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RESEARCH ARTICLE

View Article Online

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Cite this: *Inorg. Chem. Front.*, 2024, **11**, 3731

Employing a template synthesis to access diastereopure Np(IV) and U(IV) complexes and analysis of their 5f orbitals in bonding†

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Development of new synthetic methods to employ with transuranium organometallic complexes will help further the field and may provide a stronger understanding of their distinct reactivity, electronic structures, and chemical bonding. Although the organoneptunium field has grown over the last decade, just a handful of complexes have been synthesized, most of which are found in the trivalent oxidation state. Few reactions with neptunium (Np) have considered the stereochemical outcome, thus we employed a diastereoselective template protocol to synthesize *meso* (or: C_2) complexes of neptunium(IV). Following previous success with the diastereoselective synthesis of $(^{tBu_2P}ONO)UCl_2(dtbp)$ (**1**), adaptation to $NpCl_4(dme)_2$ led to the isolation of diastereopure $(^{tBu_2P}ONO)NpCl_2(dtbp)$ (**2**), $(^{tBu_2P}ONO) = 2,6\text{-bis}((di\text{-}tert\text{-butylphosphino})\text{-methanolato})pyridine$). The addition of a bulky indenide ligand, derived from the *in situ* deprotonation of 4,7-dimethyl-1,3-bis(1-methylethyl)-1*H*-indene (1,3-(*t*Pr)₂-4,7-Me₂-C₉H₃), was pursued to synthesize the organoactinide complexes $(^{tBu_2P}ONO)U(1,3\text{-}(tPr)_2\text{-4,7-Me}_2\text{-C}_9\text{H}_3)Cl$ (**3**) and $(^{tBu_2P}ONO)Np(1,3\text{-}(tPr)_2\text{-4,7-Me}_2\text{-C}_9\text{H}_3)Cl$ (**4**). Bonding analyses consisting of orbital localization and energy decomposition methods show resemblances in covalent character between uranium (**1** and **3**) and neptunium (**2** and **4**) systems. However, an increased orbital overlap was identified in the metal – indenide with respect to the metal–dtbp interaction, mainly driven by 5f electrons. This was attributed to the better symmetry match between the organic ligand and metal orbitals that allowed the participation of the 5f orbitals.

Received 5th April 2024,
Accepted 29th May 2024

DOI: 10.1039/d4qi00810c

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Introduction

Neptunium coordination chemistry has experienced rapid growth in recent years,^{1–11} which has been facilitated by its lower activity, cost, and regulatory burden relative to plutonium with respect to the most common isotopes of these elements used in synthetic chemistry. However, the specialized facilities required for the radiological hazards of ²³⁷Np and its ²³³Pa daughter are more significant than those of uranium. Although progress is reflected in the increasing number of

complexes that have been reported, the gap with respect to organoactinide chemistry is pronounced. Traversal of the actinide series is marked by a gradual energetic stabilization of the 5f shell, unlike the 6d shell which remains almost at constant energy. This, added to the more contracted nature of the 5f with respect to the 6d shell, makes the latter more prone to participate in metal–ligand interactions.^{12,13} However, by taking advantage of the unique shape of the f-orbitals, in combination with the diverse symmetry-adapted interactions that aromatic ligands such as arene or indene ligands can provide, a preferential interaction that exposes the 5f over the 6d shell can be favored, encouraging their participation in covalent interactions.

Recent developments in organoneptunium chemistry can be traced back to well-defined neptunium starting materials for air- and moisture-free synthesis, $NpCl_4(thf)_3$, $NpCl_3(py)_4$, $NpBr_3(thf)_4$, $NpI_3(thf)_4$ and $NpCl_4(dme)_2$ (*thf* = tetrahydrofuran, *dme* = 1,2-dimethoxyethane),^{14–18} all of which circumvent the need for Np⁰ metal that was required previously.¹⁹ In addition to neptunium complexes bearing cyclopentadienyl ligands,^{20–22} η⁶-arene complexes and hydrocarbyl-coordinated

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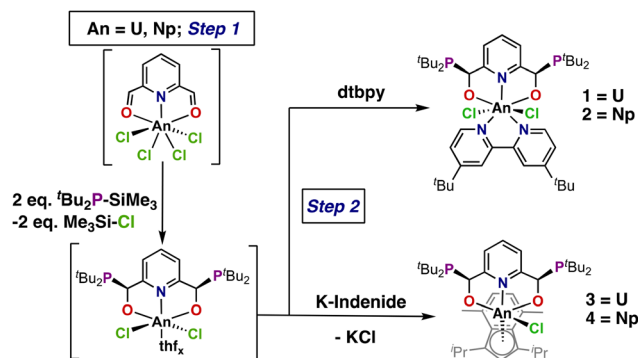


Np^{4+} centers have also been described.^{23–25} These systems have set the foundations of organoneptunium chemistry and analysis of its chemical bonds. Investigating novel coordination motifs in organoactinide complexes may provide further understanding of f-orbital participation in neptunium–ligand bonding interactions. Accordingly, we expanded upon our prior efforts and introduced template techniques to neptunium chemistry.

Template synthesis protocols are reactions that use a metal center to arrange reactive fragments prior to reactive product generation. Templating is pursued when mixtures of products, or facile deleterious side reactions, are expected during a chemical reaction. Accordingly, templating has often been employed for the formation of macromolecules and molecular knots.^{26–29} Additional work has shown that template synthesis can aid in the activation of inert pre-ligand moieties to enable reactivity that is unobserved with traditional synthesis.^{30–32} Our interests lie in developing synthetic procedures for use with actinide elements, where fundamental research efforts are often limited to small scales due to low availability and radiological hazards, persistent problems when researching the chemistry of neptunium. Here, employing template protocols to increase product yields and minimize product mixtures was sought during the requisite use of small scales of neptunium.

We previously reported a template procedure that constructed diastereopure complexes of both iron and uranium.³³ Coordination of 2,6-pyridine dicarboxaldehyde ((CHO)₂Py) to the metal center followed by the addition of two equivalents of di^{tert}-butyl(trimethylsilyl)phosphine (*t*Bu₂PSiMe₃) generated the *meso* complexes. Synthesis of the free ligand without a template generated a 63:37 *rac*:*meso* (C₂:C_S) ratio favoring slightly the formation of the *rac* isomer, which required fractional crystallization to purify the diastereoisomers following coordination to FeCl₂.³³ Remarkably, pre-coordination of (CHO)₂Py to a metal halide using either FeCl₂ or UCl₄ prior to addition of *t*Bu₂PSiMe₃ resulted in substantial stereospecificity (>90%) for the *meso* diastereoisomers of the respective products. The reaction of the uranium congener produced a pyridine-based dianionic [ONO]^{2–} complex arising from the loss of TMS-Cl and U–O bond formation (Scheme 1, top right).³³ Both the improved stereospecificity and higher yields suggested that adaptation of this method to neptunium would be successful.

Similar bond dissociation energies³⁴ for U–O (755 kJ mol^{–1}) and Np–O (731 kJ mol^{–1}) suggest a successful diastereoselective synthesis of a neptunium complex can be achieved *via* this synthetic route. The ease and high purity of this method lends itself to the generation of unique starting materials for subsequent chemistry. In this report, organometallic derivatization was pursued to establish chemistry beyond the synthesis of halide materials and to generate the respective diastereopure organoactinide complexes. An indene derivative was employed to generate the corresponding organoactinide derivatives (Scheme 1). Characterization of the resulting complexes *via* single-crystal X-ray diffraction (SC-XRD), UV-Vis-NIR, and NMR spectroscopies was performed



Scheme 1 Proposed reactivity of the template synthesis from either UCl₄ or (dme)₂NpCl₄ (dme = 1,2-dimethoxyethane) with the loss of TMS-Cl (TMS = trimethylsilyl).

alongside natural bond orbital (NBO) analysis, time-dependent density functional theory (TD-DFT) and multi-reference wavefunction calculations to explore the electronic structure and bonding of these systems. This analysis allowed for direct comparisons between homologous neptunium and uranium complexes.

