.57 ^b
$\text{Pt}(\text{NH}_3)\text{ClI}_2^-$ (Iso a, b)	X	4.09	3.90 ^a , 3.86 ^b
	A	4.14	4.30 ^a , 4.19 ^b
	B	4.55	4.39 ^a , 4.55 ^b
	C	4.71	4.77 ^a /4.79 ^a
	D	4.96	4.84 ^b
	E	5.18	5.18 ^a
	F	5.36	5.27 ^a , 5.48 ^b /5.55 ^b
	G	5.71	5.74 ^a , 5.86 ^b

^a The isomer a of PtClX_2^- , or $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Br}, \text{I}$). ^b The isomer b of PtClX_2^- , or $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Br}, \text{I}$). ^c The experimental uncertainty of VDEs is 0.02 eV.

the corresponding anions were examined (Fig. 2). In PtCl_3^- , the central Pt atom carries a positive charge of 0.23, while the apical Cl is at -0.29 and two other Cl ligands at the base possess more negative charges of -0.47 . This trend of NPA charges among three Cl atoms can be attributed to that the apical Cl ligand is adjacent to two negatively charged Cl atoms, affording a maximum reduction in electrostatic repulsion. Substituting two base Cl ligands in PtCl_3^- by two Br or two I generate iso a of PtClBr_2^- or PtClI_2^- , respectively, with a structure similar to that of PtCl_3^- . The amount of negative charges of the base halogens follow their electron negativity, in the order of $\text{Cl} > \text{Br} > \text{I}$, accompanied by a gradual reduction of the Pt positive charge. Notably, the charge at the apical Cl atom remains largely unchanged and is significantly lower than those at the base ligands. In iso b of $\text{PtClBr}_2^-/\text{PtClI}_2^-$,

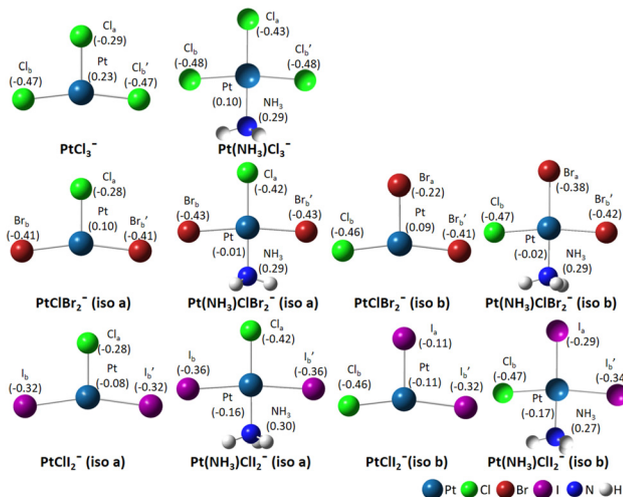


Fig. 2 The natural population analysis (NPA) charge distribution of PtClX_2^- and $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Cl}, \text{Br}, \text{I}$) anions, calculated at the B3LYP-D3(BJ)/aug-cc-pVTZ-(pp) level of theory.

the apical Br_a/I_a ligand also consistently possesses notable fewer negative charges than the other two halogens at the base. On going from PtCl_3^- to PtClBr_2^- and PtClI_2^- , the positive charge on Pt reduces progressively, and becomes even negative in PtClI_2^- , following the electron affinities of halogens in the order $\text{Cl} > \text{Br} > \text{I}$.

Upon NH_3 coordination in $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$, the charges on the two base Cl atoms remain largely unchanged, while the positive charge on the central Pt atom decreases significantly. Concomitantly, the amount of negative charge on the apical Cl atom increases noticeably, and the NH_3 ligand itself carries a partial positive charge. The above NPA charge analyses suggest that NH_3 , as an electron-donating group, transfers its electron density primarily to the central Pt and apical Cl atom. The induced NPA charge distribution change upon coordination of NH_3 in $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Br}, \text{I}$) closely resembles those observed in the $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$ case. Specifically, the halogen ligand positioned *trans* to NH_3 exhibits a significantly increased negative charge, whereas those at the base remain nearly intact.

