Influence of Halogen Bonding Interaction on Supramolecular Assembly of Coordination Compounds; Head-to-Tail N…X Synthon Repetitivity

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Supporting Information



ABSTRACT: In this study, *N*-(3-halophenyl)-2-pyrazinecarboxamide ligands, L^{3-F} , L^{3-Cl} , L^{3-Br} , and L^{3-I} , carrying a different halogen atom on the phenyl *meta*-position and *N*-phenyl-2-pyrazinecarboxamide ligand, L^{H} , have been employed for the synthesis of 12 mercury(II) complexes, $[HgCl_2(L^{H})]_{n}$, **1**, $[HgCl_2(L^{3-Cl})]_{n}$, **2**, $[Hg_2Cl_4(L^{3-Br})_2]$, **3**, $[Hg_2Cl_4(L^{3-I})_2]$, **4**, $[Hg_2Br_4(L^{H})_2]$, **5**, $[HgBr_2(L^{3-F})]$, **6**, $[HgBr_2(L^{3-Cl})]$, **7**, $[HgBr_2(L^{3-Br})]$, **8**, $[HgBr_2(L^{3-I})]$, **9**, $[Hg_2I_4(L^{H})_2]$, **10**, $[HgI_2(L^{3-Br})]$, **11**, and $[HgI_2(L^{3-I})]_{n}$, **12**. Interestingly, structural analysis clearly shows that, by the replacing of coordinated anions from chloride with bromide and iodide in each series containing the same ligand, the coordination geometry and structural motif of the resulting compounds have been dramatically affected. One of the common features in the crystal structures of these complexes is that there is a strong tendency to form halogen bonding synthons between adjacent halophenyl and pyrazine rings. The influence of these halogen bonding interactions on the supramolecular assemblies has been discussed with the help of geometrical analysis and theoretical calculations. The X…N halogen bonding distances are 2.5–9.4% shorter than the sum of the van der Waals radii of nitrogen and halogen atoms. Theoretical methods also show the halogen bonding energies within a range of -27.86 to -46.15 kJ·mol⁻¹. In all complexes synthesized here, the pyrazine ring is coordinated to the mercury(II) ion through the N atom *syn* to the carbonyl. Therefore, the second common feature of the crystal structures for complexes studied here is the selectivity of the metal ion coordination site. The halogen bond synthon repetitivity across these compounds and selectivity in the mercury(II) ion coordination site further point to application in the coordination crystal engineering research field.

INTRODUCTION

Structural assemblies in coordination compounds rely on a range of directional and nondirectional intermolecular interactions to hold the components together. The most extensively exploited of these interactions is highly directional hydrogen bonding.¹ The influence of $\pi \cdots \pi$, cation $\cdots \pi$, anion $\cdots \pi$, lone-pair $\cdots \pi$, and metal $\cdots \pi$ interactions is also well accepted among supramolecular coordination chemists.² Another interaction that has more recently become of interest as a construction tool in the field of crystal engineering is directional attractive halogen bonding interaction (hereafter referred to as XB). The XB, denoted in the general form D $\cdots X-Y$ (Y = C, N, halogen, etc.), refers to any noncovalent interaction involving a halogen as the acceptor of electron density, a Lewis acid, and an electrondonating atom, D, as a Lewis base.³ Various types of XB, including R-X…Nu, X₂…Nu, R-X1…X2-R, R-X1…X2-M, and R-X1…X2⁻, where Nu is a nucleophile, R-Xi is a haloalkyl group, X₂ is a dihalogen molecule, Xi⁻ is a halide anion, and M is a metal, have been studied in detail.⁴ Although the role of XB in metalcontaining crystal structures and analogies and differences with respect to hydrogen bonds have been reviewed by Brammer⁵ and

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Bertani and co-workers,⁶ compared to organic systems, to our knowledge, far less attention has been paid to *systematic* studies that examine the effect of halogen bonding on the supramolecular aggregation of coordination complexes. To make progress in controlling specific interactions in the solid state of coordination compounds, it is desirable to synthesize a series of coordination compounds of predetermined chemical structure, allowing for a comparison between assemblies with specific and controllable changes to their molecular structure.

Recently, the possibility of using C=O…X (X = Cl, Br, I) halogen bonds for directed assembly in copper(II) coordination compounds containing β -diketonate moieties bearing chloro, bromo, and iodo substituents has been systematically studied by Aakeröy and co-workers.⁷ In 2008, competition between coordination network formation and halogen bond network formation using copper(II) iodobenzoate units was reported by Brammer and co-workers.⁸

On the other hand, with the exploration of how weak interactions work individually or cooperatively in the supramolecular structure, the identification of reproducible synthons in the supramolecular structures is an important line of research.⁹ In this regard, even failed attempts at supramolecular assembly can offer some insight into discovering how intermolecular interactions can be ranked in terms of structural effectiveness. Assembly of discrete copper(II) and cobalt(II) coordination compounds into 1D linear chains using halogen bond interactions has been investigated by Aakeröy, and the results showed that attempts at creating desired architectures had failed.¹⁰ In Ward and co-worker's paper, the repetitivity of C–I···NC(Ru) halogen bonds in [Ru(bipy)(CN)₄]^{2–} salts containing iodinated cations has been shown, while the analogous brominated cations do not exhibit halogen bond interactions.¹¹

brominated cations has been shown, this the line and g_{1} interactions.¹¹ During our research effort on $\pi - \pi$ stacking,^{12a} C-H… π^{12b} interaction, and ligand substituent effect^{12c} in the crystal packing of mercury coordination compounds containing pyrazine carboxamide ligands, that the authors are developing, the effect of halogen bonding on the three-dimensional supramolecular architecture of coordination compounds has herein been reported. In this Article, a series of N-(3-halophenyl)-2-pyrazinecarboxamide ligands, L^{3-F} , L^{3-Cl} , L^{3-Br} , and L^{3-I} , and Nphenyl-2-pyrazinecarboxamide ligand, L^H (Scheme 1), carrying a different halogen atom in the phenyl meta-position, have been employed for the synthesis of mercury(II) complexes in order to get insights into the effect of halogen bonding interactions on the structural assembly of the complexes. Because of the freedom of C-C and C-N single bond rotations in these ligands, the pyrazine and halophenyl rings can be freely twisted to meet the requirements of the coordination geometries of metal atoms and intermolecular interactions in the assembly process. Twelve Hg(II) complexes of these ligands, $[HgCl_2(L^H)]_n$, 1, $[HgCl_2 [L^{3-Cl}]_{,v}$ 2, $[Hg_2Cl_4(L^{3-Br})_2]$, 3, $[Hg_2Cl_4(L^{3-I})_2]$, 4, $[Hg_2Br_{4^-}(L^H)_2]$, 5, $[HgBr_2(L^{3-F})]$, 6, $[HgBr_2(L^{3-Cl})]$, 7, $[HgBr_2(L^{3-Br})]$, 8, $[HgBr_2(L^{3-1})], 9, [Hg_2I_4(L^H)_2], 10, [HgI_2(L^{3-Br})], 11, and$ $[HgI_2(L^{3-1})]_n$, 12, have been prepared by the reaction of equimolar quantities of mercury(II) halides (chloride, bromide, and iodide) in a methanol solution (Scheme 1). The structural details show that the presence of Cl, Br, and I atoms in the metaposition of the N-phenyl ring significantly influences the structural assembly of the resulting mercury(II) complexes and the ligands are pointed toward the adjacent molecules to generate simple or head-to-tail $N \cdots X$ (X = Cl, Br, I) halogen bond synthons.

Scheme 1. Synthetic Route of 1–12 and Presence of Halogen Bond Synthons (Red Band)



RESULTS AND DISCUSSION

Synthesis. The ligands L^H, L^{3-F}, L^{3-Cl}, L^{3-Br}, and L^{3-I} were prepared by simply mixing the same equivalents of *meta*-haloaniline or aniline and pyrazinecarboxylic acid in pyridine in the presence of triphenyl phosphite.¹³ The reaction of equimolar