Experimental results

Applying a diastereoselective synthesis to neptunium

Scale-down reactions using UCl₄ were performed prior to work using NpCl₄(dme)₂ in order to identify optimal reaction and crystallization conditions. Coordination of (CHO)₂Py to a stirring solution of NpCl₄(dme)₂ in THF at room temperature resulted in little color change of the pale-pink solution (Fig. S18†). Following the coordination of the dialdehyde to the metal center, *t*Bu₂PSiMe₃ (2 equivalents) was added to the reaction solution, resulting in a color change to yellow (Fig. S19†). Removal of TMS-Cl was performed by holding the reaction solution under reduced pressure to a constant mass. The addition of 4,4'-di-^{tert}-butyl-2,2'-bipyridyl (dtbpy) in fluorobenzene to the tacky yellow solid resulted in the formation of a pale-peach solution (Fig. S20†). Pale-peach rectangular plates were observed after leaving the filtered reaction solution undisturbed for 10 days at –35 °C (Fig. S21†). These crystals were isolated in 46% yield and were assigned as (^{*t*}Bu₂P)₂(ONO)NpCl₂(dtbpy)₂.

Complex 2 was successfully characterized by NMR and UV-Vis-NIR spectroscopies. The ¹H NMR spectrum of 2 has a broad signal range, as expected for a formally paramagnetic 5f³ Np⁴⁺ complex, with resonances found over a 70 ppm window. The resonances attributed to the ^{*t*}Bu₂P groups were found in the ¹H NMR spectrum at δ 13.05 ppm and δ –1.86 ppm as broadened doublets with a resolved coupling of ~10 Hz. A diagnostic set of downfield resonances centered at δ 65.50 ppm and δ 54.43 ppm in 2 is reminiscent for those observed for 1 (δ 67.16 ppm and δ 24.30 ppm), which are tentatively assigned as the *m*-pyridine C–H and the two methine C–



H groups. The ^{31}P NMR spectrum of **2** contained a single resonance at δ 79.3 ppm, significantly shifted from the resonance of the free ligand, *meso*- $^{\text{OTMS}}\text{PNP}^{\text{tBu}}$ (δ 50.2 ppm), and slightly shifted from the observed resonance of **1** (δ 86.0 ppm).³³ The presence of a single phosphorus resonance suggests formation of a single diastereoisomer, a result consistent with the U-based analogue. The chemical shift difference between **1** and **2** likely arises from the inherent disparities of their expected magnetic moments based on electron count; U^{4+} is formally a $5f^2$ system with an inherently lower magnetic moment when compared to the $5f^3$ Np^{4+} center. Previously established Evans method protocols for transuranium complexes were followed,^{24,35} and the magnetic moment of **2** was found to be $2.32\mu_{\text{B}}$, lower than previously observed solution values for a $5f^3$ Np^{4+} complexes.²⁴

SC-XRD experiments were performed to unambiguously assign the identity of the complex as the desired *meso* diastereoisomer (Fig. 1). The solid-state structure of **2** was solved in the triclinic space group $P\bar{1}$ with unit cell parameters that were significantly different from our original report of **1** (solved in the monoclinic $P2_1/c$ space-group). The differences in the structures arise from a different arrangement of the solvent molecules in the unit cells. In the reported $P2_1/c$ cell of **1**, the asymmetric unit cell contained four discrete fluorobenzene molecules. The $P\bar{1}$ cell of **2** contained three disordered fluorobenzene molecules that were modelled as partially occupied. Additional crystallization attempts produced a polymorph of **1** that was solved in $P\bar{1}$, allowing for a direct comparison to **2** (Table 1, ESI section 5†).

Little difference between the core molecule of **1** was found between the two different unit cells. The bond lengths and angles between **1** and **2** are nearly identical, though the Np–O bond distances are consistently slightly contracted than the U–O bond distances as expected (Table 1), although the values remain within 3σ of ESD values (ESD = estimated standard

Table 1 Select bond distances (Å) and angles ($^\circ$) for **1**³³ ($P\bar{1}$), **1** ($P2_1/c$), and **2**

Distance (Å), \angle ($^\circ$)	1 ($P2_1/c$) ^a	1 ($P\bar{1}$)	2 ($P\bar{1}$)
M–O (avg)	2.165(4)	2.154(4)	2.150(4)
M–Cl (avg)	2.670(4)	2.653(3)	2.642(3)
M–N _{1_{pyr}}	2.481(2)	2.484(3)	2.461(4)
M–N _{dtbpy} (avg)	2.621(7)	2.602(4)	2.586(6)
Cl1–M–Cl2	163.69(7)	163.41(4)	162.45(4)
O1–M–O2	129.2(2)	129.21(10)	129.68(10)
N _{1_{pyr}} –M–O1	65.0(2)	64.85(10)	66.35(10)
N _{1_{pyr}} –M–O2	65.02(2)	64.64(10)	65.84(10)

^a Previously reported in ref. 33.

deviation). The M–N bond is found to be 0.02 Å shorter in the neptunium congener, also within 3σ of ESD values. Complex **2** represents a rare $[\text{ONO}]^{2-}$ ligand type for neptunium, with dipicolinate $[\text{ONO}]^{2-}$ representing the previously known complexes.^{36–42} Additionally, there is recently a single report of an enantioselective synthesis on neptunium bearing optically active (chiral) amidinates,⁴³ yet complex **2**, to our knowledge, describes the first diastereoselective reaction of neptunium by employing the benefits of template methods.

Beyond dtbpy: synthesis of actinide-indenyl complexes

Complexes **1** and **2** were isolated in good yields and high purity. However, the lability of the dtbpy ligand of **1** was problematic for subsequent experiments, contaminating resultant products. An alternative route to substitution was pursued to circumvent the use of dtbpy in the synthesis of these complexes. The addition of an anionic ligand derived from *in situ* deprotonation of 4,7-dimethyl-1,3-bis(1-methylethyl)-1*H*-indene ($1,3\text{-}(\text{Pr})_2\text{-}4,7\text{-Me}_2\text{-C}_9\text{H}_3$) was employed following U–O bond formation and TMS–Cl release. The indenide ligand was chosen for electronic and coordination variability and reported high-degree of crystallinity,⁴⁴ and was expected to provide steric protection and stability to the $[(\text{ONO})\text{An}]$ -core of these complexes. Furthermore, steric hindrance was expected to induce coordination to the metal *trans* to the $^{\text{tBu}}\text{P}$ groups.

