

Comparative Analysis of Mononuclear 1:1 and 2:1 Tetravalent Actinide (U, Th, Np) Complexes: Crystal Structure, Spectroscopy, and Electrochemistry

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1 Comparative Analysis of Mononuclear 1:1 and 2:1
2 Tetravalent Actinide (U, Th, Np) Complexes: Crystal
3 Structure, Spectroscopy, and Electrochemistry

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7 **ABSTRACT.** Six mononuclear tetravalent actinide complexes (**1-6**) have been synthesized using
8 a new Schiff base ligand 2-methoxy-6-(((2-methyl-1-(pyridin-2-yl)propyl)imino)methyl)phenol
9 (**HL^{Pr}**). The **HL^{Pr}** is treated with tetravalent actinide elements in varied stoichiometry to afford
10 mononuclear 1:1 complexes [$MCl_3-L^{Pr} \cdot nTHF$] (**1-3**) and 2:1 complexes [$MCl_2-L^{Pr}_2$] (**4-6**) ($M =$
11 Th^{4+} (**1** and **4**), U^{4+} (**2** and **5**) and Np^{4+} (**3** and **6**)). All complexes are characterized using different
12 analytical techniques such as IR, NMR, and absorption spectroscopy as well as crystallography.
13 UV-vis spectroscopy revealed more red-shifted absorption spectra for 2:1 complexes as compared
14 to 1:1 complexes. 1H NMR of Th(IV) complexes exhibit diamagnetic spectra whereas U(IV) and
15 Np(IV) complexes revealed paramagnetically shifted 1H NMR. Interestingly, NMR signals are
16 paramagnetically shifted between -70 to 40 ppm in **2** and **3**, but are confined within -35 to 25 ppm
17 in 2:1 complexes **5** and **6**. Single crystal structures for 1:1 complexes revealed an eight-coordinated
18 Th(IV) complex (**1**) and seven-coordinated U(IV) (**2**) and Np(IV) (**3**) complexes. Whereas, all 2:1
19 complexes **4-6** were isolated as eight-coordinated isostructural molecules. The geometry around
20 the Th^{4+} center in **1** is found to be trigonal dodecahedral and, capped trigonal prismatic around

21 U(IV) and Np(IV) centers in **2** and **3**, respectively. Whereas, An⁴⁺ centers in 2:1 complexes are
22 present in dodecahedral geometry. Importantly, 2:1 complexes exhibit increased bond distances in
23 comparison to their 1:1 counterparts as well as interesting bond modulation w.r.t. ionic radii of
24 An(IV) centers. Cyclic voltammetry displays an increased oxidation potential of the ligand by 300
25 to 500 mV, after coordination with An⁴⁺. CV studies indicates Th(IV)/Th(II) reduction beyond
26 -2.3 V whereas attempts were made to identify redox potentials for U(IV) and Np(IV) centers.
27 Spectroscopic binding studies reveal that complex stability in 1:1 stoichiometry follows the order
28 Th⁴⁺ ≈ U⁴⁺ > Np⁴⁺.

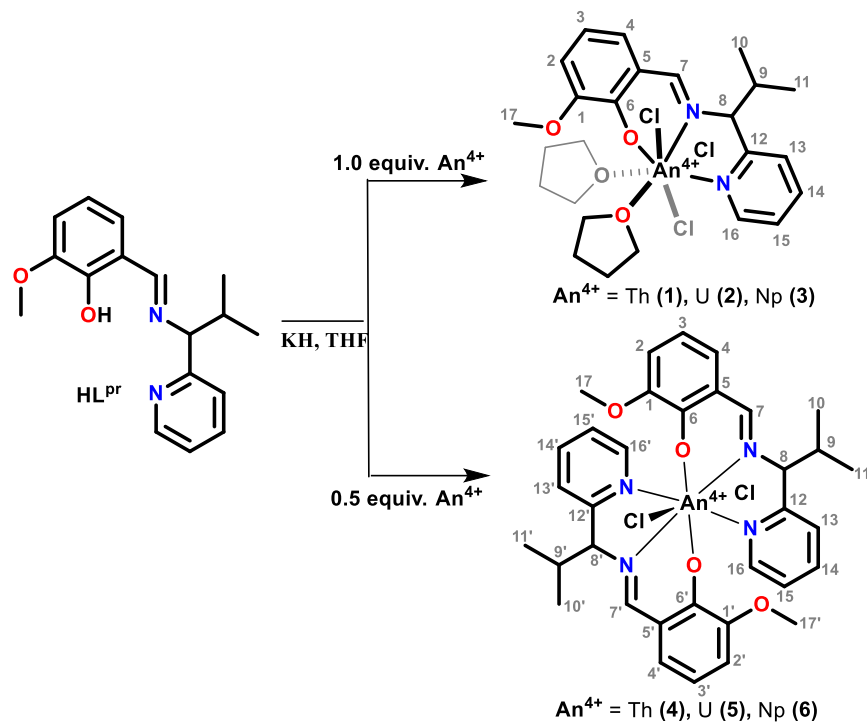
29 **Introduction**

30 In recent years, coordination chemistry of actinide elements has gained widespread attention due
31 to their underexplored coordination properties as well as interesting redox and catalytic properties.
32 [1-14] Importantly, studies on actinide chemistry mostly focus on uranyl U(VI)O₂²⁺, with
33 significantly fewer studies for any other actinide or oxidation state.[15-21] Nowadays, efforts are
34 underway to synthesize and isolate highly reactive low valent actinide compounds, after the
35 realization of their potential in small molecule activation and catalysis.[4,22-25] Meanwhile, the
36 low valent actinide complexes are dominated by cyclopentadienyl and related ligands,
37 emphasizing a demand to design and synthesize other organic ligand(s) to stabilize low valent
38 actinide complexes.[6,15,26,27] In this context, di- and mono-anionic Schiff base ligands are
39 frequently explored due to the presence of a strongly coordinating aryl-oxide donor along with
40 chelating motif. While multiple U(VI)O₂²⁺, U(IV), and Th(IV), and fewer Np(IV) complexes have
41 been reported with di-anionic Schiff base ligands, e.g. of salen-type [5,21,28-33], Uranium
42 complexes with Schiff base ligands have been shown to promote interesting electrochemical

43 processes owing to the non-innocent redox behavior of the ligand(s).[3-5] Therefore, there is a
44 need for the in-depth characterization of structure, bond properties, and redox behavior of actinide
45 complexes with non-redox innocent ligands to achieve actinide complexes of desired properties.

46 Concerning the relatively well-explored dianionic Schiff base ligands, there are no reports on the
47 isolation of mononuclear 1:1 actinide complex(es) with monoanionic Schiff base ligands. Notably,
48 monoanionic, tridentate Schiff base ligands are promising candidates to produce 1:1 actinide
49 complex with larger numbers of labile sites and thus potentially more versatile reactivity.[34,35]
50 Indeed, synthesis and isolation of 1:1 Schiff base-actinide complexes requires careful control of
51 the synthesis conditions due to the potential formation of dimerized products or other unintended
52 follow-up reactions.[36] Therefore, mono-anionic Schiff base ligands are largely reported as stable
53 bis-actinide complexes.[33,36] On the other hand, availability of such 1:1 actinide complex will
54 be extremely informative for comparative analysis of coordination, structural and electronic
55 variations between mono- (1:1) and bis-ligated (2:1) An complexes.

56 Taking this into consideration we have synthesized a new non-redox-innocent mono-ionic Schiff
57 base ligand **HL^{Pr}** and its mono-ligated (**1-3**) and bis-ligated (**4-6**) actinide complexes $[\text{AnCl}_x\text{-}$
58 $(\text{L}^{\text{Pr}})_y \cdot n\text{THF}]$ (An = Th (**1** and **4**), U (**2** and **5**) and Np (**3** and **6**); $x = 3$, $y = n = 1$ for **1-3**, except n
59 $= 2$ for **1**; $x = y = 2$ and $n = 0$ for **4-6** complexes) (Scheme 1). All complexes were comprehensively
60 characterized by FTIR, UV-vis, NMR, SC-XRD, and cyclic voltammetry to understand the
61 influence of structure and electronic properties on bonding, coordination, and redox behavior.



62

63 **Scheme 1.** Synthetic route for the preparation of actinide complexes **1-6** with **HL^{Pr}**.

64 **Experimental**

65 **Caution!**

66 Th-232sec, U-nat and Np-237+ consist of long lived α emitters with half-lives of 1.41×10^{10} , 4.47
 67 $\times 10^9$ and 2.14×10^6 years, respectively. Special precautions as well as appropriate equipment and
 68 facilities for radiation protection are required for handling these materials. All experiments were
 69 carried out in a controlled laboratory at the Institute of Resource Ecology, Helmholtz-Zentrum
 70 Dresden-Rossendorf.