Table 1. Structural Data and Refinement Parameters for Compounds 1-12

]	L	2			3		4		5	6	
formula	C ₁₁ H ₉ Cl ₂ HgN ₃ O		Cl ₂ HgN ₃ O C ₁₁ H ₈ Cl ₃ HgN		$N_{3}O = C_{22}H_{16}Br_{2}Cl_{4}Hg_{2}N_{6}O_{2}$		$C_{22}H_{16}Cl_4Hg_2I_2N_6O_2\\$		$C_{22}H$	$I_{18}Br_4Hg_2N_6O_2$	C ₁₁ H ₈ Br ₂ FH	gN ₃ O
fw	470.70		505.14		1099.18		1193.18		1119.20		577.59	
cryst syst	monoclii	monoclinic			monoclinic		monoclinic		monoclinic		orthorhombic	
space group	$P2_1/c$		$P\overline{1}$		$P2_{1}/c$		$P2_{1}/c$		$P2_1/n$		Pbca	
a (Å)	7.1019(1	1)	3.9330(8)		16.4086(2)		16.5955(12)		7.2762(10)		6.7945(4)	
b (Å)	11.2071((19)	12.012(2)		13.2074(2)		13.4945(7)		15.4995(14)		12.5409(12)	
c (Å)	16.435(2	2)	15.803(3)		13.1793(2)		13.4342(9)		12.0093(16)		33.507(2)	
α (deg)	90		109.56(3)		90		90		90		90	
β (deg)	91.144(1	2)	96.68(3)		104.4990(1	10)	104.95	104.954(6)		21(11)	90	
γ (deg)	90		98.47(3)		90		90		90		90	
$V(Å^3)$	1307.8(3	3)	684.8(2)		2765.19(7))	2906.7	7(3)	1353	.5(3)	2855.1(4)	
$D_{\text{calc}} (\text{Mg} \cdot \text{m}^{-3})$	2.391		2.450		2.640		2.727		2.746	5	2.688	
Ζ	4		2		4		4		2		8	
$\mu \text{ (mm}^{-1})$	12.167		11.816		14.402		13.076	5	17.26	6	16.385	
F(000)	872		468		2016		2160		1016		2096	
2θ (deg)	58.40		54.00		58.60		58.60		58.42	2	58.66	
R (int)	0.0616		0.1097		0.0519		0.0729)	0.100)8	0.2859	
GOF	1.12		1.08		1.04		1.12		1.20		1.38	
$R_1^a (I > 2\sigma(I))$	0.044		0.047		0.029		0.052		0.054		0.179	
$wR_2^{b} (I > 2\sigma(I))$	0.140 0.115		0.115	0.053		0.124		0.139		0.360		
final electron density (e∙Å ⁻³)	-3) 1.86, -1.97		1.63, -1.66		1.26, -1.20		2.06, -2.22		1.77, -3.76		1.74, -1.44	
CCDC No.	901972		901975		901970		91246	5	9019	71	901981	
		7		8	3	9		10		11	12	
formula		C ₁₁ H ₈ Br ₂ C	lHgN ₃ O	$C_{11}H_8Br$	₃ HgN ₃ O	C ₁₁ H ₈ Br ₂ HgII	N ₃ O	$C_{22}H_{18}Hg_2I_4N_6O$	2 (C ₁₁ H ₈ BrHgI ₂ N ₃ O	C ₁₁ H ₈ Hgl	I ₃ N ₃ O
fw		594.04		638.49		685.49		1307.20	7	732.49	779.49	
cryst syst		orthorhom	bic	monoclii	nic	monoclinic		monoclinic	1	nonoclinic	monoclini	ic
space group		Стса		C2/m		C2/m		C2/c	(C2/m	C2/m	
a (Å)		6.6796(13)		13.3681((17)	13.681(13)		7.2849(11)	1	13.609(2)	13.9063(1	(3)
b (Å)		13.257(3)		6.6595(9))	6.6901(17)		12.5265(14)	e	5.9070(10)	6.9034(7)	1
c (Å)		33.744(7)		17.1358((18)	17.162(4)		33.5265(4)	1	17.596(3)	17.5611(1	17)
β (deg)		90		103.551((9)	104.740(18)		91.415(11)	1	102.494(12)	103.402(7	7)
V (Å ³)		2988.1(10)		1483.0(3	3)	1519.0(6)		3043.7(7)	1	1614.8(4)	1640.0(3)	1
$D_{\text{calc}} (\text{Mg} \cdot \text{m}^{-3})$		2.641		2.860		2.997		2.853	3	3.013	3.157	
Ζ		8		4		4		4	4	4	4	
$\mu (\text{mm}^{-1})$		15.823		18.455		17.419		14.162	1	15.824	15.028	
F(000)		2160		1152		1224		2320	1	1296	1368	
2θ (deg)		58.50		58.64		58.74		58.56	5	58.44	58.34	
R (int)		0.1265		0.1275		0.1489		0.1118	(0.1049	0.0725	
GOF		1.17		1.15		1.19		1.07	1	1.26	1.09	
$R_1^a (I > 2\sigma(I))$		0.059		0.094		0.077		0.131	(0.063	0.047	
$wR_2^b (I > 2\sigma(I))$		0.141		0.239		0.193		0.358	(0.175	0.126	
final electron density (e	e∙Å ^{−3})	2.27, -2.54	ł	2.19, -2	.71	3.25, -2.12		1.70, -1.31	2	2.12, -4.86	2.56, -2.3	30
CCDC No.		901978		901973		901974		901980	ç	901979	901977	
${}^{a}R_{1} = \sum F_{o} - F_{c} /2$	$\sum F_{o} $. ^b w	$vR_2 = \left[\sum\right]$	$w(F_{\rm o}^{2}-F_{\rm o})$	$(2^{2})^{2})/\sum u$	$w(F_{\rm o}^{2})2]^{1/2}$							

amounts of these ligands and HgX_2 (X = Cl, Br, and I) in methanol gave the corresponding complexes. Slow evaporation of the solvent resulted in air-stable block crystals of 1, 3–5, 8, and 10–12; prism crystals of 2, 7, and 9; and needle crystals of 6, after a few days. Attempts were made to form complexes with $HgCl_2$ and HgI_2 using the L^{3-F} ligand and HgI_2 using the L^{3-Cl} ligand. Yet unfortunately, no mercury-containing species were isolated from the reaction mixtures. Crystallographic data for compounds 1– 12 are listed in Table 1. Selected bond distances and angles are summarized in Table 2.

Structural Analysis of HgCl₂ Complexes, [HgCl₂(L^H)]_n, 1, [HgCl₂(L^{3-Cl})]_n, 2, [Hg₂Cl₄(L^{3-Br})₂], 3, and [Hg₂Cl₄(L³⁻¹)₂], 4. A simple reaction between HgCl₂ and L^H, L^{3-Cl}, L^{3-Br}, and L^{3-I} in methanol afforded well-formed crystals of 1–4. The asymmetric

unit of 1 consists of one crystallographically independent Hg²⁺ ion, one *N*-phenyl-2-pyrazinecarboxamide, L^H, and two chloride ions. As depicted in Figure 1a, in this compound, the highly distorted tetrahedral geometry of the Hg(II) center can be better described as a seesaw structure. The Cl–Hg–Cl angle is 176.17(10)°, where the two Hg–Cl bonds form the plank (Table 2). The angle between the other two bonds (Table 2), which form the pivot, is 90.0(2)°. The Cl–Hg–Cl and N–Hg–O planes are nearly perpendicular, with a dihedral angle of 87.0(4)°. For this complex, the four-coordinate geometry index, τ_4 , of 0.64 fits with a seesaw description.¹⁴

Compound 2 crystallizes in the $P\overline{1}$ space group. The asymmetric unit of 2 consists of one Hg²⁺ ion, two halogen anions, and one crystallographically independent ligand. As

Table 2. Selected Bond Lengths (Å) and Angles (deg) around Mercury(II) for Complexes 1-12

				cor	nplex		
		1	2	3	4	5	6
bond distance	Hg1-X1	2.289(2)	2.3464(18)	2.3106(6)	2.295(2)	2.4147(10)	2.443(4)
	Hg1-X1		$3.047(2)^{a}$				
	Hg1-X2	2.303(2)	2.3375(19)	2.3089(7)	2.294(2)	2.4293(0)	2.462(4)
	Hg1-X2		$3.044(2)^{b}$				
	Hg1-O1	$2.748(6)^a$				$2.949(7)^{c}$	
	Hg1-N2	2.656(8)	2.575(5)	2.573(2)	2.599(7)	2.514(7)	2.49(3)
	Hg2-X3			2.3340(7)	2.300(2)		
	Hg2-X4			2.3088(7)	2.325(2)		
	Hg2-O1			2.815(2)	2.822(7)		
	Hg2–N5			2.516(2)	2.551(9)		
bond angle	X1-Hg1-X2	176.17(10)	173.26(6)	169.44(2)	169.81(9)	161.93(4)	163.90(15)
	X1-Hg1-X2		$92.71(6)^a$				
	X1-Hg1-X1		$86.82(6)^{b}$				
	X2-Hg1-X2		$92.98(6)^{b}$				
	X1-Hg1-O1	$92.0(1)^{a}$				105.8(1)	
	X2-Hg1-O1	$86.6(1)^a$				82.8(1)	
	X1-Hg1-N2	93.18(18)	93.01(15)	94.30(5)	94.34(17)	101.3(2)	98.2(7)
	X2-Hg1-N2	90.40(17)	93.73(14)	95.36(5)	95.12(17)	95.8(2)	97.9(7)
	X1-Hg1-N2		$91.5(5)^{a}$				
	X2-Hg1-N2		$93.7(1)^{b}$				
	N1-Hg1-O1	$90.0(2)^{a}$				$79.4(2)^{c}$	
	Hg1-01=C5	$125.4(5)^{a}$				$114.8(5)^{c}$	
	X3-Hg2-X4			166.36(3)	166.00(9)		
	X3-Hg2-N5			93.57(6)	99.32(17)		
	X3-Hg2-O1			98.53(5)	89.4(1)		
	X4-Hg2-N5			99.15(6)	93.77(17)		
	X4-Hg2-O1			88.49(5)	97.3(1)		
	N5-Hg2-O1			80.48(7)	82.34(16)		
	Hg2-01=C5			106.8(2)	109.0(5)		
	0			con	nplex		
		7	8	9	10	11	12
bond distance	Hg1-X1	2.4639(12)	2.459(2)	2.4569(18)	2.622(2)	2.6381(10)	2.6394(8)
	Hg1-X1				$3.512(3)^d$	$3.4957(5)^e$	$3.5182(4)^{f}$
	Hg1-X2	2.4287(14)	2.424(2)	2.420(2)	2.602(2)	2.5950(11)	25953(8)
	Hg1–N2	2.469(10)	2.480(13)	2.449(13)	2.54(3)	2.474(10)	2.484(7)
bond angle	X1-Hg1-X2	161.05(5)	160.26(9)	159.07(8)	163.17(10)	156.92(4)	155.95(3)
0	X1-Hg1-X2					$93.63(3)^{e}$	$94.46(2)^{f}$
	X1-Hg1-X1				$92.61(8)^d$	$89.84(3)^{e}$	$90.06(2)^{f}$
	X1-Hg1-N2	98.3(3)	98.2(4)	98.8(3)	98.4(7)	102.0(2)	102.3(2)
	X2-Hg1-N2	100.6(2)	101.6(4)	102.1(3)	97.6(7)	101.0(2)	101.8(2)
	X1-Hg1-N2	~ /	~ /	~ /	94.24(8)	81.3(2) ^e	79.1(2) ^f
'Symmetry codes: 1	1 + x, y, z. ^b Symmet	ry codes: $-1 + x$,	<i>y, z.</i> ^{<i>c</i>} Symmetry	codes: -x, 2 -	y, $2 - z$. ^d Symm	etry codes: 1/2 -	-x, 1/2 - y, -z

