Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2024, 15, 12754

dll publication charges for this article have been paid for by the Royal Society of Chemistry

Received 15th March 2024 Accepted 22nd June 2024

DOI: 10.1039/d4sc01767f

rsc.li/chemical-science

$PuCl_3\{CoCp[OP(OEt)_2]_3\}$: transuranic elements entering the field of heterometallic molecular chemistry†

Thomas E. Shaw, ‡a Zachary R. Jones, ‡a Sara L. Adelman, [b*a] Nickolas H. Anderson, [b*a] Eric G. Bowes, [b*a] Eric D. Bauer, *a David Dan, *a Jan Klouda, [b*a] Karah E. Knope, *b Stosh A. Kozimor, [b*a] Molly M. MacInnes, *a Veronika Mocko, *a Francisca R. Rocha, *a Harrison D. Root, *a Benjamin W. Stein, [b*a] Joe D. Thompson *a and Jennifer N. Wacker [b*a]

Recent advances enabled the discovery of heterometallic molecules for many metals: main group, d-block, lanthanides, and some actinides (U, Th). These complexes have at least two different metals joined by bridging ligands or by direct metal—metal bonding interactions. They are attractive because they can enable chemical cooperativity between metals from different parts of the periodic table. Some heterometallics provide access to unique reactivity and others exhibit physical properties that cannot be accessed by homometallic species. We envisioned that transuranic heterometallics might similarly enable new transuranic chemistry, though synthetic routes to such compounds have yet to be developed. Reported here is the first synthesis of a molecular transuranic complex that contains plutonium (Pu) and cobalt (Co). Our analyses of PuCl₃{CoCp[OP(OEt)₂]₃} showed Pu(IV) and Co(III) were present and suggested that the Pu(IV) oxidation state was stabilized by the electron donating phosphite ligands. This synthetic method – and the demonstration that Pu(IV) can be stabilized in a heterobimetallic molecular setting – provides a foundation for further exploration of transuranic multimetallic chemistry.

Introduction

Recent synthetic advances have enabled discovery of heterometallic molecules for many elements including those from the main, d-block, lanthanide, and even some actinide (uranium and thorium) elements. These heterometallic compounds are defined as having two – or more – different metals within one discrete molecule, where the metals are fixed in proximity through bridging ligands or direct metal–metal bonds. Combining properties from elements on opposite sides of the periodic table within one molecule unlocks potential for new cooperative chemistry that would otherwise be inaccessible to homometallic species. The application space for heterometallics is now quite broad and ranges from catalysis to conductive materials, photophysical applications, and molecular magnets. Alternative properties and molecular magnets.

that preparation of plutonium (Pu) heterometallic molecules could be a gateway for new and interesting chemistry, provided appropriate synthetic pathways can be developed. This notion was fueled by the isolation of heterometallic extended solids reported from the transuranic solid-state and intermetallic communities. ^{32–35} Perhaps the most notable result from those synthetic efforts was the discovery of PuCoGa₅. This intermetallic enabled plutonium superconductivity to be observed for the first time and advanced fundamental understanding of plutonium's electronic structure and physical properties. ³²

Herein, we report the isolation of the first discrete heterobimetallic molecule for plutonium, namely plutonium(IV) tri-(cyclopentadienyl)tris(diethylphosphito)cobalt(III), chloride PuCl₃{CoCp[OP(OEt)₂]₃}. This compound was prepared by reacting an acidic Pu4+(aq) residue with Na{CoCp[OP(OEt)2]3}, affectionately referred to in the literature as the Kläui ligand.36-46 Our synthetic methodology was attractive because it could be carried out without exclusion of air and moisture. It also enabled large amounts (>300 mg) of PuCl₃{CoCp[OP(OEt)₂]₃} to be isolated in modest crystalline yield (70%) and in high purity. Although many plutonium molecules have been characterized using a wide variety of techniques, limited access to plutonium and the substantial infrastructure needed to safety handle large quantities of plutonium often limits the extent to which a given plutonium compound can be studied. Access to 300 mg of

^aLos Alamos National Laboratory (LANL), P. O. Box 1663, Los Alamos, New Mexico, 87545, USA. E-mail: sadelman@lanl.gov; stosh@lanl.gov

^bDepartment of Chemistry, Georgetown University, 37th and O Streets NW, Washington, D.C. 20057, USA. E-mail: karah.knope@georgetown.edu

[†] Electronic supplementary information (ESI) available: Structural metrics and additional ¹H NMR spectra. CCDC 2323718. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d4sc01767f

[‡] Contributed equally and are joint first authors.

Edge Article Chemical Science

PuCl₃{CoCp[OP(OEt)₂]₃} facilitated rigorous characterization by single crystal X-ray diffraction, nuclear magnetic resonance (NMR) spectroscopy, variable temperature magnetometry, cyclic voltammetry, and optical spectroscopy. The results indicated that PuCl₃{CoCp[OP(OEt)₂]₃} contained discrete Pu(IV) and Co(III) metal cations within the same molecule. These data also showcased that Pu(IV) was stabilized by the electron donating properties of the bridging phosphite ligands. It is our hope that the discovery of PuCl₃{CoCp[OP(OEt)₂]₃} will enable synthetic access to higher order multi-metallic transuranic molecules (*e.g.*, tri-, tetra-, penta-metallic, and beyond) and potentially inspire others to begin studying heterometallic transuranic molecular chemistry.

Results and discussion

Synthesis

The plutonium used in this study was recovered from residues generated by previous research campaigns using a published Purecycling procedure. 47 This process generated an oxidation state pure and chemically pure stock-solution of Pu⁴⁺(aq) in HCl(aq) (6 M). Heating an aliquot from that Pu⁴⁺(aq) stock-solution on a hot block under a stream of filtered air afforded an acidic $Pu^{4+}_{(aq)}$ residue used to prepare $PuCl_3\{CoCp[OP(OEt)_2]_3\}$. The acidic nature of the Pu4+(aq) residue raised concern for further reactivity, e.g. would residual HCl(aq) decompose the CoCp[OP(OEt)2]31- fragment. To alleviate these types of concerns, numerous scoping studies were conducted that targeted previously reported MCl₃{CoCp[OP(OEt)₂]₃} (M = Ce, U) complexes using (NH₄)₂Ce(NO₃)₆ and UCl₄ surrogate starting materials dissolved in $HCl_{(aq)}$. ⁴⁸⁻⁵⁰ These studies suggested that the Na{CoCp[OP(OEt)2]3} reagent was tolerant to the acidic conditions associated with the $Pu^{4+}_{(aq)}$ acidic residue and enabled us to optimize experimental conditions for crystallization of the heterometallic product. Moving forward, we established a robust protocol that reproducibly provided single crystals of PuCl₃{CoCp[OP(OEt)₂]₃} on a small scale (3 mg scale of Pu). Subsequent efforts demonstrated that the synthesis could be scaled-up 100-fold, generating >300 mg of PuCl₃{CoCp[OP(OEt)₂]₃} in 70% crystalline yield.

