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Halogen bonding between metal centers and halocarbons†

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Metal-involving halogen bonding was detected in a series of associates of CHI $_3$ with trans-[PtX $_2$ (NCNAlk $_2$) $_2$] (X = Cl, Br). The HI $_2$ C-I··· η^1 (Pt) halogen bonding and the bifurcated HI $_2$ C-I··· η^2 (Pt-Cl) halogen bonding – the latter undergoes the thermally induced reversible HI $_2$ C-I··· η^2 (Pt-Cl) \rightleftarrows HI $_2$ C-I··· η^1 (Pt) transformation – were observed and confirmed theoretically.

Halogen bonding (XB), among various non-covalent weak interactions, is one of the most intensively studied systems in modern science. Together with hydrogen bonding, π – π -, and metallophilic interactions, it is successfully applied in crystal engineering^{1,2} and design of compounds exhibiting pharmacological significance.^{3–5} Recent applications of XB include stabilization of explosives⁶ and molecular design of materials with tunable photophysical properties.^{7,8}

In accord with the IUPAC definition, XB^9 is the real $R-X\cdots Y$ (X= halogen) contact, when, firstly, the interatomic distance between X and an appropriate nucleophilic atom of Y (Y=F, O, N, Cl, *etc.*) is less than the sum of their van der Waals radii and, secondly, the \angle ($R-X\cdots Y$) angle is close to 180° (exceptions from the linearity have been reviewed 10). These geometrical parameters are usually easily obtained using single-crystal X-ray diffraction (XRD), which is the most common method for the detection of XB. When application of the two criteria in XRD data fails, some other criteria (lengthening of R-X covalent bonds due to XB and also changes in the appropriate spectroscopic parameters 9), or

In the majority of the reported cases when XB was reliably identified, only typical Lewis basic non-metal atoms bearing lone pairs (viz. O, N, S, halogens etc.) behave as XB acceptors, whereas possible XBs with metal centers was found only for I₂. ¹⁶⁻¹⁹ Our inspection of the relevant literature indicates few experimental studies describing X···M short contacts, which were either not considered as halogen bonds, or their geometrical parameters fulfill only one of the two aforementioned structural IUPAC criteria. Therefore definitive attribution of these weak interactions to XB was not provided and the other IUPAC criteria should be applied for their identification. Indeed, Chen et al., 20 Wölper et al., 21 and Laguna et al. 22 observed X···Au^I and X···Ag^I contacts, which fulfilled only the "van der Waals criterion", but in all the three cases the corresponding R-X···M angles strongly deviate from 180° (by $30-90^{\circ}$, 20 $60-70^{\circ}$, 21 and 85° 22) thus contrasting with the second structural IUPAC criterion. Yamamoto et al.23 and Liau et al. 24 reported, respectively, the I···Au and Cl···Au contacts, which can be interpreted as X···Au^I halogen bonds, but the nature of these contacts was not verified. Finally, Blakey et al.²⁵ observed contacts between the typical XB donors such as iodoperfluorobenzenes and gold nanoparticles in water solutions, but these halogen bonds were confirmed only by indirect methods such as UV-vis difference spectroscopy, SERS, and XPS and the geometrical parameters were not obtained.

It is important that a metal atom can be an XB acceptor if it contains at least one lone pair, which could interact with empty σ^* -orbitals of an XB donor. In particular, some d^8 -Pt^{II} and d^8 -Rh^I centers may behave similarly to d^{10} -Au^I ones as these centers feature lone pairs on $4d_{z^2}$ or $5d_{z^2}$ orbitals. In particular, weak contacts were observed between some d^8 -Pt^{II} and d^8 -Rh^I species and iodine. ^{16–19} These two were the only reports on unambiguous recognition of metal-involving XB.

Taking into account that no data for XB between metal centers and halocarbons were published, we focused our efforts

application of some other indirect methods, *e.g.* UV-vis, IR and Raman spectroscopy, ^{11,12} differential anomalous X-ray scattering studies, ¹³ NMR, ^{14,15} and phosphorescence⁷ spectroscopy, were also applied for XB recognition.

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[†] Electronic supplementary information (ESI) available: Synthesis of complexes (Section EX1), their co-crystallization with iodoform, full description of XRD experiments, crystallographic data and refinement parameters (Section EX2), full description of the geometrical parameters of covalent bonds and non-covalent interactions (Sections XR1–XR3), and full description of the results of theoretical calculations and computational details (Sections TH1–TH3). CCDC 1446656–1446670. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c6cc01107a

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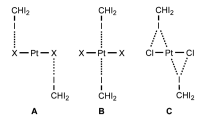


Chart 1 Three types of XB between the linear PtX₂ (X = Cl, Br) fragment and two CHI₃ molecules.

on the search of metal-involving XB and we applied platinum(II) species as potential XB donors and iodoform as an XB acceptor.