^eSymmetry codes: 3/2 - x, -1/2 + y, -z. ^fSymmetry codes: 1/2 - x, -1/2 + y, 1 - z.

depicted in Figure 1b, the coordination geometry around the metal center is a square-based pyramid (SBP) (Table 2) with a trigonality index, τ_{5} ,¹⁵ of 0.03. In this structure, the plane of the square-based pyramid is occupied by four halogen anions (Table 2). The apical position is occupied by a nitrogen atom from the pyrazine ring of the L^{3-Cl} ligand at a normal distance of 2.575(5) Å. As it is clear from the τ_5 value and geometrical parameters around the central metal atom (Table 2), the coordination geometry around Hg(II) is almost perfect SBP.

Dinuclear isostructural compounds **3** and **4** consist of two different mercury environments. In these compounds, each asymmetric unit consists of two crystallographically independent mercury centers, four chloride ions, and two neutral L^{3-Br} or L^{3-1} ligands (Figure 1c,d). The Hg1 atom is located in a three

coordination environment, chelating with two chloride atoms and one pyrazine nitrogen atom from the ligand L^{3-X} (X = Br, I) (Table 2). The Cl–Hg–Cl angle of 169.44(2)° and 169.81(9)° for 3 and 4, respectively, shows that the geometry around the metal center is slightly distorted T-shaped. In these compounds, the L^{3-X} ligand acts as an angular bridge *via* the carbonyl oxygen and the pyrazine nitrogen atoms to link Hg1 and Hg2 ions. The coordination geometry around the Hg2 atom in compounds 3 and 4 can be described as a seesaw structure with a τ_4 index of 0.67 for both complexes, with nearly linear Cl–Hg–Cl (Table 2) forming the plank, and N–Hg–O (Table 2) for 3 and 4 forming the pivot. The dihedral angles between the Cl–Hg–Cl and N– Hg–O planes are 83.78(6)° and 84.91(7)° for 3 and 4,



(d)

Figure 1. Portion of the structure of coordination compounds formed between HgCl₂ and L^H, **1** (a), L^{3-Cl}, **2** (b), L^{3-Br}, **3** (c), and L^{3-I}, **4** (d), showing coordination geometry around the central metal. Displacement ellipsoids are shown at 30% probability level. Symmetry codes: (a) (i) 1 + *x*, *y*, *z*; (b) (i) 1 + *x*, *y*, *z*; (ii) -1 + x, *y*, *z*.

respectively. We propose a simple geometry index for almost planar three-coordinate complexes, trigonal-planar index, τ_3 , eq 1

$$\tau_3 = \left| \left[(\alpha + \beta + \gamma)/360 \right] - \left| (\alpha - 120)/60 \right| \right|$$
(1)

inspired by Houser's four-cooordinate τ_4 index¹⁴ and Addison and Reedijk's five-coordinate τ_5 index.¹⁵ In this equation, α is the largest angle in the three-coordinated complex. The values of τ_3 will range from 1.00 for a perfect trigonal-planar geometry, since [(120 + 120 + 120)/360] - [(120 - 120)/60] = 1 - 0 = 1, to zero for a perfect T-shaped geometry, since [(180 + 90 + 90)/360] - [(180 - 120)/60] = 1.00 - 1.00 = 0. Therefore, this parameter can be used for almost planar three-coordinate complexes since trigonal-planar and T-shaped geometries are the extremes. Trigonal-planar index, τ_3 , for Hg1 in compound 3 and 4 is 0.17 for both complexes. In 1, the adjacent mercury atoms are linked by C=O-Hg bonds to form a 1D linear polymeric chain spanning along the *a* axis (Figure 2a). In this compound, the phenyl ring involved in



Figure 2. Representation of 1D linear polymeric chain in 1 (a), and a side view representation of 1 in the *a* direction showing the association of the adjacent molecules in the chain, through an additional head-to-tail C_{phen}-H···Cl-Hg and C=O···H-C_{phen} nonclassical hydrogen bonds and Hg···Cl contacts and Hg··· π_{phen} intermolecular interactions (b). In (b), different colors show different adjacent linear chains.

the *intramolecular* $\pi - \pi$ stacking interaction with the adjecent pyrazine rings is arranged in such a way that the angle between the plane (containing C-CO-N fragment) normal and the O-Hg vector (for geometrical definition, see ref 12a) reaches about 36.81(6)°. In $\pi_{\text{phen}} \cdots \pi_{\text{pyz}}$ interactions, the centroid-centroid distance is 3.569(5) Å. Such $\pi - \pi$ interaction effects on the primary structure directing the coordination geometry around Hg(II), which contain similar ligands to those discussed in this paper, have previously been reported in detail by some of us.^{12a} As shown in Figure 2b, these 1D linear chains are further linked to each other from one side by head-to-tail dimeric C_{phen}-H…Hg-Cl and C_{phen}-H…O=C nonclassical hydrogen bonds (Table S1, Supporting Information), and from the other side by head-to-tail Hg…Cl contacts and Hg… π_{phen} intermolecular interactions (Table 3). Coordination geometry and intermolecular interactions of compound 1 are illustrated in Scheme 2.

Within the asymmetric unit of **2**, each mercury coordinates to the nitrogen donor atom of the pyrazine ring, the N atom *syn* to the carbonyl, of the L^{3-Cl} ligand, while each chloride atom in the basal plane bridges two adjacent metal centers to generate a 1D double-chain motif in the *a* direction (Figure 3a). The planar organic ligands stack along both sides of the HgCl skeleton, and the distance between their mean planes is 3.933(1) Å (Figure 3a). These 1D chains are further linked to adjacent ones by Hg...Cl contacts to generate a double chain in the *a* direction (Table 3). The resulting double chains are further linked to adjacent chains from one side by head-to-tail dimeric C_{phen}-

complex	D–H…A	$d(D \cdots A)$ (Å)	symmetry
1	Hg1…Cl2	3.203(2)	1 - x, 2 - y, -z
	Hg1 $\cdots \pi_{\rm phen}$	3.548(2)	1 + x, $3/2 - y$, $1/2 + z$
2	Hg1…Cl1	3.198(2)	2 - x, 2 - y, 1 - z
3	Hg1…Cl2	3.288(1)	-x, 1-y, -z
	Hg2…Cl4	3.247(1)	-x, 1-y, -z
	$\pi_{\rm phen} \cdots \pi_{\rm pyz}$	3.449(4)	
	Hg1 $\cdots \pi_{\rm phen}$	3.452(1)	1- <i>x</i> , <i>y</i> , <i>z</i>
4	Hg1…Cl3	3.324(3)	1 - x, 1 - y, 1 - z
	Hg2…Cl2	3.324(3)	1 - x, 1 - y, 1 - z
	$\pi_{\mathrm{phen}} \cdots \pi_{\mathrm{pyz}}$	3.516(6)	
	Hg1 $\cdots \pi_{\rm phen}$	3.517(4)	x, y, -1 + z
5	$\pi_{\mathrm{phen}} \cdots \pi_{\mathrm{phen}}$	3.847(7)	1/2 - x, $1/2 + y$, $5/2 - z$
6	Hg1…Br2	3.378(4)	1/2 + x, $1/2 - y$, $2 - z$
	Hg1…Br2	3.454(4)	-1/2 + x, $1/2 - y$, $2 - z$
	amide $\cdots \pi_{\text{phen}}$	3.475(5)	1/2 + x, y, 5/2 - z
7	Hg1…Br1	3.401(1)	-1/2 + x, $1/2 - y$, $-z$
	amide $\cdots \pi_{\text{phen}}$	3.373(1)	1/2 - x, y, 1/2 - z
8	Hg1…Br1	3.411(1)	1/2 - x, $-1/2 + y$, $1 - z$
	amide $\cdots \pi_{\text{phen}}$	3.363(1)	1/2 - x, $-1/2 + y$, $2 - z$
9	Hg1…Br1	3.443(1)	1/2 - x, -1/2 + y, -z
	amide $\cdots \pi_{\text{phen}}$	3.390(1)	1/2 - x, $-1/2 + y$, $1 - z$
10	$\pi_{\rm phen} \cdots \pi_{\rm pyz}$	3.661(8)	-x, y, 1/2 - z

Table 3. Hg···X, Hg··· π , Amide··· π , and π ··· π Distances for Compounds 1–10

H···O=C and C_{pyz} -H···Cl-Hg nonclassical hydrogen bonds (Figure 3b; Table S1, Supporting Information). The coordination geometry and the intermolecular interactions of compound **2** are illustrated in Scheme 2.

