Accepted Manuscript

Synthesis, structural, DFT calculations and biological studies of rhodium and iridium complexes containing azine Schiff-base ligands

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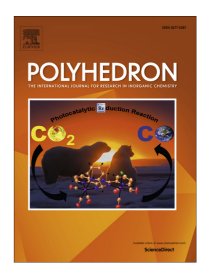
PII: S0277-5387(16)30218-2

DOI: http://dx.doi.org/10.1016/j.poly.2016.06.001

Reference: POLY 12039

To appear in: Polyhedron

Received Date: 4 May 2016 Accepted Date: 1 June 2016



Please cite this article as: S. Adhikari, D. Sutradhar, S.L. Shepherd, R.M. Phillips, A.K. Chandra, K. Mohan Rao, Synthesis, structural, DFT calculations and biological studies of rhodium and iridium complexes containing azine Schiff-base ligands, *Polyhedron* (2016), doi: http://dx.doi.org/10.1016/j.poly.2016.06.001

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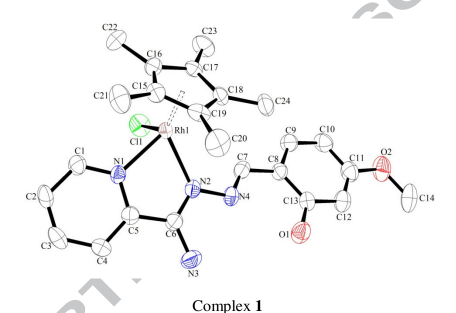
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Graphical abstract

Half-sandwich Cp*Rh(III) and Cp*Ir(III) complexes have been synthesized with N-N' azine Schiff-base ligands and characterized by spectroscopic techniques. The molecular structures of some of the representative complexes have been confirmed by single crystal X-ray analysis.

Chemo-sensitivity activities of the complexes were evaluated against HT-29 (human colorectal

cancer) cell line and non-cancer cell line ARPE-19 (human retinal epithelial cells).



Abstract

The reaction of $[Cp*MCl_2]_2$ (M = Rh/Ir) with N-N' azine Schiff-base ligands (L1-L4)
leads to the formation of mononuclear cationic half-sandwich complexes having the general
formula $[Cp*M(L)Cl]^+$ (1–8), (M = Rh/Ir and L = (2-hydroxy-4-methoxybenzylidene)2-
pyridylamidrazone (L1), (2-hydroxybenzylidene)2-pyridylamidrazone (L2), (1-(2-
hydroxyphenyl)ethylidene)2-pyridylamidrazone (L3) and (1-phenylethylidene)2-
pyridylamidrazone (L4). All these complexes were isolated as their hexafluorophosphate salts
and fully characterized by spectroscopic and analytical techniques. The molecular structure of
complexes (1), (3), (4), (7) and (8) have been determined by single crystal X-ray crystallographic
studies which displayed the coordination of the ligand to the metal in a bidentate $N\cap N$ fashion
through nitrogen atom of pyridine and one azine nitrogen. The chemo-sensitivity activities of the
complexes were evaluated against HT-29 (human colorectal cancer) cell line and non-cancer cell
line ARPE-19 (human retinal epithelial cells) which revealed that the complexes are moderately
cytotoxic to cancer cells over human cells although complex 5 was the most potent among all the
compounds. Theoretical studies carried out using DFT and TD-DFT at B3LYP level shows good
agreement with the experimental results.

Keywords: Rhodium, Iridium, Azine Schiff-base ligands, Cytotoxicity

1. Introduction

The chemistry of half-sandwich organometallic complexes has evolved as a versatile
subject of research during the past few decades due to its wide application in biological and
medicinal fields [1-4]. Organometallic half-sandwich compounds of the general formula
[Cp*MCl(LL')] (M = Rh, Ir and LL' = N,N or N,O donor ligands) have been extensively studied
for their cytostatic activity, DNA binding, cellular uptake and as DNA intercelators [5-9].
Rhodium and iridium complexes have also been investigated as an alternative to platinum based
drugs mainly because of their water solubility and lability towards ligand exchange [10, 11].
Recently Therrien et.al reported dinuclear dithiolato bridged rhodium and iridium complexes
which exhibit cytotoxicity against human ovarian cancer cells lines (A2780 and A2780cisR)
[12]. C-H activated cyclometallated Rh(III) and Ir(III) complexes can effectively bind to DNA
and protein through electrostatic and hydrophobic interactions [13]. Iridium complexes of
dihydroxybipyridine are active catalysts for homogenous water oxidation under mild reaction
conditions [14]. Rh(III) and Ir(III) polypyridyl complexes exhibits strong antiproliferative
activity towards human cancer cell lines and are also capable of binding to DNA [15]. A number
of half-sandwich Ir(III) complexes have been reported by Sadler et al with chelating C, N and
pyridine ligands and N, N donor ligands which showed strong antiproliferative activity [16, 17].
Pyridyl azines represent an important class of organic compounds with interesting
properties having wide applications in various areas [18]. Open chain diazine Schiff base ligands
linked by a single N-N bond are of great interest due to its rotational flexibility around the N-N
bond and potential donor sites which can give rise to a rich variety of coordination compounds
with different binding modes [19]. The N-N bridging ligand plays a crucial role in
communicating the metal centers to form mononuclear, dinuclear or polynuclear complexes [20].

The diazine ligand has been employed into several transition metal azido and thiocyanato systems namely Mn(II)-azido, Cd(II)-NCS to obtain several 1D, 2D and 3D polymers which exhibit interesting magnetic properties [21, 22]. Dinuclear transition metal complexes of Cu, Zn, Mn and Ni have been reported with bridging N-N diazine ligands which give rise to strong ferromagnetic and antiferromagnetic coupling [23]. In the recent years our group has reported many half-sandwich Ru(II), Rh(III) and Ir(III) complexes with azine ligands [24, 25]. In continuation with our interest of these ligands herein we report four new azine Schiff base ligands derived from 2-pyridylamidrazone and its corresponding rhodium and iridium half-sandwich metal complexes. The complexes were tested for their cytotoxic property to selectively kill HT-29 cancer cell line against normal ARPE-19 cells.

2. Experimental Section

79 2. 1. Physical methods and materials

All the reagents were purchased from commercial sources and used as received. Starting materials RhCl₃.nH₂O, IrCl₃.nH₂O were purchased from Arora Matthey limited. 2-cyanopyridine, 2-hydroxybenzaldehyde, 2-hydroxyacetophenone, were obtained from Aldrich, acetophenone and 2-hydroxy-4-methoxybenzaldehyde were obtained from Alfa-Aesar. The solvents were purified and dried according to standard procedures [26]. All the reactions were carried out under normal conditions. The starting precursor metal complexes [Cp*MCl₂]₂ (M = Rh/Ir) were prepared according to the literature methods [27]. Infrared spectra were recorded on a Perkin-Elmer 983 spectrophotometer by using KBr pellets in the range of 400-4000 cm⁻¹. ¹H NMR spectra were recorded on a Bruker Avance II 400 MHz spectrometer using DMSO-d₆ and CDCl₃ as solvents. Absorption spectra were recorded on a Perkin-Elmer Lambda 25 UV/Visible spectrophotometer in the range of 200-800 nm at room temperature in acetonitrile. Elemental

- analyses of the complexes were performed on a Perkin-Elmer 2400 CHN/S analyzer. Mass
- 92 spectra were recorded using Q-Tof APCI-MS instrument (model HAB 273). All these
- 93 mononuclear metal complexes were synthesized and characterized by using FT-IR, ¹H NMR,
- 94 UV-Vis, and Single-crystal X-ray diffraction techniques.
- 95 2. 2. Single-crystal X-ray structures analyses
- The orange crystals of complexes (1), (3), (7) and (8) were obtained by slow diffusion of 96 hexane into acetone or DCM solution and yellow crystals of complex (4) was obtained by 97 diffusing hexane into DCM solution. Single crystal X-ray diffraction data for all the complexes 98 (1), (3) (4), (7) and (8) were collected on a Oxford Diffraction Xcalibur Eos Gemini 99 diffractometer at 293 K using graphite monochromated Mo-K α radiation ($\lambda = 0.71073$ Å). The 100 strategy for the data collection was evaluated using the CrysAlisPro CCD software. Crystal data 101 were collected by standard "phi-omega scan" techniques and were scaled and reduced using 102 CrysAlisPro RED software. The structures were solved by direct methods using SHELXS-97 103 and refined by full-matrix least squares with SHELXL-97 refining on F² [28, 29]. The positions 104 of all the atoms were obtained by direct methods. Metal atoms in the complex were located from 105 the E-maps and non-hydrogen atoms were refined anisotropically. The hydrogen atoms bound to 106 the carbon were placed in geometrically constrained positions and refined with isotropic 107 temperature factors, generally 1.2 U_{eq} of their parent atoms. Crystallographic and structure 108 refinement details for the complexes are summarized in Table 1, and selected bond lengths and 109 bond angles are presented in Table S1. Figures 1-3 were drawn with ORTEP3 program. Figure 4 110 and Figures S3-S6 were drawn with MERCURY3.6 program [30]. 111
- 112 2.3. Biological studies