71 **General remarks**

72 All preparations were performed under the rigorous exclusion of moisture and oxygen in nitrogen
 73 filled glove boxes or using Schlenk techniques. The used solvents were dried using solvent

74 purification system MBraun SPS 5 and stored over molecular sieve (3 Å) prior to use. Chemicals
75 were sourced from Sigma-Aldrich and were used as received. Thin layer chromatography (TLC)
76 was carried out on aluminum plates coated with silica gel mixed with fluorescent indicator sourced
77 from Merck, Germany. NMR spectra were recorded inside a controlled laboratory on a Varian
78 Inova 400 spectrometer with an ATB indirect probe equipped with z-gradients operating at a ^1H
79 frequency of 399.89 MHz and a ^{13}C frequency of 100.56 MHz. Special precautions were taken to
80 avoid contamination. All spectra were recorded with a Varian AutoX ID probe head with z
81 gradient. Deuterated solvents were purchased at Deutero GmbH and dried over potassium mirror
82 prior to use. FTIR spectra were measured on an Agilent Cary 630 FT-IR spectrometer equipped
83 with a single-reflection attenuated total reflection (ATR) accessory made of diamond. The
84 measurements were performed in an inert glove box filled with N_2 . The spectra were recorded
85 between 4000 and 650 cm^{-1} with a resolution of 2 cm^{-1} . UV/visible/NIR spectra were recorded
86 with a J&M Analytik AG TIDAS 100 spectrometer connected with optical fibers to a cuvette
87 housing in the glove box. The spectra were recorded between 200 and 1025 nm in 1 cm quartz
88 cuvettes.

89 **Cyclic and Differential Pulse Voltammetry (CV/DPV):** CV and DPV were carried out using a
90 computer controlled potentiostat (CHI 650C), and a standard three electrode arrangement that
91 consisted of glassy carbon, platinum, and saturated calomel (SCE) containing Ag/AgCl in AgNO_3
92 acetonitrile solution as working, auxiliary, and reference electrodes, respectively.[37,38] All
93 electrochemical measurements were carried out in dry and Ar-purged acetonitrile with $n\text{-Bu}_4\text{NPF}_6$
94 as the supporting electrolyte. All the potentials are calculated in reference to $\text{Fc}^{0/+}$ which was
95 measured separately.

96 **X-ray Crystallography:** The data for **1-6** were analyzed on a Bruker D8 Venture single-crystal
97 X-ray diffractometer with micro-focused Mo K α radiation ($\lambda = 0.71073 \text{ \AA}$) and a PHOTON 100
98 CMOS detector. The frames for **1, 2,** and **4-6** were collected at 100 K, and for **3** at 293 K. Data
99 treatment was performed with the Bruker APEX 3 program suite including the Bruker SAINT
100 software package for integration [39], and empirical absorption corrections was applied by using
101 the spherical harmonic incorporated in the SCALE3 ABSPACK scaling algorithm.[40] The
102 structures were solved and refined with full-matrix least-squares data on F^2 using the Bruker
103 SHELXTL [41] software package and SHELXL-2018 [42] in the WinGX module. [43] All
104 hydrogen atoms were fixed at the calculated positions and refined isotropically. Complex **2**
105 exhibits a Flack parameter of 0.380 suggesting potential racemization. [45] For complexes **4** and
106 **6,** some electron density, potentially corresponding to disordered THF or toluene molecules, could
107 not be resolved and was therefore masked using the solvent masking ‘Squeeze’ command in
108 PLATON. [44] In complex **5,** highly disordered toluene molecules are modelled using OLEX 2
109 software. Details of the crystallographic data collection and structural solution parameters are
110 provided in Table S1.

111 **Synthesis**

112 *2-methoxy-6-(((2-methyl-1-(pyridin-2-yl)propyl)imino)methyl) phenol (HL^{Pr})*. In a 25 ml round
113 bottom flask, 2-methyl-1-(pyridin-2-yl)propan-1-amine dihydrochloride (0.17 g, 0.78 mmol) was
114 dissolved in deionized water (5ml) and neutralized by solid Na₂CO₃ (0.2 g, 1.95 mmol). The
115 solution was stirred for 20 minutes followed by the dropwise addition of methanolic solution of o-
116 vanillin (0.1 g, 0.65 mmol) resulting in a yellow color solution. The reaction was further stirred
117 for another 30 minutes resulting in the formation of the yellow oily product. The reaction was
118 stopped and dichloromethane (25 ml) was added to the reaction mixture dissolving the oily

119 product. The yellow colored organic layer was isolated using a separating funnel. The organic
120 portion was washed multiple times with water followed by removal of DCM under reduced
121 pressure to afford an oily product. Yield = 96% (179 mg). ¹H NMR (400 MHz, CDCl₃): δ = 14.13
122 (d, 1H, -(H)C=N), 8.39 (s, 1H, -CH_{py}), 7.67 (m, 1H, -CH_{py}), 7.41 (d, 1H, -CH_{phenoxy}), 7.18 (m, 1H,
123 -CH_{py}), 6.91-6.88 (m, 2H, -CH_{py} & -CH_{phenoxy}), 6.78 (t, 1H, -CH_{phenoxy}), 4.32 (d, 1H, -
124 (H)CCH(CH₃)₂), 3.87 (s, 3H, -OCH₃), 2.39 (m, 1H, -HC(CH₃)₂), 0.87 (d, 6H, -CH₃). ¹³C NMR
125 (125 MHz, CDCl₃): δ = 165.45, 161.07, 151.71, 148.71, 148.38, 137.09, 124.46, 123.05, 122.34,
126 121.88, 118.39, 113.99, 80.80, 55.99, 34.30, 19.72, 17.67. FTIR spectrum (ATR, selected peaks,
127 cm⁻¹): 1626 (C=N), 1586 (C=N_{py}). Absorption spectrum [λ_{\max} , nm, THF]: 334, 440.

128 [*ThCl₃-L^{Pr}.2THF*] (**1**). To a scintillation vial, charged with 1 ml solution of **HL^{Pr}** (0.02g, 0.070
129 mmol) in dry THF, was added with excess of KH, leading to the immediate evolution of molecular
130 hydrogen gas. The reaction mixture was stirred for 15 min and the clear supernatant liquid was
131 separated from unreacted solid KH by centrifugation. To a clear supernatant solution,
132 ThCl₄·2DME (0.038g, 0.070 mmol) solution in dry THF (1ml) was added dropwise under constant
133 stirring resulting in formation of yellow solution. The reaction mixture was stirred for another 1h
134 and was centrifuged to remove salt impurity. The resulting yellow color solution was left for
135 evaporation to afford deposition of crystalline material at the bottom of vial. Yield = 86% (41 mg).
136 Anal. calc. for C₂₅H₃₅Cl₃N₂O₄Th: C, 37.42; H, 4.57; N, 3.45. Found: C, 39.20; H, 4.61; N, 3.66.
137 ¹H NMR (400 MHz, THF-*d*₈): δ = 9.95 (d, 1H), 8.44 (s, 1H), 7.89 (t, 1H), 7.46 (d, 1H), 7.40 (t,
138 1H), 7.08 (m, 1H), 7.00 (d, 1H), 6.74 (t, 1H), 4.43 (m, 1H), 3.78 (s, 3H), 3.32(m, 1H), 0.92-0.85
139 (dd, 6H). ¹³C NMR (125 MHz, THF-*d*₈): δ = 168.48, 162.24, 153.21, 150.41, 138.91, 127.29,
140 124.80, 123.47, 118.51, 116.53, 88.86, 55.67, 34.56. FTIR spectrum (ATR, selected peaks, cm⁻¹):
141 1612 (C=N), 1559 (C=N_{py}). Absorption spectrum [λ_{\max} , nm, THF]: 305, 370, 430.

142 $[UCl_3-L^{Pr}\cdot THF]$ (**2**). The synthesis of **2** was performed in the similar manner to **1** except using
143 UCl_4 (0.026 g, 0.070 mmol) as the uranium precursor. After the dropwise addition of UCl_4
144 solution, the reaction was stirred for 30 min followed by centrifugation to remove insoluble salt
145 impurities. The resulting green colour solution was diffused with toluene to afford green crystal
146 after 72 h. Yield = 83% (41 mg). Anal. calc. for $C_{21}H_{27}Cl_3N_2O_3U\cdot 2THF$: C, 38.90; H, 4.57; N,
147 3.63. Found: C, 37.0; H, 3.89; N, 3.59. 1H NMR (400 MHz, $THF-d_8$): δ = 38.45 (s, 1H), 36.48 (s,
148 1H), 35.90 (s, 1H), 31.87 (s, 1H), 23.87 (s, 1H), 20.11 (s, 3H), 3.19 (s, 1H), 0.94 (s, 1H), -0.27 (s,
149 1H), -15.43 (s, 3H), -15.57 (s, 1H), -24.69 (s, 3H), -29.23 (s, 1H), -71.67 (s, 1H). FTIR spectrum
150 (ATR, selected peaks, cm^{-1}): 1612 (C=N), 1563 (C=N_{py}). Absorption spectrum [λ_{max} , nm, THF]:
151 309, 370, 470.