The organometallic synthesis was optimized on uranium, with adjustments to the synthetic protocol of **1** to include the use of $[\text{K}][1,3\text{-}(\text{Pr})_2\text{-}4,7\text{-Me}_2\text{-C}_9\text{H}_3]$. $(\text{CHO})_2\text{Py}$ (1 equivalent) was initially added to UCl_4 to form a tan slurry in THF. The addition of two equivalents of $^{\text{tBu}}\text{P}(\text{SiMe}_3)_3$ to the reaction slurry resulted in the formation of a vibrant green solution and the loss of TMS–Cl. Removal of all volatiles under reduced pressure left a tacky, green material, which was then redissolved in THF. A THF solution of $[\text{K}][1,3\text{-}(\text{Pr})_2\text{-}4,7\text{-Me}_2\text{-C}_9\text{H}_3]$ was added dropwise to the uranium solution, resulting in a rapid color change to dark red. After work-up, crystalline material was isolated overnight from a concentrated pentane solution at -35 °C in 61% crystalline yield and assigned as the desired product $(1,3\text{-}(\text{Pr})_2\text{-}4,7\text{-Me}_2\text{-C}_9\text{H}_3)(^{\text{tBu}}\text{P})_2\text{ONO}(\text{U})\text{Cl}$ (**3**). Analysis of the ^1H NMR spectrum suggested a paramagnetic complex, the spectrum comprising broadened, featureless peaks over a wide range ($\sim\delta$ –64 ppm to δ 79 ppm). The reso-

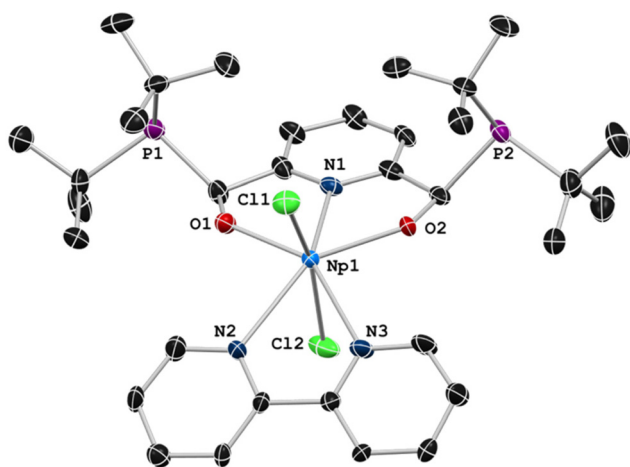


Fig. 1 Solid-state structure of **2** presented at 50% probability ellipsoids. Hydrogen atoms, disordered fluorobenzene solvent molecules (Fig. S25†), and *para*-*tert*-butyl groups of the dtbpy ligand have been omitted for clarity.



nances attributed to the ${}^t\text{Bu}_2\text{P}$ groups were found with a broad singlet at δ 32.21 ppm and a broadened doublet ($J = 6$ Hz) at δ -2.97 ppm, also with diagnostic downfield resonances centered at δ 78.90 ppm and δ 44.75 ppm. The ${}^{31}\text{P}$ NMR spectrum of **3** contained a single resonance centered at δ 81.8 ppm, slightly shifted from that of complex **1** (δ 86.0 ppm).³³ Magnetic studies performed *via* Evans method NMR gave a magnetic moment of $2.71\mu_{\text{B}}$, for **3** consistent with both complex **1** ($2.33\mu_{\text{B}}$) and other previously reported U^{4+} complexes.^{33,45}

Crystals of **3** grew as well-defined blocks but the crystals produced diffraction patterns suggestive of twinning. Following several attempts and careful crystal splitting, smaller single crystals produced sufficient, high-quality data with suggestions of minor twinning of the data. SC-XRD analysis of these crystals confirmed the assignment of the product as $(1,3\text{-}({}^t\text{Pr})_2\text{-}4,7\text{-Me}_2\text{-C}_9\text{H}_3)^{({}^t\text{Bu}_2\text{P})}\text{ONO}(\text{U})\text{Cl}$ (Fig. 2). Complex **3** was solved in the monoclinic $P2_1/n$ space group, where the asymmetric unit cell contained a single molecule of **3**, as well as pentane solvent disordered over two positions.

The coordination environment around the uranium center consists of a single η^5 -indenide ligand, the $[\text{ONO}]^{2-}$ chelate, and one Cl^- . As anticipated, the installation of the indenide occurred on the sterically accessible side, and the remaining Cl^- is nestled between the steric bulk of the ${}^t\text{Bu}_2\text{P}$ moieties. The solid-state metrical parameters of **1** and **3** show little change to the chelate between the two metals, with the U–N and U–O bond distances found slightly contracted in **3**. The O–U–O angle of the chelate is effectively unchanged, found at $129.2(2)^\circ$ in **1** and $130.66(2)^\circ$ in **3**. The average U–centroid bond of **3** ($2.477(1)$ Å) is consistent with, if slightly contracted, tris(indenyl)uranium chloride, $\text{U}(\text{Cl})(\text{C}_9\text{H}_7)_3$, which is reported at 2.49 Å.⁴⁶ Additional metrical parameters can be found in Table 2. Compound **3** joins a small number of U–indenide complexes that have been reported^{47–53} in the larger established field of organouranium chemistry.

Table 2 Select bond distance (Å) and angles ($^\circ$) for **3** and **4**

Distance (Å), \angle ($^\circ$)	3	4	4 _{Calc.} ^a
M–O1 (avg)	2.130(2)	2.126(11)	2.146
M–Cl1	2.603(1)	2.574(3)	2.585
M–N1 _{pyr}	2.465(1)	2.449(6)	2.479
M–Centroid	2.477(1)	2.443(6)	2.458
Cl1–M–Cent.	118.96(1)	116.33(15)	112
O1–M–O2	130.66(2)	130.6(4)	128
N1 _{pyr} –M–O1	65.63(2)	64.8(4)	65
N1 _{pyr} –M–O2	65.44(2)	66.4(4)	65

^a Structural data obtained from a geometry optimization employing scalar ZORA-B3LYP/TZP level of theory.

Synthesis of the Np-indenide homologue

The synthesis employed for **3** was adapted to neptunium using $(\text{dme})_2\text{NpCl}_4$. Following TMS–Cl elimination and formation of the $[\text{ONO}]^{2-}$ chelate, a THF solution of $[\text{K}][[1,3\text{-}({}^t\text{Pr})_2\text{-}4,7\text{-Me}_2\text{-C}_9\text{H}_3]]$ was added. The color of the solution rapidly turned dark from the yellow color (Fig. S22[†]). Following work-up, the dark residue was taken into pentane and crystallized at -35 °C. Dark, diamond-shaped crystals formed and were assigned as $(1,3\text{-}({}^t\text{Pr})_2\text{-}4,7\text{-Me}_2\text{-C}_9\text{H}_3)^{({}^t\text{Bu}_2\text{P})}\text{ONO}(\text{Np})\text{Cl}$, **4** (Fig. S23[†]). Characterization of **4** using NMR and UV-Vis-NIR spectroscopies and SC-XRD conclusively show the generation of the desired complex **4** in low yields (12% crystalline yield).