All complexes (1-8) were dissolved in DMSO at 100 mM and stored at -20 °C until needed. The complexes were tested against cancer cell line HT-29 (human colorectal cancer), and one non-cancer cell line ARPE-19 (human retinal epithelial cells). Cells were seeded into 96 well plates at 1 x 10³ cells per well and incubated at 37 °C in a CO₂ enriched (5%), humidified atmosphere overnight to adhere. The cells were exposed to a range of drug concentrations in the range of 0-100 µM for four days before cell survival was determined using the MTT assay [31]. To each well MTT (0.5 mg/ml) was added and was further incubated at 37 °C for 4 h. After this the MTT was removed from each well and the formazan crystals formed were dissolved in 150 µM DMSO. The absorbance of the resulting solution was recorded at 550 nm using an ELISA spectrophotometer. The percentage of cell inhibition was calculated by dividing the absorbance of treated cell by the control value absorbance (exposed to 0.1 % DMSO). The results were expressed in terms of IC₅₀ values (concentration required to kill 50 % cell) and all studies were performed in triplicate. The results were also expressed in terms of a 'selectivity index' defined as the IC₅₀ of the non-cancer cell line ARPE divided by the IC₅₀ of cancer cell lines [32]. Values greater than 1 demonstrate that the compound is preferentially active against tumor cell compared to normal cell lines.

2.4. Computational methodology

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All the electronic structure calculations of the metal complexes (1-8) were carried out using the Gaussian 09 suite of program [33]. The geometries of the rhodium and iridium complexes were optimized in the gas phase employing the DFT-based B3LYP method with 6-31G** basis set for (H, C, N, O, Cl, F and P atoms and LANL2DZ [34, 35] for (Rh and Ir) atoms. Harmonic frequency calculations were carried out at the same level of theory to ensure that the optimized geometries were true minima on the potential energy surface (PES). Natural

- Bond Orbital (NBO) analysis [36] was used to obtain the charge distribution on individual atoms 136 and the d-orbital occupations of the metal present in the complexes. Time dependent-Density 137 Functional Theory (TD-DFT) [37] has been employed to evaluate the absorption spectra and the 138 electronic transitions of the metal complexes. In order to incorporate the effect of the solvent 139 around the molecule, the Polarizable Continuum Model (PCM) [38] was used in TD-DFT 140 calculations. The percentage contribution of molecular orbital analysis was carried out using 141 Chemissian software package [39]. 142 143
 - 2.4. General procedure for preparation of ligands 1-4
- 2.4.1. The azine Schiff base ligands (L1-L4) were prepared by two step procedure. 144
- In the first step 2-pyridylamidrazone was prepared, by following a reported procedure 145 [40]. 2-cyanopyridine and hydrazine hydrate were dissolved and stirred in absolute ethanol 146 overnight to give 2-pyridylamidrazone as yellow crystalline solid which was used in the next 147 step without further purification (Scheme-1). In the second step (5 mmol) of aldehyde or ketone 148 and 2-pyridylamidrazone (5 mmol) was refluxed in 10 ml ethanol for 5 hours (Scheme-2). The 149 products obtained after cooling the solution were filtered off washed with cold methanol and 150 diethyl ether and dried in vacuum. 151
- Data for ligands (L1-L4) 152
- 2.4.2. (2-hydroxy-4-methoxybenzylidene)2-pyridylamidrazone (L1) 153
- Color: Yellow needles; Yield: 88%; IR (KBr, cm⁻¹): 3487(s), 3380(s), 3333(m), 2964(m), 154
- 1627(s), 1587(m), 1566(m), 1394(m), 1340(s); ¹H NMR (400 MHz, CDCl₃): $\delta = 11.82$ (s, 1H, 155
- OH), 8.60 (s, 1H, $CH_{(imine)}$), 8.57 (d, 1H, J = 4.0 Hz, $CH_{(pv)}$), 8.34 (d, 1H, J = 8.0 Hz, $CH_{(pv)}$), 156
- 7.76 (t, 1H, $CH_{(py)}$), 7.35 (t, 1H, $CH_{(py)}$), 7.20 (d, 2H, J = 8.0 Hz, $CH_{(Ar)}$), 6.46-6.50 (m, 3H, NH_2 , 157
- $CH_{(Ar)}$), 3.80 (s, 3H, OMe); HRMS-APCI (m/z): 271.11 [M+H]⁺; UV-Vis {Acetonitrile, λ_{max} , 158

- 159 nm ($\epsilon/10^{-4}$ M⁻¹ cm⁻¹)}: 218 (0.84), 314 (0.68), 342 (0.92), 355 (0.94); Anal. Calc. for $C_{14}H_{14}N_4O_2$
- 160 (270.29): C, 62.21; H, 5.22; N, 20.73. Found: C, 62.36; H, 5.35; N, 20.86%.
- 2.4.3. (2-hydroxybenzylidene)2-pyridylamidrazone (L2)
- 162 Color: Yellow needles; Yield: 92%; IR (KBr, cm⁻¹): 3477(s), 3363(s), 3340(s), 3043(m), 1626(s),
- 163 1576(m), 1567(m), 1473(m), 1337(m); ¹H NMR (400 MHz, CDCl₃): δ = 11.61 (s, 1H, OH), 8.59
- 164 (s, 1H, $CH_{(imine)}$), 8.54 (d, 1H, J = 4.0 Hz, $CH_{(py)}$), 8.28 (d, 1H, J = 8.0 Hz, $CH_{(py)}$), 7.74 (t, 1H,
- 165 $CH_{(py)}$), 7.33 (t, 1H, $CH_{(py)}$), 7.24-7.27 (m, 3H, NH_2 , $CH_{(Ar)}$), 6.96 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$),
- 166 6.87 (t, 2H, CH_(Ar)); HRMS-APCI (m/z): 241.10 [M+H]⁺; UV–Vis {Acetonitrile, λ_{max} , nm ($\epsilon/10^{-4}$
- 167 M^{-1} cm⁻¹)}: 219 (0.84), 247 (0.55), 349 (1.30), 361 (1.29); Anal. Calc. for $C_{13}H_{12}N_4O$ (240.26):
- 168 C, 64.99; H, 5.03; N, 23.32. Found: C, 65.12; H, 5.18; N, 23.44%.
- 2.4.4. (1-(2-hydroxyphenyl)ethylidene)2-pyridylamidrazone (L3)
- 170 Color: Yellow crystalline solid; Yield: 95%; IR (KBr, cm⁻¹): 3482(s), 3339(s), 3056(m),
- 3003(m), 1615(s), 1562(m), 1507(m), 1300(m); ¹H NMR (400 MHz, CDCl₃): $\delta = 13.73$ (s, 1H,
- OH), 8.59 (d, 1H, J = 4.0 Hz, $CH_{(py)}$), 8.36 (d, 1H, J = 8.0 Hz, $CH_{(py)}$), 7.79 (t, 1H, $CH_{(py)}$), 7.58
- 173 (t, 1H, $CH_{(py)}$), 7.21-7.28 (m, 3H, NH_2 , $CH_{(Ar)}$), 6.98 (d, 2H, J = 8.0 Hz, $CH_{(Ar)}$), 6.89 (t, 1H,
- 174 $CH_{(Ar)}$), 2.62 (s, 3H, CH₃); HRMS-APCI (m/z): 255.12 [M+H]⁺; UV–Vis {Acetonitrile, λ_{max} , nm
- 175 $(\epsilon/10^{-4} \text{ M}^{-1} \text{ cm}^{-1})$: 217 (1.21), 303 (0.74), 344 (0.95); Anal. Calc. for $C_{14}H_{14}N_4O$ (254.29): C,
- 176 66.13; H, 5.55; N, 22.03. Found: C, 66.25; H, 5.68; N, 22.21%.
- 177 *2.4.5.* (1-phenylethylidene)2-pyridylamidrazone (L4)
- 178 Color: Yellow crystalline solid; Yield: 92%; IR (KBr, cm⁻¹): 3450(s), 3331(s), 3056(m),
- 3009(m), 1604(s), 1568(m), 1445(m), 1362(m); ¹H NMR (400 MHz, CDCl₃): δ = 8.58 (d, 1H, J
- $=4.0~{\rm Hz},~{\rm CH_{(py)}}),~8.23~({\rm d},~1{\rm H},~J=8.0~{\rm Hz},~{\rm CH_{(py)}}),~7.71~({\rm t},~1{\rm H},~{\rm CH_{(py)}}),~7.30~({\rm t},~1{\rm H},~{\rm CH_{(py)}}),~7.21-{\rm t},~{\rm t},~{\rm$
- $7.28 \ (m, \, 3H, \, NH_2, \, CH_{(Ar)}), \, 6.93 \ (m, \, 3H, \, CH_{(Ar)}), \, 6.89 \ (t, \, 1H, \, CH_{(Ar)}), \, 2.39 \ (s, \, 3H, \, CH_3); \, HRMS-100 \ (t, \, 1H, \, CH_{(Ar)}), \, 2.39 \ (s, \, 2H, \, CH_3); \, HRMS-100 \ (t, \, 2H, \, 2H_3); \, HRMS-100 \ (t, \, 2H,$