152 $[NpCl_3-L^{Pr}\cdot THF]$ (**3**). The synthesis of **3** was performed in the similar manner to **1** using
153 $NpCl_4\cdot 2DME$ (0.029g, 0.070 mmol) as the neptunium precursor. The dropwise addition of Np^{4+}
154 solution resulted in the formation of brown color solution. Crystal suitable for X-ray diffraction
155 were obtained by diffusing pentane to the complex solution in THF to afford dark brown crystals
156 within 24h. Yield = 85% (42 mg). 1H NMR (400 MHz, $THF-d_8$): δ = 41.37 (s, 1H), 37.48 (s, 1H),
157 35.12 (s, 1H), 27.19 (s, 1H), 21.44 (s, 1H), 16.23 (s, 3H), 5.02 (s, 1H), 0.31 (s, 1H), -15.51 (s, 4H),
158 -20.14 (s, 3H), -30.54 (s, 1H), -65.23 (s, 1H). FTIR spectrum (ATR, selected peaks, cm^{-1}):
159 1613(C=N), 1559 (C=N_{py}). Absorption spectrum [λ_{max} , nm, THF]: 373, 474.

160 $[ThCl_2-(L^{Pr})_2]$ (**4**). The synthesis of **4** was performed in the similar manner to **1** using 0.5 equiv.
161 $ThCl_4\cdot 2DME$ (0.015g, 0.035 mmol) as the Thorium precursor. The dropwise addition of Th^{4+}
162 solution resulted in the formation of colorless solution. Crystal suitable for X-ray diffraction were
163 obtained by diffusing pentane to the complex solution in THF to afford colorless crystals within
164 24h. Yield = 83% (51 mg). Anal. calc. for $C_{34}H_{38}Cl_2N_4O_4Th$: C, 46.96; H, 4.40; N, 6.44. Found:

165 C, 46.25; H, 4.18; N, 6.72. ^1H NMR (400 MHz, THF- d_8): δ = 9.66 (s, 2H), 8.49 (s, 2H), 7.98 (s,
166 2H), 7.60 (s, 2H), 7.41 (s, 2H), 7.0 (s, 4H), 6.71 (s, 2H), 4.50 (s, 2H), 3.75 (s, 2H), 3.39 (s, 6H),
167 0.88 (s, 12H). FTIR spectrum (ATR, selected peaks, cm^{-1}): 1610(C=N_{py}), 1551 (C=N). Absorption
168 spectrum [λ_{max} , nm, THF]: 442.

169 $[\text{UCl}_2\text{-}(\mathbf{L}^{\text{Pr}})_2]$ (**5**). The synthesis of **5** was performed in the similar manner to **2** except using 0.5
170 equiv. UCl_4 (0.013 g, 0.035 mmol) as the uranium precursor. The dropwise addition of U^{4+}
171 solution resulted in the formation of brown solution. The solvent was removed under reduced
172 pressure and the resulting solid compound was washed with pentane to afford brown precipitates.
173 The precipitates were redissolved in toluene followed by diffusion with pentane to afford brown
174 crystals after 48 h. Yield = 80% (49 mg). Anal. calc. for $\text{C}_{34}\text{H}_{38}\text{Cl}_2\text{N}_4\text{O}_4\text{U}$: C, 46.64; H, 4.37; N,
175 6.40. Found: C, 46.94; H, 4.82; N, 6.59. ^1H NMR (400 MHz, THF- d_8): δ = 23.93 (s, 1H), 21.40
176 (s, 1H), 15.95 (s, 1H), 15.26 (s, 1H), 14.66 (s, 1H), 13.96 (s, 1H), 12.89 (s, 1H), 12.78 (s, 1H),
177 12.28 (s, 1H), 11.68 (s, 3H), 10.03 (s, 3H), 2.24 (s, 3H), 1.02 (s, 3H), 0.38 (s, 1H), -0.61 (s, 1H), -
178 1.26 (s, 1H), -5.11 (s, 3H), -10.73 (s, 3H), -11.34 (s, 3H), -13.84 (s, 1H), -14.0 (s, 1H), -14.60 (s,
179 3H), -23.89 (s, 1H), -26.80 (s, 1H), -29.93 (s, 1H), -33.40 (s, 1H). FTIR spectrum (ATR, selected
180 peaks, cm^{-1}): 1613 (C=N_{py}), 1560 (C=N). Absorption spectrum [λ_{max} , nm, THF]: 442, 523.

181 $[\text{NpCl}_2\text{-}(\mathbf{L}^{\text{Pr}})_2]$ (**6**). The synthesis of **6** was performed in the similar manner to **5** using 0.5 equiv.
182 $\text{NpCl}_4\cdot 2\text{DME}$ (0.015 g, 0.035 mmol) as the neptunium precursor to afford wine-red colored
183 complex solution. Crystals suitable for X-ray diffraction were obtained in a similar manner like **5**.
184 Yield = 84% (45 mg). ^1H NMR (400 MHz, THF- d_8): δ = 22.37 (s, 1H), 18.32 (s, 1H), 16.46 (s,
185 1H), 15.25 (s, 1H), 14.79 (s, 1H), 13.04 (s, 3H), 11.52 (s, 3H), 10.99 (s, 1H), 10.24 (s, 1H), 9.21
186 (s, 1H), 8.49 (s, 1H), 1.86 (s, 3H), 1.28 (s, 1H), 0.85 (s, 1H), 0.51 (s, 1H), -0.46 (s, 1H), -6.26 (s,
187 3H), -9.85 (s, 3H), -10.54 (s, 1H), -11.65 (s, 1H), -12.26 (s, 3H), -13.31 (s, 3H), -21.21 (s, 1H), -

188 22.86 (s, 1H), -26.39 (s, 1H), -28.77 (s, 1H). FTIR spectrum (ATR, selected peaks, cm^{-1}):
189 1613($\text{C}=\text{N}_{\text{py}}$), 1551 ($\text{C}=\text{N}$). Absorption spectrum [λ_{max} , nm, THF]: 452, 530.

190 **Results and discussion**

191 Condensation of o-vanillin with 2-methyl-1-(pyridine-2-yl)propane-1-amine dihydrochloride
192 results in the isolation of a yellow oily product **HL^{Pr}** in high yield. The formation of **HL^{Pr}** was
193 confirmed by the presence of the characteristic (H)C=N proton signal at 8.51 ppm in ^1H NMR
194 (Figure S1, ESI). Complexes **1-6** were synthesized by treating (**L^{Pr}**)⁻, deprotonated with KH, with
195 An^{4+} ($\text{An} = \text{Th}, \text{U}, \text{Np}$) to afford mononuclear 1:1 (**1-3**) and 2:1 (**4-6**) complexes. Interestingly,
196 addition of Th^{4+} or U^{4+} in different stoichiometry leads to the observation of differently colored
197 solutions during the preparation of 1:1 and 2:1 complexes (see experimental section for detail).
198 FT-IR measurements exhibit superimposable spectra of all the complexes suggesting the
199 coordination of metal ions with the ligand in a similar fashion (Figure S2). All complexes exhibit
200 (H)C=N $\nu_{\text{C}=\text{N}}$ and pyridyl $\nu_{\text{C}=\text{N}}$ stretching modes as expected, bathochromically shifted by 14 cm^{-1}
201 and $23\text{-}31\text{ cm}^{-1}$ in relation to **HL^{Pr}**, respectively, indicating a complexation involving (H)C=N and
202 pyridyl nitrogen atoms (Table 1). M-Cl bond vibrations appear below 650 cm^{-1} and thus could not
203 be identified. [46] The absorption spectrum of the ligand **HL^{Pr}** exhibits an absorption maximum
204 at approx. 334 nm (λ_{max}) along with a weak band at 440 nm (Figure S3). This ligand-based
205 absorption feature at 334 nm (in **HL^{Pr}**) is red shifted by 30-40 nm in 1:1 complexes **1-3**, whereas
206 it is red shifted by 110-120 nm for the 2:1 complexes **4-6**. Moreover, weak absorption bands
207 between 580 and 1025 nm, indicating $f-f$ transitions, are observed for complexes **2, 3, 5, and 6**
208 (Figure S3). [45]

209

Table 1. Absorbance and selected IR stretches in **HL^{Pr}** and complexes **1-6**.