In isostructural **3** and **4**, the two independent dinuclear moieties are closely packed through head-to-tail Hg…Cl contacts (Table 3) to generate dimeric units (Figure 4a). Adjacent dimeric units are further linked to each other by head-to-tail N…Br/I halogen bonds (Figure 4b). In the packing of these complexes, the overall supramolecular structure results from the linkage of neighboring units of Hg… π_{phen} , C_{phen}-H…O=C, and C_{phen}-H…Cl-Hg with nonclassical hydrogen bonds (Figure 4c). Coordination geometries and intermolecular interactions of compounds **3** and **4** are illustrated in Scheme 2.

Structural Analysis of HgBr₂ Complexes, $[Hg_2Br_4(L^H)_2]$, 5, $[HgBr_2(L^{3-F})]$, 6, $[HgBr_2(L^{3-Cl})]$, 7, $[HgBr_2(L^{3-Br})]$, 8, $[HgBr_2(L^{3-I})]$, 9. The reaction of 1 equiv of HgBr₂ with 1 equiv of L^H in methanol produces a dinuclear mercury complex, 5, with a 12-membered ring (Figure 5a). This compound displays 2/*m* point symmetry in the solid state, making half of the molecules crystallographically unique. Metal is in a seesaw geometry, with a four-coordinate geometry index, τ_4 , of 0.64, coordinated by two Br atoms (Table 2) at the plank and one pyrazine nitrogen atom of the L^H ligand and one carbonyl oxygen atom of the second L^H ligand (Table 2), which form the pivot. The recent L^H ligand acts as an angular bridge *via* carbonyl oxygen and pyrazine nitrogen

Scheme 2. Representation of Coordination Geometry and Intermolecular Interactions of Compounds 1-12





Figure 3. Representation of 1D double chain (a), and a side view representation of 2 (b), showing the association of the adjacent molecules in the chains, through head-to-tail C_{phen} -H···O=C and C_{pyz} -H···Cl-Hg nonclassical hydrogen bonds and head-to-tail N···Cl halogen bonds. Different colors show different adjacent linear chains. Halogen bonds are shown in red (b).

atoms to link Hg1 and Hg1ⁱ (symmetry code: (i) x, 2 - y, 2 - z) ions. The Br-Hg-Br and N-Hg-O planes are nearly perpendicular, with a dihedral angle of $80.17(9)^\circ$.

The coordination geometries around the Hg atom in **6**–**9** are similar and made up of discrete neutral HgBr₂L^{3-X} units. Figure 5b–e depicts the representative molecular structure showing the arrangement about the Hg(II) center for **6**–**9**, and selected bond distances and angles are listed in Table 2. The Hg atom lies in a T-shaped structure defined by two bromide atoms and one the pyrazine nitrogen atom of the L^{3-X} ligand (Table 2). Trigonalplanar indexes, τ_3 , for these compounds are 0.27, 0.32, 0.33, and 0.35, respectively (Table 4). The angle between the Br–Hg–Br plane and the pyrazine ring for **6** is 6.87(9)° and for 7–**9** is almost 0.00°.

In 5, the structure is stabilized by an ordered network of intermolecular $\pi \cdots \pi$ interactions that exist between the pyrazine and phenyl rings (Figure 6, Table 3). In these $\pi_{phen} \cdots \pi_{pyz}$ interactions, the centroid–centroid distance is 3.847(5) Å. As shown in Figure 6, these $\pi \cdots \pi$ interactions cooperate with C_{phen} – $H \cdots Br$ –Hg and C_{pyz} – $H \cdots Br$ –Hg to further link to each other from one side by head-to-tail dimeric Hg–Cl \cdots H– C_{phen} nonclassical hydrogen bonds (Table S1, Supporting Information). The coordination geometry and intermolecular interactions of compound 4 are illustrated in Scheme 2.

In compounds **6–9**, discrete units are linked to adjacent ones by Hg…Br contacts to generate a double-stranded stair motif (Figures 7a, 8a, and 9a). In these compounds, the Hg…Br contacts of 3.378(4) and 3.454(4) Å for **6** and 3.401(1), 3.411(1), and 3.443(1) Å for **7–9**, respectively (Table 3), are comparable to those previously reported (3.385(1) Å) by Wu et al.^{16a} The resulting double chains are further linked to adjacent chains by π_{phen} —NHCO– interactions (Figures 7a and 8a and Table 3). In 6, these chains are further linked to adjacent chains by C_{phen} -H···Br-Hg and C_{pyz} -H···F-C intermolecular interactions (Table S1, Supporting Information), while in 7–9, cooperation of N_{pyz} ···Cl/Br/I halogen bonds and C_{phen} -H···O=C nonclassical hydrogen bonds are main factors in the generation of supramolecular assemblies (Figures 7b, 8b, and 9b). It is notable that compounds 8 and 9 have isostructural coordination geometries, and intermolecular interactions of compounds 6–9 are illustrated in Scheme 2.

Structural Analysis of Hgl₂ Complexes, [Hg_2I_4(L^H)_2], 10, $[HgI_2(L^{3-Br})]_n$, **11, and [HgI_2(L^{3-I})]_n, 12.** In compound **10**, the metal center has the same seesaw geometry observed in compounds **1** and **5**, with two iodide atoms as the plank, and one iodide and one pyrazine nitrogen atom in the pivot position (Table 2). For this complex, the four-coordinate geometry index, τ_4 , is 0.70. In this compound, each discrete molecular unit consists of a centrosymmetric dinuclear iodo-bridge Hg_2I_2 fragment. This bridged Hg_2I_2 forms a unsymmetric tetra-atomic rhombohedral plane (Figure 10a).

In 1D structures of **11** and **12**, each asymmetric unit consists of one crystallographically independent Hg center, two iodide ions, and one neutral L^{3-Br} or L^{3-1} ligand. In both structures, the coordination geometry around the Hg center is square-based pyramid (SBP) (Figure 10b,c), respectively, with a trigonality index, τ_5 , of 0.09 and 0.03, respectively. In both structures of **11** and **12**, the plane of square-based pyramid is occupied by iodide anions (Table 2). The apical position in both structures is occupied by a nitrogen atom from the pyrazine ring of the L ligand at a normal distance of 2.474(10) and 2.484(7) Å for **11** and **12**, respectively. As it is clear from the τ_5 values and geometrical parameters around the central metal atoms (Table 2), Hg has a slightly distorted SBP environment in both compouds.

In 10, the dinuclear units are further linked to adjacent ones from one side by C_{phen} —H···I—Hg nonclassical hydrogen bonds (Figure 11a) and from the other side by π_{pyz} ··· π_{phen} intermolecular interactions (Figure 11b and Table 3). The coordination geometry and intermolecular interactions of compound 10 are illustrated in Scheme 2.

In 11 and 12, the arrangement of the asymmetric units defines the well-known double-stranded stair motif¹⁶ (Figure 12a). In this stair 1D polymer, the translation axis is parallel to the bdirection. In these compounds, the Hg-I bond distance of 3.4957(5) and 3.5182(4) Å is comparable to that previously reported (3.3773(5) Å) by Popovic et al.^{16b} The I1 atom adopts a μ_3 mode to bridge three Hg atoms, while I2 is coordinated to metal centers as a terminal ligand. The iodide anion in the stair is tricoordinated with distorted T-shaped geometry (diagonal I---I = 4.3733(9) and 4.4005(6) Å for 11 and 12, respectively). The organic ligand moieties approximately preserve their planarity and stack along both sides of the HgI skeleton (Figure 12a). These 1D chains are further linked by head-to-tail N_{pvz}…Br/I halogen bonds in the other direction (Figure 12b). Coordination geometries and intermolecular interactions of compounds 11 and 12 are illustrated in Scheme 2.

Influence of Halide Counteranion and Halogen Bonding on Coordination Geometry and Supramolecular Assembly. Progress in controlling the structural assemblies of coordination complexes in the solid state requires investigation of the families of compounds designed so that the effects of different factors on the resultant supramolecular and structural properties may be systematically delineated. In this regard, the effect of halogen bonding and the effect of different halogen

Article



Figure 4. Side view representation of 3, showing the generation of dimeric moieties through the Hg…Cl short contacts (a), and head-to-tail N…Br halogen bonds between adjacent dimeric units; halogen bonds are shown in red (b). A representation of part of 3 (c), showing the cooperation of head-to-tail N…Br halogen bonds, Hg… π_{phen} , C_{phen}-H…O=C, and C_{phen}-H…Cl-Hg nonclassical hydrogen bonds in generation of three-dimensional packing. Different colors show different adjacent moieties; halogen bonds are shown in red.

anions on the coordination geometry and supramolecular architecture of the 12 coordination compound of mercury(II) have been studied. Much of the difference in the structural motifs can be explained by the changing of the halogen atom on the halophenyl ring, but the role of the different halogen ions should not be neglected. Therefore, in the following discussion, first, we will focus on controlling the effect of the coordinated anion on the structural motif of the resulting coordination compounds.