- APCI (m/z): 239.13 [M+H]⁺; UV–Vis {Acetonitrile, λ_{max} , nm (ϵ /10⁻⁴ M⁻¹ cm⁻¹)}: 225 (0.21), 327
- 183 (0.29); Anal. Calc. for C₁₄H₁₄N₄ (238.29): C, 70.57; H, 5.92; N, 23.51. Found: C, 70.72; H, 6.03;
- 184 N, 23.62%.
- 185 2.5. General procedure for preparation of metal complexes (1-8)
- A mixture of metal precursor $[Cp*MCl_2]_2$ (M = Rh/Ir) (0.1 mmol), azine Schiff-base ligands (L1-
- L4) (0.2 mmol) and 2.5 equivalents of NH₄PF₆ in dry methanol (10 ml) was stirred at room
- temperature for 8 hours (Scheme-3). The solvent was evaporated under reduced pressure, and the
- 189 residue was dissolved in dichloromethane and filtered over celite to remove excess salt. The
- 190 filtrate was reduced to 2 ml and diethyl ether was added to induce precipitation. The yellow
- 191 colored precipitate, which formed, was filtered and washed with diethyl ether and dried in
- 192 vacuum.
- 193 $2.5.1. [Cp*Rh(L1)Cl]PF_6(1)$
- 194 Yield: 56 mg (40%); IR (KBr, cm⁻¹): 3460(m), 3237(m), 2926(w), 1630(s), 1595(m), 1296(m),
- 195 846(s); ¹H NMR (400 MHz, CDCl₃): $\delta = 10.5$ (s, 1H, OH), 9.02 (s, 1H, CH(imine)), 8.76 (d, 1H, J
- 196 = 4.0 Hz, $CH_{(py)}$), 8.54 (d, 1H, J = 4.0 Hz, $CH_{(py)}$), 8.13 (t, 1H, $CH_{(py)}$), 7.76 (t, 1H, $CH_{(py)}$), 7.41
- 197 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$), 7.38 (s, 2H, NH₂), 6.53 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$), 6.50 (s, 1H,
- 198 $CH_{(Ar)}$), 3.81 (s, 3H, OMe), 1.58 (s, 15H, $CH_{(Cp^*)}$); HRMS-APCI (m/z): 507.12 [M-PF₆-HCl]⁺;
- 199 UV–Vis {Acetonitrile, λ_{max} , nm ($\epsilon/10^{-4}$ M⁻¹ cm⁻¹)}: 233 (0.98), 277 (0.57), 352 (0.42); Anal.
- 200 Calc. for C₂₄H₂₉ClF₆N₄O₂PRh (688.84): C, 41.85; H, 4.24; N, 8.13. Found: C, 41.96; H, 4.16; N,
- 201 8.23%.
- 202 2.5.2. [Cp*Ir(L1)Cl]PF₆ (2)
- 203 Yield: 70 mg (45%); IR (KBr, cm⁻¹): 3447(m), 3241(m), 2925(m), 1630(s), 1610(m), 1293(m),
- 204 846(s); ¹H NMR (400 MHz, CDCl₃): $\delta = 10.4$ (s, 1H, OH), 9.02 (s, 1H, CH_(imine)), 8.77 (d, 1H, J

- 205 = 4.0 Hz, $CH_{(py)}$), 8.51 (d, 1H, J = 4.0 Hz, $CH_{(py)}$), $8.17 \text{ (t, 1H, } CH_{(py)}$), $7.78 \text{ (t, 1H, } CH_{(py)}$), $7.42 \text{ (t, 2H, } CH_{(py)})$
- 206 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$), 7.39 (s, 2H, NH₂), 6.56 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$), 6.54 (s, 1H,
- 207 $CH_{(Ar)}$), 3.87 (s, 3H, OMe), 1.62 (s, 15H, $CH_{(Cp^*)}$); HRMS-APCI (m/z): 597.18 [M-PF₆-HCl]⁺;
- 208 UV–Vis {Acetonitrile, λ_{max} , nm ($\epsilon/10^{-4}$ M⁻¹ cm⁻¹)}: 266 (0.36), 347 (0.29); Anal. Calc. for
- 209 C₂₄H₂₉ClF₆N₄O₂PIr (778.14): C, 37.04; H, 3.76; N, 7.20. Found: C, 37.19; H, 3.89; N, 7.31%.
- 210 2.5.3. [Cp*Rh(L2)Cl]PF₆ (3)
- 211 Yield: 52 mg (39%); IR (KBr, cm⁻¹): 3422(m), 3310(w), 2923(w), 1636(s), 1603(m), 1457(m),
- 212 845(s); ¹H NMR (400 MHz, CDCl₃): $\delta = 10.1$ (s, 1H, OH), 9.11 (s, 1H, CH_(imine)), 8.78 (d, 1H, J
- 213 = 4.0 Hz, $CH_{(py)}$), 8.49 (d, 1H, J = 8.0 Hz, $CH_{(py)}$), 8.14 (t, 2H, $CH_{(py)}$), 7.78 (t, 1H, $CH_{(Ar)}$), 7.60
- 214 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$), 7.38 (s, 2H, NH₂), 6.92-7.01 (m, 2H, $CH_{(Ar)}$), 1.58 (s, 15H, $CH_{(Cp^*)}$);
- 215 HRMS-APCI (m/z): 477.12 [M-PF₆-HCl]⁺; UV–Vis {Acetonitrile, λ_{max} , nm ($\epsilon/10^4$ M⁻¹ cm⁻¹)}:
- 216 235 (1.55), 283 (0.79), 348 (1.00); Anal. Calc. for C₂₃H₂₇ClF₆N₄OPRh (658.81): C, 41.93; H,
- 217 4.13; N, 8.50. Found: C, 42.08; H, 4.25; N, 8.68%.
- 218 $2.5.4. [Cp*Ir(L2)Cl]PF_6$ (4)
- Yield: 52 mg (34%); IR (KBr, cm⁻¹): 3479(s), 3329(s), 2924(w), 1642(s), 1618(m), 1602(m),
- 220 842(s); ¹H NMR (400 MHz, CDCl₃): $\delta = 10.1$ (s, 1H, OH), 9.13 (s, 1H, CH_(imine)), 8.80 (d, 1H, J
- = 4.0 Hz, $CH_{(py)}$), 8.56 (d, 1H, J = 8.0 Hz, $CH_{(py)}$), 8.18 (t, 2H, $CH_{(py)}$), 7.80 (t, 1H, $CH_{(Ar)}$), 7.64
- 222 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$), 7.40 (s, 2H, NH₂), 6.99-7.35 (m, 2H, $CH_{(Ar)}$), 1.63 (s, 15H, $CH_{(Cp^*)}$);
- 223 HRMS-APCI (m/z): 567.17 [M-PF₆-HCl]⁺; UV–Vis {Acetonitrile, λ_{max} , nm ($\epsilon/10^4$ M⁻¹ cm⁻¹)}:
- 224 291 (0.62), 344 (0.78); Anal. Calc. for C₂₃H₂₇ClF₆N₄OPIr (748.12): C, 36.93; H, 3.64; N, 7.49.
- 225 Found: C, 37.11; H, 3.83; N, 7.62%.
- 226 $2.5.5. [(Cp*Rh(L3)Cl]PF_6(5)]$