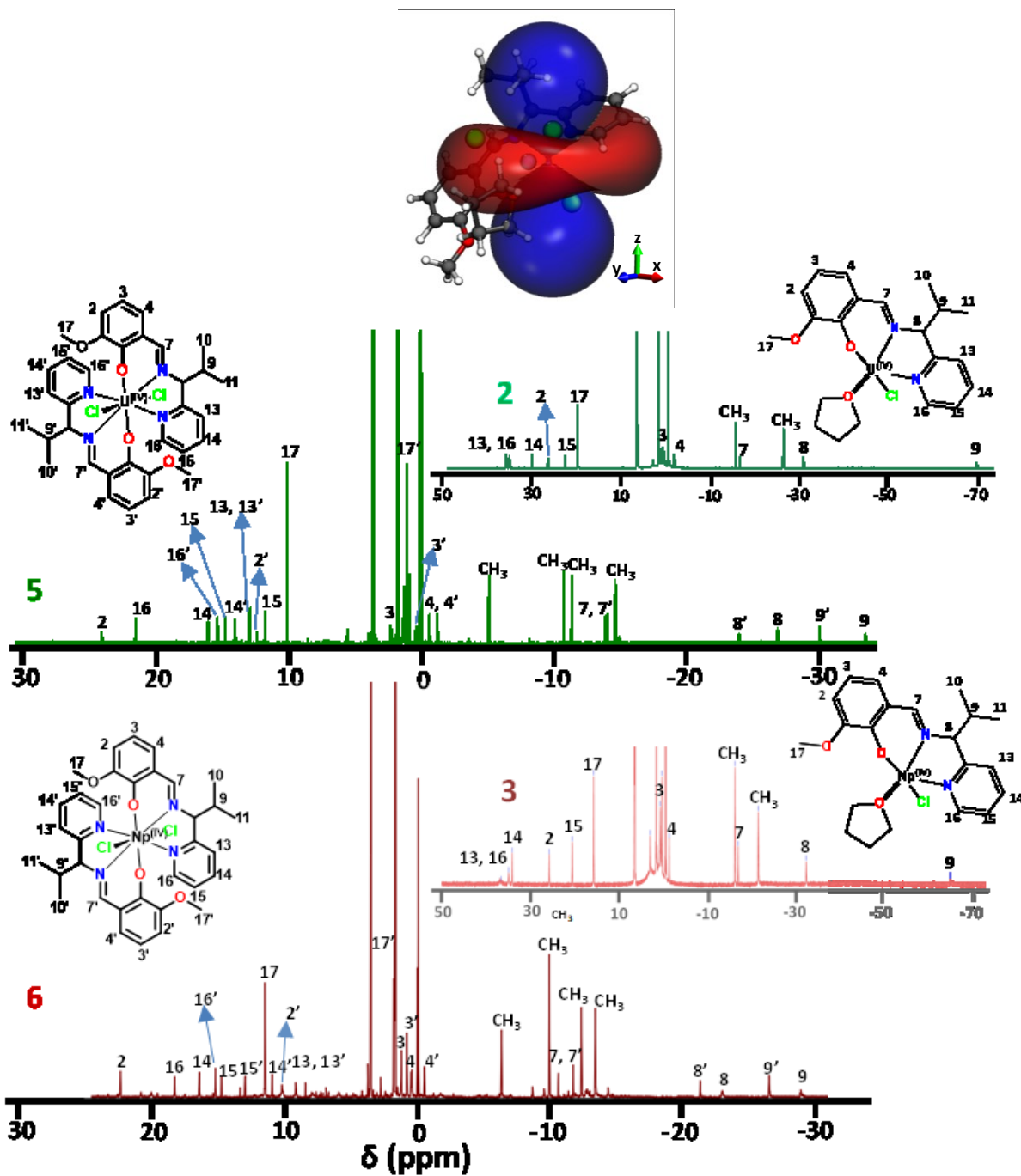
	HL^{Pr}	1	2	3	4	5	6
Absorbance (nm)							
	334, 440	366,	370, 473	370, 490	442	442	452
		436,				523	530
FTIR vibrations (cm ⁻¹)							
$\nu_{C=N}$ (HC=N)	1586	1563	1563	1559	1551	1560	1551
$\nu_{C=N}$ (pyridyl)	1626	1612	1612	1613	1610	1613	1613

210 NMR Spectroscopy

211 To analyze the molecular structure in solution, NMR spectroscopy was used for all complexes
212 (Figures S4-S12). Complexes **1** and **4** exhibit diamagnetically shifted ¹H NMR spectra, whereas
213 highly shifted paramagnetic ¹H NMR spectra were observed for complexes **2**, **3**, **5**, and **6** (Table
214 2). The proton signals in **1** and **4** were observed downfield shifted as compared to free ligand
215 (Figures S4 and S8). Interestingly, the change of coordination environment around the An center
216 from 1:1 to 2:1 complex exhibits small but distinct shifts of these diamagnetic ¹H NMR signals.
217 For example, the (H7)C=N proton signal in 2:1 complex **4** is comparatively more upfield shifted
218 by 0.30 ppm than in 1:1 complex **1**. Whereas, proton signals for (H8)C and (H9)C in **4** are
219 downfield shifted by 0.07 ppm and 0.33 ppm respectively, as compared to their 1:1 counterparts.
220 Apart from that, pyridyl protons appear between 6.71 ppm to 7.97 ppm for both complexes, **1** and
221 **4**. We believe that the presence of the additional ligand results in a reduced interaction of Th(IV)
222 with both ligands leading to the upfield shift in the proton signals in **4**. On the other hand, we
223 observe the presence of additional NMR signals corresponding to every ¹H signal in both **1** and **4**.
224 Since we worked with the ligand having both ‘R’ and ‘S’ configuration at C8 center, we tentatively

225 assign these signals as belonging to the complex moiety with the ligand in the opposite
226 configuration. Based on integration of both types of NMR signals, we can estimate an enantiomeric
227 excess for one of the conformations by ~50% in our starting material.

228 Due to the presence of unpaired $5f$ electrons in U(IV) and Np(IV), complexes **2**, **3**, **5**, and **6** exhibit
229 paramagnetically shifted ^1H NMR signals (Figures S5, S9 and S10). Notably, unpaired electron
230 density mainly interacts with the NMR observed nuclear spins either due to spin dipolar
231 interactions in form of pseudo contact shifts (PCS) or by Fermi contact interactions (FCS) which
232 may arise from molecular orbitals (MO) containing unpaired electron density originating from
233 metal contributions to the MO and featuring significant levels of s -contribution at the observed
234 nucleus. Typically, only sizeable PCS contributions are detected on nuclei remote from the metal
235 center. From previous studies, a prolate density of unpaired electrons is expected at the An^{4+}
236 centers [28-30, 45, 47-49]. This results in a PCS field that leads to shielding PCS shifts in the
237 direction of the z -axis of the magnetic tensor and to de-shielding PCS shifts in the x,y -plane. Thus
238 nuclei with resonances found in the negative scale are expected to be located in z -direction relative
239 to the metal center and those found shifted to positive ppm values are expected close to the x,y -
240 plane. Apart from that, the change in coordination environment from 1:1 to 2:1 around An^{4+} center
241 is also very prominent in the paramagnetically shifted NMR. [50] Interestingly, Np(IV) complexes
242 **3** and **6** exhibit similar NMR patterns as their U(IV) counterparts **2** and **5**, respectively. Moreover,
243 the 1:1 Np(IV) complex **3** is paramagnetically more shifted as compared to its U(IV) counterpart
244 **2** whereas, in contrast, 2:1 U(IV) complex **5** is comparatively more shifted than its Np(IV)
245 counterpart **6** (see Table 2). The ^1H NMR signals for 1:1 complexes **2** and **3** appear in the range
246 between -70 to 40 ppm, whereas these signals are confined within -35 to 25 ppm for 2:1
247 complexes **5** and **6** (Figure 1).



248

249 **Figure 1.** Comparative ^1H NMR spectra for 1:1 complexes 2 and 3 and 2:1 complexes 5 and 6. A

250 schematic representation of possible PCS cones is shown at the top for the 1:1 complex. The

251 corresponding figure for the 2:1 complexes is discussed in the SI.

252 Notably, while complexes **2** and **3** exhibit six signals on the negative side and seven signals on the
253 positive side of the spectra, complexes **5** and **6** display twice the number of signals on the same
254 side of spectrum, due to the presence of two enantiomers of the ligand. More importantly, proton
255 signals (H and H') for both ligands in the 2:1 complexes exhibit distinct positions suggesting
256 different electronic interactions of each ligand with the An^{4+} center.[50] The highly shifted
257 isopropyl 'H9' proton at -69.74 ppm is found for the 1:1 U(IV) complex **2** whereas, in 2:1 U(IV)
258 complex **5**, H9 and H9' protons appear separately at -33.40 ppm and -29.93 ppm, respectively.
259 These values are shifted by approximately 35 to 40 ppm when changing from 1:1 to 2:1 complexes.
260 The paramagnetic shift is considerably reduced to 5-10 ppm for H8 (-26.78 ppm) and H8'
261 (-23.89 ppm) protons in **5** as compared to its position in **2** (-30.59 ppm). The resonances of the
262 whole isopropyl group appear in the strongly shielded region of the NMR spectrum and thus should
263 be located in the z-axis region of the PCS field. [51] In contrast, the pyridyl proton 'H13' in **2** is
264 observed as most downfield shifted signal at 36.45 ppm, while a phenoxide 'H2' (23.90 ppm) from
265 one of the ligands, was found to be the most de-shielded proton in **5**. Moreover, phenoxide protons
266 (H2, H3/H3', H4/H4') are shifted to the similar extent in both the complexes **2** and **5**, except H2'
267 (12.27 ppm) in **5**, which appeared highly shielded as compared to H2 (27.16 ppm) in complex **2**.
268 This indicates that these nuclei and thus the respective aromatic rings are located in the x,y-plane
269 of the PCS field. Variations of the magnitude of shielding in the respective resonances may account
270 for slight differences in position in the PCS field or originate from FCS contributions due to the
271 differences in the respective interactions of the donors with the metal center. Both PCS and FCS
272 are strongly influenced by the coordination geometry around the metal center and thus depend on
273 the size of the metal center itself. In conclusion, we observe a quite symmetric complexation in
274 the 2:1 ligands that resembles the coordination environment in the 1:1 complexes, with the ligands

275 having coordinating heteroaromatic rings in the x,y-plane region and the isopropyl groups in the
 276 z-direction (magnetic principal axis) of the PCS field. Due to the observed one coordination
 277 environment in the 1:1 complexes and two sets of resonances in the 2:1 complexes, most likely
 278 due to the presences of both enantiomers of the ligand, we assume that in the time and ensemble
 279 average of the NMR spectra, we only observe a single (averaged) coordination environment. In
 280 case of the 2:1 complexes, the magnetic principal axis coincides with a (pseudo-) symmetry axis,
 281 most likely coinciding with the O-U-O bond. It is evident from the NMR spectra that the structure
 282 of the 2:1 complexes in solution differs from that found in the solid state (see below). Assigning
 283 magnetic axes or even “PCS cones” to the molecule is further complicated by the molecules’ low
 284 symmetry, which causes all three quadrupole moments to be unequal. Details regarding this
 285 assignment can be found in the SI.