To visualize the impact of the *trans* effect on electron transfer upon NH_3 coordination, ETS-NOCV calculations were performed. The results indicate that the NH_3 coordination induces electron transfer predominately from NH_3 (blue, negative orbital density) to its opposite halogen ligand (green, positive orbital density), while the adjacent ligands remain largely unaffected (Fig. 3). This observation is consistent with the NPA charge distribution analysis, in which the negative charge on the *trans* halogen ligand substantially increases after NH_3 coordination, independent of the halogen type, highlighting the dominate role of the *trans* effect in this process. Furthermore, the *trans* effect induced by NH_3 coordination significantly impacts the polarity of the *trans* Pt–halogen bonds. As shown in Fig. 4, the ionic character of the *trans* Pt–halogen bond increases by more than 16% after NH_3 coordination, whereas the ionic character of Pt to the adjacent halogen ligands changes less than 1.5%. Detailed data on the ionic

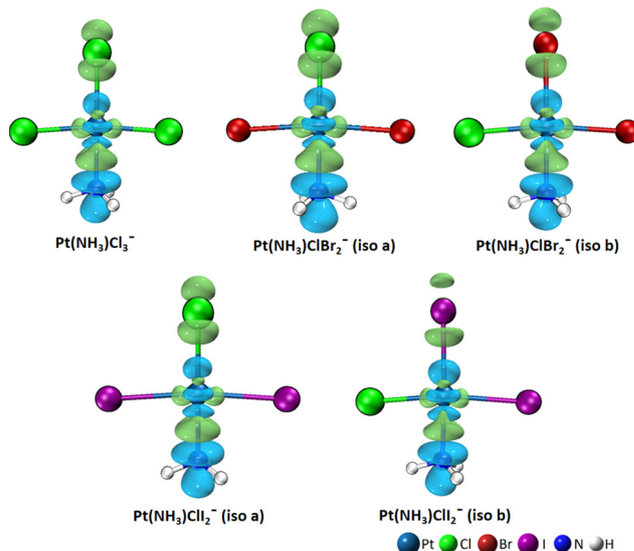


Fig. 3 The calculated ETS-NOCV of $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$, $\text{Pt}(\text{NH}_3)\text{ClBr}_2^-$ (iso a and b) and $\text{Pt}(\text{NH}_3)\text{ClI}_2^-$ (iso a and b) with the isovalue of 0.005 at the B3LYP-D3(BJ)/aug-cc-pVTZ-(pp) level of theory. The colour green represents the positive orbital density and blue represents negative. The corresponding orbital interactions induce electron transfer from the blue region to the green region.

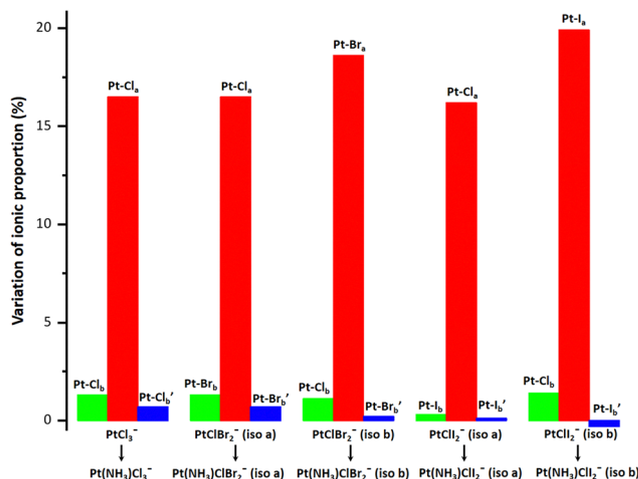


Fig. 4 The difference in ionic proportion of the corresponding Pt–halogen bonds in PtClX_2^- and $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Cl}, \text{Br}, \text{I}$). The calculations were performed at the B3LYP-D3(BJ)/aug-cc-pVTZ-(pp) level of theory.

character are provided in Table S1 (ESI[†]). These results demonstrate the pronounced *trans* effect brought by coordinating NH_3 in platinum trihalides.