- Yield: 58 mg (43%); IR (KBr, cm⁻¹): 3452(s), 3318(s), 2924(m), 1648(s), 1600(m), 1566(m),
- 228 1489(m), 842(s); ¹H NMR (400 MHz, DMSO-d₆): δ = 12.5 (s, 1H, OH), 8.96 (d, 1H, J = 4.0 Hz,
- 229 $CH_{(py)}$), 8.33-8.38 (m, 3H, $CH_{(py)}$), 7.91 (t, 1H, $CH_{(Ar)}$), 7.86 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$), 7.48 (t,
- 230 1H, J = 8.0 Hz, $CH_{(Ar)}$), 7.01-7.06 (m, 3H, NH_2 , $CH_{(Ar)}$), 2.48 (s, 3H, CH_3), 1.59 (s, 15H,
- 231 $CH_{(Cp^*)}$; HRMS-APCI (m/z): 491.14 [M-PF₆-HCl]⁺; UV–Vis {Acetonitrile, λ_{max} , nm ($\epsilon/10^4$ M⁻¹
- 232 cm⁻¹)}: 229 (0.95), 268 (0.59), 332 (0.32); Anal. Calc. for C₂₄H₂₉ClF₆N₄OPRh (672.84): C,
- 233 42.84; H, 4.34; N, 8.33. Found: C, 42.98; H, 4.26; N, 8.48%.
- 234 2.5.6. $[Cp*Ir(L3)Cl]PF_6$ (6)
- 235 Yield: 65 mg (42%); IR (KBr, cm⁻¹): 3460(m), 3237(m), 2926(w), 1630(s), 1595(m), 1296(m),
- 236 846(s), 3456(m), 3369(m), 2925(m), 1649(s), 1618(m), 1598(m), 1306(m), 845(s); ¹H NMR
- 237 (400 MHz, DMSO-d₆): δ = 12.3 (s, 1H, OH), 8.94 (d, 1H, J = 4.0 Hz, CH_(DV)), 8.44 (d, 1H, J =
- 238 4.0 Hz, $CH_{(pv)}$), 8.35 (t, 2H, $CH_{(pv)}$), 7.90 (t, 1H, $CH_{(Ar)}$), 7.86 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$), 7.48 (t,
- 239 1H, $CH_{(Ar)}$), 7.01-7.06 (m, 3H, NH_2 , $CH_{(Ar)}$), 2.46 (s, 3H, CH_3), 1.58 (s, 15H, $CH_{(Cp^*)}$); HRMS-
- 240 APCI (m/z): 581.19 [M-PF₆-HCl]⁺; UV–Vis {Acetonitrile, λ_{max} , nm ($\epsilon/10^{-4}$ M⁻¹ cm⁻¹)}: 209
- 241 (1.27), 263 (0.66), 330 (0.36); Anal. Calc. for $C_{24}H_{29}ClF_6N_4OPIr$ (762.15): C, 37.82; H, 3.84; N,
- 242 7.35. Found: C, 37.96; H, 3.96; N, 7.44%.
- 243 2.5.7. $[(Cp*Rh(L4)Cl)PF_6(7)]$
- Yield: 54 mg (41%); IR (KBr, cm⁻¹): 3441(s), 3137(m), 2961(w), 1640 (s), 1593(m), 1464(m),
- 245 841(s); ¹H NMR (400 MHz, DMSO-d₆): $\delta = 8.96$ (d, 1H, J = 4.0 Hz, $CH_{(py)}$), 8.33-8.37 (m, 2H,
- 246 $CH_{(py)}$), 7.88 (t, 1H, $CH_{(py)}$), 7.81 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$), 7.46 (t, 1H, $CH_{(Ar)}$), 7.23-7.28 (m,
- 247 2H, $CH_{(Ar)}$), 6.97-7.02 (m, 3H, NH_2 , $CH_{(Ar)}$), 2.47 (s, 3H, CH_3), 1.59 (s, 15H, $CH_{(Cp^*)}$); HRMS-
- 248 APCI (m/z): 511.12 [M-PF₆]⁺; UV–Vis {Acetonitrile, λ_{max} , nm ($\epsilon/10^{-4}$ M⁻¹ cm⁻¹)}: 229 (1.37),

- 249 265 (0.37), 400 (0.22); Anal. Calc. for C₂₄H₂₉ClF₆N₄PRh (656.84): C, 43.89; H, 4.45; N, 8.53.
- 250 Found: C, 44.02; H, 4.39; N, 8.61%.
- 251 2.5.8. [Cp*Ir(L4)Cl]PF₆ (8)
- 252 Yield: 65 mg (43%); IR (KBr, cm⁻¹): 3458(s), 3383(s), 2922(m), 1643(s), 1603(m), 1567(m),
- 253 1447(m), 844(s); ¹H NMR (400 MHz, DMSO-d₆): $\delta = 8.97$ (d, 1H, J = 4.0 Hz, CH_(pv)), 8.31-8.34
- 254 (m, 2H, $CH_{(pv)}$), 7.85 (t, 1H, $CH_{(pv)}$), 7.79 (d, 1H, J = 8.0 Hz, $CH_{(Ar)}$), 7.44 (t, 1H, $CH_{(Ar)}$), 7.19-
- 255 7.23 (m, 2H, $CH_{(Ar)}$), 6.99-7.03 (m, 3H, NH_2 , $CH_{(Ar)}$), 2.46 (s, 3H, CH_3), 1.59 (s, 15H, $CH_{(Cp^*)}$);
- 256 HRMS-APCI (m/z): 601.17 [M-PF₆]⁺; UV–Vis {Acetonitrile, λ_{max} , nm (ϵ /10⁻⁴ M⁻¹ cm⁻¹)}: 256
- 257 (0.53), 361 (0.20); Anal. Calc. for $C_{24}H_{29}ClF_6N_4Plr$ (746.15): C, 38.63; H, 3.92; N, 7.51. Found:
- 258 C, 38.74; H, 4.03; N, 7.63%.
- 259 3. Results and discussion
- 260 3.1. Synthesis of ligands and complexes
- The azine Schiff-base ligands (L1-L4) were prepared by the reaction of 2-
- 262 pyridylamidrazone and the respective aldehyde or ketone in absolute ethanol medium. The
- complexes (1-8) were synthesized by the reaction of Rh/Ir metal precursors with the azine
- Schiff-base ligands. The cationic complexes were isolated with PF₆ counter ion. All these metal
- complexes were obtained in good yields and are yellow in color. They are stable in air as well as
- 266 in solid state, and are non-hygroscopic. These complexes are soluble in common organic
- solvents such as dichloromethane, acetonitrile and acetone but insoluble in diethyl ether and
- 268 hexane. All the synthesized ligands and complexes were fully characterized by spectroscopic
- techniques.
- 270 *3.2. Spectroscopic characterization of ligands*

The infrared spectra of the free ligand shows characteristic stretching frequencies for NH₂, OH, C=N and C=C groups. The NH₂ and OH stretching frequencies for the azine ligand appeared in the range of 3300-3500 cm⁻¹. The C=C and C=N stretching frequencies were observed in the range of 1550-1626 cm⁻¹. The proton NMR spectra of the ligands displayed signals in the range of 7.30-8.57 ppm assignable to the protons of the pyridine ring. The imine protons for L1 and L2 are located at 8.60 and 8.59 ppm respectively. The methoxy proton signal was observed as a singlet for L1 at 3.80 ppm. The methyl protons of L3 and L4 were observed as a singlet at 2.62 and 2.39 ppm respectively. The hydroxyl proton resonance for the ligands appeared in the range of 11.5-11.9 ppm. The aromatic protons of the ligand appeared as doublet, triplet and multiplet in the range of 6.21-7.29 ppm. The [M+H]⁺ molecular ion peak for the ligands are shown in the experimental section which are found to be in good agreement with the expected range. The electronic spectra of the free ligands are shown in (Figure S1). The electronic spectra of the free ligands show absorption bands in the range of 210-360 nm. The band in the range of 210-250 nm can be assigned as π - π * and n- π * transition. The band around 300-370 nm is due to the intermolecular charge transfer transition within the whole molecule [41].

3.3. Spectroscopic characterization of complexes

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The IR spectra of the complexes show sharp bands around 842-846 cm⁻¹ due to the P-F stretching frequency of the counter ion [42]. The OH and NH₂ stretching vibrations in the complexes were found around 3300-3500 cm⁻¹. The retaining of the OH and NH₂ stretching frequencies indicates that they are not involved in bonding to the metal center. The strong absorption band for $V_{C=N}$ around 1630-1650 cm⁻¹ at higher wave numbers as compared to the free

ligand around 1615-1626 cm⁻¹ suggest that the coordination to the metal occurs through the imine and pyridine nitrogen.