Influence of Halide Anion on Coordination Geometry and Supramolecular Assembly. It is well-known that anions play an important role in the crystal engineering of self-assembled coordination polymers.¹⁷ Some of us have recently demonstrated that cooperative noncovalent interactions can lead to the formation of one- or two-dimensional architectures under anion-directed self-assembly.¹⁸

Interestingly, structural analysis clearly shows that, by the replacing of coordinated anions from chloride with bromide and iodide in each series containing the same ligand, the coordination geometry and structural motif of the resulting compounds have been dramatically affected. It seems that the size of the coordinated halides would make a substantial contribution to the local mercury coordination environment and can also affect the extended structures.^{17a,18,19} In compounds **1**, **5**, and **10**, where the carboxamide ligand, L^H, is similar, all complexes have the same seesaw coordination geometries (Table 4, Scheme 2). Compound **1** shows a 1D linear chain structure. The situation for

compounds 5 and 10 (where the chloride anion is replaced by bromide and iodide) is quite different from that of compound 1. The different structural motifs between 1, 5, and 10 are seemingly attributed to the anion size effect. Unlike the formation of the 1D linear chain in 1, the bulkier Br and I ions in 5 and 10 are sterically hindered to form a larger aggregate. By replacing the coordinated anion from chloride to bromide and iodide, the supramolecular structure changed from 1D linear chains to dinuclear units (Table 4). Similar results were also observed for complexes 2 and 7. For complexes containing the L^{3-Cl} ligand with HgCl₂ and HgBr₂, 2 and 7, the coordination sphere changes from SBP for 2 to T-shape for 7 (Table 4, Scheme 2). The structural motifs in these compounds are also quite different, and the 1D double chain in 2 is changed to a discrete monomer structure in 7.

For HgX₂ adducts, while the carboxamide ligand is L^{3-Br}, from 3 to 8 and to 11 (where counteranions are chloride, bromide, and iodide, resepectively), the coordination sphere changes from the seesaw/T-shape (for two crystallographically independent metal ions) in 3, to a T-shape in 8 and SBP in 11 (Table 4, Scheme 2). As listed in this table, compounds 3 and 8 show dinuclear and monuclear structures, respectively, whereas compound 11 has a 1D ladder chain structure. In the case of 11, although increasing the size of the anion results in the formation of a larger molecular complex, the study of Hg–I distances indicates that two of the four Hg–I bond lengths in the basal plane, connecting Hg metal



Figure 5. Structure of coordination compounds formed between HgBr₂ and L^H, **5** (a), L^{3-F}, **6** (b), L^{3-Cl}, **7** (c), L^{3-Br}, **8** (d), and L^{3-I}, **9** (e), showing coordination geometry around the central metal. Displacement ellipsoids are shown at 30% probability level. Symmetry codes: (i) -x, 2 - y, 2 - z.

centers through a Hg–I–Hg bridge, are much longer than the others, 3.496(1) Å, compared to the remaining Hg–I bond lengths, and deviate considerably from the average bond length (Table 2). It is noteworthy that a search of the Cambridge Structural Database²⁰ for the Hg–I distance shows that there are only 8 out of a total of 468 hits that have values longer than 3.496 Å. In the recent series, the structural analysis confirms that, as the size of the counteranion increases, the extended structures change significantly. It can be anticipated that the halide steric factor cannot be the only factor dictating the coordination environment and extended structures but certainly plays an important role in this regard.

As compounds 3 and 4, 8 and 9, and 11 and 12 are isostructural, for HgX_2 adducts, while the carboxamide ligand is L^{3-1} (compounds 4, 9, and 12), the same trend as L^{3-Br} (compounds 3, 8, and 11) is observed (Table 4, Scheme 2).

Influence of N···X Halogen Bonding Interaction on Supramolecular Assembly. The compounds studied here were designed to probe the influence of halogen bonding interactions on the supramolecular assembly of similar coordination compounds. Organic ligands with flexible torsions can indeed generate dissimilar conformations with different halogen bond patterns and packing modes. The common feature in the crystal structures of complexes obtained from the reaction between HgX₂ and L^{3-Cl}, L^{3-Br} and L³⁻¹ ligands is the existence of a N···X halogen bond. Thus, a systematic evaluation of supramolecular synthons consisting of N···X halogen bonding interactions in a series of conformationally flexible mercury complexes containing halophenyl pyridazinamides is interesting.

In these compounds, since the halophenyl ring is flexible in rotation around the $N_{amide}-C_{phen}$ bond, this allows subtle conformational adaptation of L^{3-X} to produce additional synthons via the rotation of the halophenyl ring. In all Hg(II) complexes reported here, the coordination of the pyrazine ring occurred through the N atom syn to the carbonyl and the N atom anti to the carbonyl is involved in classical intramolecular N-H…N hydrogen bonds. Because of the presence of such intramolecular hydrogen bonds, the pyrazine-CONH moiety is approximately planar, and C5=O1 and C4-N1 bonds are in the *anti* position. Therefore, the conformational variation of L^{3-X} can be discussed by a dihedral angle of O1=C5-C8-X, which is listed in Table 5 for compounds 1-12. As in compounds 3 and 4, each asymmetric unit consists of two crystallographically independent L^{3-X} ligands; two different values for this angle are listed in Table 5. According to the direction of the C=O and C-X bonds, the L^{3-X} ligands can be arranged in anti and syn conformations (Scheme 3). In the presence of this flexibility, the halogen atom of the halophenyl ring is pointed toward the adjacent molecule to generate different synthons. Two basic halogen bond synthons (Scheme 3) were observed in the mercury complexes examined. Compound 7 adopts the syn conformation while the others, 2, 3, 4, 8, 9, 11, and 12, adopt anti conformations. It is notable that these two different conformations play different roles in the formation of packing. The syn-conformational L^{3-Cl} ligand generates a single Cl...N halogen bond in 7, whereas in the complexes of HgX2 with L^{3-Br} and \tilde{L}^{3-I} , and also HgCl₂ with L^{3-Cl}, the anti-conformational ligands generate a head-to-tail halogen bond synthon (Table 5).

Table 5 provides a summary of the geometrical parameter of the halogen bonding synthons observed in 2-4, 7-9, 11, and 12. These results show that there is a strong tendency to form halogen bonding synthons between adjacent halophenyl and pyrazine rings, in order to stabilize the packing of these compounds. As it is clear from Table 5, halogen bonds show geometries consistent with a donor-acceptor binding model in which the pyrazine nitrogen atom serves as the electron donor and the halocarbon C-X bond serves as the acceptor. In complexes containing the L^{3-Cl} ligand, 2 and 7, contact distances of Cl...N were 3.190(6) and 3.217(5) Å, respectively (Table 5), indicating a 3.3% and 2.5% reduction of the sum of van der Waals radii.²¹ The Br…N halogen bonding distances in compounds 3, 8, and 11, complexes containing the L^{3-Br} ligand, were found between 3.100(3) and 3.251(9) Å, which are 8.8-4.4% shorter than the sum of the van der Waals radii of nitrogen and bromide atoms.²¹

Tab	le 4.	Coordination	Geometries	and	τ_i	Va	lues	of	Compounds	1-	-12	2
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compound	coordination geometry	$ au_i$	polymeric/nonpolymeric
1, $[HgCl_2(L^H)]_n$	seesaw	$\tau_4 = 0.64$	1D linear chain
2 , $[HgCl_2(L^{3-Cl})]_n$	SBP	$\tau_{5} = 0.03$	1D double chain
3, $[Hg_2Cl_4(L^{3-Br})_2]$	T-shape/seesaw	$\tau_3 = 0.17 / \tau_4 = 0.67$	dinuclear
4, $[Hg_2Cl_4(L^{3-I})_2]$	T-shape/seesaw	$\tau_3 = 0.17 / \tau_4 = 0.67$	dinuclear
5, $[Hg_2Br_4(L^H)_2]$	seesaw	$\tau_4 = 0.64$	cyclic dinuclear
6 , $[HgBr_2(L^{3-F})]$	T-shape	$\tau_{3} = 0.27$	monomer
7, [HgBr ₂ (L ^{3-Cl})]	T-shape	$\tau_3 = 0.32$	monomer
8, $[HgBr_2(L^{3-Br})]$	T-shape	$\tau_3 = 0.33$	monomer
9, $[HgBr_2(L^{3-I})]$	T-shape	$\tau_3 = 0.35$	monomer
10 , $[Hg_2I_4(L^H)_2]$	seesaw	$ au_4 = 0.70$	dinuclear
11, $[HgI_2(L^{3-Br})]_n$	SBP	$\tau_{5} = 0.09$	1D ladder chain
12 , $[HgI_2(L^{3-I})]_n$	SBP	$\tau_5 = 0.03$	1D ladder chain



Figure 6. Side view representation of **5**, showing the cooperation of $\pi_{pyz} \cdots \pi_{phen}$ interactions and C_{phen} -H···Br-Hg and C_{pyz} -H···Br-Hg nonclassical hydrogen bonds in generation of three-dimensional packing. Different colors show different adjacent moieties.



Figure 7. Side view representation of **6**, showing the generation of double-stranded stair motifs through the Hg…Br short contacts, and π_{phen} …amide interactions between adjacent chains (a). A representation of part of **6** in the *a* direction (b), showing the cooperation of C_{phen}–H…Br–Hg and C_{pyz}–H…F–C intermolecular interactions in generation of three-dimensional packing. Different colors show different adjacent moieties.



Figure 8. Side view representation of 7, showing the generation of double-stranded stair motifs through the Hg…Br short contacts, and head-to-tail π_{phen} …amide interactions between adjacent chains (a). A representation of part of 7 in the *a* direction (b), showing the cooperation of N_{pyz}…Cl halogen bond and C_{phen}-H…O=C nonclassical hydrogen bonds in generation of three-dimensional packing. Different colors show different adjacent moieties; halogen bonds are shown in red.