In the current work, we report that the dialkylcyanamide d^8 -Pt^{II} complexes trans-[PtX₂(NCNR₂)₂] (R₂ = Me₂, X = Cl **1a**, Br 1b; Et₂ 2, (CH₂)₄ 3, (CH₂)₅ 4) easily co-crystallize with iodoform taken as the XB donor. We observed that CHI3 tends to co-crystallize with 1a-b and 4 forming HI₂C-I···Pt metal-involving halogen bonds (Chart 1, B) that fulfill both structural IUPAC criteria for halogen bonding. The existence of metal-involving halogen bonding has also been confirmed theoretically by the combination of DFT and AIM²⁶ methods. Within the AIM theory, the (3, -1)critical points (bond critical points, BCPs) on the surface of the theoretical electron density function $\rho(r)$ unambiguously reveal all bonding intermolecular interactions, including non-covalent interactions. Energies for the corresponding contacts $E_{\rm int}$ can be determined using semi-empirical correlations between local energy densities in BCPs and interaction strength. 27,28 Moreover, we found and confirmed theoretically by combination of DFT and AIM methods the unique bifurcated $HI_2C-I \cdot \cdot \cdot \eta^2_{(Pt-Cl)}(C)$ XB that undergoes thermally induced reversible $HI_2C-I \cdots \eta^2_{(Pt-Cl)} \rightleftharpoons$ $HI_2C-I\cdots\eta^1_{(Pt)}$ transformation.

Complexes 1a and 1b form isostructural associates 1a: CHCl₃· $1\frac{1}{2}$ CHI₃, $1a \cdot 2$ CHI₃, and $1b \cdot 2$ CHI₃ upon crystallization from 1a: $CHI_3 = 1:2$ (molar ratio) solutions in chloroform or dichloromethane, or from 1b:CHI₃ = 1:2 (molar ratio) solution in dichloromethane, respectively. Apart from numerous HI₂C-I···X-Pt halogen bonds (Chart 1, A), they exhibit the HI₂C-I···Pt short contacts, which can be interpreted as metal-involving XB accordingly to both structural IUPAC criteria as the distances I···Pt (3.4276(5) Å, 3.4389(5) Å, and 3.4023(5) Å) are less than sums of Bondi's²⁹ vdW radii $(R_{vdW}(Pt) + R_{vdW}(I) = 3.73 \text{ Å})$ and the corresponding angles (164.84(17)°, 169.8(3)°, and 172.7(2)°) are close to 180° (for more details see Section XR2 of the ESI†). We also confirmed the formation of the I···Pt non-covalent interaction in $1a \cdot \frac{1}{2}$ CHCl₃· $1\frac{1}{2}$ CHI₃ and $1b \cdot 2$ CHI₃ systems theoretically for the 1a·(CHI₃)₆ and 1b·(CHI₃)₆ model structures by using Bader's AIM analysis (Fig. 1 and 2; for details see Section TH1 in the ESI†). The estimated energies (E_{int}) of these contacts are in the ranges of 2.4–2.8 kcal mol⁻¹ and 2.7–3.1 kcal mol⁻¹, respectively. The strength of these weak interactions has been defined according to the procedures proposed by Espinosa et al.27 and Vener et al.,28 these approaches considered explore linear relationships between the local electronic potential V(r) and kinetic G(r) densities at the bond critical points and energies of appropriate contacts.

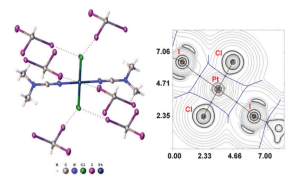


Fig. 1 Left: View of fragment 1a.(CHI₃)₆ from 1a. \(\frac{1}{2}\)CHCl₃. \(\frac{1}{2}\)CHCl₃. Thermal ellipsoids are shown with 50% probability and short contacts are indicated by dotted lines. Coordinates of this fragment were applied in further DFT calculations. Right: Contour line diagram of the Laplacian distribution $\nabla^2 \rho(r)$ (cross-section along the plane Pt–Cl–I) was received as a result of the topological analysis of the electron density distribution in the model structure. Bond paths are shown as brown lines, selected zero-flux surfaces – as blue lines, bond critical points (3, -1) are shown in blue, nuclear critical points (3, -3) – in pale brown, length unit – Å.