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The proton NMR spectra of the metal complexes show that the ligand resonance signals are shifted downfield as compared to that of the free ligand. These signals are shifted downfield because of the ligand coordination to the metal atom. The imine proton signal was observed in the range of 9.0-9.13 ppm for complexes (1-4). The hydroxyl proton resonance for the complexes appeared in the range of 10.1-12.5 ppm respectively. The appearance of the hydroxyl proton signal indicates that the hydroxyl group is not involved in bonding to the metal atom. The pyridine ring protons also showed downfield signals comprising of doublet and triplet in the range of 7.75-8.96 ppm. The NH₂ protons were observed as a singlet for complexes (1-4) in the range of 7.35-7.37 ppm respectively. The methoxy proton resonance for complexes (1 and 2) appeared as a singlet at 3.81 and 3.83 ppm. The aromatic proton signals for complexes appeared in the range of 6.50-7.86 ppm as doublet, triplet and multiplet. The methyl proton signal for complexes (5-8) appeared as a singlet around 2.46-2.48 ppm respectively. In addition to the signals for the ligand protons, a sharp singlet was observed for all the complexes between 1.58-1.63 ppm respectively corresponding to the methyl protons of the Cp* ring. In the mass spectra of the complexes (1–6) the peaks at m/z: 507.12, m/z: 597.18, m/z: 477.12, m/z: 567.17, m/z: 491.13 and m/z: 581.20 can be assigned as [M-PF₆-HCl]⁺ ion peaks respectively. Whereas, the mass spectra of the complex 7 and 8 displayed molecular ion peaks at m/z: 511.12 and 601.17 which corresponds to the $[M-PF_6]^+$ ion.

The electronic spectra of the complexes were recorded in acetonitrile at 10⁻⁴ M concentration at room temperature and the plot is shown in (Figure S2). The electronic spectra of complexes display two absorption band in the higher energy region around 210-330 nm. The

bands in the higher energy UV region can be assigned as ligand centered or intra ligand π - π * and n- π *transition. The Rh(III) and Ir(III) complexes provides filled d π (t_{2g}) orbitals which can interact with low lying π * orbitals (C=N) of the ligand. The band in the lower energy region around 345-405 nm can be assigned as Rh (d π) or Ir (d π) to π * ligand metal to ligand charge transfer (MLCT) transition [43].

3.4. Molecular structures of complexes

The molecular structures of some of the respective complexes have been elucidated by single crystal X-ray analysis. Suitable single crystals were attached to a glass fibre and transferred into the Oxford Diffraction Xcalibur Eos Gemini diffractometer. The crystallographic details and structure refinement details are summarized in Table 1. The geometrical parameters around the metal atom involving ring centroid are listed in Table S1. In all these complexes the ligand is coordinated to the metal atom in a similar manner with $N\cap N$ binding mode. Complex (1) and (8) crystallized in triclinic system with space group PT. Complex (8) crystallized with one PF₆ and one chloride counter ion. Complex (3) and (4) crystallized in monoclinic system with space group P2₁/c whereas complex (7) crystallized in monoclinic system with space group P2₁.

All these complexes display a typical three-legged piano stool geometry around the metal center with coordination sites occupied by one chloride group, two σ bonded nitrogen atoms from chelating azine ligand and the pentamethylcyclopentadienyl (Cp*) ring in η^5 manner. The metal atom in all these complexes is situated in a pseudo-octahedral arrangement with the azine ligand coordinating through the pyridine and azine nitrogen atoms forming a five membered metallocycle. In complexes (1), (3), (4) and (7) the M-N bond length {2.088(5), 2.099(3), 2.098(4) and 2.102(4) Å} from pyridine is comparatively shorter than the azine nitrogen-metal

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distances {2.135(5), 2.116(3), 2.105(4) and 2.159(4) Å}, which are similar to those, reported with similar complexes [24, 44]. However in complex (8) the metal-nitrogen distance from pyridine {2.102(5) Å} is comparatively larger than azine nitrogen-metal distance, which is {2.096(5) Å}. The C=N bond length of the coordinated nitrogen in complex (1), (3), (4) and (8) is longer than that of the uncoordinated C=N (Table S1) which could be due to the back bonding of electron from metal $(d\pi)$ to π^* orbital of the ligand. But in complex (7), a reverse pattern has been observed where the C=N bond length of the coordinated nitrogen {1.346(7) Å} is shorter than uncoordinated C=N {1.358(7) Å} bond. The average M-C distances are {2.159 (1), 2.1534 (3), 2.1616 (4), 2.1528 (7) and 2.1726 (8) Å} while the distance between the metal to Cp* centroid ring is in the range of 1.758–1.793 Å respectively. The M-Cl bond lengths {2.3976(15) (1), 2.4172(9) (3), 2.4190(12) (4), 2.4242(16) (7) and 2.4220(17) (8) shows no significant differences and is comparable to previously reported values (Table 1) [45-48]. The bite angle N(1)-Rh(1)-N(2) values are 75.10(19) (1), 75.09(11) (3), and 75.44(17) (7) whereas in complex (4) and (8) the bite angle values are N(1)-Ir(1)-N(2) values are 74.99(14) (4) and 75.26(18) respectively which probably indicates an inward bending of the coordinated pyridyl and azine group [49]. The bond angles N(1)-M-Cl(1) and N(2)-M-Cl(1) in complexes are comparable to the piano stool arrangement about the metal atom and is comparable to reported values for closely related systems [50-52]. Further the crystal packing in complex (1) is stabilized by weak intermolecular hydrogen bonding C-H·····O (2.702 Å) between the hydrogen atom from methoxy group and oxygen atom of the hydroxyl group and C-H·····Cl (2.793 Å) interaction between CH₃ group of Cp* and chloride atom (Figure S3). These interactions play a significant role in the formation of supramolecular motifs.

On the other hand in the crystal structure of complex (3) and (4) two types of intramolecular hydrogen bonding has been observed; the first one between the uncoordinated nitrogen atom of the azine linkage with the hydrogen atom of the hydroxyl group O-H····N (1.916 and 1.908 Å) and the second between the hydrogen atom from NH₂ and uncoordinated azine nitrogen atom N-H·····N (2.323 and 2.328 Å) (Figure 4). The selected hydrogen bonding distances and angles for complex (3) and (4) are given in (Table 2). Also the crystal packing in complex (3) and (4) is further stabilized by two different C-H·····Cl interaction between the Cl atom attached to metal M (where M = Rh/Ir) with hydrogen atom of pyridine ring and NH₂ (Figure S4). Complex (7) shows C-H····· π (2.832 and 2.937 Å) interactions between the methyl hydrogen atom and Cp* moiety and between pyridine ring and hydrogen atom of Cp* group respectively (Figure S5). Interestingly the crystal packing in complex (8) leads to a dimeric unit via intermolecular C-H·····Cl interaction between the chloride counter ion and hydrogen atom from pyridine ring, NH₂ and Cp* group (Figure S6).

3.5. Chemosensitivity studies

The complexes (1-8) were tested for their cytotoxicity against cancer cell line HT-29 (human colorectal cancer), and non-cancer cell line ARPE-19 (human retinal epithelial cells). The response of the cell lines HT-29 to the test complexes and cisplatin (1-8) is presented in graphical form in Figure 5 and in tabular form in Table 3. All the complexes tested were found to be active against HT-29 cancer cell line (IC₅₀< 30 μ M). Complex (5) was the most potent among all the complexes with (IC₅₀ value of 96.93 \pm 5.31 μ M). However all the complexes were less potent than cisplatin (IC₅₀ value of 0.25 \pm 0.11 μ M against HT-29). The selectivity index (SI) defined as the ratio of IC₅₀ values in ARPE19 cells divided by the IC₅₀ value of cancer cell line demonstrates that all the complexes are effective against cancer cell with SI values ranging from

1.01 to 2.11 (Table S2). Moreover although complex (**5**) showed more selectivity than other complexes for HT-29 cancer cell, however its selectivity was significantly lower than cisplatin where SI value is 25.64 (Figure 6).

3.6. *Optimized geometry*

The comparison of the geometric parameters (selected bond lengths and bond angles) of the optimized structures and the crystal structures of the complexes (1, 3, 4, 7 and 8) are listed in Table S3. All the metal complexes are found to be closed shell structures. The calculated bond lengths and the bond angles of the complexes are in good agreement with the experimental data indicating the reliability of the theoretical method (B3LYP/6-31G**/LanL2DZ) used in the present study. It should be noted that for complexes (3, 4, 7 and 8), the M(1)-N(2) (where M = Rh/Ir) bond length is slightly longer than the M(1)-N(1) bond length whereas for complex (1), a reverse pattern has been observed (Table S3).