Since iodide is the most polarizable atom, it would have a higher positive charge than bromide and chloride. This should also affect the intermolecular bond strength, which is indeed reflected in the observed distance between I and N (of C–I···N) (Table 5). The C–I···N intermolecular distances in compounds 4, 9, and 12, containing the L³⁻¹ ligand, were found to be between 3.198(7) and 3.300(9) Å, which are 9.4-6.5% shorther than the sum of the van der Waals radii of nitrogen and iodide atoms.²¹

As expected,²² Table 5 shows that the strength of the halogen bond increases from chloride to bromide and further to iodide. Besides the N…X distance, another contributing factor to halogen bond strength is the halogen bond angle C–X…N. The C–X…N angles are all close to linearity, which is consistent with the theory of electron donation into the C–X σ^* orbitals of the X-bond donor.²³

The Cambridge Structural Database (CSD) was analyzed for halogen bonding of 2-4, 7-9, 11, and 12 to determine the common range of NM–X···N interactions, where NM is a nonmetal and nonhalogen and X is Cl, Br, or I (Figure 13).²⁰



Figure 9. Side view representation of **8** and **9**, showing the generation of double-stranded stair motifs through the Hg…Br short contacts (a), and cooperation of N_{pyz}…Br/I halogen bond and C_{phen}-H…O=C nonclassical hydrogen bonds in generation of three-dimensional packing (b). Different colors show different adjacent moieties; halogen bonds are shown in red.



Figure 10. Portion of the structure of coordination compounds formed between HgI₂ and L^H, **10** (a), L^{3-Br}, **11** (b), and L^{3-I}, **12** (c), showing coordination geometry around the central metal. Displacement ellipsoids are shown at 30% probability level. Symmetry codes: (a) (i) 1/2 - x, 1/2 - y, -z; (b) (i) 3/2 - x, -1/2 + y, -z; (ii) 3/2 - x, 1/2 + y, -z; (c) (i) 1/2 - x, -1/2 + y, 1 - z; (ii) 1/2 - x, 1/2 + y, 1 - z.



Figure 11. Side view representation of **10** showing the association *via* $C-H_{phen}\cdots I-Hg$ nonclassical hydrogen bonds (a), and $\pi_{pyz}\cdots\pi_{phen}$ intermolecular interactions (b), in the generation of three-dimensional packing. Different colors show different adjacent linear moieties.



Figure 12. Side view representation of **11** and **12**, showing the generation of double-stranded stair motifs through the Hg–I bonds (a), and N_{pyz} ...Br/I halogen bond in generation of three-dimensional packing (b). Different colors show different adjacent moieties; halogen bonds are shown in red.

Similar plot correlations have been reported for C-I...N halogen bonds.²⁴ This figure reveals that all parameters for determining these interactions are in the normal range. Therefore, it seems that the driving force for the crystal formation of these compounds is a halogen bond between the pyrazine nitrogen atom and the halogen atom on the halophenyl ring. It was thought of interest to further investigate the XB energy in these compounds by using theoretical methods. The binding energies obtained from DFT on two relative fragments (from complexes 2-4, 7-9, 11, and 12) provide us an opportunity to evaluate the complexes involving XB interactions. Calculations were performed with the experimental structures as the starting point at the LDA-ZORA-TZP level. The outcomes obtained from DFT methods are listed in Table 5. From these data, it is explicit that halogen bonding energies vary within a range of -27.86 to -46.15 kJ·mol⁻¹. In view of the directionality and

complex	X…N	X…N (Å)	C–X…N (deg)	reduction of the sum of the van der Waals radii (%)	symmetry code	XB synthon	L ^{3-X} conformation	O1=C5-C8-X torsion angle (deg)	calculated XB energy (kJ/mol) (LDA-Zora-TZP)
					X = Cl				
2 , $[HgCl_2(L^{3-Cl})]_n$	Cl…N	3.190(6)	172.7(3)	3.3	1-x,-y,-z	Ι	anti	4.16(6)	-46.15
7, [HgBr ₂ (L ^{3-Cl})]	Cl…N	3.217(5)	163.5(5)	2.5	1 - x, $1/2 + y$, $1/2 - z$	II	syn	0.00	-27.86
					X = Br				
3, $[Hg_2Cl_4(L^{3-Br})_2]$	Br…N	3.100(3)	173.6(1)	8.8	-x, 1-y, 1-z	Ι	anti	6.64(5)	-39.84
		3.188(3)	170.9(1)	6.2	-x, 1-y, 1-z	Ι	anti	29.72(8)	
8, $[HgBr_2(L^{3-Br})]$	Br…N	3.201(17)	171.0(8)	6.1	-x, y, 2-z	Ι	anti	0.00	-34.06
11, $[HgI_2(L^{3-Br})]$	Br…N	3.251(9)	173.1(5)	4.4	1 - x, y, 1 - z	Ι	anti	0.00	-32.35
					X = I				
4, $[Hg_2Cl_4(L^{3-1})_2]$	I…N	3.198(7)	175.3(2)	9.4	-x, 1-y, 2-z	Ι	anti	6.49(8)	-44.02
		3.288(7)	170.7(2)	6.9	-x, 1-y, 2-z	Ι	anti	32.44(9)	
9 , [HgBr ₂ (L ^{3-I})]	I…N	3.248(18)	174.3(5)	8.0	-x, y, 1-z	Ι	anti	0.00	-41.59
12, $[HgI_2(L^{3-I})]$	I…N	3.300(9)	174.2(3)	6.5	1 - x, y, -z	Ι	anti	0.00	-39.48

Table 5. Halogen Bond, Conformational Parameters, and Calculated XB Binding Energies for Compounds 2–4, 7–9, 11, and 12

Scheme 3. Halogen Bonding Synthons Exhibited in Hg(II) Complexes



strength of the halogen bonds, the N_{pyz} ···Cl/Br/I interaction can be recognized as an effective driving force to determine the alignment of molecules in supramolecular assemblies, which enables the design and development of new functional solid material.

The decomposition behavior of these compounds was also investigated in a static air atmosphere from ambient conditions to 500 °C. Stacked TG curves of compounds **2**, **3**, **4**, **7**, **8**, **9**, **11**, and **12** are given in Figure S1 (Supporting Information). The thermogravimetric analyses reveal that these compounds have identical single-step decomposition patterns.

CONCLUSION

The coordination geometry and supramolecular architecture of the 12 coordination compounds of mercury(II) halides based on L^{3-X} ligands (L = N-(3-halophenyl)-2-pyrazinecarboxamide) carrying a different halogen atom in the phenyl meta-position and L^{H} (L = N-phenyl-2-pyrazinecarboxamide) have been studied. Structural analysis clearly shows that the size of the coordinated halides would make a substantial contribution to the local mercury coordination environment and can also affect the extended structures. In each series containing the same ligand, the bulkier Br and I ions are sterically hindered to form a larger molecule. The crystal packing of the studied compounds is determined mostly by $C-H\cdots O/X$ (X = F, Cl, Br, I) nonclassical hydrogen bonds and $\pi \cdots \pi/\text{Hg}/\text{amide}$ and $\text{Hg} \cdots \text{X}$ intermolecular interactions. Since the halophenyl ring is flexible in rotation around the $N_{amide}-C_{phen}$ bond, it allows for subtle conforma-tional adaptation of L^{3-X} (X = Cl, Br, I) to produce additional halogen bond synthons via the rotation of the halophenyl ring. In view of the directionality and strength of the halogen bonds, the N_{pvz}…Cl/Br/I interaction can be recognized as an effective



Figure 13. Scatter plot of the halogen bonding distance X···N versus the angle of NM–X···N (where NM is a nonmetal and nonhalogen). X is Cl, Br, or I shown in deep green, brown, and purple, respectively. Reported data were obtained from CSD, version 2011, filtered to include only error-free, nondisordered structures with *R*, 0.075. Correlation values: chloride, 0.243; bromide, 0.438; iodide, 0.489.

driving force to determine the alignment of molecules in supramolecular assemblies. The N···X halogen bonding distances are 2.5-3.3, 4.4-8.8, and 6.5-9.4% shorter than the sum of the van der Waals radii of nitrogen and halogen in complexes containing L^{3-Cl}, L^{3-Br}, and L^{3-I} ligands, respectively. Theoretical methods show that the halogen bonding energies within a range of -27.86 to -46.15 kJ·mol⁻¹ are comparable to classical hydrogen bonding. The presence of a strong tendency to form halogen bonding synthons between adjacent halophenyl and pyrazine rings is one of the common features in the crystal structures of these complexes. Structural analysis of complexes studied here has clearly resulted in new sets of ligands that have been especially developed for selective and efficient metal coordination. Designing ligands that can provide capabilities for selective metal ion coordination is one of the important factors in controlling the architecture of self-assembled species. In all complexes synthesized here, the pyrazine ring is coordinated to the mercury(II) ion through the N atom syn to the carbonyl. Accordingly, the second nitrogen of the pyrazine ring, the N atom anti to the carbonyl, is now available to serve as a halogen bond acceptor to interact with the halogen atom. Therefore, the second common feature of the crystal structures of complexes studied here is the selectivity of the metal ion coordination site. The similarity of the halogen bond synthon across these compounds and selectivity in the mercury(II) ion coordination site further point to their application in coordination crystal engineering especially in the clever design of such special metal species and the crystal structure prediction research field.