Table 2. Selected ¹H NMR shift (ppm) for complexes **1-6**.

Proton	HL ^{pr}	1	2	3	4	5	6
2	6.86	6.99	27.16	27.26	7.0	23.90, 12.27	22.37, 10.24
3	6.74	6.74	1.10	2.05	6.71	2.245, 0.38	1.28, 0.85
4	7.49	7.46	-1.35	0.31	7.60	-0.63, -1.24	0.51, -0.46
7	8.52	9.95	-16.26	-16.45	9.66	-14.0, -13.86	-10.54, -11.65
8	4.32	4.43	-30.59	-31.90	4.50	-26.78, -23.89	-22.86, -21.20
9	2.39	3.32	-69.74	-64.66	3.75	-33.39, -29.93	-28.77, -26.39
10, 11	0.87	0.92-0.85	-15.29, -25.99	-15.72, -21.02	0.88	-14.60, -11.33, -10.72, -5.11	-13.31, -12.26, -9.85, -6.26
13	6.89	7.08	36.45	41.37	7.0	12.79, 12.91	9.21, 8.49
14	7.65	7.89	30.73	35.23	7.98	15.96, 13.96	10.99, 16.46

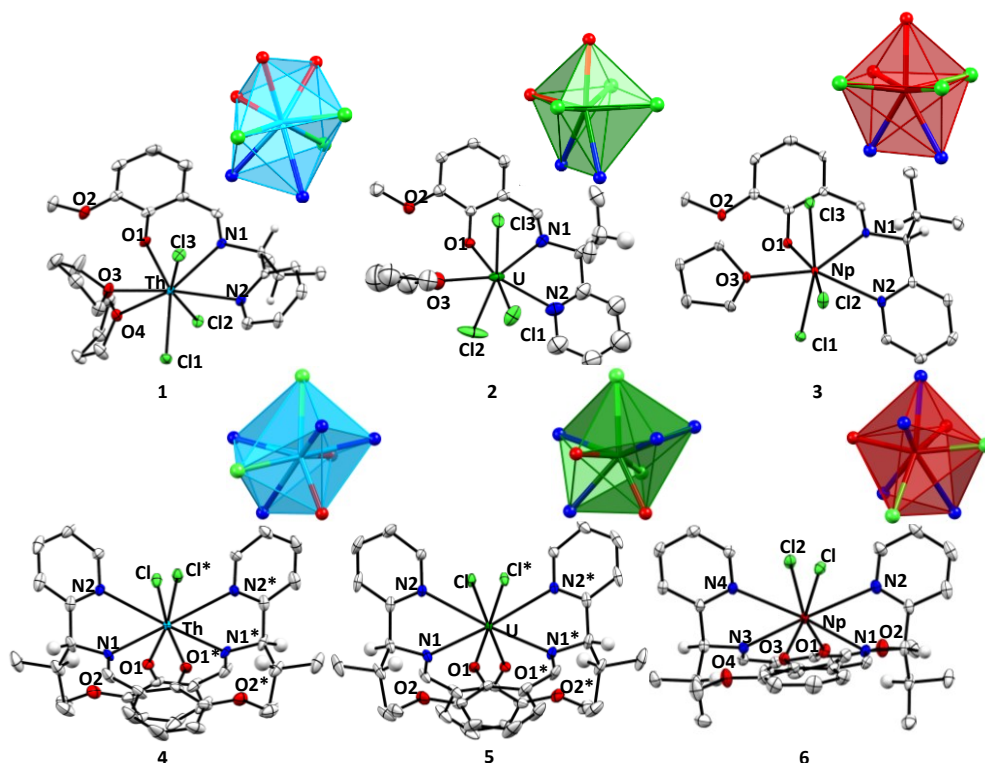
15	7.15	7.39	23.29	21.49	7.41	14.65, 11.67	14.79, 13.04	286
16	8.39	8.44	35.81	37.38	8.49	21.40, 15.27	18.32, 15.25	
17	3.85	3.78	20.40	16.27	3.39	10.03, 1.02	11.52, 1.86	

287 **Molecular structures of 1-6**

288 All complexes have been characterized crystallographically and exhibit mononuclear molecular
289 structures with 1:1 and 2:1 L:An stoichiometry for **1-3** and **4-6**, respectively. The isolated
290 complexes were found with the chemical compositions $[\text{AnCl}_3\text{L}^{\text{Pr}}\cdot(\text{THF})_n]$ for **1-3** and
291 $[\text{AnCl}_2\text{L}^{\text{Pr}_2}]$ for **4-6** [An = Th, U and Np; n = 1 or 2] (Figures 2 and 3, and Table 3). Complexes **1-**
292 **3** contain one ligand unit, three chlorine atoms and one or two THF molecule(s) within their
293 coordination sphere. Whereas 2:1 complexes **4-6** were isolated as isostructural eight-coordinated
294 tetravalent complexes containing two ligand units and two chloride ions. The crystal structures of
295 **2** and **3** exhibit seven-coordinated U^{4+} and Np^{4+} centers coordinated by the ligand via phenoxide
296 oxygen (O_{Ph}), imine nitrogen ($\text{N}_{\text{C}=\text{N}}$), pyridyl nitrogen (N_{py}), as well as three chloride ions and a
297 THF molecule. However, complex **1** exhibits an eight-coordinated Th^{4+} due to the presence of an
298 additionally coordinated THF molecule. The geometry around the U^{4+} and Np^{4+} center in **2** and **3**
299 can be best described as capped trigonal prismatic with distortion of 6.96 % and 13.968 %
300 respectively (Figure 2 inset and Table S3).[52] In **1**, however, an additionally coordinating THF
301 molecule results in the formation of an eight-coordinated trigonal dodecahedral geometry
302 (distortion = 5.92 %) around the Th^{4+} center (Figure 2, Inset). Gratifyingly, all the 2:1 complexes
303 **4-6** were found having a dodecahedral geometry around the An(IV) centers deviating by 5.62 %,
304 5.46 % and 5.12 %, respectively (see Table S3).

305 The An– O_{Ph} , An– $\text{N}_{\text{C}=\text{N}}$, An– N_{py} and An–Cl bond distances were found to be in the range 2.199–
306 2.234 Å, 2.518–2.673 Å, 2.559–2.640 Å and 2.626–2.734 Å, respectively for complexes **1-3**, and

307 2.235–2.167 Å, 2.653–2.615 Å, 2.743–2.669 Å and 2.681–2.766 Å, respectively for complexes 4-
308 **6** (Table 3). The An–O_{Ph} and An–N_{C=N} distances in the 2:1 complexes **4-6** are comparable with
309 previously reported An(IV) complexes, while bond distances for 1:1 complexes **1-3** are
310 comparatively shorter by approximately 0.5 Å (Table S2). Interestingly, the bond distances in bis-
311 ligated complexes **4-6** are larger as compared to their 1:1 counterpart **1-3**, except for Th–N_{C=N} in
312 **4** (2.653 Å) which is shorter by 0.02 Å than Th–N_{C=N} in **1** (2.673 Å), potentially due to the presence
313 of an additional THF molecule in **1**. Notably, comparative analysis of mono-ligated complex **1**
314 with bis-ligated complexes **4** exhibit marginal change of approximately 0.03 Å for Th–O_{Ph}, Th–
315 N_{C=N}, Th–N_{py} and Th–Cl. Whereas U(IV) and Np(IV) complexes **5** and **6** exhibit considerably
316 larger increases of nearly 0.05 Å (An–O_{Ph}), 0.08 Å (An–N_{C=N}), 0.12 Å (An–N_{py}) and 0.07 Å (An–
317 Cl) as compared to their 1:1 counterpart **2** and **3**. These values indicate a weakening of An-ligand
318 interactions in 2:1 complexes as compared to the 1:1 complexes. Moreover, bond distance
319 comparison among 1:1 complexes **1-3** and 2:1 complexes **4-6** exhibits largest changes from Th(IV)
320 (**1/4**) to U(IV) (**2/5**), with noticeable decreases in An–O_{Ph}, An–N_{C=N} and An–N_{py} distances
321 by -0.07 Å, 0.15 Å and 0.12 Å, respectively from **1** to **2**, and by 0.06 Å, 0.05 Å, and 0.02 Å,
322 respectively from **4** to **5**. Similarly, An–Cl bond distances also exhibit decreases of 0.08-0.10 Å
323 from **1** to **2** and 0.06 Å from **4** to **5**, while no considerable change in bond distances is observed
324 among U to Np complexes. Such a trend in decreasing bond distances from Th⁴⁺ (0.94 Å) > U⁴⁺
325 (0.89 Å) > Np⁴⁺ (0.87 Å) while traversing the An series is well describe in literature. [29]
326 Interestingly, the 1:1 complexes exhibit more pronounced decreases in An–N_{C=N/py} bond distances
327 as compared to An–O_{ph} suggesting a potential covalent contribution in An–N bonds which is not
328 present in An–O_{ph}.