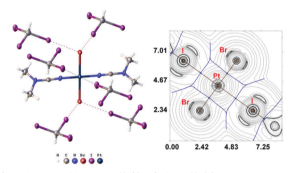


Fig. 2 Left: View of fragment 1b·(CHI₃)₆ from 1b·2CHI₃. Thermal ellipsoids are shown with 50% probability and short contacts are indicated by dotted lines. Coordinates of this fragment were applied in further DFT calculations. Right: Contour line diagram of the Laplacian distribution $\nabla^2 \rho(r)$ (cross-section along the plane Pt-Br-I) was received as a result of the topological analysis of the electron density distribution in the model structure. Bond paths are shown as brown lines, selected zero-flux surfaces – as blue lines, bond critical points (3, -1) are shown in blue, nuclear critical points (3, -3) – in pale brown, length unit – Å.

Complex 4 was co-crystallized with CHI₃ in a 1:2 molar ratio from chloroform solution forming 4.2CHI3. Apart from two types of HI₂C-I···Cl-Pt XB (Chart 1, A), this solid also contains the bifurcated $HI_2C-I\cdots\eta^2_{(Pt-Cl)}$ metal-involving XB (C). Both distances $(d(I \cdot \cdot Pt) = 3.7060(7) \text{ Å and } d(I \cdot \cdot Cl) = 3.6974(19) \text{ Å})$ are slightly less than the corresponding sums of Bondi's²⁹ vdW radii $(R_{vdW}(Pt) + R_{vdW}(I) = 3.73 \text{ Å}; R_{vdW}(Cl) + R_{vdW}(I) = 3.73 \text{ Å})$ and both contact angles are not much different from 160° $(\angle(C-I\cdots Pt) = 158.4(2)^{\circ}, \angle(C-I\cdots Cl) = 163.4(2)^{\circ})$ (see Section XR2 of the ESI†). The formation of the I···Pt and I···Cl noncovalent interactions was confirmed theoretically for the 4-(CHI₃)₆ model structure by using Bader's AIM analysis (Fig. 3, see also TH1 of the ESI \dagger). The estimated energies E_{int} of these contacts are in good agreement with the energy range for XB $(1.3-1.4 \text{ kcal mol}^{-1} \text{ for HI}_2\text{C-I}\cdots\text{Cl and } 1.6 \text{ kcal mol}^{-1} \text{ for}$ HI₂C-I···Pt). To the best of our knowledge, these weak ChemComm

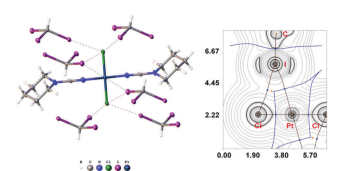


Fig. 3 Left: View of fragment $4\cdot (\text{CHI}_3)_6$ from $4\cdot 2\text{CHI}_3$. Thermal ellipsoids are shown with 50% probability and short contacts are indicated by dotted lines. Coordinates of this fragment were applied in further DFT calculations. Right: Contour line diagram of the Laplacian distribution $\nabla^2 \rho(r)$ (cross-section along the plane Pt–Cl–I) was received as a result of the topological analysis of the electron density distribution in the model structure. Bond paths are shown as brown lines, selected zero-flux surfaces – as blue lines, bond critical points (3,-1) are shown in blue, ring critical points (3,+1) – in orange, nuclear critical points (3,-3) – in pale brown, length unit – Å.

interactions are the first example of metal-involving bifurcated XB. It is noteworthy that examples of metal-free bifurcated XB 6 and metal-involving bifurcated hydrogen bonding 30 are known. Taking into account that the observed distances within the $HI_2C\text{--}I\cdots\eta^2_{(Pt\text{--}Cl)}$ moiety are only slightly less than their van der Waals separations, more examples of metal-involving bifurcates are needed to exclude completely the packing effects on the bonding.

Variable temperature XRD experiments for the same singlecrystal of 4·2CHI₃ were performed at 100, 150, 200, 250, and 300 K. An inspection of the obtained data indicates that the heating leads only to the elongation of the intermolecular distances, whereas the covalent bond lengths remain almost the same. The most significant changes were detected for the bifurcated $HI_2C-I \cdot \cdot \cdot \eta^2_{(Pt-Cl)}$ contacts (by 0.08 Å upon an increase in temperature from 100 to 300 K), whereas the HI₂C-I···Cl-Pt halogen bonds were lengthened by only 0.05 and 0.03 Å, correspondingly (see also XR3 of the ESI†). Bader's AIM analysis (TH2 of ESI†) for the 4·(CHI₃)₆ model systems at each temperature indicated above showed the gradual reduction of the estimated energies of all halogen bonds. Apart from that one of the bond critical points (between Cl and I) in the bifurcated XB disappeared on going from 250 to 300 K. The reversibility of the thermal transformation was proved by three consecutive XRD experiments for the same single-crystal at 100 K, 300 K, and again at 100 K. This experiment gives evidence supporting the thermally induced reversible $HI_2C-I \cdots \eta^2_{(Pt-Cl)} \rightleftharpoons$ $HI_2C-I \cdot \cdot \cdot \eta^1_{(Pt)}$ transformation.