EXPERIMENTAL SECTION

General. All chemicals were purchased from Aldrich or Merck and used without further purification. The synthesis and recrystallization of $L^{\rm H}$, $L^{3-\rm F}$, $L^{3-\rm Cl}$, $L^{3-\rm Br}$, and $L^{3-\rm I}$, and mercury(II) complexes 1–12 were carried out in air. The ¹H NMR spectrum was recorded on a Bruker AC-300 MHz spectrometer at ambient temperature in CD₃OD. All chemical shifts are quoted in parts per million (ppm) relative to tetramethylsilane. Infrared spectra (4000–250 cm⁻¹) of the solid sample were taken as 1% dispersion in CsI pellets using a BOMEM-MB102 spectrometer. Elemental analysis was performed using a Heraeus CHN-O Rapid analyzer. Melting points were obtained by a Bamstead Electrothermal type 9200 melting point apparatus and corrected. All ligands were prepared according to the reported procedure.¹³

Single-Crystal Diffraction Studies. For crystals 1-9, 11, and 12, intensity data were collected using a Bruker SMART APEX-II CCD diffractometer, equipped with a fine focus 1.75 kW sealed tube with Mo K α radiation (λ , 0.71073 Å). The total number of images was based on the results from the program COSMO.^{25a} Cell parameters were retrieved using APEX II software^{25b} and refined using SAINT on all observed reflections. Data reduction was performed using the SAINT software,^{25c} which corrects for the Lorentz and polarizing effects. Scaling and absorption corrections were applied using the SADABS^{25d} multiscan technique, supplied by George Sheldrick. X-ray data for compound 10 were collected on a STOE IPDS-II diffractometer with graphite monochromated Mo K α radiation (λ , 0.71073 Å). Data were collected in a series of ω scans in 1° oscillations and integrated using the Stöe X-AREA^{26a} software package. A numerical absorption correction was applied using the X-RED^{26b} and X-SHAPE^{26c} software. All structures were solved by direct methods using SHELXS-97²⁷ and refined with full-matrix least-squares on F^2 using the SHELXL-97²⁷ program package. All non-hydrogen atoms were refined anisotropically. Hydrogen atoms were added at ideal positions and constrained to ride on their parent atoms, with $U_{iso}(H) = 1.2U_{eq}$. All refinements were performed using the X-STEP32 crystallographic software package.²⁸ Structural illustrations have been drawn with ORTEP-329 and MERCURY.³⁰ Crystallographic data for compounds 1-12 are listed in Table 1. Selected bond distances and angles are summarized in Table

2. As expected, the highest peaks and deepest holes for these mercury complexes are slightly high, but the residual densities are close to heavy mercury atoms (with a maximum distance of 1.21 Å from Hg1 in complex 1). For compounds 5 and 11, which have high residual densities, the highest peaks and deepest holes are close to the Hg atoms with a distance of 0.86 and 0.79 Å for the highest peaks and 0.89 and 0.85 Å for deepest holes, respectively.

Synthesis of *N*-(3-Halophenyl)pyrazine-2-carboxamide ligands, L^H, L^{3-F}, L^{3-CI}, L^{3-Br}, and L^{3-I}. These ligands were prepared by using the previously reported method.¹³

Synthesis of [HgCl₂(L^H)]_n, **1.** To a solution of HgCl₂ (0.027 g, 0.1 mmol) in 3 mL of methanol was added a solution of L^H (0.019 g, 0.1 mmol) in 3 mL of methanol while stirring. The mixture was heated at 40 °C for about 1 h and then filtered. Upon slow evaporation of the filtrate at room temperature, suitable yellow cystals of complex 1 for X-ray analysis were obtained after 5 days (yield 41%, mp = 153–155 °C). Anal. Calcd for C₁₁H₉Cl₂HgN₃O: C, 28.07; H, 1.93; N, 8.93. Found: C, 28.05; H, 1.95; N, 8.94. FT-IR (CsI pellet, cm⁻¹): 3337 (N-H str. vib.), 1666 (C=O amide str. vib.), 1606, 1540 (C-N str. and N-H bend vib.), 1434(C=C str. vib.), 1387 (C-N str. vib.), 1016, 877, 685 (O=C-N group bend vib.), 572 (O=C-N group bend vib), 439. ¹H NMR (CD₃OD, δ from TMS): 10.71 (s, 1H-pyrazine), 9.30 (s, amidic H), 8.93 (d, 1H-pyrazine), 8.81–8.83 (m, 1H-pyrazine), 7.89 (d, 2H-phenyl), 7.37 (t, 2H-phenyl), and 7.14 (t, 1H-phenyl). **Synthesis of [HgCl₂(L^{3-C})**]_n, **2.** The procedure was similar to the

Synthesis of [HgCl₂(L^{3-Cl})]_n, 2. The procedure was similar to the synthesis of 1 except that L^{3-Cl} (0.023 g, 0.1 mmol) was used instead of L^H. Colorless crystals of **2** were formed after 5 days (yield 72%, mp = 184–186 °C). Anal. Calcd for C₁₁H₈Cl₃HgN₃O: *C*, 26.15; H, 1.60; N, 8.32. Found: C, 26.13; H, 1.58; N, 8.30. FT-IR (CsI pellet, cm⁻¹): 3356 (N-H str. vib.), 1692 (C=O amide str. vib.), 1600, 1527 (C-N str. and N-H bend vib.), 1474 (C=C str. vib.), 1420, 1394 (C-N str. vib.), 1122 (C-Cl str. vib.), 1049, 844, 777, 678 (O=C-N group bend vib.), 579 (O=C-N group bend vib), 426. ¹H NMR (CD₃OD, *δ* from TMS): 10.96 (s, 1H-pyrazine), 9.30 (s, amidic H), 8.95 (d, 1H-pyrazine), 8.82–8.84 (m, 1H-pyrazine), 8.09 (s, 1H-phenyl), 7.86 (d, 1H-phenyl), 7.40 (t, 1H-phenyl), and 7.20 (d, 1H-phenyl). **Synthesis of [Hg₂Cl₄(L^{3-B1}₂)₂], 3.** The procedure was similar to the

Synthesis of [Hg₂Cl₄(L^{3-Br})₂], 3. The procedure was similar to the synthesis of 1 except that L^{3-Br} (0.027 g, 0.1 mmol) was used instead of L^H. Colorless crystals of **3** were formed after 7 days (yield 69%, mp = 196–198 °C). The complex **3** is relatively stable in the solid state when stored in the mother liquor, but decomposes slowly when exposed to open air. Therefore, the diffraction data for this compound were collected at 173 K. Anal. Calcd for C₂₂H₁₆Br₂Cl₄Hg₂N₆O₂: C, 24.04; H, 1.47; N, 7.65. Found: C, 24.01; H, 1.45; N, 7.63. FT-IR (CsI pellet, cm⁻¹): 3350 (N-H str. vib.), 1692 (C=O amide str. vib.), 1659 (C=O amide str. vib.), 1573 (C-N str. and N-H bend vib.), 1467 (C=C str. vib.), 1394 (C-N str. vib.), 1122 (C-Br str. vib.), 857, 685 (O=C-N group bend vib.), 579 (O=C-N group bend vib.), 579 (d, 1H-pyrazine), 9.29 (s, amidic H), 8.95 (d, 1H-pyrazine), 8.81–8.84 (m, 1H-pyrazine), 8.24 (s, 1H-phenyl), 7.90 (d, 1H-phenyl), 7.56 (d, 1H-phenyl), and 7.35 (t, 1H-phenyl).

Synthesis of [Hg₂Cl₄(L³⁻¹)₂], 4. The procedure was similar to the synthesis of 1 except that L³⁻¹ (0.032 g, 0.1 mmol) was used instead of L^H. Light yellow crystals of 4 were formed after 2 weeks (yield 55%, mp = 200–202 °C). Anal. Calcd for C₂₂H₁₆Cl₄Hg₂I₂N₆O₂: C, 22.15; H, 1.35; N, 7.04. Found: C, 22.14; H, 1.35; N, 7.04. Found: C, 22.14; H, 1.35; N, 7.04. For L(CSI pellet, cm⁻¹): 3337 (N-H str. vib.), 1692 (C=O amide str. vib.), 1675 (C=O amide str. vib.), 1573 (C-N str. and N-H bend vib.), 1533, 1467 (C=C str. vib.), 1394 (C–N str. vib.), 1116 (C-I str. vib), 1016, 850, 685 (O=C-N group bend vib.), 585 (O=C-N group bend vib), 435. ¹H NMR (CD₃OD, δ from TMS): 10.83 (s, 1H-pyrazine), 9.28 (s, amidic H), 8.93 (d, 1H-pyrazine), 8.79–8.82 (m, 1H-pyrazine), 8.38 (s, 1H-phenyl), 7.90 (d, 1H-phenyl), 7.50 (d, 1H-phenyl), and 7.17 (t, 1H-phenyl).

Synthesis of [Hg_2Br_4(L^H)_2], 5. The procedure was similar to the synthesis of 1 except that $HgBr_2$ (0.036 g, 0.1 mmol) was used instead of $HgCl_2$. Colorless crystals of **5** were formed after 5 weeks (yield 69%, mp = 146–148 °C). Anal. Calcd for $C_{22}H_{18}Br_4Hg_2N_6O_2$: C, 23.61; H, 1.62; N, 7.51. Found: C, 23.60; H, 1.60; N, 7.50. FT-IR (CsI pellet, cm⁻¹):

3343 (N-H str. vib.), 1672 (C=O amide str. vib.), 1546 (C-N str. and N-H bend vib.), 1447 (C=C str. vib.), 1401 (C-N str. vib.), 1016, 744, 685 (O=C-N group bend vib.), 579 (O=C-N group bend vib), 453. ¹H NMR (CD₃OD, *δ* from TMS): 10.72 (s, 1H-pyrazine), 9.29 (s, amidic H), 8.91 (d, 1H-pyrazine), 8.79–8.83 (m, 1H-pyrazine), 7.88 (d, 2H-phenyl), 7.36 (t, 2H-phenyl), and 7.16 (t, 1H-phenyl).