The $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ heterobimetallic complex was prepared by a salt metathesis reaction that occurred when the acidic $Pu^{4+}_{(aq)}$ residue was reacted with crystalline $Na\{CoCp[OP(OEt)_2]_3\}$ in THF (Scheme 1). In this reaction,

a chloride ligand associated with the $Pu^{4+}_{(aq)}$ residue was displaced by the $CoCp[OP(OEt)_2]_3^{1-}$ monoanion, which consequently outcompeted neutral THF and H_2O solvents for complexation of Pu(rv). During the synthesis the $Pu^{4+}_{(aq)}$ residue did not completely dissolve in THF. It formed a suspension instead. Only after combining that orange slurry with the THF solution of $Na\{CoCp[OP(OEt)_2]_3\}$ did homogeneity occur. Stirring the resulting solution overnight generated a brown solution and a white precipitate. That solid – presumedly NaCl – was removed in two steps. First, the mixture was filtered through a Celite column and the solvent was removed. Second, dichloromethane was added to the resulting residue and that mixture was filtered through another Celite column. The title complex was then isolated as a dark green powder following the removal of the solvent.

Structure

Green, rectangular prisms of PuCl₃{CoCp[OP(OEt)₂]₃} suitable for X-ray diffraction were reproducibly obtained from dichloromethane solutions layered with a toluene counter solvent in a 1:1 ratio. Data were collected at room temperature to avoid icing imposed from the tertiary containment method required for safe handling of radioactive single crystal samples.47 Despite having significantly disordered chloride and ethoxy functional groups, the room temperature single crystal X-ray diffraction data from PuCl₃{CoCp[OP(OEt)₂]₃} were modeled in the monoclinic $P2_1/n$ space group (see Fig. 1 and ESI†). Within this heterobimetallic compound were the duo of Pu(IV) and Co(III) cations bridged by three phosphite ligands. The inner coordination sphere around Pu(IV) was comprised of three chloride ligands and three oxygen atoms from the bridging phosphites. These chlorine and oxygen atoms were arranged in a distorted octahedron with approximate C_{3v} symmetry around Pu(IV). With respect to Co(III), the structure was best described using the "three-legged piano stool" analogy often invoked to describe molecules with one cyclopentadienide and three monodentate ligands.

Defining metrics from the $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ structure were the $Co\cdots Pu$, Pu-Cl, $Pu-O_{phosphite}$, $Co-P_{phosphite}$, and $Co-C_{Cp}$ bond distances. Comparing these bond distances with those reported previously from $MCl_3\{CoCp[OP(OEt)_2]_3\}$ $\{M = Ce, Ti, Tc\}^{48,51,52}$ showed regularity and predictability when the six-coordinate metal ionic radii of the M(IV) cations

Scheme 1 Reaction that generated PuCl₃{CoCp[OP(OEt)₂]₃}.

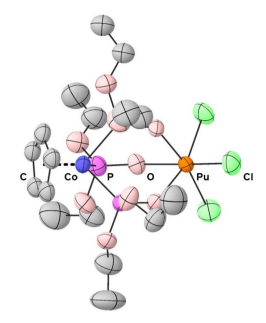


Fig. 1 Thermal ellipsoid plot (30% probability) generated from the single crystal X-ray diffraction data from $PuCl_3\{CoCp[OP(OEt)_2]_3\}$. Hydrogen atoms have been omitted.

were accounted for (Table 1). For example, a Co···M distance of 3.3 Å was obtained for each complex when the six-coordinate metal ionic radii were subtracted from the Co···M distances. This value of 3.3 Å was also similar to the

ionic radius corrected Co···U distance in the structurallyrelated UCl₃(THF){CoCp[OP(OEt)₂]₃} compound. Comparison with the later compound, UCl₃(THF){CoCp[OP(OEt)₂]₃}, further indicated that increasing the M(IV) coordination number by adding a neutral ligand had no impact on the Co···M distance. 49 All of these Co···M distances were far too long than what would be anticipated for Co-M bonds. As testament, the Co···Pu distance in PuCl₃{CoCp[OP(OEt)₂]₃} was over 1 Å longer than the Co-U bond distance observed by Bart, Thomas, and coworkers in their related UX[CoI(Ph2PNiPr)]3 (X = Ph₂PNiPr, I) structures.⁵³ Other structural metrics worth noting were the three equivalent Pu-O_{phosphite} bond distances that ranged from 2.216(5) to 2.231(4) Å and the three equivalent Pu-Cl bond distances that ranged from 2.567(7) to 2.576(5) Å. These Pu-O_{phosphite} and Pu-Cl distances averaged 2.222 ± 0.008 Å and 2.572 ± 0.005 Å, respectively. Uncertainty for all average bond distances have been reported as the standard deviation from the mean at 1σ and denoted by the " \pm " symbol.

The three Co–P_{phosphite} distances ranged from 2.160(2) to 2.166(18) Å, averaged 2.162 \pm 0.003 Å, and were equivalent within the measurement uncertainty. These distances were slightly shorter than the Co–P_{phosphite} distances in the Na{CoCp[OP(OEt)₂]₃} precursor: 2.187(2) and 2.190(2) Å. 54 They were also equivalent to the average Co–P_{phosphite} distances previously reported in the other MCl₃{CoCp[OP(OEt)₂]₃} complexes (Table 1). The five Co–C_{Cp} distances in PuCl₃{CoCp[OP(OEt)₂]₃} ranged from 2.064(12) to 2.093(12) Å and had a Co–C_{centroid} distance equal to 1.699 Å. These distances also matched the Co–C_{Cp} distances reported

Table 1 Solid-state structural metrics from $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ compared with other relevant solid-state structures from $(Me_4N)_2$ $PuCl_6$, 55 $MCl_3\{CoCp[OP(OEt)_2]_3\}$, where M = Ce, 48 Ti, 51 Tc, 52 and $UCl_3(THF)\{CoCp[OP(OEt)_2]_3\}$. 49 Only one set of disordered chlorides from $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ are reported in this table