3.7. Charge distribution

The charges on the individual atoms for the metal complexes obtained from NBO analysis are listed in Table S4. The charges on the Rh atom in the complexes (1), (3), (5) and (7) are 0.136, 0.200, 0.216 and 0.214 e whereas the charges on Ir for complexes (2), (4), (6) and (8) are 0.186, 0.252, 0.268 and 0.214 e respectively. These NBO charges on Rh and Ir are comparatively lower than their formal charge of +3 which suggests that the ligand transfers their negative charge to the respective rhodium and iridium metal on complex formation. In metal complexes (1-8), the charge on Cl ranges between -0.439 e (Complex-1) to -0.394 e (Complex-4). In isolated ligands, the charge on N(1) ranges between -0.416 and -0.417 e whereas for N(2) it ranges between -0.324 e and -0.348 e. It should be noted that for isolated ligands as well as for complexes (1-8), the negative charges on N(1) (-0.385, -0.381, -0.372, -0.373, -0.369, -0.398, -0

0.368 and -0.373 e) are slightly higher than the charges on N(2) (-0.258, -0.253, -0.284, -0.283, -0.305, -0.297, -0.311 and -0.305 e). On complex formation, the negative charge on the N(1) and N(2) reduces slightly giving an indication of the charge transfer on Rh and Ir in metal complexes. The population of the 4d ($4d_{xy}$, $4d_{xz}$, $4d_{yz}$, $4d_{x}^2$, $4d_{x}^2$ and $4d_{z}^2$) orbital of Rh complexes and 5d orbital of Ir complexes are shown in Table S5. The orbital occupations of each orbital (nd_{xy}, nd_{yz}, nd_{yz}, nd_x² and nd_z²) for all the complexes are comparatively higher in rhodium complexes than iridium complexes. In free Rh(III) and Ir(III) state, the population of nd_{xy}, nd_{xz} and nd_{yz} are 2.0, 2.0 and 2.0 e and the other two orbitals remain vacant. But on complex formation, the population on nd_{xy}, nd_{xz} and nd_{yz} orbital gets reduced whereas the nd_{x2-y2} and nd_z² orbitals gain some population as indicated in Table S5. For most of the complexes, the population of 4d and 5d orbital containing the same ligand follow similar pattern of filling, except for the complexes containing ligand L1 where the nd_{xz} orbital population is slightly lower and nd_x²-y² is higher as compared to the other complexes.

3.8. Frontier molecular orbitals and absorption spectra

The molecular orbital representation of the complexes along with their HOMO, LUMO energies and HOMO-LUMO energy gaps are shown in Figure 7. The HOMO-LUMO energy gap can be used as an important parameter in analyzing the chemical reactivity and kinetic stability of a molecule. This energy gap is also related to the hardness/softness of a chemical species [53]. The lower HOMO-LUMO energy gap is a suitable condition where a molecule can be excited easily and thereby increasing its reactivity and decreasing its kinetic stability whereas higher energy gap can lead to more kinetic stability but less reactivity. The HOMO-LUMO energy gaps for all the complexes (1-8) are found to be 3.20, 2.98, 3.63, 3.46, 3.61, 3.60, 3.68 and 3.59 eV respectively. The gap is slightly lower for the iridium complexes as compared to rhodium

complexes containing the same ligand indicating the reactivity of Ir complexes over the
complexes containing Rh metal. The % contribution of molecular orbital analysis as shown in
Table S6, predicts that the most percentage of HOMO is located on the ligand itself except for
complex (2) and (8) where as it is mostly present on the Ir metal. On the other hand, LUMO is
located on the ligand for complexes (1) (about 97%), (2) (91%), (4) (89%), (6) (92%) and (8)
(69%) whereas for complexes (3) (40%), (5) (35%) and (7) (38%), it is located on the Rh metal.

The electronic absorption spectra were calculated using the TD-DFT method in acetonitrile solvent employing PCM model. The calculated and the experimental absorption data, HOMO-LUMO energy gaps, and the character of electronic transitions are listed in Table 4. The H \rightarrow L transitions for complexes (1), (4) and (6) occurring at 417, 444 and 441 nm corresponds to ILCT character, for complexes (2) and (8) at 463 and 440 nm corresponds to MLCT character whereas for complexes (3), (5) and (7) at 532, 519 and 518 nm corresponds to LMCT character. These MLCT character can be assigned for $d\pi(M)\rightarrow\pi*(L)$ transitions whereas the ILCT character are for $\pi\rightarrow\pi*$ transitions. It should be noted that all LMCT transitions are occurring at higher wavelength regions (i.e. > 500 nm). In good agreement with the experimental data, the TD-DFT calculations shows few MLCT transitions at 358 nm complex (2), 332 nm, complex (4), 334 nm complex (6) and 372, 358 nm complex (8). However, in the range between 340-400 nm, few LMCT, ILCT and LLCT transitions have also been observed (Table 4).

4. **Conclusion**

In summary, we have synthesized four new azine Schiff-base ligands and its rhodium and iridium half-sandwich complexes. All these complexes and ligands were full characterized by various spectroscopic techniques. The ligands under study preferably bind to the metal in a bidentate $N\cap N$ fashion using pyridine and one azine nitrogen atom. Our attempt to synthesize

dinuclear rhodium and iridium complexes with NN' and NO bonding was however unsuccessful irrespective of molar ratio of metal to ligand where as in the presence of base, it leads to decomposition of the reaction. These complexes possess some important intramolecular and intermolecular hydrogen bonding and also possess some weak non-covalent interactions, particularly C-H·····Cl and C-H···· π interactions. Chemosensitivity activity of the complexes against HT-29 cancer cell demonstrates that the complexes are active however complex (5) was found to be the most potent among all other complexes. Theoretical studies reveal that the HOMO-LUMO energy gap is lower for iridium complexes indicating better reactivity over the rhodium complexes. TD-DFT calculations were carried out in order to evaluate the electronic transitions occurring in the metal complexes, which are in good agreement with the experimental results. The charge distribution analysis (using NBO analysis) of these complexes helps to understand how the charges on nitrogen atom (which are coordinating to the metal) are delocalized on complex formation. Especially, the NBO charges, on rhodium and iridium confirm that the ligands transfer their negative charge to the respective metal on complex formation. The lower HOMO-LUMO energy gap leads to greater chemical reactivity but lesser kinetic stability and vice versa. Furthermore, the nature of HOMO and LUMO illustrate the electronic origin of the lowest energy transition and the resulting electronic reorganization. Moreover, the molecular orbital analysis was helpful to understand and locate the % contribution of HOMO and LUMO on different fragments of the complexes, which is otherwise not possible to predict from experimental data.

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Sanjay Adhikari and Dipankar Sutradhar thanks UGC, New Delhi, India for providing financial assistance in the form of university fellowship (UGC-Non-Net). We thank DST-PURSE

476	SCXF	RD, NEHU-SAIF, Shillong, India for providing Single crystal X-ray analysis and other
477	spectr	ral studies. AKC thanks Computer centre, NEHU, for computational facilities.
478	Supp	lementary material
479		CCDC 1477976 (1), 1477977 (3), 1477978 (4), 1477979 (7) and 1477980 (8) contains
480	the su	pplementary crystallographic data for this paper. These data can be obtained free of charge
481	via <u>w</u>	www.ccdc.cam.ac.uk/data_request/cif, by e-mailing data_request@ccdc.cam.ac.uk, or by
482	contac	cting The Cambridge Crystallographic Data Centre, 12, Union Road, Cambridge CB2 1EZ,
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Scheme-1 Synthesis of 2-pyridylamidrazone

$$R_2$$
 NH_2
 NH_2
 R_3
 R_2
 R_1
 R_2
 R_2
 R_1
 R_2
 R_3
 R_2
 R_3
 R_4
 R_5
 R_4
 R_5
 R_5
 R_7
 R_7
 R_8

Scheme-2 Preparation of ligands (L1-L4)

Scheme-3 Preparation of metal complexes (1-8)

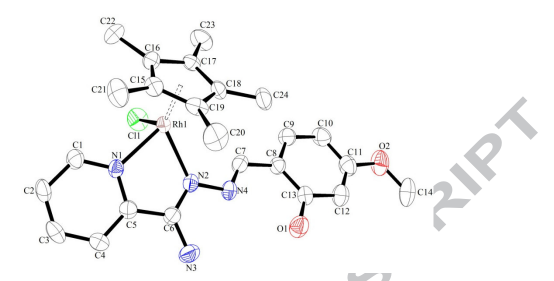


Figure 1 ORTEP diagram of complex [Cp*RhCl(L1)Cl]PF₆ (1) with 50% probability thermal ellipsoids. Hydrogen atoms and counter ions are omitted for clarity.

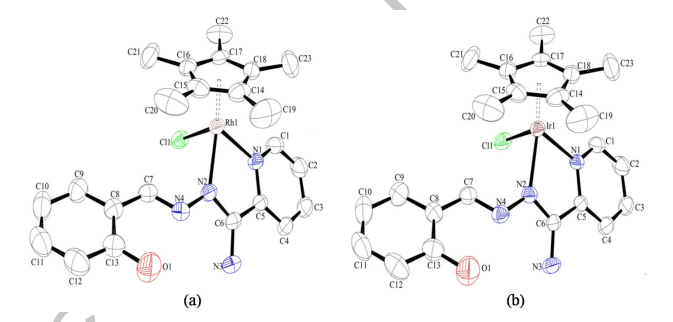


Figure 2 (a) ORTEP diagram of complex [Cp*RhCl(L2)Cl]PF₆ (**3**) and (b) ORTEP diagram of complex [Cp*IrCl(L2)Cl]PF₆ (**4**) with 50% probability thermal ellipsoids. Hydrogen atoms and counter ions are omitted for clarity.