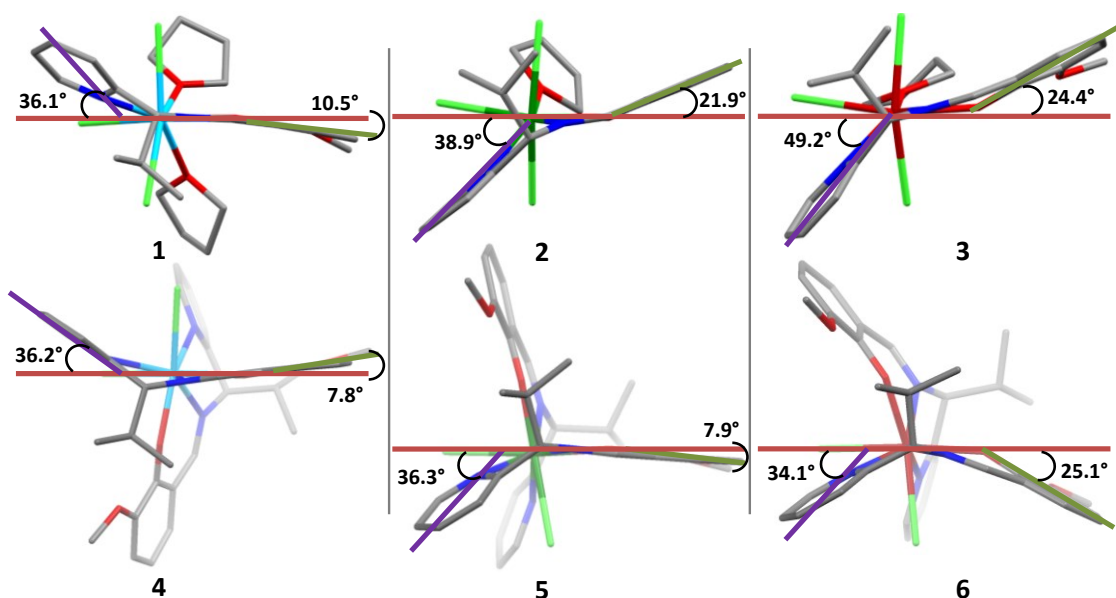


329

330 **Figure 2.** Ellipsoidal representation of complexes **1-6**. Ellipsoids are drawn at 30% probability
 331 level. The hydrogen atoms and lattice THF molecules are omitted for clarity. Inset: polyhedral
 332 representation of coordination environment around central metal ion.

333 Apart from that, change in the number of coordinating ligand to the various An^{4+} ions leads to
 334 differences in the alignment of the aromatic rings relative to the arbitrary plane comprising of
 335 An^{4+} , O_{Ph} , and $(H)C=N$ (Figures 3 and S13). In the solid-state structures of **1-3**, the aromatic rings
 336 are arranged in a pseudo-trans manner relative to the plane containing An^{4+} , O_{Ph} , and $(H)C=N$.
 337 While the phenoxide ring makes angles of 10.5° in **1**, 21.9° in **2** and 24.4° in **3** with the plane, the
 338 pyridyl ring is inclined by 36.1° , 38.9° , and 49.2° in **1**, **2**, and **3**, respectively (see Figure 3). With
 339 respect to the plane, the angle for the phenoxide and pyridyl rings increases in the order **3** > **2** > **1**
 340 suggesting increasingly constrained coordination with decreasing ionic radii. For complexes **4-6**,
 341 a change in the alignment of ligands with respect to the plane is seen. The complex appears to be

342 constrained in having two ligands in a manner that both the aromatic rings of a ligand are aligned
343 on the same side of the plane. Interestingly, for complexes **4** and **5**, pyridyl and phenoxide rings
344 are aligned with the horizontal plane in similar fashion (pyridyl 36.2° (**4**), 36.3° (**5**); phenoxide
345 7.8° (**4**), 7.9° (**5**)). Whereas, in complex **6**, pyridyl and phenoxide rings are inclined to angles of
346 34.1° and 25.1° w.r.t the plane. The maximum deviation w.r.t plane is observed in Np⁴⁺ complexes
347 **3** and **6** potentially due to Np's smallest ionic radius among the three An⁴⁺ under investigation.
348 This change in arrangement of the ligands depending on the size of the metal center once again
349 highlights the considerable spatial flexibility of L^{Pr} in the complexes. This is in agreement with
350 the observed slight differences between the solid-state structures, which may be affected by
351 packing effects and weak intermolecular interactions, and the more symmetrical solution structure,
352 which was suggested by our NMR results. In the end, we believe that An-Ligand coordination is
353 comparatively relaxed and does not show considerable changes in the structural properties while
354 changing the An⁴⁺ whereas coordination in the 2:1 complexes is comparatively strained and
355 consequently exhibits greater structural rearrangements with small changes in ionic radii (0.02 Å
356 from U⁴⁺ to Np⁴⁺) of An⁴⁺. [28]



357
 358 **Figure 3.** Capped stick representation of complexes **1-6**, displaying the angle formed by phenoxide
 359 ring (green line), pyridyl ring (magenta line) with the horizontal plane (brown line). The hydrogen
 360 atoms and lattice THF molecules are omitted for clarity.

Table 3. Selected bond distance (Å) parameters for complexes **1-6**.

Bond	1	2	3 ^a	4	5	6
An–O _{Ph}	2.199(4)	2.125(12)	2.122 (2)	2.235(2)	2.178(1)	2.167(1), 2.184(1)
An–N _{C=N}	2.673(5)	2.528(14)	2.514(2)	2.653(2)	2.606(1)	2.579(2), 2.615(2)
An–N _{Py}	2.725(5)	2.610(16)	2.559(2)	2.743(2)	2.720(1)	2.667(2), 2.669(2)
An–O _{THF}	2.543(5), 2.601(4)	2.455(16)	2.446(2)	-----	-----	----
An–Cl ₁	2.733(14)	2.635(5)	2.626(1)	2.766(7)	2.708(1)	2.681(1)
An–Cl ₂	2.709(14)	2.632(6)	2.629(1)	2.766(7)	2.708(1)	2.698(1)
An–Cl ₃	2.734(15)	2.662(5)	2.635(1)	----	----	----

^avalues are average of three molecular structures in the unit cell. Ionic radii (CN = 6): Th(IV) = 0.94 Å; U(IV) = 0.89 Å; Np(IV) = 0.87 Å.

361 Electrochemistry

362 To understand the redox properties of all synthesized compounds, cyclic voltammetry experiments
363 were performed in acetonitrile solvent (Figures 4 and S14 and Table 4) and values are calculated
364 vs $\text{Fc}^{0/+}$. [53] The free ligand **HL**^{Pr} exhibits three oxidative signals at, 0.47, 0.84, and 1.73 V vs
365 $\text{Fc}^{0/+}$ and two reductive responses at 0.74 and 0.34 V vs $\text{Fc}^{0/+}$ (Figure S14). The oxidative responses
366 at 0.47 V and 0.84 V are quasi-reversibly coupled with reductive responses at 0.34 V and 0.74 V,
367 respectively. The $E_{1/2}$ values for these redox couples are 0.40 V and 0.79 V, respectively with peak-
368 to-peak separations (ΔE_p) of 130 mV and 100 mV, respectively. Notably, the cathodic wave at
369 0.06 V (E_{pc}) and its anodic couple at 0.19 V (E_{pa}) are formed during the scan at the expense of
370 features corresponding to both $E_{1/2}$'s. The reverse cycle exhibits the formation of an additional
371 reductive response at -1.10 V potentially due to the reduction of a previously oxidized imine group
372 but producing a chemically different species. The differential pulse voltammetry (DPV)
373 experiment further confirms the presence of redox couples at 0.40 V, 0.76 V and 1.65 V.