| Bond | Pu(ɪv) this work (Å) | $PuCl_6^{2-}$ (Å) | Ce(IV) (Å) | U(IV) (Å) | Ti(rv) (Å) | Tc(IV) (Å) |
|---|----------------------|-------------------|------------|--------------|------------|------------------------|
| M-O(1) | 2.218(4) | _ | 2.226(3) | 2.2358^{b} | 1.956(6) | 2.035(2) |
| M-O(2) | 2.216(5) | _ | 2.233(3) | 2.2607^{b} | 1.962(7) | 2.013(2) 2.027(2) |
| M-O(3) | 2.231(4) | _ | 2.244(3) | 2.2607^{b} | 1.979(6) | 2.028(2) 2.026(2) |
| М⋯Со | 4.1958(9) | | 4.2044(7) | 4.289(5) | 3.973(2) | 2.016(2) 3.9269(5) |
| M-O _{mean} ^a | 2.222(8) | _ | 2.234(9) | 2.25(1) | 1.97(1) | 3.9518(5) 2.029(4) |
| M-O _{mean} ^a corrected | 1.362(8) | _ | 1.365(9) | 1.36(1) | 1.36(1) | 2.019(8) 1.378(4) |
| M-Cl(1) | 2.567(7) | 2.5757 | 2.5749(10) | 2.6480^b | 2.305(3) | 1.374(8) 2.3036(7) |
| M-Cl(2) | 2.568(4) 2.575(5) | 2.5935 | 2.5903(11) | 2.6679^{b} | 2.304(3) | 2.3171(7) 2.3098(7) |
| M-Cl(3) | 2.572(7) 2.569(6) | 2.5943 | 2.5840(12) | 2.6679^{b} | 2.322(3) | 2.3105(7) 2.3136(7) |
| M-Cl _{mean} ^a | 2.568(5) 2.571(4) | 2.59(1) | 2.583(8) | 2.66(1) | 2.31(1) | 2.3290(7) 2.309(5) |
| | 2.570(2) | | . , | | | 2.319(9) |
| M-Cl _{mean} ^a corrected | 1.711(4) 1.710(2) | 1.734(7) | 1.713(8) | 1.77(1) | 1.71(1) | 1.664(5) 1.674(9) |
| | | | | | | |

^a The uncertainty for average distances were reported as the standard deviation of the mean at 1σ . ^b Uncertainty for these uranium distances were difficult to extract from the reported primary literature.

Edge Article Chemical Science

previously in the $MCl_3\{CoCp[OP(OEt)_2]_3\}$ complexes shown in Table 1. Regularities in the Pu-ligand and Co-ligand bond distances within $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ and the other reported $MCl_3\{CoCp[OP(OEt)_2]_3\}$ complexes highlighted ubiquitous steric saturation and stability within this heterobimetallic framework for Co(III), Pu(IV), and the other Ce(IV), Ti(IV), and Tc(IV) cations.

Nuclear magnetic resonance spectroscopy (NMR)

Solution-phase ¹H and ³¹P NMR data from PuCl₃{CoCp[OP(OEt)₂]₃} and Na{CoCp[OP(OEt)₂]₃} were provided in Fig. 2 and in the ESI.† These data indicated that PuCl₃{CoCp[OP(OEt)₂]₃} was stable when dissolved in deuterated chloroform. In contrast to the diamagnetic Na{CoCp[OP(OEt)₂]₃} precursor, spectra from PuCl₃{CoCp[OP(OEt)₂]₃} exhibited a paramagnetic contact shift associated with the Pu(IV) cation (Fig. 2). For instance, the ¹H NMR spectrum from Na{CoCp[OP(OEt)₂]₃} contained three resonances: a singlet from the Cp¹⁻ ligand at 5.18 ppm, a broad multiplet from the methylene protons on the phosphite functional group at 4.09 ppm, and a triplet corresponding to the methyl protons at 1.27 ppm. In the paramagnetic product, the Cp¹⁻ singlet shifted downfield by 1.53 ppm to 6.71 ppm. It also shifted the methyl triplet upfield to 0.73 ppm and the methylene resonance downfield to 3.03 ppm. Another defining feature for the PuCl₃{CoCp[OP(OEt)₂]₃} spectrum was that the methylene resonance was split into a complicated multiplet, which we described as a doublet of pentets (I = 92.0, 7.8 Hz). These methylene protons were diasteriotopic and experienced geminal proton-proton coupling, vicinal proton-proton coupling, and coupling to adjacent phosphorous atoms. Consistent with this interpretation were the relative peak intensities (5 for Cp1-, 12 for the methylene, and 18 for the methyl) and similarity between chemical shift values from PuCl₃{CoCp[OP(OEt)₂]₃} and the previously reported uranium analogue, UCl₃{CoCp[OP(OEt)₂]₃}. The spectrum from

UCl₃{CoCp[OP(OEt)₂]₃} had Cp¹⁻, methylene, and methyl resonances at 6.45, 3.04, and 0.50 ppm, respectively.⁵⁰

The ³¹P NMR spectrum from PuCl₃{CoCp[OP(OEt)₂]₃} was impacted by the paramagnetic Pu(IV) cation, as well. It displayed a single phosphite resonance at 24.5 ppm. This value was substantially shifted upfield from the 114.61 ppm resonance associated with the diamagnetic Na{CoCp[OP(OEt)₂]₃} precursor. The ³¹P phosphite chemical shift from PuCl₃{CoCp[OP(Oet)₂]₃} was quite different than that observed for the Ce(IV) (diamagnetic) and U(IV) (paramagnetic) analogues, which displayed phosphite resonances at 124.2 and 191.4 ppm, respectively. ^{48,50} The ³¹P NMR data also suggested that there was only one phosphorous containing compound in solution, namely PuCl₃{CoCp[OP(OEt)₂]₃}. Evaluating this data alongside the ¹H NMR data provided no evidence of CoCp[OP(OEt)₂]₃¹⁻ decomposition or ligand redistribution and highlighted the solution-phase stability of PuCl₃{CoCp[OP(OEt)₂]₃}.

Magnetometry

Variable-temperature magnetic susceptibility measurements were made on PuCl₃{CoCp[OP(OEt)₂]₃} to determine to the 5f-and 3d-electron occupancies for the Pu and Co metals. Data collected from the paramagnetic PuCl₃{CoCp[OP(OEt)₂]₃} complex was compared with data from a known Pu(IV) standard, (Me₄N)₂PuCl₆ (Fig. 3).⁵⁵ It is often difficult to collect reproducible magnetic data on radioactive samples because magnetic measurements require extreme precision regarding sample mass. Unfortunately, many actinide synthetic campaigns are carried out on the 1 to 10 mg scale and it is hard to accurately transfer those small quantities into holders that provide radiological containment for magnetic measurements. We reduced the impact from this variable by developing synthetic methods and sample handling procedures that were compatible

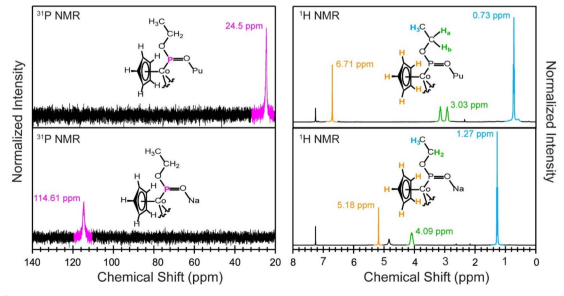


Fig. 2 (Left) 31 P NMR spectra from PuCl₃(CoCp[OP(OEt)₂]₃) (top) and Na{CoCp[OP(OEt)₂]₃} (bottom) dissolved in CDCl₃. (Right) 1 H NMR spectra from PuCl₃(CoCp[OP(OEt)₂]₃) (top) and Na{CoCp[OP(OEt)₂]₃) (bottom) dissolved in CDCl₃. The molecular fragments illustrate our interpretation of the NMR data (methyl – blue; methylene – green; cyclopentadienyl – orange; phosphorus – pink).