Additionally, the *trans* effect also influences the frontier molecular orbitals (FMOs) of the system. Taking PtCl_3^- and $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$ as an example, NH_3 coordination does not significantly alter the orbital shapes but upshifts their energies, making them easier to ionize. Molecular orbitals with a higher contribution from the apical Cl atom in PtCl_3^- (HOMO–2, HOMO–4) exhibit greater energy upshifts upon the *trans* NH_3 binding in $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$, which can even cause orbital reordering, *i.e.*,

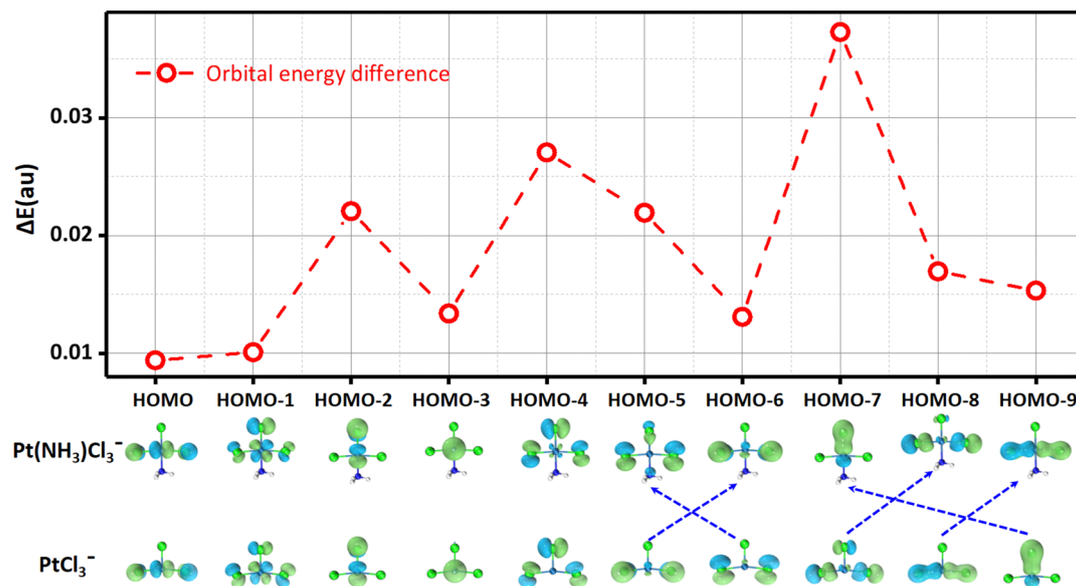


Fig. 5 The HOMO energy difference upon NH_3 coordination and orbital rearrangement between PtCl_3^- and $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$ with the pictures of the corresponding MOs illustrated. $\Delta E = \epsilon(\text{FMO})(\text{Pt}(\text{NH}_3)\text{Cl}_3^-) - \epsilon(\text{FMO})(\text{PtCl}_3^-)$ for $\text{FMO} = \text{HOMO}-1, -2, -3, -4$; and the correlated orbital differences (lined by blue lines) for deeper FMOs (HOMO-5, -6, ..., -9). The calculations were performed at the B3LYP-D3(BJ)/aug-cc-pVTZ-(pp) level of theory.

HOMO-6 and HOMO-9 of PtCl_3^- become HOMO-5 and HOMO-7 of $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$, respectively (Fig. 5). Consistent trends are observed from PtClX_2^- ($X = \text{Br}, \text{I}$) to $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Br}, \text{I}$) (Fig. S5–S8, ESI[†]).