The ${}^1\text{H}$ NMR spectrum of **4** exhibited a wide chemical shift range (~ 60 ppm) that is consistent with those of **1**, **2**, and **3**. Resonances attributed to the ${}^t\text{BuP}$ moieties were found at δ 8.80 and δ 4.56 ppm as doublets ($J = 7$ Hz, 5 Hz, respectively), and again containing resonances downfield at δ 51.91 and δ 24.67 ppm. The ${}^{31}\text{P}$ NMR spectrum reveals a single resonance at δ 72.71 ppm, similar to the observed ${}^{31}\text{P}$ NMR spectrum of both **2** and **3**. Evans method NMR was used to calculate the solution phase magnetic moment of **4**, which was found at $1.21\mu_{\text{B}}$, uncharacteristically low for the expected magnetic moment for a $5f^3$ ion of Np^{4+} of $3.62\mu_{\text{B}}$.⁵⁴ The ${}^1\text{H}$ NMR spec-

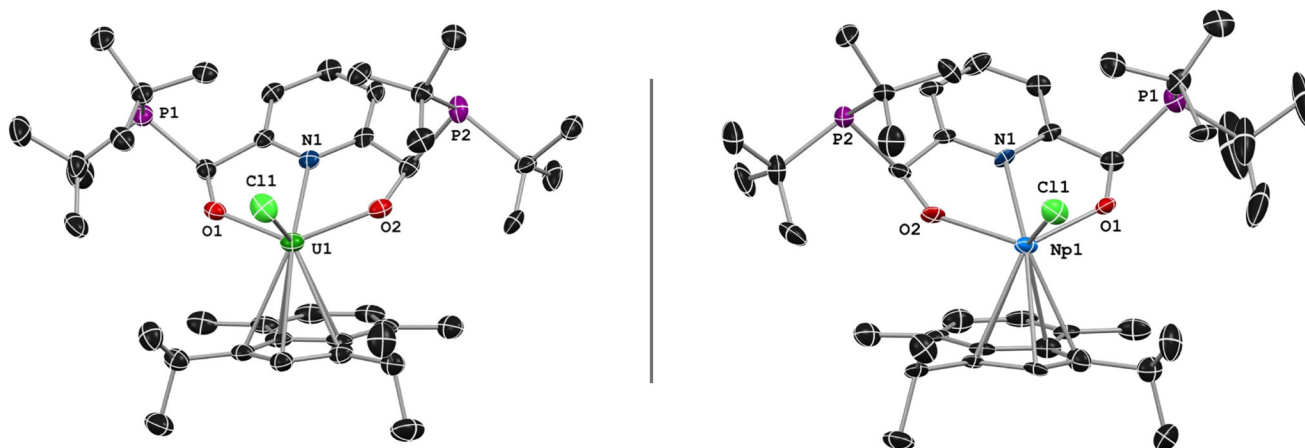


Fig. 2 General solid-state structure of **3** (left) and **4** (right) presented at 50% probability ellipsoids. Hydrogen atoms and pentane solvent molecule have been omitted for clarity.



trum of **4** gave evidence for the presence of diamagnetic indene integrating to roughly 40% as compared to **4**, likely causing the significantly low magnetic value and complicating the ^1H NMR characterization of the complex. This diamagnetic material is attributed to the difficulties in small scale synthesis, sensitivity of the isolated complex (decomposition) or unreacted starting material, though the exact cause remains unknown.

SC-XRD studies of **4** were marred by significant twinning. The most promising crystal mounted during the SC-XRD experiments contained twinned data, which was modelled as a two-component, non-merohedral twin. Complex **4** was solved in the monoclinic $P2_1/n$ space group, with unit cell parameters consistent with those of **2** (Fig. 2). As in **3**, a disordered pentane molecule was present in the unit cell, and one of the $^t\text{Bu}_2\text{P}$ -units was disordered. Many of the bond distances between **3** and **4** were within 3σ of the ESD values, but in certain cases, such as the An-centroid distances, **4** displayed the contraction of the bonds expected for neighbors in the actinide series (Table 2). Despite the technical challenges in the synthesis and characterization of **4**, we were able to confirm the first synthesis of an organometallic, diastereopure neptunium complex.

Electronic structure studies of uranium- and neptunium-ONO complexes

The twinning issue observed with crystals of **4** hampered precise metrical analysis between complexes **3** and **4**. Accordingly, **1–4** were studied computationally, whereby both geometry optimizations and electronic structure studies were performed to assist with the comparison of the differences in geometry, electronic structure, and bonding. Detailed descriptions of the metal-to-supporting ligand bonding and the differences in f-orbital participation between uranium and neptunium was pursued. UV-Vis-NIR and computational geometry optimizations provide further insights for the comparison of the uranium and neptunium homologues.

UV-Vis-NIR studies. The UV-vis-NIR spectra (measured in C_6H_6 : C_6D_6 solutions with ratio 60:40) of **2** and **4** are relatively similar, both exhibiting several sharp, weak features from 500 nm through 1100 nm. These features are likely attributable to Laporte forbidden $5f \rightarrow 5f$ transitions and $5f \rightarrow 6d$ transitions for a Np^{4+} ion.²⁴ The molar absorptivity of the spectral features of **2** are smaller than those observed for **4**, consistent with the observed dark coloration of **4** (Fig. S14 and S17[†]). The spectral features observed for **3** are more pronounced and contain similar features to those of the previously reported spectrum of **1** (Fig. S15[†]).³³ These results suggest minimal change in the electronic structure of the An^{4+} center upon substitution of dtbpy for 1,3-(^tPr)₂-4,7-Me₂-C₉H₃.

Geometry optimizations. The hybrid B3LYP functional with scalar relativistic effects was used to optimize the structures, showing good agreement with respect to the experiment (Tables 2, S1 and S2[†]). It is not surprising to observe the actinide contraction when moving across the series comparing **1** vs. **2**, and **3** vs. **4**. Notably, however, a decrease of the An-

[ONO]²⁻ bond lengths was experimentally and theoretically observed when substituting the dtbpy with the indenide ligand comparing **1** and **3**. This decrease was more pronounced in the An-O bonds with differences of ~ 0.04 Å. An analysis of the steric effects was conducted to delve deeper into this unusual decrease in the same metal-ligand motif (An-[ONO]²⁻) in both complexes. Geometry optimization of a truncated ligand model of di(methyl)phosphine was performed (details in Table S3[†]). The model complexes result in shorter An-[ONO]²⁻ bonds, like those observed in **3**. It suggests that the origin of this decrease from **1** to **3** is in fact because of smaller steric effects between [ONO]²⁻ motif with the indenide ligand compared to dtbpy, rather than an intrinsic electronic effect or chemical bonding. These conclusions also apply to the neptunium systems, **2** and **4**.

Electronic structure studies. The electronic structures of complexes **1–4** were analyzed *via* multi-reference wavefunction methods with a relativistic Hamiltonian, in order to provide an accurate treatment of relativistic and correlation effects (see ESI Computational descriptions[†]). The crystal-field splitting, interelectronic repulsion, and spin-orbit coupling are of comparable magnitude in the actinides, making either the Russell-Saunders (RS) or the j-j coupling models incomplete to characterize these systems. In this context, the intermediate coupling is the most suitable option.⁵⁵ In this formalism, states with the same value of total angular momentum, J, mix, resulting in electronic states that can be described by linear combinations of RS terms.

The U^{4+} and Np^{4+} systems have formal $5f^2$ and $5f^3$ configurations, respectively and according to Hund's rule, they possess ^3H and ^4I ground terms. Under the intermediate coupling scheme, the ground state with spin-orbit coupling (SO-GS) is expected to be dominated by these terms, with additional contributions from some excited terms. This was supported by SO-CAS($n,7$)/SCF/NEVPT2 ($n = 2, 3$ for U^{4+} , Np^{4+} , respectively) calculations, where the ground-state term contributed between ~ 76 and 86% to the total composition of the SO-GS (Tables S4–S7[†]). The substitution of dtbpy with the indenide ligand causes negligible differences in the composition of the SO-GS wavefunction in both the uranium and neptunium systems. For instance, the SO-GS of the uranium systems **1** and **3** respectively have ~ 88 and 84% contribution from the ^3H RS term, and ~ 8 and 7% from the excited ^1G . An extension of the active space was performed by including indenide orbitals in complexes **3** and **4** to evaluate correlation effects between the metal and ligand orbitals (see ESI for Computational descriptions[†]). Similarities were observed in terms of the SO-GS composition (Tables S6 and S7[†]), where excitations involving ligand orbitals only contribute about $\sim 1\%$. Comparable observations were found for the spin-orbit excited states (SO-ES) of all systems **1–4** (Tables S4–S7[†]).