Complexes 2 and 3 were co-crystallized with CHI $_3$ in a 1:2 molar ratio from chloroform solutions forming 2·2CHI $_3$ and 3·2CHI $_3$, respectively. These associates were found to form only three types of HI $_2$ C-I···Cl-Pt XB in each case (Fig. 4 and 5). In all cases ($1a\cdot\frac{1}{2}$ CHCl $_3\cdot1\frac{1}{2}$ CHI $_3$, $1b\cdot2$ CHI $_3$, 2·2CHI $_3$, 3·2CHI $_3$, and 4·2CHI $_3$), the formation of halogen bonds with the halide ligands was confirmed theoretically by using Bader's AIM

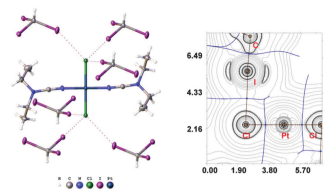


Fig. 4 Left: View of fragment $2\cdot (\text{CHI}_3)_6$ from $2\cdot 2\text{CHI}_3$. Thermal ellipsoids are shown with 50% probability and short contacts are indicated by dotted lines. Coordinates of this fragment were applied in further DFT calculations. Right: Contour line diagram of the Laplacian distribution $\nabla^2 \rho(r)$ (cross-section along the plane Pt–Cl–I) was received as a result of the topological analysis of the electron density distribution in the model structure. Bond paths are shown as brown lines, selected zero-flux surfaces – as blue lines, bond critical points (3,-1) are shown in blue, nuclear critical points (3,-3) – in pale brown, length unit – Å.

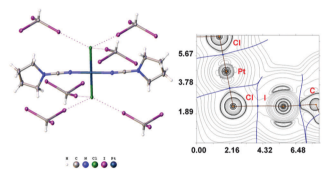


Fig. 5 Left: View of fragment ${\bf 3}\cdot ({\sf CHI}_3)_6$ from ${\bf 3}\cdot {\sf 2CHI}_3$. Thermal ellipsoids are shown with 50% probability and short contacts are indicated by dotted lines. Coordinates of this fragment were applied in further DFT calculations. Right: Contour line diagram of the Laplacian distribution $\nabla^2 \rho(r)$ (cross-section along the plane Pt–Cl–I) was received as a result of the topological analysis of the electron density distribution in the model structure. Bond paths are shown as brown lines, selected zero-flux surfaces – as blue lines, bond critical points (3,-1) are shown in blue, nuclear critical points (3,-3) – in pale brown, length unit – Å.

analysis (XR2 and TH1 of the ESI†). The estimated energies ($E_{\rm int}$) of this type of weak interaction are in the range of 1.9–3.5 kcal mol⁻¹. Only the HI₂C-I···Cl⁻ and the HI₂C-I···Br⁻ weak interactions were previously described,³¹ but no single example of the XB between metal-coordinated chloride or bromide and CHI₃ has been previously reported.

The elongation of the Pt–X (X = Cl, Br) bonds in the CHI₃-associates was detected when XRD data of the associates were compared with those of CHI₃-free 1a, 32 1b, and 4^{33} (see XR1 in the ESI†). The Pt–Cl distances in $2\cdot 2$ CHI₃ and $3\cdot 2$ CHI₃ are longer than in CHI₃-free complexes 1a and 4. Although these differences are not too large (0.01–0.03 Å), the observed elongation indirectly point out (Chart 2) to the presence of the HI₂C–I···X–Pt and HI₂C–I···Pt contacts (see XR2 and TH1 in the ESI†) additionally supporting our conclusions.

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Chart 2 The Pt–X bond elongation caused by the $HI_2C-I\cdots X-Pt$ and $HI_2C-I\cdots Pt$ weak interactions.

In this communication, we reported the first examples of definite and unambiguous identification of metal-involving XB with halocarbons, *viz.* between Pt^{II} and CHI₃. Various Pt^{II} centers are known to participate in donor–acceptor interactions with Lewis acids (B^{III} and S^{IV} compounds^{34,35}), but in all cases these donor–acceptor bonds are too strong to be treated as non-covalent bonds. The most relevant example of XB is weak contacts between the Pt^{II} and Rh^I centers and iodine.^{16–19} Examples of the X···M short contacts given in the Introduction have not been clearly identified as metal-involving XB. We plan to continue our studies in the area of metal-involving XB and to expand the range of studied systems to other X's and X-containing species (*e.g.* Br) and metal centers (*e.g.* Pd^{II}) and studies in these directions are underway in our group.

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