Previous studies have established that cisplatin (*cis*- $\text{Pt}(\text{NH}_3)_2\text{Cl}_2$) exhibits potent anticancer activity, whereas its *trans* analogue transplatin (*trans*- $\text{Pt}(\text{NH}_3)_2\text{Cl}_2$) is inactive.^{38–40} The structural difference between the two lies in the spatial arrangement of the two NH_3 ligands: in cisplatin, each NH_3 ligand is positioned *trans* to a Cl atom, while in transplatin, two NH_3 ligands (also two Cl) face each other from the opposite side. During the drug activation process inside the cell, the Cl ligand opposite to the NH_3 ligand in cisplatin undergoes dissociation. Table 4 presents the theoretical bond dissociation energies (BDE) of $\text{Pt}-\text{Cl}_a$ and $\text{Pt}-\text{Cl}_b$. In PtCl_3^- , the $\text{Pt}-\text{Cl}_a$ bond has the shortest bond length of 2.247 Å and highest BDE of 3.61 eV compared to that of $\text{Pt}-\text{Cl}_b$ (2.310 Å, 3.29 eV). Upon NH_3 combination, the *trans* effect elongates $\text{Pt}-\text{Cl}_a$ in $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$ to 2.318 Å and substantially weakens this bond with BDE = 2.90 eV, rendering it the weakest bond prone to dissociate. More BDE calculations in PtClX_2^- and $\text{Pt}(\text{NH}_3)\text{ClX}_2^-$ ($X = \text{Br}, \text{I}$) (Table S2, ESI[†]) reveal the same trend, further corroborating the significant role of the *trans* effect in facilitating the *trans* halide dissociation by binding a NH_3 ligand.

Based on these findings, we speculate that in cisplatin and other *cis*-structured metal-based anticancer drugs, the leaving

group ligands are those positioned opposite to the NH_3 ligand. In contrast, transplatin has halide ligands all at the adjacent, not *trans* position to NH_3 , rendering them less susceptible to this effect. Although the drug activation mechanism occurs *in vivo* and under liquid-phase environments, our gas-phase study provides valuable insights into the underlying molecular mechanisms that govern such metal-based anticancer drug activities and offers benchmark references for future research aimed at designing therapeutic agents with greater efficacy.

To further investigate the *trans* effect in tetracoordinated platinum halide complexes with different ligands, we optimized the structures of Zeise's salt analogs, $\text{Pt}(\text{C}_2\text{H}_4)\text{ClX}_2^-$ ($X = \text{Cl}, \text{Br}, \text{I}$), using the same level of theory (B3LYP-D3(BJ)/aug-cc-pVTZ-(pp)) (Table S4, ESI[†]). Structural comparisons between PtClX_2^- and $\text{Pt}(\text{C}_2\text{H}_4)\text{ClX}_2^-$ reveal a pronounced *trans* effect induced by ethylene coordination, characterized by a significant elongation of the $\text{Pt}-\text{X}$ bond *trans* to the C_2H_4 ligand. Taking $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$ and $\text{Pt}(\text{C}_2\text{H}_4)\text{Cl}_3^-$ as representative examples, we compared their structural parameters and ligand dissociation energies. Upon coordination, the $\text{Pt}-\text{Cl}_a$ bond *trans* to NH_3 elongates to 2.318 Å with a corresponding dissociation energy of 2.90 eV. In contrast, in $\text{Pt}(\text{C}_2\text{H}_4)\text{Cl}_3^-$, the *trans* $\text{Pt}-\text{Cl}_a$ bond elongates further to 2.338 Å, with a slightly lower dissociation energy of 2.82 eV (Table S5, ESI[†]). These observations demonstrate that the *trans* effect exerted by C_2H_4 is more pronounced than that of NH_3 , as

Table 4 The dissociation energies of different dissociation channels of PtCl_3^- and $\text{Pt}(\text{NH}_3)\text{Cl}_3^-$. All calculations were carried out at the B3LYP-D3(BJ)/aug-cc-pVTZ-(pp) level of theory