374 Complex **1** displays a broad oxidative response (E_{pa}) at 0.86 V, 1.34 V, and 1.52 V along with an
375 observable reductive couple (E_{pc}) at 0.75 V, 1.03 V and 1.44 V respectively, corresponding to
376 ligand-based signals (Figure 4). The oxidative signals (E_{pa}) are positively shifted by ca. 390 mV
377 and ca. 500 mV compared to the free ligand indicating an influence of the coordinated Th^{4+} in **1**
378 on the ligand's oxidative potential. Moreover, cathodic waves (E_{pc}) at -0.92 V, -1.42 V and
379 -1.69 V could be assigned to ligand-based signals due to potential formation of a C-C bond after
380 ligand reduction. Such electrochemical behavior for An-(Schiff base) complexes was previously
381 reported by Clément et. al. [36]. Interestingly, the bis-ligated Th(IV) complex **4** displays
382 voltammogram similar to its 1:1 counterpart **1**. The two oxidative responses were observed at
383 approximately 0.92 V and 1.20 V and four reductive responses at -1.40, -1.63, -2.12, and

384 -2.21 V. Since thorium is not expected to show any redox response within the potential range
385 under discussion, these signals most likely corresponds to ligand-based responses.[54] We
386 anticipate signals corresponding to Th(IV)/Th(III/II) beyond -2.3 V, [55,56] and thus not within
387 the range of our experiments.

388 The U(IV) complex **2** exhibits five redox couples at $E_{1/2} = 0.45, 0.76, 0.98, 1.25,$ and 1.52 V on
389 the positive side of voltammogram (Figure 4). While redox couples at $E_{1/2} = 0.76, 1.25,$ and 1.52 V
390 best matches to ligand-based responses, the responses at $E_{1/2} = 0.45$ V ($E_{pa} = 0.49$ V) and 0.98 V
391 ($E_{pa} = 1.05$ V) could be tentatively assigned to metal-based U(IV)/U(V) and U(V)/U(VI) redox
392 couples, respectively. Interestingly, ligand based oxidative responses (E_{pa}) at 0.82 V and 1.37 V
393 are positively shifted by ca. 350 mV and 530 mV in comparison to **HL**^{Pr}. Moreover, similar to
394 what was observed for complex **1** ($E_{pc} = -1.68$ V), a ligand-based reductive feature at -1.76 V
395 (E_{pc}) is observed in **2**. Whereas the feature at -2.09 V most likely corresponds to a U(IV)/U(III)
396 reductive process. [33] On the other hand, the 2:1 U(IV) complex **5** exhibits four prominent
397 oxidative features at $E_{pa} = 0.73, 1.18, 1.61,$ and 1.86 V along with the four noticeable reductive
398 responses at $E_{pc} = -1.22, -1.89, -2.05,$ and -2.22 V. The oxidative features at $0.73, 1.61$ and
399 1.86 V correlate to ligand-based responses. However, in contrast to observations of two metal-
400 centered oxidative responses in **2**, only one oxidative feature at 1.18 V is observed in **5** which we
401 tentatively assigned as U(IV)/U(V/VI). In addition, reductive features at $E_{pc} = -1.22, -1.89$ and
402 -2.22 V [33] correspond to ligand-based responses, whereas the feature at -2.05 V could be
403 assigned to a U(IV)/U(III) response.

404 For complex **3**, broad oxidative (E_{pa}) and reductive signals (E_{pc}) indicating that metal-based
405 responses have closely matched potentials with that of ligand. However, a closer look at the
406 voltammogram shows the presence of four oxidative responses 0.97 V, 1.05 V, 1.28 V, and

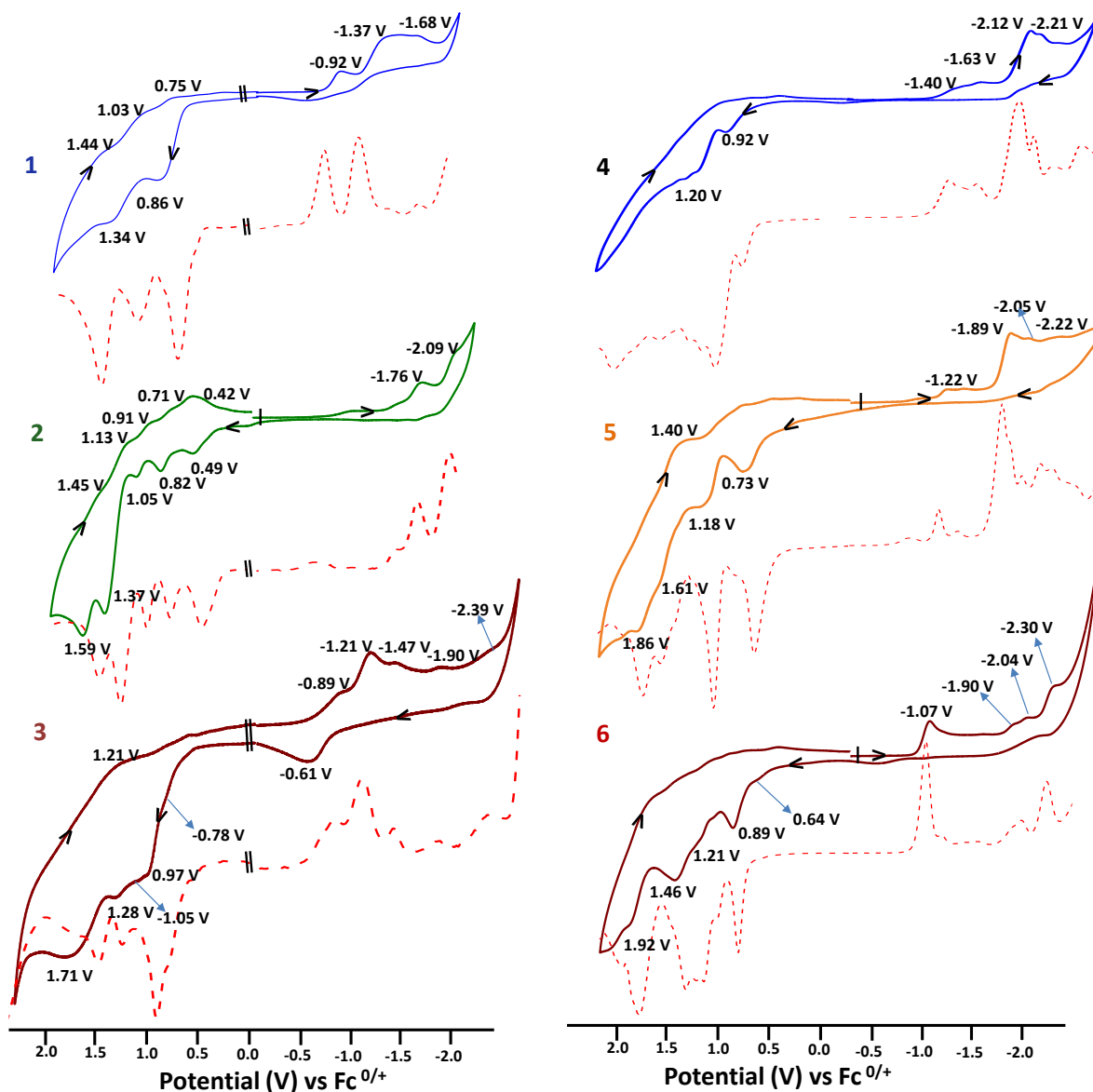
407 1.71 V. In addition, a weak signal at approx. 0.78 V is observed. Oxidative responses at 0.97 V,
408 1.28 V, and 1.71 V are closely matched with the ligand-based responses in complexes **1** and **2**,
409 whereas signals at approx. 0.78 V and 1.05 V could be tentatively assigned as Np(IV)/Np(V) and
410 Np(V)/Np(VI) processes, respectively. Importantly, ligand-based signals at $E_{pa} = 0.97$ V and
411 1.28 V are positively shifted by 500 mV and 440 mV, respectively, in comparison to the free
412 ligand, and are similar to those of complexes **1** and **2** containing Th^{4+} and U^{4+} ions. The reductive
413 responses at the negative side of voltammogram could be assigned as ligand-based responses.
414 Based on one reference available in the literature, we anticipate the observation of Np(IV) based
415 reduction responses beyond our analytical range of -2.5 V.[57]. On the other hand, the 2:1
416 counterpart **6** displays broad oxidative signals, similar to **3**, having four measurable features at
417 0.79, 1.21, 1.46, and 1.92 V along with a weak trace at 0.64 V. Considering our observation in
418 other complexes (**1-5**), we tentatively assign 0.89, 1.46 and 1.92 V to the ligand-based responses,
419 whereas, features at 0.64 V and 1.21 V could be assigned as Np(IV)/Np(V) and Np(V)/Np(VI)
420 responses, respectively. Moreover, four reductive responses at $E_{pc} = -1.07, -1.90, -2.04$ and -2.30 V
421 could be assigned to ligand-based responses. Gratifyingly, DPV experiments further corroborate
422 the observations for these complexes.

423

Table 4. Redox potentials values for **HL^{pr}** and complexes **1-6** determined by CV.