Spectroscopy. TD-DFT, as a natural approach, was first employed to assign the bands observed in the solution UV-Vis-NIR spectra. In general, the region between 300 and 500 nm shows good agreement with the experimental spectra for all complexes with little dependence on the choice of DFT func-



tional, such as GGA (GGA = generalized gradient approximation) and hybrid functionals. However, the low energy region, where the dipole-forbidden $f-f$ transitions are usually observed, is not well reproduced and the results are very sensitive to the DFT functional used (Fig. S3–S6†). Therefore, the multi-reference SO–CASSCF/NEVPT2 methodology was applied to model the Laporte forbidden $f-f$ transitions.

Charge transfer bands and $5f \rightarrow 6d$ transitions. TD-DFT calculations show that the region between 300 and 500 nm is dominated by inter-ligand, charge transfer (CT) and $5f \rightarrow 6d$ bands in all the complexes. The most intense spectroscopic features are ascribed to LMCT and MLCT transitions, which are mixed together making it difficult to assign their categorization by energy ranges (see energy level orbital diagrams in Fig. 3). The $6d$ orbitals are split between ~ 250 – 500 nm showing a significant mixing with ligand orbitals. Therefore, the $5f \rightarrow 6d$ transitions that are observed in this range are not pure and are found overlapping with CT transitions. In the uranium systems **1** and **3**, most of these excitations involve the dtbpy ligand and the $[\text{ONO}]^{2-}$ chelate (Tables S8 and S10†). Similar observations are found in the neptunium complexes (**2** and **4**), although a shift associated to the stabilization of the $5f$ orbitals is found, consistent with a bathochromic shift observed in some of the calculated transitions (Tables S9 and S11†).

$f-f$ transitions. The assignment of the absorption bands was based on the analysis of low-lying excited states calculations (500 and 2000 nm range for complexes **1** and **3**; 500 and 1500 nm range for complexes **2** and **4**). The energy of each

state along with its composition is shown in Tables S4–S7.† For the sake of clarity, each band in the experimental and theoretical spectra was labeled accordingly (Fig. 4). Generally speaking, the substitution of dtbpy by the indenide ligand does not significantly affect the positions of the absorption peaks for both the uranium and neptunium systems. Small differences were observed in terms of the splitting and intensities of some bands and attributed to the differential ligand fields.

Eight well defined absorption bands were predicted between 500 and 2000 nm for the uranium complexes (**1** and **3**, Fig. 4a and c). The first absorption band predicted (peak A) was not observed experimentally, although previous reports on $[\text{UCl}_5(\text{THF})]^-$ have identified a band in similar energy ranges.^{56,57} Due to the limited resolution in the experimental spectrum of complex **1**, the band assignment was performed only for the predicted bands (Fig. 4a). Conversely, a comparison between experimental and theoretical data was possible in complex **3**. Whereas the results are in good agreement with the experimental spectrum, the calculated spectrum appears blue shifted with respect to experiment (Fig. 4c and Fig. S5†). This shift is likely due to an incomplete treatment of the electronic correlation as consequence of a limited active space. Previous reports show comparable spectroscopic features with same assignments for U^{4+} complexes.^{56–58}

The region between ~ 680 and 1600 nm was assigned in terms of major contributions of RS terms to each state for the neptunium systems **2** and **4**, as done above with the uranium

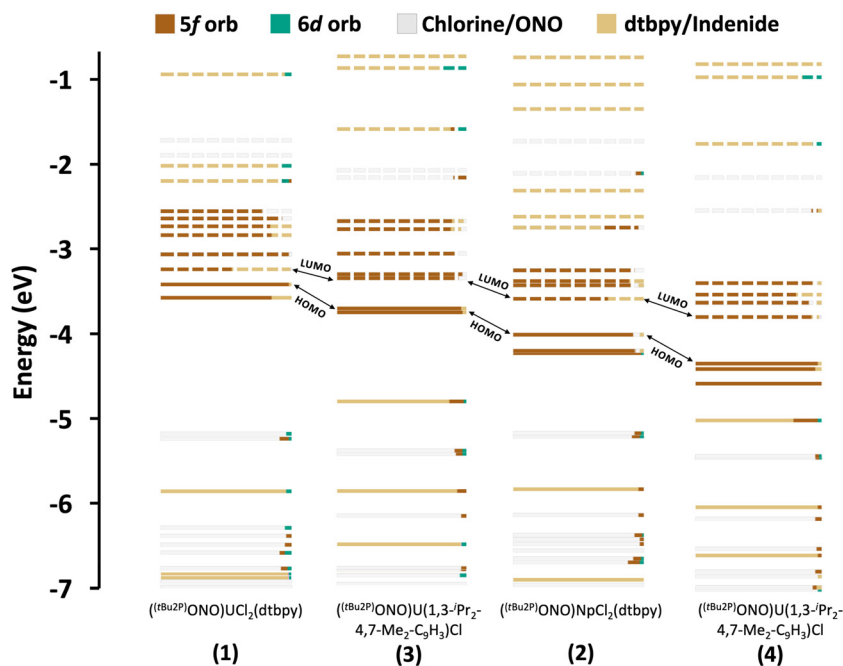


Fig. 3 Energy level diagrams of systems $(t\text{Bu}_2\text{P})\text{ONO})\text{UCl}_2(\text{dtbpy})$ (**1**), $(t\text{Bu}_2\text{P})\text{ONO})\text{NpCl}_2(\text{dtbpy})$ (**2**), $(t\text{Bu}_2\text{P})\text{ONO})\text{U}(1,3-(\text{Pr})_2-4,7-\text{Me}_2-\text{C}_9\text{H}_3)\text{Cl}$ (**3**) and $(t\text{Bu}_2\text{P})\text{ONO})\text{Np}(1,3-(\text{Pr})_2-4,7-\text{Me}_2-\text{C}_9\text{H}_3)\text{Cl}$ (**4**) calculated at ZORA/B3LYP–TZP level of theory. The solid and dashed lines indicate occupied and empty orbitals, respectively. Each orbital is depicted in different colours, depending on its most predominant contributing fragment. Since the percentage of metal participation in the orbitals found in the range of -4 to -7 eV is small, the raw data with all the compositions and energies is available in the ESI.†