Channel no.	Dissociation channels	E (eV)	Dissociation channels	E (eV)
1	$\text{PtCl}_3^- \rightarrow \text{PtCl}_2 + \text{Cl}_a^-$	3.61	$\text{Pt}(\text{NH}_3)\text{Cl}_3^- \rightarrow \text{Pt}(\text{NH}_3)\text{Cl}_2 + \text{Cl}_a^-$	2.90
2	$\text{PtCl}_3^- \rightarrow \text{PtCl}_2 + \text{Cl}_b^-$	3.29	$\text{Pt}(\text{NH}_3)\text{Cl}_3^- \rightarrow \text{Pt}(\text{NH}_3)\text{Cl}_2 + \text{Cl}_b^-$	2.93
3	$\text{PtCl}_3^- \rightarrow \text{PtCl}_2 + \text{Cl}_b'^-$	3.29	$\text{Pt}(\text{NH}_3)\text{Cl}_3^- \rightarrow \text{Pt}(\text{NH}_3)\text{Cl}_2 + \text{Cl}_b'^-$	2.93

reflected in both greater bond elongation and reduced bond strength for the *trans*-positioned halide. To further assess the coordination strength of the two ligands, we calculated the binding energies (BEs) of NH₃ and C₂H₄ with PtClX₂[−] (X = Cl, Br, I), incorporating both basis set superposition error (BSSE) and zero-point energy (ZPE) corrections. The results show that the Pt–C₂H₄ interaction is consistently stronger than that of Pt–NH₃, suggesting that the C₂H₄-coordinated complexes are thermodynamically more stable. Detailed structural parameters, dissociation energies, and binding energies are provided in Tables S4–S6 (ESI[†]). Previous work by Hou *et al.*⁴¹ reported significant back-donation from the Pt center to the ethylene ligand in Zeise's salt complexes, which inspired us to investigate whether a similar interaction occurs in Pt(NH₃)Cl₃[−]. However, as shown in Fig. S9 (ESI[†]), an examination of the LUMO to LUMO+10 orbitals in Pt(NH₃)Cl₃[−] reveals no evidence of back-donation from Pt to the NH₃ ligand, further supporting its role as a classical σ-donor.

Conclusions

In this contribution, we report a combined photoelectron spectroscopic and theoretical study on PtClX₂[−] in comparison to their corresponding NH₃ complexes Pt(NH₃)ClX₂[−] (X = Cl, Br, I), aimed to provide a molecular level understanding on the widely accepted 'trans effect' proposed nearly 100 years ago. Our results show that introducing an electron-donating NH₃ ligand leads to a decrease in the eBE of the platinum trihalide anions, and facilitates electron transfer primarily along the H₃N–Pt–X_a axis to the *trans* positioned halogen ligand. Furthermore, our study demonstrates that the most stable apex halide ligand undergoes a substantial reduction in BDE upon NH₃ coordination, manifesting an auspicious *trans* effect. At the electronic state level, this research elucidates how the *trans* effect influences the physical and chemical properties of platinum trihalide anions when coordinated with NH₃. Finally, we also conducted a comparison with the widely utilized Zeise's salt system. The results reveal that PtClX₂[−] (X = Cl, Br, I) coordinated with C₂H₄ similarly exhibits a pronounced *trans* effect, and the influence of this effect on the stability of the *trans* ligand is slightly greater than that observed with the NH₃ ligand. These findings provide valuable insights into transition-metal systems and establish a theoretical foundation for understanding platinum-based anticancer drugs, aiding in their design and synthesis.