Entry	Ligand-based				Metal-based			
	E_{pa}	E_{pc}	$E_{1/2}$	DPV	E_{pa}	E_{pc}	$E_{1/2}$	DPV
HL^{pr}	0.47, 0.84, 1.73	0.34, 0.74	0.40, 0.79	0.40, 0.76, 1.65	----	----	----	----
1^a	0.86, 1.34, 1.52	0.75, 1.03, 1.44, -0.92, -1.37, -1.68	0.80, 1.18	0.70, 1.09, 1.44, -0.75, -1.09	----	> 2.1	----	----
2	0.82, 1.37, 1.59	0.71, 1.13, 1.45, -1.76	0.76, 1.25, 1.52	0.76, 1.24, 1.45, -1.70	0.49, 1.05	0.42, 0.91, -2.09	0.45, 0.98	0.44, 0.98, -2.02
3	0.97, 1.28, 1.71	-0.89, -1.21	-----	0.89, 1.22, 1.46	0.78, 1.05	-1.47, -1.90, -2.40	-----	0.51, 0.77, -1.44, -1.85, -2.39
4	0.92, 1.20	-1.40, -1.63, -2.21	-----	0.77, 1.03, 2.05, -1.31, -2.00, -2.60	----	----	----	-2.14
5	0.73, 1.61, 1.86	1.40, -1.22, -1.89, -2.22	----	0.67, 1.07, 1.59, 1.78, -1.86, -1.81, -2.19	1.18	-2.05	----	-1.90
6	0.89, 1.46, 1.92	-1.07, -1.90, -2.30		0.83, 1.05, 1.19, 1.36, 1.82, - 1.03, -2.25	0.64, 1.21	-2.04	----	-1.99

^a values could not be assigned exactly.



424

425 **Figure 4.** Cyclic (thick line) and Differential Pulse (dotted line) voltammogram of 1:1
 426 complexes **1** (—), **2**(—) and **3** (—) and 2:1 complexes **4** (—), **5**(—) and **6** (—) vs $\text{Fe}^{0/+}$ in
 427 acetonitrile solvent. Conditions: Ligand/complex approx. 1 mM, $n\text{-Bu}_4\text{NPF}_6$, supporting
 428 electrolyte approx. 100 mM, glassy carbon working electrode, Pt wire auxiliary electrode, Ag/Ag^+
 429 reference electrode; scan rate: 100 mV s^{-1} .

430

431 **Binding Studies**

432 We performed concentration dependent absorption titrations to understand the binding affinity of
433 $(\text{L}^{\text{Pr}})^-$ with different metal ions (Figures S15-S17). In complex **1-3**, absorption spectral titration
434 between Th^{4+} , U^{4+} , and Np^{4+} with $(\text{L}^{\text{Pr}})^-$ exhibited an increase in absorption feature at 450 nm.
435 The changes in the absorbance reached maxima on addition of 1.0 equiv. metal ion indicating a 1:1
436 stoichiometry of $(\text{L}^{\text{Pr}})^-$ with respect to the actinide ion, in agreement with the SC-XRD results.
437 The binding coefficients were calculated (at 450 nm for **1-3**) using the Benesi–Hildebrand eq. 1.
438 [58-60]

$$439 \quad \frac{1}{(A - A_0)} = \frac{1}{K(A_{\text{max}} - A_0)[\text{An}^{n+}]^x} + \frac{1}{(A_{\text{max}} - A_0)} \quad (1)$$

440 Here, ' A_0 ' is the absorbance of the ligand $(\text{L}^{\text{Pr}})^-$, ' A ' is the change in absorbance after addition
441 of An ions, ' A_{max} ' is the absorbance value after adding excess amount of An ions, ' K ' is the
442 association constant (M^{-1}), $[\text{An}^{n+}]$ is the concentration of the An ions added (M), and ' x ' is the no.
443 of equivalents. The linear regression plot between absorption intensity, $1/[A - A_0]$ vs $1/[\text{An}^{n+}]^x$
444 varied linearly as a function of $1/[\text{An}^{n+}]$ ($x = 1$), confirming a 1:1 stoichiometry (Figures S15-S17).
445 The binding constants (K) were calculated as $(2.3 \pm 0.4) \times 10^3 \text{ M}^{-1}$ for **1**, $(1.54 \pm 0.3) \times 10^3 \text{ M}^{-1}$ for
446 **2**, and $(2.01 \pm 0.4) \times 10^2 \text{ M}^{-1}$ for **3**. These numbers suggest that Th^{4+} and U^{4+} have similar binding
447 affinities to $(\text{L}^{\text{Pr}})^-$ whereas Np^{4+} has an approx. ten-fold weaker binding affinity.

448 **Conclusions**

449 Herein we report the synthesis and characterization of six mono-nuclear tetravalent actinide
450 complexes (**1-6**) comprising mono-ligated (**1-3**) and bis-ligated complexes (**4-6**) with a novel
451 Schiff base-type ligand. Their comparative analysis exhibits the influence of change in

452 coordination environment on the (electro-)chemical properties of actinide complexes. Notably, the
453 effect of the actinide on the electronic properties of the complexes is more pronounced in the 1:1
454 complexes than for their 2:1 counterparts. This is most obvious in NMR spectroscopy, where ^1H
455 NMR for 1:1 Th(IV) complex (**1**) is shifted upfield from its 2:1 counterpart **4** and ^1H NMR signals
456 for 1:1 U(IV) and Np(IV) complexes are highly paramagnetically shifted between -70 to 40 ppm.
457 These signals appeared within -35 to 25 ppm for 2:1 complexes **5** and **6**. Single crystal structures
458 reveal an increase in the An-ligand bond distances when moving from 1:1 to 2:1 stoichiometry,
459 which also confirms our interpretation of NMR data as indicating weak An-ligand interactions in
460 2:1 complexes. Apart from that, bond distances in both 1:1 or 2:1 complex decrease with the
461 decreasing ionic radii of An(IV) center while traversing the An series. These structural changes
462 are accompanied by changes in the arrangement of the aromatic rings in the complexes, which
463 emphasizes the limited structural flexibility in this system. The maximum deviation of phenoxide
464 and pyridyl ring(s) is observed in Np^{4+} complexes **3** and **6**. This goes along with binding studies
465 suggesting that Np^{4+} has a significantly weaker binding affinity to $(\text{L}^{\text{Pr}})^-$ than Th^{4+} and U^{4+} . Cyclic
466 voltammetry studies revealed an increase in the ligand-based oxidation potential by 200 - 300 mV
467 in all the complexes. In addition, several redox transitions could be tentatively assigned to actinide
468 reductions and oxidations. While these processes need to be verified, cyclic voltammetry suggests
469 $(\text{L}^{\text{Pr}})^-$ may be suitable to stabilize actinides in both low and high oxidation states. Our results add
470 to the very limited database of structures of actinide-organic complexes and sheds new light on
471 the interplay of molecular and electronic structure in these compounds.

472

473 **Supporting Information.**

474 The Supporting Information is available free of charge at XXXXXXXXXX.

- 475 • Additional details related to NMR, UV-vis, FTIR and crystal refinement parameters for all
476 the synthesized molecules.

477 **Accession Codes**

478 CCDC 2075052-2075054 for complexes **1-3** and 2152165-2152167 for complexes **4-6** contain the
479 supplementary crystallographic data for this paper. These data can be obtained free of charge via
480 www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by
481 contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ,
482 UK; fax: +44 1223 336033.

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487 **Notes**

488 The authors declare no competing financial interest.

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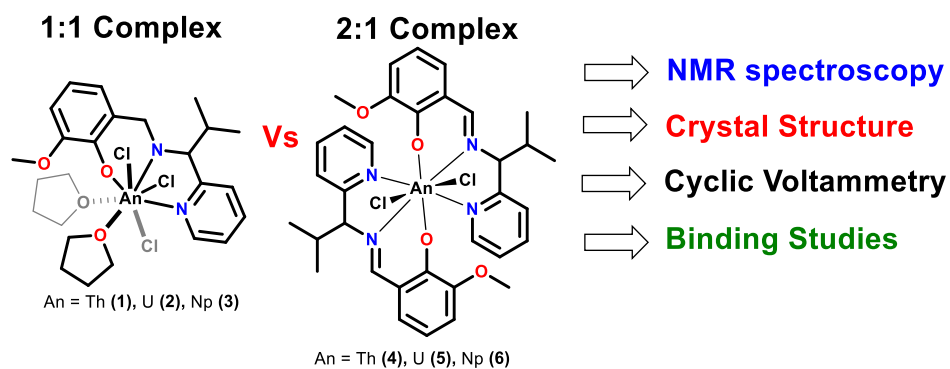
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625 independently. Whereas 2:1 complexes (**4-6**) display stable cyclic voltammogram for at least
626 one full cycle of measurement. Notably, though we have observed electronic influence of
627 chirality only in Th(IV) complexes **1** and **4**. We believe that the redox signals contain responses
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650 **SYNOPSIS.** A series of mononuclear tetravalent actinide complexes (**1-6**) have been synthesized
651 using a new monoanionic Schiff base ligand (**HL^{Pr}**). Comparative analysis between 1:1 complexes
652 [**MCl₃-L^{Pr}.nTHF**] (**1-3**) and 2:1 complexes [**MCl₂-L^{Pr}₂**] (**4-6**) shows intriguing influence of
653 coordinating ligands in coalescence with coordinating properties of An⁴⁺ centers on the spectral,
654 structural and electrochemical properties.



655