Synthesis of [HgBr₂(L^{3-F})], 6. The procedure was similar to the synthesis of 5 except that L^{3-F} (0.021 g, 0.1 mmol) was used instead of L^H. Colorless crystals of 6 were formed after 7 days (yield 48%, mp = 156–158 °C). Anal. Calcd for C₁₁H₈Br₂FHgN₃O: C, 22.87; H, 1.40; N, 7.27. Found: C, 22.87; H, 1.40; N, 7.29. FT-IR (CsI pellet, cm⁻¹): 3339 (N-H str. vib.), 1694 (C=O amide str. vib.), 1617, 1544 (C-N str. and N-H bend vib.), 1445 (C=C str. vib.), 1407 (C-N str. vib.), 1272, 1146 (C-F str. vib), 860, 680 (O=C-N group bend vib.), 580 (O=C-N group bend vib.), 459. ¹H NMR (CD₃OD, δ from TMS): 10.98 (s, 1H-pyrazine), 9.31 (s, amidic H), 8.95 (d, 1H-pyrazine), 8.81–8.83 (m, 1H-pyrazine), 7.85 (d, 1H-phenyl), 7.75 (d, 1H-phenyl), 7.41 (dd, 1H-phenyl), and 6.97 (t, 1H-phenyl).

Synthesis of [HgBr₂(L^{3-CI})], 7. The procedure was similar to the synthesis of 5 except that L^{3-CI} (0.023 g, 0.1 mmol) was used instead of $L^{\rm H}$. Colorless crystals of 7 were formed after 5 days (yield 45%, mp = 185–187 °C). Anal. Calcd for C₁₁H₈Br₂ClHgN₃O: C, 22.24; H, 1.36; N, 7.07. Found: C, 22.23; H, 1.36; N, 7.06. FT-IR (CsI pellet, cm⁻¹): 3350 (N-H str. vib.), 1686 (C=O amide str. vib.), 1586, 1533 (C-N str. and N-H bend vib.), 1440 (C=C str. vib.), 1414 (C-N str. vib.), 1129 (C-Cl str. vib.), 1049, 857, 784, 685 (O=C-N group bend vib.), 572 (O=C-N group bend vib), 453. ¹H NMR (CD₃OD, δ from TMS): 10.95 (s, 1H-pyrazine), 9.31 (s, amidic H), 8.95 (d, 1H-pyrazine), 8.81–8.83 (m, 1H-pyrazine), 8.09 (s, 1H-phenyl), 7.85 (d, 1H-phenyl), 7.40 (t, 1H-phenyl), and 7.20 (d, 1H-phenyl).

Synthesis of [HgBr₂(L^{3-Br})], 8. The procedure was similar to the synthesis of 5 except that L^{3-Br} (0.027 g, 0.1 mmol) was used instead of L^H. Colorless crystals of 8 were formed after 2 weeks (yield 63%, mp = 190–192 °C). Anal. Calcd for C₁₁H₈Br₃HgN₃O: C, 20.69; H, 1.26; N, 6.58. Found: C, 20.69; H, 1.26; N, 6.58. FT-IR (CsI pellet, cm⁻¹): 3350 (N-H str. vib.), 1699 (C=O amide str. vib.), 1573, 1540 (C-N str. and N-H bend vib.), 1474 (C=C str. vib.), 1410 (C-N str. vib.), 1129 (C-Br str. vib), 850, 771, 681 (O=C-N group bend vib.), 579 (O=C-N group bend vib), 433. ¹H NMR (CD₃OD, δ from TMS): 10.96 (s, 1H-pyrazine), 9.30 (s, amidic H), 8.94 (d, 1H-pyrazine), 8.80–8.83 (m, 1H-pyrazine), 8.23 (s, 1H-phenyl), 7.89 (d, 1H-phenyl), 7.56 (d, 1H-phenyl), and 7.34 (t, 1H-phenyl).

Synthesis of [HgBr₂(L³⁻¹)], **9.** The procedure was similar to the synthesis of **5** except that L³⁻¹ (0.032 g, 0.1 mmol) was used instead of L^H. Colorless crystals of **9** were formed after 5 days (yield 61%, mp = 191–193 °C). Anal. Calcd for C₁₁H₈Br₂HgIN₃O: C, 19.27; H, 1.18; N, 6.13. Found: C, 19.27; H, 1.17; N, 6.13. FT-IR (CsI pellet, cm⁻¹): 3337 (N-H str. vib.), 1686 (C=O amide str. vib.), 1586, 1533 (C-N str. and N–H bend vib.), 1460 (C=C str. vib.), 1402 (C-N str. vib.), 1122 (C-I str. vib), 1009, 850, 784, 685 (O=C-N group bend vib.), 585 (O=C-N group bend vib), 413. ¹H NMR (CD₃OD, δ from TMS): 10.88 (s, 1H-pyrazine), 9.31 (s, amidic H), 8.95 (d, 1H-pyrazine), 8.79–8.83 (m, 1H-pyrazine), 8.31 (s, 1H-phenyl), 7.89 (d, 1H-phenyl), 7.51 (d, 1H-phenyl), and 7.18 (t, 1H-phenyl).

Synthesis of [Hg₂I₄(L^H)₂], 10. The procedure was similar to the synthesis of 1 except that HgI₂ (0.045 g, 0.1 mmol) was used instead of HgCl₂. Colorless crystals of **10** were formed after 4 days (yield 59%, mp = 140–142 °C). Anal. Calcd for C₂₂H₁₈Hg₂I₄N₆O₂: *C*, 20.21; H, 1.39; N, 6.43. Found: *C*, 20.19; H, 1.38; N, 6.42. FT-IR (CsI pellet, cm⁻¹): 3360 (N-H str. vib.), 1695 (C=O amide str. vib.), 1533 (C-N str. and N-H bend vib.), 1445 (C=C str. vib.), 1387 (C-N str. vib.), 1308, 1155, 883, 685 (O=C-N group bend vib.), 587 (O=C-N group bend vib), 433. ¹H NMR (CD₃OD, δ from TMS): 10.73 (s, 1H-pyrazine), 9.28 (s, amidic H), 9.28 (d, 1H-pyrazine), 8.79–8.83 (m, 1H-pyrazine), 7.87 (d, 2H-phenyl), 7.37 (t, 2H-phenyl), and 7.15 (t, 1H-phenyl).

Synthesis of [Hgl₂(L^{3-B})], 11. The procedure was similar to the synthesis of 10 except that L^{3-Br} (0.027 g, 0.1 mmol) was used instead of L^H. Colorless crystals of 11 were formed after 2 weeks (yield 51%, mp = 200–202 °C). Anal. Calcd for C₁₁H₈BrHgI₂N₃O: C, 18.04; H, 1.10; N, 5.74. Found: C, 18.03; H, 1.10; N, 5.72. FT-IR (CsI pellet, cm⁻¹): 3350

(N-H str. vib.), 1692 (C=O amide str. vib.), 1540 (C-N str. and N-H bend vib.), 1474 (C=C str. vib.), 1401 (C-N str. vib.), 1175, 1122 (C-Br str. vib.), 1056, 850, 671 (O=C-N group bend vib.), 585 (O=C-N group bend vib), 433. ¹H NMR (CD₃OD, δ from TMS): 10.93 (s, 1H-pyrazine), 9.29 (s, amidic H), 8.95 (d, 1H-pyrazine), 8.79–8.83 (m, 1H-pyrazine), 8.23 (s, 1H-phenyl), 7.88 (d, 1H-phenyl), 7.56 (d, 1H-phenyl), and 7.33 (t, 1H-phenyl).

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Computational Details. DFT calculations were performed using the ORCA quantum chemistry suite.³¹ The local density approximation (LDA) exchange correlation potential was used with the local density approximation of the correlation energy.³² Gradient-corrected geometry optimizations³³ were performed by using the generalized gradient approximation.³⁴ Large atom basis sets TZP were used to ascribe all of the atoms. Scalar relativistic effects were taken into account by using the zeroth-order regular approximation (ZORA).³⁵

Thermal Analysis (TG). TG experiments were carried out using a Rheometric Scientific (STA/1500) with a heating rate of 10 °C/min in the range of 25-500 °C, under an air atmosphere. The measurements were carried out using around 10 mg of a powdered sample sealed in aluminum pans with a mechanical crimp.

ASSOCIATED CONTENT

S Supporting Information

TG curves for compounds 2, 3, 4, 7, 8, 9, 11, and 12; hydrogen bond geometries, and X-ray crystallographic files in CIF format for structural determination of 1 (CCDC No. 901972), 2 (CCDC No. 901975), 3 (CCDC No. 901970), 4 (CCDC No. 901976), 5 (CCDC No. 901971), 6 (CCDC No. 901981), 7 (CCDC No. 901978), 8 (CCDC No. 901973), 9 (CCDC No. 901974), 10 (CCDC No. 901980), 11 (CCDC No. 901979), and 12 (CCDC No. 901977). This material is available free of charge *via* the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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