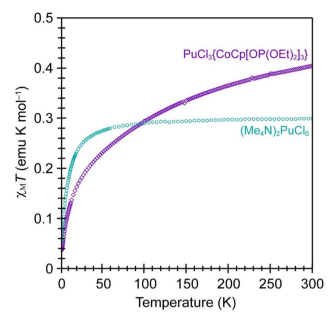


Fig. 3 Variable temperature magnetic susceptibility in a 5 T field from $PuCl_{\pi}\{CoCp[OP(OEt)_{2}]_{\pi}\}$ (purple trace) and $(Me_{\alpha}N)_{\alpha}PuCl_{\pi}$ (teal trace).

with large (hundreds of milligrams) crystalline sample sizes. As such, the masses for the magnetometry samples used for both compounds shown in Fig. 3 were determined with confidence and concerns regarding errors from mass dependence of the measurements were low.

Data from PuCl₃{CoCp[OP(OEt)₂]₃} were quite similar to those from $PuCl_6^{2-}$. Both compounds exhibited a drop in $\chi_M T$ as the temperature was lowered, decreasing to approximately $0.04 \text{ emu K mol}^{-1}$ at low temperature (near 5 K). These data suggested that the number of unpaired 5f-electrons in PuCl₃{CoCp[OP(OEt)₂]₃} was identical to that in (Me₄N)₂PuCl₆, both having the 5f4 electron configuration associated with Pu(IV). Note, that $\chi_M T$ approached an approximate value of 0.4 emu K mol⁻¹ at high temperature for $PuCl_3\{CoCp[OP(OEt)_2]_3\}$. This value was similar to the high-temperature $\chi_M T$ data obtained from (Me₄N)₂PuCl₆, which saturated at 0.3 emu K mol⁻¹. These values were substantially suppressed from what would be expected for a free Pu(IV) cation (as expected), where $\chi_{\mathbf{M}}T$ would saturate at 0.9 emu K mol⁻¹ (the Currie constant) and the effective magnetic moment (μ_{eff}) would equal 2.68 μ_{B} . Like $(Me_4N)_2PuCl_{6,55}$ the reduced Currie constant (saturated χ_MT) in PuCl₃{CoCp[OP(OEt)₂]₃} was most naturally explained by ligand field splitting of the 5f4 multiplet. Overall, these comparisons suggested that PuCl₃{CoCp[OP(OEt)₂]₃} contained diamagnetic Co(III) and paramagnetic Pu(IV).

Optical spectroscopy (UV-vis-NIR spectroscopy)

The single crystal UV-vis-NIR absorption spectrum from $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ validated conclusions based on the magnetometry data and indicated that the compound contained Pu(IV) and Co(III) (Fig. 4). Confidence in the oxidation state assignments came through comparisons with UV-vis data we collected on the $Na\{CoCp[OP(OEt)_2]_3\}$ precursor (see ESI†)

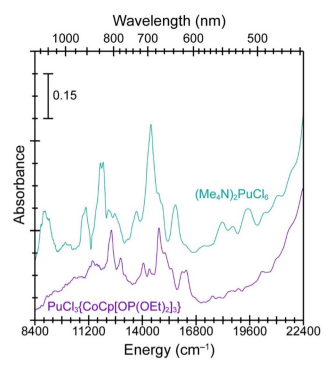


Fig. 4 The solid-state UV-vis-NIR absorption spectra from $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ (purple trace, bottom) vs. $(Me_4N)_2PuCl_6$ (teal trace, top).

and the aforementioned Pu(IV) standard, $(Me_4N)_2PuCl_6$ (Fig. 4). The spectrum from $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ was similar to $PuCl_6^{2^-}$ in that both data sets contained absorbance features characteristic of Pu(IV), most notably between 10 000 and 16 000 cm⁻¹ (600 to 1000 nm). To Coxidation state purity for the $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ sample was clear, as the UV-vis spectrum displayed no evidence of lower [Pu(III)] or higher $(PuO_2^{1^+}; n=1,2)$ oxidation state plutonium species. We inferred that $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ must have Co(III) to maintain charge neutrality, which was also consistent with the magnetometry analyses described above.

Comparing single crystal UV-vis data from (Me₄N)₂PuCl₆ with $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ provided an experimental glimpse into how the Pu(IV) ligand field was impacted by changing from Pu-Cl to Pu-Ophosphite bonding interactions. For example, moving from a homoleptic and octahedral chloride ligand environment in PuCl₆²⁻ to the heteroleptic and pseudooctahedral environment in PuCl₃{CoCp[OP(OEt)₂]₃} had a substantial impact on the Pu(IV) absorbance peak energies. We observed a hypsochromic energy shift for the PuCl₃{CoCp[OP(OEt)₂]₃} absorbance peaks between 10 000 and 16 000 cm⁻¹ (600 to 1000 nm) by approximately 430 cm⁻¹ vs. $PuCl_6^{2-}$. A possible explanation was that substituting three Pu-Cl bonds in PuCl₆²⁻ for three Pu-O_{phosphite} bonds in PuCl₃{CoCp[OP(OEt)₂]₃} increased the valence orbital energy splitting. That change could increase the Pu(IV) 5f \rightarrow 5f and 5f \rightarrow 6d absorption transition energies between 10 000 and 16 000 cm⁻¹ (600 to 1000 nm) and likely resulted from increased Pu ← Ophosphite vs. Pu ← Cl electron donation.

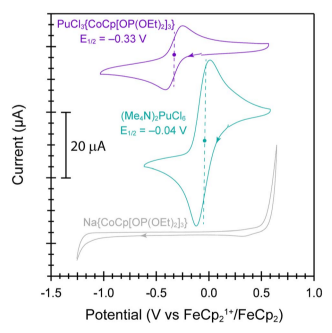
This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

Open Access Article. Published on 15 July 2024. Downloaded on 12/9/2025 12:52:10 PM

Edge Article

Chemical Science

Further evidence that the $Pu \leftarrow O_{phosphite}$ electron donation was larger than Pu ← Cl came from electrochemical analyses of PuCl₃{CoCp[OP(OEt)₂]₃} vs. (Me₄N)₂PuCl₆ (Fig. 5). These data were collected in acetonitrile solutions, with a [Bu₄N][PF₆] supporting electrolyte and using a standard three electrode cell (glassy carbon working, platinum counter, and platinum pseudo reference electrodes). Data were referenced to the FeCp₂¹⁺/FeCp₂ internal standard redox couple. Cyclic voltammograms from PuCl₃{CoCp[OP(OEt)₂]₃} showed a single quasi-reversible wave attributable to the Pu(IV)/Pu(III) redox event with a half-potential (E_1) of -0.33 V and an anodic to cathodic peak separation (ΔE_p) of 0.163 V (scan rate = 0.2 V s⁻¹). The Pu(IV)/Pu(III) redox event from PuCl₃{CoCp[OP(OEt)₂]₃} was substantially shifted negatively (>0.3 V) in comparison to the Pu(IV)/Pu(III) redox event observed from $PuCl_6^{2-}$: $E_{\frac{1}{2}}$ was near zero (-0.04 V) and ΔE_p equaled 0.095 V (scan rate = 0.2 V s^{-1}). These data highlighted that the Pu-Ophosphite bond stabilized electron deficient and high-oxidation state Pu(IV) over electron rich and low-oxidation state Pu(III). Identifying ligand environments that stabilize Pu(IV) in organic solvents is special. Although there exist examples where Pu(IV) remains when complexed in organic media,58-63 many ligand scaffolds induce auto-reduction of Pu(IV) to Pu(III). Examples include - but are not limited to - dithiophosphinate,64 oximes, 65-68 hydroxamic acids, 69-72 hydroxylamines, 73,74 and many