Author contributions

Qixu Zhao: investigation (equal); writing – original draft (equal); formal analysis (equal). Jian Zhang: investigation (equal); writing – original draft (equal); formal analysis (equal). Xueying Li: resources (equal); data curation (supporting). Peng Tang: resources (equal); investigation (supporting). Fan Yang: data curation (lead). Junyang Ma: visualization (equal). Zhubin Hu: software (supporting); visualization (equal). Haitao Sun: software (lead). Xue-Bin Wang: writing – review and editing (lead); conceptualization (equal). Zhenrong Sun: writing – review and editing (equal); funding acquisition

(lead); conceptualization (equal). Yan Yang: writing – review and editing (equal); conceptualization (lead); validation (lead).

Data availability

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

Conflicts of interest

The authors have no conflicts to disclose.

Acknowledgements

The work was partly supported by the National Natural Science Foundation of China (No. 92461301, 12034008, and 12250003) and the Program of Introducing Talents of Discipline to Universities 111 Project (B12024). X.-B. W. was supported by the US Department of Energy (DOE), Office of Science, Office of Basic Energy Sciences, Division of Chemical Science, Geosciences, and Biosciences, Condensed Phase and Interfacial Molecular Science program, FWP 16248.

Notes and references

- 1 F. R. Hartley, *Chem. Soc. Rev.*, 1973, **2**, 163–179.
- 2 R. G. Pearson, *Inorg. Chem.*, 1973, **12**, 712–713.
- 3 B. J. Coe and S. J. Glenwright, *Coord. Chem. Rev.*, 2000, **203**, 5–80.
- 4 M. H. Baik, R. A. Friesner and S. J. Lippard, *J. Am. Chem. Soc.*, 2003, **125**, 14082–14092.
- 5 D. V. Deubel, *J. Am. Chem. Soc.*, 2006, **128**, 1654–1663.
- 6 Y. Mantri, S. J. Lippard and M. H. Baik, *J. Am. Chem. Soc.*, 2007, **129**, 5023–5030.
- 7 S. Gonell, M. D. Massey, I. P. Moseley, C. K. Schauer, J. T. Muckerman and A. J. M. Miller, *J. Am. Chem. Soc.*, 2019, **141**, 6658–6671.
- 8 S. Gonell, E. A. Assaf, K. D. Duffee, C. K. Schauer and A. J. M. Miller, *J. Am. Chem. Soc.*, 2020, **142**, 8980–8999.
- 9 W. C. Zeise and O. K. Dan, *Vidensk. Selsk. Forh.*, 1825–1826, **6**, 13–14.
- 10 G. L. Hou, H. Wen, K. Lopata, W. J. Zheng, K. Kowalski, N. Govind, X. B. Wang and S. S. Xantheas, *Angew. Chem., Int. Ed.*, 2012, **51**, 6356–6360.
- 11 G.-L. Hou, N. Govind, S. S. Xantheas and X.-B. Wang, *J. Phys. Chem. A*, 2018, **122**, 1209–1214.
- 12 A. M. Amado, S. M. Fiuza, M. P. M. Marques and L. A. E. Batista de Carvalho, *J. Chem. Phys.*, 2007, **127**, 185104.
- 13 C. Zhang, E. B. Naziga and L. Guidoni, *J. Phys. Chem. B*, 2014, **118**, 11487–11495.
- 14 J. Warneke, M. Rohdenburg, Y. Zhang, J. Orszagh, A. Vaz, I. Utke, J. T. M. De Hosson, W. F. van Dorp and P. Swiderek, *J. Phys. Chem. C*, 2016, **120**, 4112–4120.