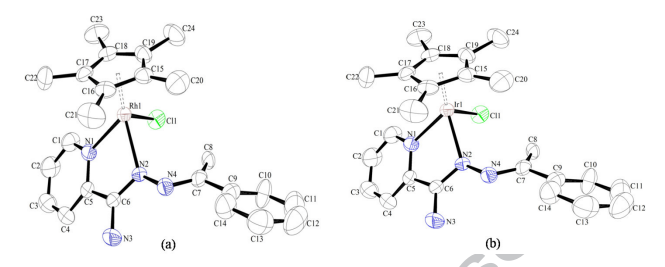


Figure 3 (a) ORTEP diagram of complex [Cp*Rh(L4)Cl]PF₆ (7) and (b) ORTEP diagram, of complex [Cp*IrCl(L4)Cl]PF₆ (8) with 50% probability thermal ellipsoids. Hydrogen atoms and counter ions are omitted for clarity.

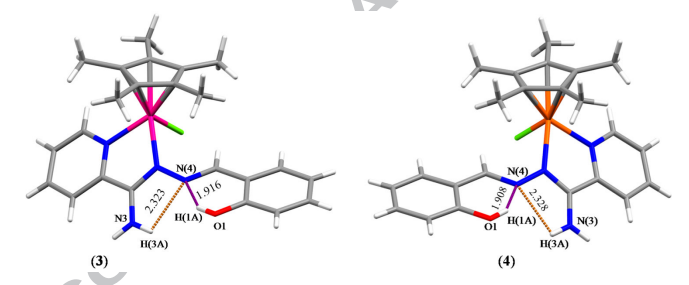


Figure 4 Crystal structure of complexes (3) and (4) showing intramolecular hydrogen bonding.

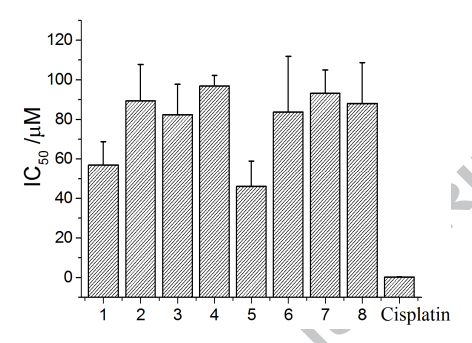


Figure 5 Response of HT-29 (human colorectal cancer) to compounds (1-8) and cisplatin. Cell were exposed to compounds (1-8) for 96 hours. Each value represents the mean ± standard deviation from three independent experiments.

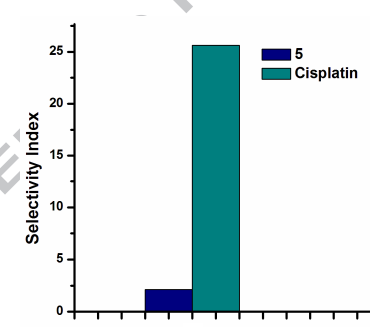
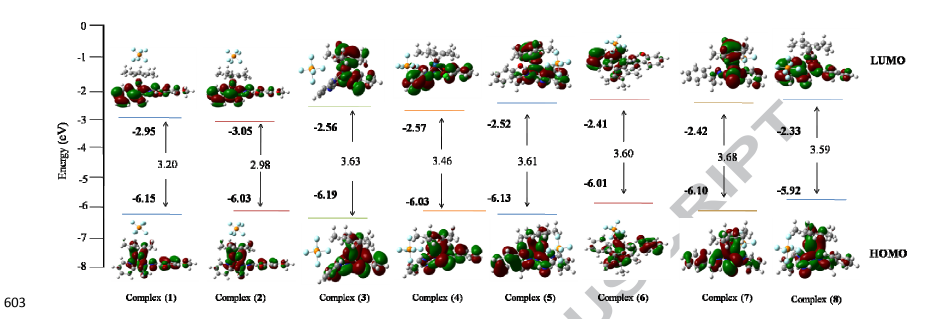


Figure 6 Graph showing selectivity index of complex **5** and cisplatin against HT-29 cancer cell line. The selectivity index is defined as the IC_{50} of ARPE19 cell divided by the IC_{50} of tumour cell line.



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Figure 7 HOMO, LUMO energies and their energy gap of complexes (1–8)

Table 1. Crystal structure data and refinement parameters of complexes.

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Complexes	[1] PF ₆	[3] PF ₆	[4] PF ₆	[7] PF ₆	[8] PF ₆ Cl
Empirical formula	$C_{24}H_{29}ClN_4O_2F_6PRh$	C ₂₃ H ₂₇ ClF ₆ N ₄ OPRh	C ₂₃ H ₂₇ ClF ₆ N ₄ OPIr	$C_{24}H_{29}ClF_6N_4PRh$	$C_{24}H_{29}Cl_2F_6N_4PIr$
Formula weight	688.84	658.82	748.11	656.84	781.58
Temperature (K)	298(2)	293(2)	293(2)	293(2)	293(2)
Wavelength (Å)	0.71073	0.71073	0.71073	0.71073	0.71073
Crystal system	triclinic	monoclinic	monoclinic	monoclinic	triclinic
Space group	PT	$P2_{I}/c$	$P2_{I}/c$	$P2_1/c$	PT
a (Å)/α (°)	8.3893(7)/89.370(6)	10.6710(6)/90	10.7019(5)/90	38.850(5)/90	7.9976(4)/87.496
b (Å)/β (°)	10.5533(7)/86.439(6)	17.0730(8)/92.708(4).	17.0860(9)/93.062(4)	7.9488(5)/98.027(4)	12.4774(4)/82.086(4)
c (Å)/γ (°)	16.6554(11)/71.182(7)	14.5390(8)/90	14.6118(9)/90	28.562(4)/90	14.6442(6)/72.596(4)
Volume (Å ³)	1393.00(18)	2645.8(2)	2668.0(2)	1344.83(10)	1381.15(10)
Z	2	4	4	2	2
Density (calc) (Mg/m ⁻³)	1.642	1.654	1.862	1.622	1.879
Absorption coefficient (μ) (mm ⁻¹)	0.836	0.874	5.231	0.857	5.148
F(000)	696	1328	1456	664	762
Crystal size (mm ³)	0.23 x 0.21 x 0.21	0.21 x 0.19 x 0.04	0.23 x 0.23 x 0.21	0.22 x 0.20 x 0.120	0.19 x 0.12 x 0.09
Theta range for data collection	3.174 to 28.654°.	3.33 to 26.73°.	3.31 to 26.37°.	3.386 to 28.842°.	3.23 to 26.37°.
Index ranges	-11<=h<=10, -12<=k<=13, -	-13<=h<=10, -10<=k<=21, -	-13<=h<=7, -21<=k<=19, -	-9<=h<=9, -12<=k<=22, -	-8<=h<=9, -15<=k<=15, -
	22<=l<=20	12<=l<=18	16<=l<=18	14<=l<8	17<=l<18
Reflections collected	10811	9506	10081	5614	7889
Independent reflections	6286 [R(int) = 0.0717]	5375 [R(int) = 0.0268]	5422 [R(int) = 0.0277]	4000 [R(int) = 0.0268]	5335 [R(int) = 0.0296]
Completeness to theta = 25.00°	99.57 %	99.5 %	99.2 %	99.2 %	94.8 %
Absorption correction	Semi-empirical from	Semi-empirical from	Semi-empirical from	Semi-empirical from	Semi-empirical from
	equivalents	equivalents	equivalents	equivalents	equivalents
Refinement method	Full-matrix least-squares on	Full-matrix least-squares on	Full-matrix least-squares on	Full-matrix least-squares on	Full-matrix least-squares on
	F^2	F^2	F^2	F^2	F^2
Data/restraints/parameters	6286/0/362	2375/0/330	5422/0/340	4000/1/340	5335/0/349
Goodness-of-fit on F ²	1.197	1.063	1.026	1.041	1.046
Final R indices [I>2sigma(I)]	R1 = 0.0703, $wR2 = 0.1706$	R1 = 0.0440, $wR2 = 0.0895$	R1 = 0.0340, $wR2 = 0.0630$	R1 = 0.0394, $wR2 = 0.0822$	R1 = 0.0368, $wR2 = 0.0875$
R indices (all data)	R1 = 0.0855, $wR2 = 0.1772$	R1 = 0.0592, $wR2 = 0.0968$	R1 = 0.0500, $wR2 = 0.0683$	R1 = 0.0462, $wR2 = 0.0858$	R1 = 0.0431, $wR2 = 0.0912$
Largest diff. peak and hole (e.Å ⁻³)	0.583 and -0.461	0.520 and -0.543	1.102and -1.143	0.512 and -0.478	1.828 and -1.071
CCDC No.	1477976	1477977	1477978	1477979	1477980

Structures were refined on F_0^2 : $wR_2 = \left[\sum [w(F_0^2 - F_c^2)^2] / \sum w(F_0^2)^2\right]^{1/2}$, where $w^{-1} = \left[\sum (F_0^2) + (aP)^2 + bP\right]$ and $P = \left[\max(F_0^2, 0) + 2F_c^2\right]/3$.