Cyclic voltammograms from acetonitrile solutions that contained $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ (purple trace), $(Me_4N)_2PuCl_6$ (teal trace), or $Na\{CoCp[OP(OEt)_2]_3\}$ (gray trace) dissolved alongside $[Bu_4N][PF_6]$ (0.1 M) in MeCN (2 mL). The dashed vertical lines and solid circles indicate the position of the reduction potential ($E_{\frac{1}{2}}$) for the Pu⁴⁺/ Pu³⁺ redox events. Arrows show the scanning direction. The working electrode was a glassy carbon disk electrode, the counter electrode was a platinum wire, and a separate platinum wire was used as the pseudo-reference electrode. The potentials were referenced internally to the FeCp₂¹⁺/FeCp₂ redox couple. Both voltammograms were collected at a scan rate of 0.200 V s⁻¹

others. In contrast, the data reported here showcased that bridging phosphite ligands in CoCp[OP(OEt)₂]₃¹⁻ formed stable bonds with Pu(IV) in organic solvents, more so than the homoleptic Cl1- ligand environment within PuCl62-. At this time, it remains difficult to predict ligand sets that provide Pu(IV) stability vs. those that incite reduction of Pu(IV). We wonder if the Co(III) is contributing inductively in some way toward that Pu(IV) stability and are currently investigating ways to enhance electronic and magnetic communication pathways between the two metals in this system.

Outlook

We report the synthesis and characterization of the first heterobimetallic transuranic molecule. Large quantities of PuCl₃{CoCp[OP(OEt)₂]₃} were prepared in modest, crystalline yield (70%) via the salt metathesis reaction between $Na\{CoCp[OP(OEt)_2]_3\}$ and an acidic $Pu^{4+}_{(aq)}$ residue suspended in THF. The synthetic procedure could be carried out under ambient conditions (without the exclusion of air and moisture) and on a large scale (hundreds of milligrams). As a result, appreciable amounts of PuCl₃{CoCp[OP(OEt)₂]₃} were accessible for thorough characterization by X-ray diffraction, NMR, UV-vis-NIR spectroscopy, cyclic voltammetry, and magnetic susceptibility measurements. These data indicated that PuCl₃{CoCp[OP(OEt)₂]₃} possessed two separate metals -Pu(IV) and Co(III) - and that the Pu(IV) oxidation state was stabilized in organic media. We assumed this stability was related to the highly electron donating phosphite ligands, but could not disregard the possibility a stabilizing inductive effect imparted by the Co(III) cation. Future efforts are focused on better understanding the electronic structure of $PuCl_3\{CoCp[OP(OEt)_2]_3\}$ and identifying ways to enhance Pu(IV) and Co(III) electronic and magnetic communication pathways. This PuCl₃{CoCp[OP(OEt)₂]₃} heterobimetallic complex has potential to serve as a precursor for higher order multi-metallic species, which is currently being pursued. More broadly speaking, developing synthetic routes to plutonium heterometallic molecules will establish a novel platform for scientific exploration within the photophysical, electron transfer, and magnetic experimental spaces and holds potential for improving understanding of actinide electronic structure and bonding. Such developments could lead to new chemistries and unveil unique properties for plutonium that would otherwise be difficult to observe and characterize.

Experimental

General consideration

All manipulations were carried out without exclusion of air and moisture. Optima grade aqueous solutions of nitric acid [HNO_{3(aq)}] and hydrochloric acid [HCl_(aq)] were obtained commercially (Fisher Scientific). Water was deionized and passed through a Barnstead water purification system until a resistivity of 18.2 M Ω cm⁻¹ was achieved. Tetrahydrofuran (THF; Sigma Aldrich, anhydrous), Celite 545 (Fischer), dichloromethane (CH2Cl2; Sigma Aldrich), toluene (Sigma Aldrich), acetonitrile (MeCN; Sigma Aldrich), tetrabutylammonium hexafluorophosphate ([Bu₄N][PF₆], Sigma Aldrich), deuterated chloroform (CDCl₃, Cambridge Isotopes), and heptane (Sigma Aldrich) were obtained commercially and used as received. The sodium (cyclopentadienyl)tris(diethylphosphito)cobalt(III) salt, Na(Cp)Co[OP(OEt)₂]₃, was prepared from bis(cyclopentadienyl)cobalt(III) and diethyl phosphite, as described previously. The tetramethyl ammonium plutonium(IV) hexachloride [(Me₄N)₂PuCl₆]⁵⁵ was prepared as previously described.

The plutonium used in this study contained a blend of ²³⁸Pu [half-life $(t_{\frac{1}{2}}) = 87.7(1)y$], ²³⁹Pu $[t_{\frac{1}{2}} = 24110(30)y]$, ²⁴⁰Pu $[t_{\frac{1}{2}} = 6561(7)y]$, 241 Pu $[t_{\frac{1}{2}} = 14.329(29)y]$, and 242 Pu $[t_{\frac{1}{2}} = 3.75(33)]$ × 10⁵ y].⁷⁵ Relative isotopic ratios by mass were approximately 0.015% for ²³⁸Pu, 95.65% for ²³⁹Pu, 6.1% for ²⁴⁰Pu, 0.224 for ²⁴¹Pu, and 0.025% for ²⁴²Pu. Caution! These radionuclides and their progeny isotopes present serious health threats owing to their radioactive properties and α -, β -, and γ -emissions. Hence, all studies that involved manipulation of these isotopes were conducted in a radiation laboratory equipped with HEPA filtered hoods, continuous air monitors, negative pressure gloveboxes, and monitoring equipment appropriate for α -, β -, and γ-particle detection. Free-flowing solids were handled within negative pressure gloveboxes equipped with HEPA filters. These boxes were used to protect the worker from loose contamination and did not contain inert atmospheres. Entrance to the laboratory space was controlled with a hand and foot monitoring instrument for α -, β -, and γ -emitting isotopes and a full-body personnel contamination monitoring station.

Spectroscopic measurements: UV-vis-NIR, NMR

Single crystal absorption ultraviolet-visible-near infrared (UV-vis-NIR) measurements were made on single crystals dispersed on a quartz slide under oil using a Craic Technologies microspectrophotometer. Data were collected from 9091 to 40 000 cm⁻¹ (1100 nm to 250 nm). Solution-phase absorption UV-vis-NIR measurements were recorded on a Varian Cary 6000i spectrophotometer in screw cap quartz cuvettes. Nuclear magnetic resonance (NMR) experiments were conducted on a Bruker Avance III 400 MHz NMR spectrometer, with samples that had been doubly contained in a Teflon liner (as previously described).⁷⁶ Spectra were processed using MestReNova v10.