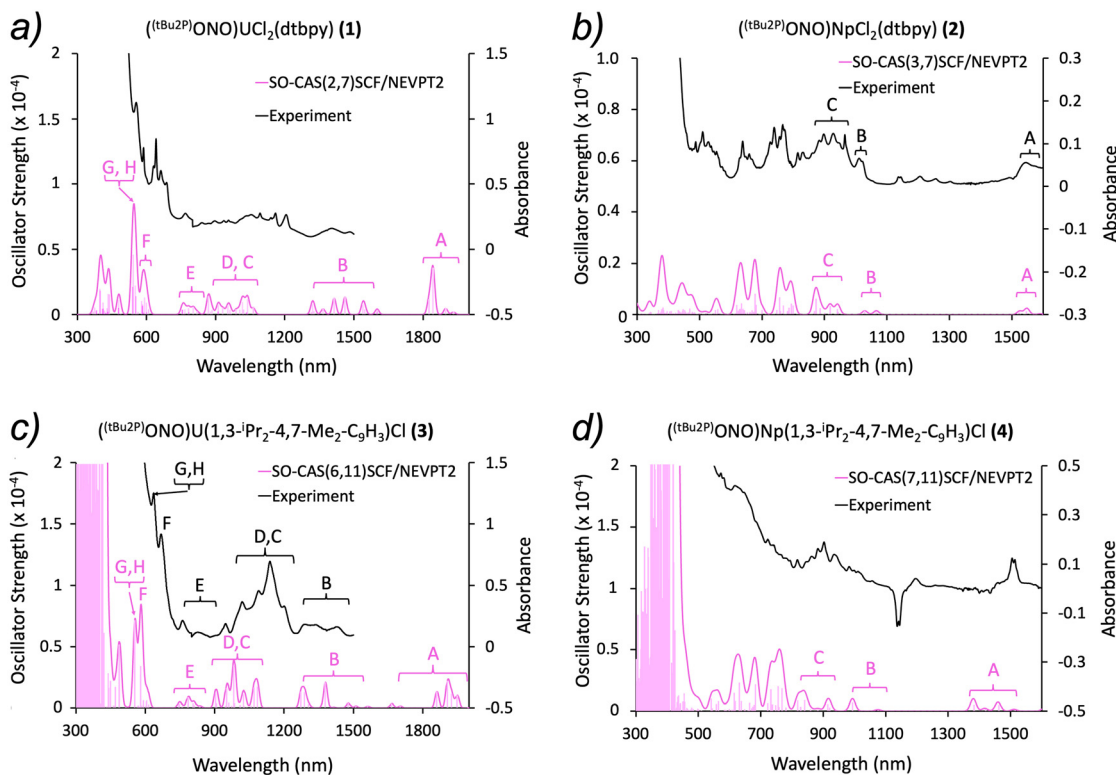


Fig. 4 Experimental (black) and simulated (pink) UV-vis-NIR spectra for complexes 1–4. The vertical bars correspond to vertical transitions calculated at SO-CASSCF/NEVPT2 level of theory. (a) The simulated spectrum for **1** was labelled from A to H to show the different assignments of the bands. (b) The experimental and simulated spectra for **2** were labelled from A to C. The remaining states and respective energies are detailed in terms of *J* in Table S6.† (c) The experimental and simulated spectra for **3** were labelled from A to H to show the different assignments of the bands. (d) The simulated spectra for **4** was labelled from A to C; note the large negative and positive features at ~1130 nm and 1510 nm are known artefacts from the instrument. The remaining states and respective energies are detailed in terms of *J* in Table S7.†

complexes. Theoretical results are in good agreement with the experimental peaks, particularly in **2**, where the spectrum is better resolved (Fig. 4b and d). A continuum of states was observed from ~680 nm and up, which rendered the analysis of its composition difficult. Since all these states appear close in energy to each other, some of them overlap, causing multiple contributions to the same spectroscopic band. Additionally, finding the predominant RS term for a specific state became even more challenging since the contribution of other terms increased. Therefore, the assignment was performed only in terms of *J* in the range of ~500 to 680 nm. Theoretical studies on Np⁴⁺ spectra are scarce, though the prediction and assignment of the spectra in complexes **2** and **4** are in good agreement with reported NpF₄ and [NpCl₆]²⁻ systems.^{59,60}

Bonding. The natural localized molecular orbital (NLMO) representation has proven valuable in analyzing An–ligand interactions^{61–64} and was first employed here to gain a localized chemical bonding picture of complexes 1–4. The goal was to study the effect of coordinating the indenide or the dtbpy ligand on the *f*-orbital participation of the metal. B3LYP and PBE densities were used as input for these calculations, both functionals leading to similar descriptions. The results for the B3LYP functional are presented in Fig. 5, while the results using the PBE functional can be found in Tables S12–S15.†

In a second step, the extended transition state (ETS) method for the energy decomposition analysis (EDA) combined with the natural orbitals for chemical valence (NOCV) theory was employed. The EDA allows to quantify contributions to the total energy in terms of electrostatic and orbital interactions along with the Pauli repulsion. Two fragments, corresponding to the metal (charge = 4+) and surrounding ligands (charge = 4–), were considered in this analysis. Thus, each component of the total interaction energy can be quantified between the defined fragments, and the orbital interactions can be further explored through the deformation energies obtained from the NOCV analysis.

Overall, the NLMO analysis describes the An–ligand interaction as predominantly ionic with a non-negligible degree of covalency (see below). It was also noticed that the An–indenide interaction displays a greater overlap compared to the An–dtbpy species. These results are consistent with conclusions coming from the EDA (Table 3). Here, it was observed that the dominance of the electrostatic component is common for all the systems, as expected considering the charged nature of the fragments employed in this calculation. When comparing uranium and neptunium analogs (**1** vs. **2** and **3** vs. **4**), similarities were observed in terms of the contribution of the electrostatic and orbital components. The most prominent