Table-2. Selected hydrogen bonding distances (Å) and angles (°) of complexes 3 and 4.

Complexes	D-H·····A	H····A (Å)	DA (Å)	DH (Å)	∠D—H····A(°)
3	O(1)-H(1A)·····N(4)	1.916	2.638	0.820	146.39
	N(3)-H(3A)N(4)	2.323	2.624	0.860	100.76
4	O(1)- $H(1A)$ $N(4)$	1.908	2.634	0.820	146.90
	N(3)-H(3A)N(4)	2.328	2.629	0.860	100.78

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Table-3 Response of HT-29 (human colorectal cancer) to complexes (1-8) and cisplatin. Each

value represents the mean \pm standard deviation from three independent experiments.

Complexes	IC ₅₀ (μM)	60	
	HT-29	ARPE-19	
1	56.95 ± 11.76	85.31 ± 14.86	
2	89.42 ± 18.33	93.45 ± 11.34	
3	82.32 ± 15.55	83.03 ± 14.76	
4	96.93 ± 5.31	>100	
5	46.17 ± 12.78	97.39 ± 4.53	
6	83.74 ± 28.17	>100	
7	93.16 ± 11.84	>100	
8	88.09 ± 20.63	>100	
Cisplatin	0.25 ± 0.11	6.41 ± 0.95	

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Table 4. The energy gap, theoretical and experimental absorption bands, electronic transitions and dominant excitation character for various singlet states of the complexes (1-8) calculated with TD-DFT method.

The most	Calculated	Energy	Oscillator	Dominant excitation	Experimental
important orbital	λ (nm)	gap E	strength	Character	λ (nm)
excitations		(eV)	(f)		
			Complex (1)		
H→L	417.16	3.20	0.2051	L1→L1(ILCT)	
H-2→L	359.64	3.53	0.0542	$Cl\rightarrow L1(LLCT)$	352.21
$H\rightarrow L+2$	355.41	4.03	0.0120	$L1\rightarrow L1(ILCT)$	
$H-4\rightarrow L+2$	338.40	4.89	0.0139	$L1\rightarrow L1(ILCT)$	
$H-1\rightarrow L+4$	278.91	4.73	0.0248	$L1\rightarrow L1(ILCT)$	276.0
H-6→L	282.81	4.54	0.0073	$L1\rightarrow L1(ILCT)$	
$H-6\rightarrow L+2$	275.41	5.37	0.0050	$L1\rightarrow L1(ILCT)$	
H-11→L	235.62	5.08	0.0216	$Rh\rightarrow L1(MLCT)$	233.3
H-5→L+4	233.62	5.74	0.0480	Cl→L1(LLCT)	

H-6→L+3	232.46	5.46	0.0105	L1→L1(ILCT)	
			Complex (2)	, ,	
H→L	462.76	2.98	0.0644	Ir→L1(MLCT)	
$H\rightarrow L+3$	358.38	4.64	0.0191	$Ir \rightarrow Cp*(MLCT)$	347.0
H-4→L	340.55	4.01	0.0075	$L1 \rightarrow L1(ILCT)$	
$H-5\rightarrow L+1$	273.73	5.11	0.0470	$Cp*\rightarrow L1(LLCT)$	266.0
$H-2\rightarrow L+4$	266.79	5.33	0.1540	$L1 \rightarrow Ir(LMCT)$	
			Complex (3)		
H→L	532.04	3.63	0.0087	L2→Rh(LMCT)	
$H-2\rightarrow L+1$	348.95	4.11	0.0369	L2→L2(ILCT)	344.10
$H\rightarrow L+2$	345.72	3.84	0.0128	L2→Rh(LMCT)	
$H-1\rightarrow L+2$	344.68	4.10	0.0089	Cl+L2→Rh(LMCT)	
H-3→L	336.07	4.24	0.0047	Cl→Rh(LMCT)	
$H\rightarrow L+4$	289.42	4.85	0.0063	L2→L2(ILCT)	286.1
H-6→L	285.32	5.14	0.0163	$Cl+L2 \rightarrow Rh(LMCT)$	
$H-5\rightarrow L+1$	282.59	4.98	0.0391	L2→L2(ILCT)	
$H-4\rightarrow L+4$	237.16	5.74	0.0422	L2→L2(ILCT)	234.30
$H-5\rightarrow L+3$	234.58	5.74	0.0161	L2→L2(ILCT)	
-			Complex (4)		
H→L	444.09	3.46	0.0355	L2→L2(ILCT)	
H-2→L	362.56	3.91	0.0077	L2→L2(ILCT)	346.1
$H-1\rightarrow L+1$	332.29	3.80	0.0076	Rh+L2→L2(MLCT/ILCT)	
$H\rightarrow L+3$	329.46	4.48	0.0468	L2→L2(ILCT)	
H-4→L	324.35	4.53	0.0265	$Cl+Cp*\rightarrow L2(LLCT)$	
$H-4\rightarrow L+2$	294.35	5.47	0.0164	$Cl+Cp*\rightarrow L2(LLCT)$	292.21
$H-1\rightarrow L+2$	288.71	4.74	0.0048	Rh+L2→Rh+L2	
$H-8\rightarrow L+3$	212.19	6.47	0.0387	L2→L2(ILCT)	210.9
$H-6\rightarrow L+4$	210.71	6.56	0.0399	Cl→L2(LLCT)	
$H-2\rightarrow L+5$	210.22	6.26	0.0669	L2→L2(ILCT)	
-		. \ /	Complex (5)	, ,	
H→L	519.01	3.61	0.0076	L3→Rh(LMCT)	
$H-2\rightarrow L+2$	338.46	4.30	0.0120	Cl→L3(LLCT)	332.0
$H-4\rightarrow L+1$	326.16	4.56	0.0052	$L3\rightarrow Rh(LMCT)$	
$H-1\rightarrow L+4$	271.36	5.01	0.265	L3→L3(ILCT)	268.0
H-2→L+4	267.30	5.21	0.0177	Cl→L3(LLCT)	
H-5→L+4	232.84	5.83	0.0498	L3→L3(LLCT)	229.0
H-10→L+2	229.44	6.14	0.0127	$Rh+L3\rightarrow L3(MLCT/ILCT)$	
			Complex (6)		
H→L	441.76	3.60	0.0160	L3→L3(ILCT)	
H-2→L+1	334.33	4.45	0.0109	$Ir \rightarrow L3(MLCT)$	330.0
$H-1\rightarrow L+1$	328.24	4.33	0.1160	L3→L3(ILCT)	
H-7→L	268.58	5.42	0.0159	$Ir \rightarrow L3(MLCT)$	256.0
H-6→L	264.64	5.25	0.0304	Cl→L3(LLCT)	
H-1→L+4	260.56	5.17	0.0350	L3→L3(ILCT)	
			Complex (7)		
H→L	518.07	3.68	0.0071	L4→Rh(LMCT)	
H-1→L	448.90	4.05	0.0132	Cl→Rh(LMCT)	400.0
$H-1\rightarrow L+1$	397.25	4.09	0.0148	$Cl \rightarrow Rh(LMCT)$	
$H-1\rightarrow L+4$	269.94	5.23	0.0297	Cl→L4(LLCT)	265.0
$H-2\rightarrow L+3$	262.52	5.20	0.0465	Cl→L4(LLCT)	
-				. ,	

H→L+5	231.04	5.66	0.0547	L4→L4(ILCT)	229.0
			Complex (8)		
H→L	439.65	3.59	0.0196	Ir→L4(MLCT)	
$H-1\rightarrow L+2$	371.55	4.79	0.0381	$Ir \rightarrow L4(MLCT)$	361.0
H-1→L	358.22	4.0	0.0349	$Ir \rightarrow L4(MLCT)$	
$H-2\rightarrow L+3$	257.57	5.29	0.0359	Cl→Cp*(LLCT)	256.0
H-4→L+3	255.65	5.61	0.0142	$L4\rightarrow Cp*(LLCT)$	