Single crystal X-ray diffraction measurements

Single crystal X-ray diffraction studies were performed at room temperature ($T=288~\mathrm{K}$) on a Bruker Quest D8 diffractometer equipped with an IµS 3.0 microfocus sourceTM (Mo Kα radiation, $\lambda=0.71073~\mathrm{\mathring{A}}$) and a CPAD Photon IITM area detector. Green, rectangular prisms of PuCl₃{CoCp[OP(OEt)₂]₃} suitable for X-ray diffraction were reproducibly obtained from dichloromethane solutions layered with a toluene counter solvent in a 1:1 ratio. Single crystals of PuCl₃{CoCp[OP(OEt)₂]₃} were mounted on MiTeGen MicromountsTM in mineral oil. Three layers of containment were employed to encapsulate the plutonium crystals, as has been previously described.⁴⁷ Data were collected at room temperature to avoid icing imposed from

the tertiary containment method required for safe handling of highly radioactive single crystal samples. Data was collected with the Apex3 software package using omega and phi scans.⁷⁷ Frame integration, including Lorentz-polarization corrections, and final cell parameter calculations were carried out using SAINT software.⁷⁷ A multi-scan absorption correction was applied to the data using the SADABS program.⁷⁷ The structure was solved using Intrinsic Phasing methods. The structural refinement was performed using SHELXL⁷⁸ within the ShelXle⁷⁹ software interface. See Tables S1–S5† for crystallographic parameters. Figures were produced using VESTA and Olex2.^{80,81}

All non-hydrogen atoms were located in the Fourier difference map and refined anisotropically, resulting in a structural model that consisted of a Pu(IV) bound to three chlorides and one tridentate CoCp[OP(OEt)₂]₃¹⁻ anion per asymmetric unit. The chlorides, cyclopentadienyl ring, and all but one ethoxide group were disordered over two positions and their respective site occupancies allowed to refine freely. The disordered groups were refined with both positional and displacement restraints; the distances between pairs and groups of atoms were restrained to be similar, the atomic displacement parameters of disordered atoms near one another were refined to be similar, and several atoms were restrained to be approximately isotropic. Finally, the displacement parameters of bonded atoms within the entire structure were restrained to be alike. Hydrogen atoms were placed in calculated positions and their U_{eq} values were assigned values 1.2 times that of their parent atom. CCDC deposit number: 2323718.

Electrochemistry

A CH Instruments 620C potentiostat was used for all voltametric measurements. In a HEPA-filtered fume hood, a threeelectrode cell was used, consisting of a glassy carbon disk working electrode (0.3 cm diameter), a platinum wire pseudo reference electrode (0.1 cm diameter), and a platinum-wire counter electrode (0.1 cm diameter). All electrodes were purchased from CH Instruments and the working electrode was cleaned and polished with a felt polishing pad and a mixture of alumina powder (0.05 μm) suspended in water. The cell consisted of a glass cylindrical body and a PTFE cap with holes to hold the electrodes in place. A small section of PTFE tubing (22 gauge) was inserted through a hole in the PTFE cap and used to flow argon into the cell. An electrolyte stock solution of [Bu₄N][PF₆] (0.1 M) was prepared in MeCN. For the voltametric measurements, an aliquot from the electrolyte solution (2 mL) was added to the cell and the solution was de-aerated with argon (purge 5 min). Argon was continuously flowed through the headspace of the cell during all subsequently described voltametric measurements. After characterizing the potential window for the electrolyte solution and confirming that no electron transfer processes occurred within that window, the electrolyte solution was replaced with either a green solution containing a few crystals of PuCl₃{CoCp[OP(OEt)₂]₃} and (Bu₄N)(PF₆) (77 mg, 0.2 mmol) dissolved in MeCN (2 mL) or an orange solution containing a few crystals of (Me₄)₂PuCl₆ and $[Bu_4N][PF_6]$ (77 mg, 0.2 mmol) dissolved in MeCN (2 mL).

Homogeneity was achieved by stirring and sparging the solution with argon (5 min) prior to collection of the cyclic voltammograms. The potentials from all data were referenced to the ferrocenium/ferrocene (FeCp $_2^{1+}$ /FeCp $_2$) redox couple by adding ferrocene to the PuCl $_3$ {CoCp[OP(OEt) $_2$] $_3$ } electrolyte solution after measurements were taken.

Sample preparation for magnetometry

The magnetic susceptibility measurements were conducted on PuCl₃{CoCp[OP(OEt)₂]₃} (120.9 mg, 0.150 mmol). The sample was weighed in a negative pressure glovebox used for radiological controls. Then, crystallites of the sample were loaded into a plastic soda straw and sandwiched between two plugs of quartz wool (99.6 mg). The top of the straw was equipped with a Delrin plug and the bottom of the straw was equipped with a Delrin plug that had a porous titanium frit (2 μm). Both plugs were sealed with Stycast (2850FT) epoxy. The holder provided a single layer of containment for the radioactive material and the frit enabled the helium gas (in the magnetometer) to contact the sample, which facilitated temperature equilibration during the magnetic susceptibility measurements. Magnetic susceptibility measurements were performed from 2.5 K to 300 K in a magnetic field of H = 5 T in a Quantum Design Magnetic Properties Measurement System-3. Data included a dominant contribution from the sample as well as a weak diamagnetic contribution from the quartz wool. The latter of which was well defined - as it had been measured independently - and was subtracted from the total susceptibility. The molar susceptibility in units of emu/mol is given by: $\chi_{\rm M} = N_{\rm A}(g\mu_{\rm B})^2 J(J+1)/3k_{\rm B}T$, where N_A is Avagadro's number, g is the Lande g-factor, J is total electronic momentum, and $k_{\rm B}$ is Boltzmann's constant. In c.g.s. units (which gives $\chi_{\rm M}$ in units of emu mol⁻¹), $3k_{\rm B}/N_{\rm A}\mu_{\rm B}^2=8$, so $\chi_{\mathbf{M}}T = p^2/8 = C$, where p is the effective magnetic moment $p = g[J(J+1)]^{1/2}$ and C is the Curie constant. The Lande factor $g = 3/2 + \frac{1}{2}[S(S+1) - L(L+1)]/[J(J+1)]$, where S is the spin quantum number and L is the angular momentum quantum number. By Hund's rules, the Curie constant is, for f^4 : S=2, L = 6 and J = 4, then g = 0.6 and p = 2.68; $\chi_{\rm M} T = ({\rm p}^2/8) = 0.9$ emu K mol⁻¹.