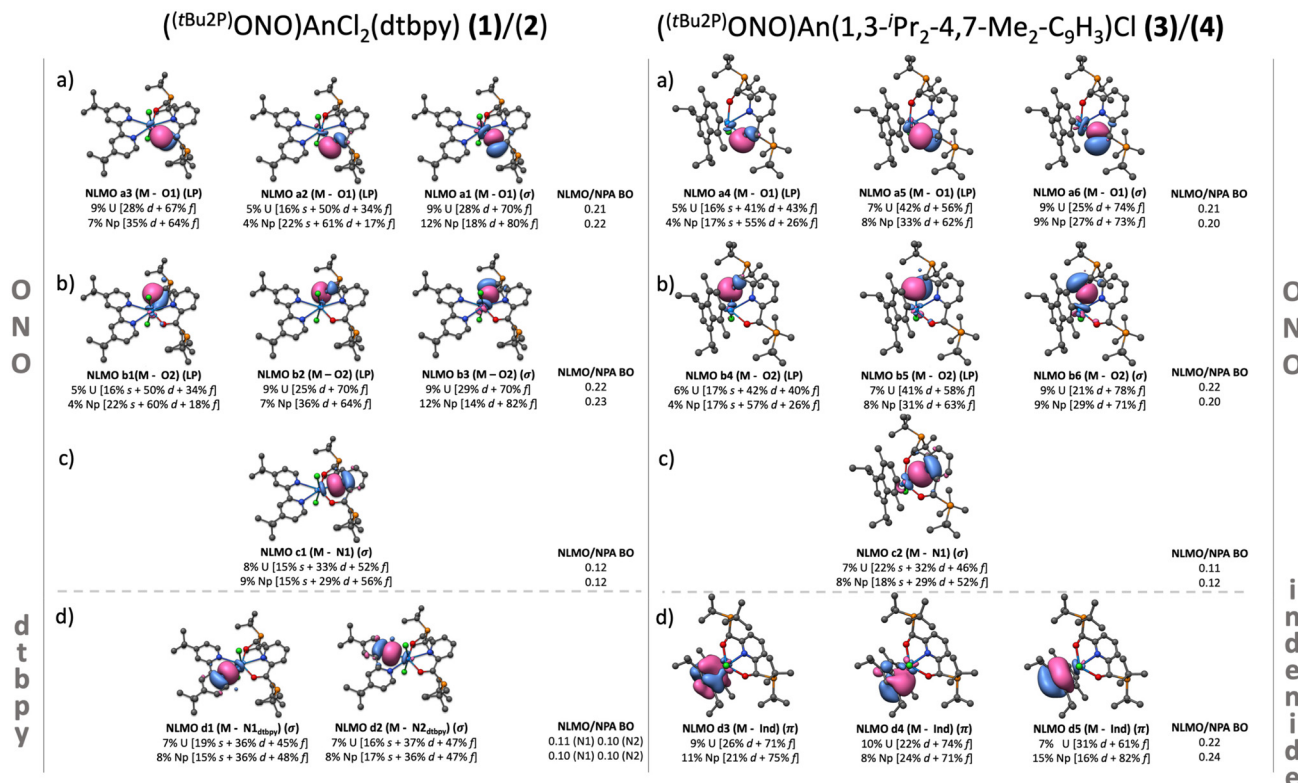


Fig. 5 Left: NLMO composition of the most important interactions describing the An–ONO (above the gray dashed line) and An–dtbpy (below the gray dashed line) bonding in **1** and **2**. The level of theory employed for these calculations was ZORA–B3LYP/TZP (isovalue = 0.03). Right: NLMO composition of the most important interactions describing the An–ONO (above the gray dashed line) and An–indenide (below the gray dashed line) bonding in **3** and **4**. All NLMOs were plotted using an isovalue = 0.03. Additionally, the type of interaction σ , π , or lone-pair (LP) plus the NLMO/NPA bond orders (NLMO/NPA BO) for the M–O, M–N and M–C bonds are specified. In the case of the indenide ligand, the M–C bond-orders correspond to an average of all carbons in the 5-member ring. The plots in this figure correspond to systems **1** (left) and **3** (right).

Table 3 Energy decomposition analysis for the M–L interactions in complexes **1–4**. For the sake of simplicity, values in parentheses correspond to individual contributions to the total stabilizing interaction of the electrostatic and orbital components. All values are in kcal mol^{−1}

Energy component	1	2	3	4
Electrostatic	−1911.1 (71%)	−1913.1 (71%)	−1696.2 (66%)	−1702.9 (65%)
Pauli	388.2	376.6	368.8	365.7
Steric	−1522.9	−1536.5	−1327.4	−1335.2
Orbital	−777.9 (29%)	−789.1 (29%)	−885.8 (34%)	−899.8 (35%)
Total	−2300.8	−2325.5	−2213.1	−2230.9

differences emerge from changes in the ligand environment, where the highest orbital contributions to the total energy were found for the indenide systems **3** and **4** (~6% more than in the dtbpy compounds).

Since the general picture of the bonding shows complementary conclusions between both methods, a more detailed discussion of the metal–ligand interactions is presented below. To facilitate this discussion, an analysis based on the orbital composition as well as the associated stabilization energies

was performed. Because the conclusions reached from both the NOCV and the perturbative analysis analyses are similar, the results of the former are discussed while the ones coming from the perturbative analysis in NBO basis can be found in Tables S20–S23.† To simplify the comparison across the four systems (**1–4**), the discussion is organized with a focus on each type of metal–ligand interaction.

An–chlorine bonding. The An–Cl interaction is represented through four NLMOs, three of which display Cl → An donation. One NLMO describes a σ (An–Cl) covalent bond, while the other three correspond to electron lone-pairs (LP) (Tables S12–S15†). This σ bond is the strongest ligand–metal donation for all the systems, judging by its bond order (0.29–0.30) and orbital composition. In complexes **1** and **2**, the average metal contribution to the An–Cl bond is ~10%, primarily by 5f orbitals, while the rest comes from the chlorine atom 3p orbitals. The metal contribution to the bond stayed constant in complexes **3** and **4**, however the average 5f contribution decreased ~6% with respect to **1** and **2**, which was mainly counterbalanced by a small 7s and 6d increment. These observations are consistent with the analysis of the deformation densities, where σ donations involving the Cl(3p) and An(5f/6d) orbitals were found to provide the most pro-



nounced stabilization energies (between -48 and -52 kcal mol $^{-1}$) (Fig. S7, a1 and b1 \dagger).

An-[ONO] $^{2-}$ bonding. In terms of individual interactions, each An-O bond is represented by three NLMOs displaying O \rightarrow An donation (Fig. 5, 5a and 5b). One corresponds to a distorted σ (An-O) bond, while the other two correspond to lone pairs. Conversely, the An-N interaction was represented by one NLMO displaying σ N \rightarrow An donation (NLMO 6a, Fig. 5). These σ bonds are the second strongest ligand-metal donations among all the systems. The average metal contribution to the bond is between 7–9% while the orbital contributions showed a negligible change between 1 and 3, and 2 and 4. The deformation densities show stabilization energies in the range of -49 and -28 kcal mol $^{-1}$, where the most stabilizing interactions are σ in nature, similar for both 1 and 3, and 2 and 4. These results suggest that the An-[ONO] $^{2-}$ core is not significantly affected by the coordination of the dtbpy or indenide ligand (Fig. S7, a2–a4 and b3–b6 \dagger).

An-dtbpy bonding. Two σ (An-N) bonds (Fig. 5d, left) describe the interactions between the actinide metal and dtbpy, with 7% and 8% metal contribution to the bond for complexes 1 and 2, respectively. A bond order of 0.10 was found for both 1 and 2. Besides these interactions, corresponding to donations from the dtbpy ligand to the An center, an interaction consistent with a small degree of back-donation was found in 1 (Fig. 6). Since no significant differences in bond lengths and bond orders were found between 1 and 2, this back-bonding was attributed to orbital mixing due to a better energy match between the virtual orbitals in the dtbpy ligand and uranium f-orbitals (Fig. 6). From the NOCV analysis, σ donations from the 2p (dtbpy) orbitals were found at lower energies (~ -30 and -21 kcal mol $^{-1}$), compared to those of the other ligands (Fig. S7, a5 and a6 \dagger). Additionally, the back-bonding interaction in 1 was found to provide a small stabilization energy of ~ 2 kcal mol $^{-1}$.

An-indenide bonding. In complexes 3 and 4, the metal is located perpendicular to the plane of the 5-member ring in the

indenide ligand. The metal preference towards this ring over the 6-member ring is attributed to the better symmetry match between its π orbitals and the 5f orbitals of the metal. The metal-ring interaction is *via* three NLMOs displaying donation towards the metal center. These correspond to π -type bonds with significant metal and 5f contributions (Fig. 5d right, Fig. 6). The uranium complex 3 exhibits an average metal contribution of $\sim 9\%$, out of which $\sim 68\%$ corresponds to 5f orbitals. In the neptunium analog, 4, a small 2% increase in the metal contribution was observed. Interestingly, this increase was accompanied by an increment in the 5f instead of 6d contribution, the more common occurrence in actinide-ligand bonding. The increased 5f contribution, along with the calculated bond orders, indicates a differential and more covalent Np-indenide interaction. In terms of the deformation densities obtained from the NOCV analysis, the stabilization energies were found between -57 and -46 kcal mol $^{-1}$, a similar energy range where the An-Cl interactions were also identified (Fig. S7, b1–b3 \dagger), but stronger than the An-dtbpy motif.