- 15 M. Rohdenburg, P. Martinović, K. Ahlenhoff, S. Koch, D. Emmrich, A. Götzhäuser and P. Swiderek, *J. Phys. Chem. C*, 2019, **123**, 21774–21787.
- 16 B. Pinter, V. Van Speybroeck, M. Waroquier, P. Geerlings and F. De Proft, *Phys. Chem. Chem. Phys.*, 2013, **15**, 17354–17365.
- 17 P. Tang, J. Zhang, X. Li, F. Yang, Q. Zhao, J. Ma, Z. Hu, H. Sun, X.-B. Wang, Z. Sun and Y. Yang, *J. Phys. Chem. A*, 2024, **128**, 5500–5507.
- 18 Q. Zhao, J. Zhang, X. Li, P. Tang, F. Yang, J. Ma, Z. Hu, H. Sun, X.-B. Wang, Z. Sun and Y. Yang, *J. Chem. Phys.*, 2024, **161**, 214305.
- 19 D. Hanstorp and M. Gustafsson, *J. Phys. B:At., Mol. Opt. Phys.*, 1992, **25**, 1773.
- 20 J. Zhang, Z.-R. Sun and X.-B. Wang, *J. Phys. Chem. A*, 2015, **119**, 6244–6251.
- 21 N. Mehta, M. Casanova-Páez and L. Goerigk, *Phys. Chem. Chem. Phys.*, 2018, **20**, 23175–23194.
- 22 J. Joseph, K. Pradhan, P. Jena, H. Wang, X. Zhang, Y. Jae Ko and K. H. Bowen, *J. Chem. Phys.*, 2012, **136**, 194305.
- 23 Y. Shi, S. Bian, Y. Ma, Y. Wang, J. Ren and X. Kong, *J. Phys. Chem. A*, 2018, **123**, 187–193.
- 24 R. Wesendrup and P. Schwerdtfeger, *Inorg. Chem.*, 2001, **40**, 3351–3354.
- 25 K. A. Peterson, B. C. Shepler, D. Figgen and H. Stoll, *J. Phys. Chem. A*, 2006, **110**, 13877–13883.
- 26 D. Figgen, K. A. Peterson, M. Dolg and H. Stoll, *J. Chem. Phys.*, 2009, **130**, 164108.
- 27 R. A. Kendall, T. H. Dunning, Jr. and R. J. Harrison, *J. Chem. Phys.*, 1992, **96**, 6796–6806.
- 28 D. E. Woon and T. H. Dunning, Jr., *J. Chem. Phys.*, 1993, **98**, 1358–1371.
- 29 E. Runge and E. K. U. Gross, *Phys. Rev. Lett.*, 1984, **52**, 997–1000.
- 30 T. Yanai, D. P. Tew and N. C. Handy, *Chem. Phys. Lett.*, 2004, **393**, 51–57.
- 31 Y. Wei, B.-W. Wang, S.-W. Hu, T.-W. Chu, L.-T. Tang, X.-Q. Liu, Y. Wang and X.-Y. Wang, *J. Phys. Org. Chem.*, 2005, **18**, 625–631.
- 32 M. Otsuka, H. Mori, H. Kikuchi and K. Takano, *Comput. Theor. Chem.*, 2011, **973**, 69–75.
- 33 J. C. Gómez Martín, O. Gálvez, M. T. Baeza-Romero, T. Ingham, J. M. C. Plane and M. A. Blitz, *Phys. Chem. Chem. Phys.*, 2013, **15**, 15612–15622.
- 34 M. P. Mitoraj, A. Michalak and T. Ziegler, *J. Chem. Theory Comput.*, 2009, **5**, 962–975.
- 35 T. Lu and F. Chen, *J. Comput. Chem.*, 2012, **33**, 580–592.
- 36 W. Humphrey, A. Dalke and K. Schulten, *J. Mol. Graph.*, 1996, **14**, 33–38.
- 37 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, D. Williams-Young, 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, Revision B.01*, Gaussian, Inc., Wallingford, CT, 2016.
- 38 L. Bica, P. J. Crouch, R. Cappai and A. R. White, *Mol. Biosyst.*, 2009, **5**, 134–142.
- 39 P. Faller, *ChemBioChem*, 2009, **10**, 2837–2845.
- 40 P. Faller and C. Hureau, *Dalton Trans.*, 2009, 1080–1094.
- 41 T. Yang, Z. Li, X. B. Wang and G. L. Hou, *Chem. Phys. Chem.*, 2023, **24**, e202200835.