Preparation of the Pu⁴⁺(aq) stock solution

Plutonium was obtained as a residue used previously in other research campaigns. This residue was dissolved in a small volume of $HNO_{3(aq)}$ (5 to 10 mL; 16 M), and subsequently heated on a hot plate under a stream of filtered flowing air until a soft dryness was reached. The resulting residue was dissolved again in $HNO_{3(aq)}$ (5 to 10 mL; 16 M) and the process was repeated two more times. This plutonium nitrate residue was purified as described previously.^{47,55} At a high-level, this two-step purification involved (1st) precipitating the $Pu^{3+}_{(aq)}$ using $HF_{(aq)}$ (28 M) and (2nd) purification by anion exchange chromatography. On occasion we have observed small amounts of plutonium reduction on the anion exchange resin that generates a small amount of $Pu^{3+}_{(aq)}$. When this occurred, the blue $Pu^{3+}_{(aq)}$ contaminant was separated easily from the reddish-orange $Pu^{4+}_{(aq)}$ product because $Pu^{3+}_{(aq)}$ passes through the anion

exchange column at a faster rate than $Pu^{4+}_{(aq)}$ during the elution process. For example, passing slightly acidic $HCl_{(aq)}$ (50 mL of H_2O to five drops of $HCl_{(aq)}$; 12 M) through the column pushed the blue $Pu^{3+}_{(aq)}$ fraction off the column first. Then, the reddishorange $Pu^{4+}_{(aq)}$ fraction was independently isolated. The eluate $HCl_{(aq)}$ concentration was then adjusted to approximately 6 M by doubling the $Pu^{4+}_{(aq)}$ eluate solution volume and adding $HCl_{(aq)}$ (12 M). For example, 7 mL of $HCl_{(aq)}$ (12 M) was added if the eluted $Pu^{4+}_{(aq)}$ fraction was 7 mL in volume. An aliquot (100 μ L) of the final stock solution was added to a quartz cuvette charged with $HClO_{4(aq)}$ (1 M, 1 mL), and assayed by UV-vis-NIR absorption spectroscopy. The absorption peak at 21 277 cm⁻¹ (470 nm; $\varepsilon_{470\,\mathrm{nm}} = 56.5$ L mol^{-1} cm⁻¹) was used to quantify the concentration of the $Pu^{4+}_{(aq)}$ stock solution, which was found to be 0.127 M.

Plutonium(IV) trichloride (cyclopentadienyl) tris(diethylphosphito)cobalt(III), PuCl₃{CoCp[OP(OEt)₂]₃}

Under ambient conditions and with no efforts to exclude air or moisture, a polyethylene falcon cone (50 mL) was charged with an aliquot of the aforementioned Pu4+(aq) stock solution (0.127 M in Pu⁴⁺, 151.2 mg, 0.633 mmol) in HCl_(aq) (5 mL, 6 M). The solution was heated on a hot plate under a stream of filtered flowing air until a soft dryness was achieved. The resulting residue was suspended in THF (7.5 mL). Most of the Pu⁴⁺_(aq) residue dissolved. The resulting red slurry was transferred into a Pyrex falcon cone (50 mL) that contained an orange solution of Na{CoCp[OP(OEt)₂]₃} (340.0 mg, 0.610 mmol) dissolved in THF (7.5 mL). Combining the plutonium mixture and the Na{CoCp[OP(OEt)₂]₃} solutions resulted in a homogenous brown solution. A white solid, presumably sodium chloride (NaCl), precipitated as the solution stirred overnight. Insoluble byproducts were removed by passing the solution through a filter stick charged with Celite. The eluent was exposed to a stream of filtered air until a soft dryness was achieved. The majority of the resulting residue dissolved in CH₂Cl₂ (5 mL) however, a stubborn white solid (likely NaCl) lingered. Removal of this persistent precipitate was achieved only after the mixture was filtered through two additional Celite columns. The resulting homogenous dark green solution was subsequently evaporated to soft dryness at room temperature under a stream of filtered air, which left a green solid. This solid was dissolved in a small amount of CH₂Cl₂ (5 mL) and layered with toluene (5 mL). Dark green and rectangular single crystals of PuCl₃{CoCp[OP(OEt)₂]₃} formed after this solution sat at room temperature for 48 h. After the mother liquor was decanted, the crystals were washed with heptane, and dried under air. This enabled the title compound, PuCl₃{CoCp[OP(OEt)₂]₃}, to be isolated (0.3735 g, 0.4619 mmol) in 70% crystalline yield; based on $Na\{CoCp[OP(OEt)_2]_3\}$.

¹H NMR, (CDCl₃, 20 °C, 400.13 MHz): δ 6.71 ppm (s, 5H); δ 3.03 (dp, J = 91, 7.8 Hz, 12H); δ 0.73 ppm (t, J = 6.9 Hz, 18H). ³¹P (CDCl₃, 20 °C, 202.46 MHz): δ 24.5 ppm (s, 1P).

UV-vis-NIR (cm⁻¹): 12 406.95, 12 866.60; 14 064.70 sh; 14 388.49 sh; 14 880.95 sh; 163 313.21 sh; 16 077.17 sh; 22 123.89 CT.

Data availability

Crystallographic data for PuCl₃{CoCp[OP(OEt)₂]₃} has been deposited at the CCDC (deposit number: 2323718). The rest of the data supporting this article have been included as part of the main text or within the ESI.†

Author contributions

All of the authors contributed to writing, reviewing, and editing the document. Thomas E. Shaw conducted electrochemical studies and formal analyses. Zachary R. Jones conducted the initial synthetic explorations and project design. Sara L. Adelman conducted experiment design. Nickolas H. Anderson contributed to synthesis and experimental design. Eric G. Bowes helped conduct the plutonium synthesis and sample preparation. Eric D. Bauer led the magnetic measurements. David Dan conducted magnetic measurements. Jan Klouda conducted electrochemical studies. Karah E. Knope was the contributed to structural analysis and experimental design. Stosh A. Kozimor provided experimental design, conceptualization, formal analyses, plutonium reprocessing. Molly M. MacInnes conducted electrochemical experiments. Veronika Mocko conducted electrochemical measurements. Francisca R. Rocha conducted electrochemical measurements. Harrison D. Root conducted synthetic experimental studies and NMR studies. Benjamin W. Stein oversaw the spectroscopic measurements. Joe D. Thompson conducted magnetic measurements. Jennifer N. Wacker conducted crystallographic measurements and solved the crystal structure.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We thank the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, Heavy Element Chemistry program (2020LANLE372) for funding the majority of this work. Additional support came from LANL's LDRD projects for electrochemical measurements (20220054DR and 20190154ER) and magnetic measurements (20200045ER). We thank the Glenn T. Seaborg Institute (Jones), the Natural Sciences and Engineering Research Council of Canada Postdoctoral Fellowship (Bowes), and the DOE, Office Of Science, Office of Workforce Development for Teachers and Scientists, Office of Science Graduate Student Research (SCGSR) program, which is administered by the Oak Ridge Institute for Science and Education (ORISE) and managed by ORAU for the DOE (contract number DE-SC0014664) (Wacker) for graduate student funding. KEK was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, Early Career Research Program under Award DE-SC0019190. LANL is an affirmative action/ equal opportunity employer managed by Triad National Security, LLC, for the National Nuclear Security Administration of the U.S. DOE.