In summary, the bonding analysis suggests that the An-ligand interactions are mostly ionic with a minor component of covalency that is consistent with the conclusions obtained from previous reports on other An(IV) compounds.^{65–67} The dtbpy ligand shows a better energy match with respect to the metal orbitals. However, based on orbital composition and bond orders, the metal-dtbpy interactions also display the weakest overlap, consistent with the experimentally observed lability of this ligand. Conversely, the indenide derivative exhibits a greater orbital overlap, which overall suggests stronger interactions. Expanding the discussion to other An-L interactions, it is noteworthy that both An-indenide and An-Cl bonds show similar magnitudes of metal contribution, an interesting result considering that organic ligands are historically considered more covalent than halides when bound to metals. Though, a slight but important difference was observed for these two interactions in terms of the 5f involvement being higher in the An-indenide bond with respect to

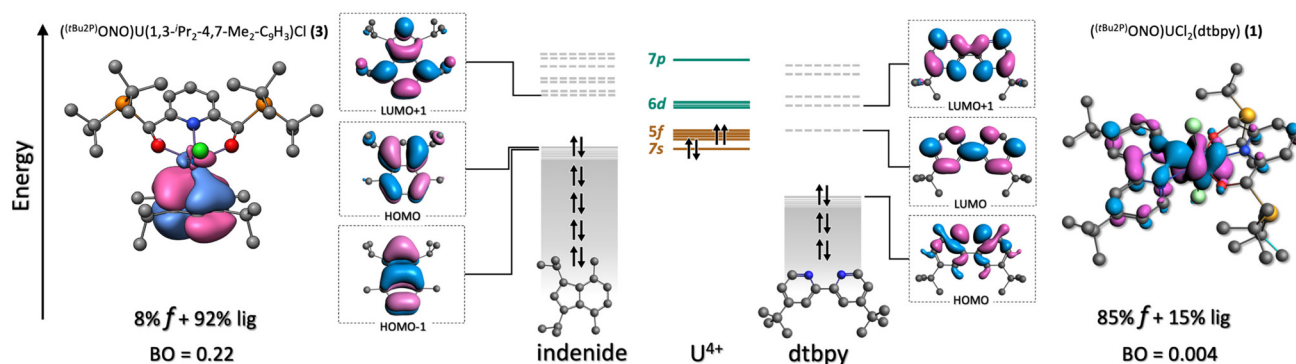


Fig. 6 Qualitative molecular orbital diagram of 1 and 3. The brown lines/blocks indicate occupied orbitals while the green lines correspond to empty orbitals. As observed, the LUMO and LUMO+1 of the dtbpy ligand are close in energy to the 5f manifold. In consequence, an interaction displaying a small degree of back-donation (depicted in the right side) takes place. In the case of 3, the energetics is different and no back-bonding is observed. However, focusing on the HOMO of the indenide ligand, it is noteworthy how the 5-member ring orbital symmetry has a better match with the 5f orbitals than that of the 6-member ring. This interaction forms the dative interaction depicted in the left side. Isovalue = 0.03.



the An–Cl bond. This was compensated by an increase in 7s and 6d contribution in the latter. The differences associated with coordination of the indenide highlights the role of organic ligands in enhancing the 5f participation in chemical bonding, which is already established with cyclooctatetraene (COT) or cyclopentadienyl ligands.^{68,69} This feature emerges from the diversity of symmetry-permitted combinations between these orbitals and the π system of the ligand; by including organic π -based orbitals of appropriate symmetries, the 5f orbitals can better participate in bonding interactions.

Finally, a synergistic combination of wavefunction and DFT approximations has enabled an accurate analysis of the electronic structure, spectroscopy, and bonding of the complexes 1–4. This combination of methods has proven successful when applied to the f-block, yet a careful analysis and interpretation of the results is required. In this case, the different approaches led to complementary conclusions shedding light on the similarities between uranium and neptunium complexes, and differences in f-orbital participation arising from the coordination of dtbpy or indenide ligands.

Conclusions

This report shows the benefits of pursuing template methodology for preferentially forming diastereopure products with uranium and neptunium, increasing the versatility of this method for the small-scale synthesis necessary for neptunium chemistry. The template method employed here is selective for the *meso* diastereoisomer. This is a stark improvement over the *rac*:*meso* mixture of diastereoisomers that form in a 63:37 ratio when the free-ligand synthesis is performed without a metal template. The *meso*-[ONO]²⁻ supported products stabilize the metal center and allow for further chemistry. The use of 1,3-(ⁱPr)₂-4,7-Me₂-C₉H₃ provided homologous organometallic uranium and neptunium complexes for comparison. Not only is this the first diastereoselective synthesis for neptunium, but also the first indenide-coordinated neptunium structure reported, contributing to the small but growing number of established organoneptunium complexes.

Detailed computational exploration of complexes 1–4 provides a complete picture of the electronic structure, spectroscopy, and bonding of these compounds. The geometry optimization process shed light on the importance of the steric effects, concluding that the An-[ONO]²⁻ contraction observed upon coordination of the indenide ligand was a consequence of lower steric hindrance, allowing for tighter metal-ligand coordination. Spectroscopic analysis performed through the CASSCF/NEVPT2 methodology was able to reproduce the UV-Vis-NIR spectra of the systems, where similarities in terms of position of the bands were detected after replacing the dtbpy by the indenide ligand. Differential splitting and intensities were identified in some bands, which was attributed to the intrinsic field produced by these ligands.

Finally, the NLMO analysis revealed similarities in composition and bond orders between the An-[ONO]²⁻ and An-dtbpy

interactions for both uranium and neptunium complexes. The most pronounced differences resulted from the substitution of the dtbpy by the indenide ligand, where an increased metal and 5f orbital contribution associated to the An-indenide interaction was found. In general terms, the bonding analysis suggests that the dtbpy ligand has a better energy match with respect to the metal orbitals, while the indenide ligand possesses a better orbital overlap. This is consistent with experimental observations describing the dtbpy ligand as extremely labile. The ETS-NOCV and second order perturbation theory analysis in NBO basis are consistent with the conclusions discussed above, suggesting a differential coordination for the indenide ligand. Additionally, this work highlights the importance of organic ligands in exposing the 5f orbital participation in An–L bonding.

Rational experimental design targeting homologous actinide complexes and implementation of multidisciplinary investigations resulted in the study of the electronic structure and bonding of diastereopure organoactinide complexes. The template protocol employed warrants further investigation in terms of generating new organometallic complexes of uranium and neptunium, as the specificity for *meso*-complexes is rather rare, and the singular steric profile may provide unique reactivity beyond the regioselectivity observed here. Further comparison of novel homologous complexes of uranium and neptunium based on complexes arising from the use of template protocols may provide further insight into differences in the electronic structure and bonding of these metals.

Author contributions

S.H.C and M.J.B.-L. contributed equally to this work, respectively performing the technical synthetic and computational tasks. A.M.T. conceived the synthetic chemistry and supervised the synthetic effort. E.R.B. and P.Y. supervised the computational effort. A.J.G. assisted with the execution of neptunium experimental work, providing guidance to S.H.C. and A.M.T. throughout. S.S. provided the necessary organic ligands for the organometallic chemistry. M.L.T. performed supplementary uranium chemistry, while J.T.M. assisted with data analysis and resolving twinned SC-XRD data. All authors contributed to manuscript preparation.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was conceived and executed at Los Alamos National Laboratory. Los Alamos National Laboratory is operated by Triad National Security, LLC, for the National Nuclear Security Administration of the U.S. Department of Energy (contract no. 89233218CNA000001). A.M.T., A.J.G., P.Y., E.R.B., S.H.C., M.L.T.,



and M.J.B.-L. acknowledge the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, Heavy Element Chemistry program (2020LANLE372, DE-AC52-06NA25396). S.S. was supported by an LDRD Early Career Research project awarded to A.M.T (20190570ECR). We are grateful for postdoctoral support provided in part by the Glenn T. Seaborg Institute for S.H.C. and M.J.B.-L.

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