References

- 1 J. Campos, Nat. Rev. Chem., 2020, 4, 696.
- 2 F.-F. Chen, Z.-Q. Chen, Z.-Q. Bian and C.-H. Huang, *Coord. Chem. Rev.*, 2010, **254**, 991.
- 3 T. Lathion, A. Fürstenberg, C. Besnard, A. Hauser, A. Bousseksou and C. Piguet, *Inorg. Chem.*, 2020, **59**, 1091.
- 4 P. Sharma, D. R. Pahls, B. L. Ramirez, C. C. Lu and L. Gagliardi, *Inorg. Chem.*, 2019, 58, 10139.
- 5 J. Shen, T. Rajeshkumar, G. Feng, Y. Zhao, S. Wang, L. Maron and C. Zhu, *Angew. Chem., Int. Ed.*, 2023, **62**, e202303379.
- 6 G. Feng, M. Zhang, P. Wang, S. Wang, L. Maron and C. Zhu, Proc. Natl. Acad. Sci. U.S.A., 2019, 116, 17645.
- 7 J. A. Hlina, J. R. Pankhurst, N. Kaltsoyannis and P. L. Arnold, J. Am. Chem. Soc., 2016, 138, 3333.
- 8 R. S. Sternal, C. P. Brock and T. J. Marks, *J. Am. Chem. Soc.*, 1985, **107**, 8270.
- 9 A. J. Ayres, A. J. Wooles, M. Zegke, F. Tuna and S. T. Liddle, *Inorg. Chem.*, 2019, 58, 13077.
- 10 A. L. Ward, W. W. Lukens, C. C. Lu and J. Arnold, J. Am. Chem. Soc., 2014, 136, 3647.
- 11 C. Z. Ye, I. D. Rosal, M. A. Boreen, E. T. Ouellette, D. R. Russo, L. Maron, J. Arnold and C. Camp, *Chem. Sci.*, 2022, 14, 861.
- 12 A. M. Chapman, M. F. Haddow and D. F. Wass, *J. Am. Chem. Soc.*, 2011, **133**, 8826.
- 13 M. Carmona, R. Pérez, J. Ferrer, R. Rodríguez, V. Passarelli, F. J. Lahoz, P. García-Orduña and D. Carmona, *Inorg. Chem.*, 2022, 61, 13149.
- 14 D. W. Stephan, J. Am. Chem. Soc., 2021, 143, 20002.
- 15 I. Čorić and P. L. Holland, *J. Am. Chem. Soc.*, 2016, **138**, 7200.
- 16 B. H. Hoffman, D. Lukoyanov, Z.-Y. Yang, D. R. Dean and L. C. Seefeldt, *Chem. Rev.*, 2014, **114**, 4041.
- 17 T. Le Borgne, E. Rivière, J. Marrot, J.-J. Girerd and M. Ephritikhine, *Angew. Chem., Int. Ed.*, 2000, **39**, 1647.
- 18 D. Patel, F. Moro, J. McMaster, W. Lewis, A. J. Blake and S. T. Liddle, *Angew. Chem., Int. Ed.*, 2011, 50, 10388.
- 19 C. Camp, D. Toniolo, J. Andrez, J. Pécaut and M. Mazzanti, *Dalton Trans.*, 2017, **46**, 11145.
- 20 B. M. Gardner, J. McMaster, F. Moro, W. Lewis, A. J Blake and S. T. Liddle, *Chem.-Eur. J.*, 2011, 17, 6909.
- 21 B. M. Gardner, D. Patel, A. D. Cornish, J. McMaster, W. Lewis, A. J. Blake and S. T. Liddle, *Chem.-Eur. J.*, 2011, 17, 11266.
- 22 V. Mougel, L. Chatelain, J. Pécaut, R. Caciuffo, E. Colineau, J.-C. Griveau and M. Mazzanti, *Nat. Chem.*, 2012, **4**, 1011.
- 23 L. Cahtelain, J. P. S. Walsh, J. Pécaut, F. Tuna and M. Mazzanti, *Angew. Chem., Int. Ed.*, 2014, 53, 13434.
- 24 P. Yang, E. Zhou, G. Hou, G. Zi, W. Ding and M. D. Walter, *Chem.-Eur. J.*, 2016, **22**, 13845.
- 25 E. Lu, A. J. Wooles, M. Gregson, P. J. Cobb and S. T. Liddle, Angew. Chem., Int. Ed., 2018, 57, 6587.
- 26 M. A. Boreen, T. D. Lohrey, G. Rao., D. Britt, L. Maron and J. Arnold, J. Am. Chem. Soc., 2019, 141, 5144.
- 27 O. Kahn, Adv. Inorg. Chem., 1995, 43, 179.

28 B. G. Cooper, J. W. Napoline and C. M. Thomas, *Catal. Rev.*, 2012, 54, 1.

Edge Article

- 29 P. Buchwalter, J. Rosé and P. Braunstein, *Chem. Rev.*, 2015, 115, 28.
- 30 S. Sinhababu, Y. Lakliang and N. P. Mankad, *Dalton Trans.*, 2022, **51**, 6129.
- 31 A. Lachguar, A. V. Pichugov, T. Neumann, Z. Dubrawski and C. Camp, *Dalton Trans.*, 2024, 53, 1393.
- 32 N. J. Curro, T. Caldwell, E. D. Bauer, L. A. Morales, M. J. Graf, Y. Bang, A. V. Balatsky, J. D. Thompson and J. L. Sarrao, *Nature*, 2005, 434, 622.
- 33 N. Budantseva, G. Andreev, M. Sokolova and A. Fedoseev, *Inorg. Chem.*, 2021, **60**, 18395.
- 34 C. Tamain, B. A. Chapelet, M. Rivenet, F. Abraham, R. Caraballo and S. Grandjean, *Inorg. Chem.*, 2013, **52**, 4941.
- 35 J. N. Cross, P. M. Duncan, E. M. Villa, E. V. Alekseev, C. H. Booth and T. E. Albrecht-Schmitt, J. Am. Chem. Soc., 2013, 135, 2769.
- 36 W. Kläui, H. O. Asbahr, G. Schramm and U. Englert, *Chem. Ber./Recl*, 1997, **130**, 1223.
- 37 I. Goldberg, H. Shinar, G. Navon and W. Kläui, *J. Incl. Phenom.*, 1987, 5, 181.
- 38 H. Shinar, G. Navon and W. Kläui, J. Am. Chem. Soc., 1986, 108, 5005.
- 39 H. Werner, H. Neukomm and W. Kläui, *Helv. Chim. Acta*, 1977, **60**, 326.
- 40 M. Glaum, W. Kläui, B. W. Skelton and A. H. White, *Aust. J. Chem.*, 1997, **50**, 1047.
- 41 W. Kläui and T. Hardt, J. Organomet. Chem., 1998, 553, 241.
- 42 W. Kläui, M. Glaum, E. Hahn and T. Lugger, Eur. J. Inorg. Chem., 2000, 21.
- 43 W. Kläui, W. Peters, N. Liedtke, S. Trofimenko, A. L. Rheingold and R. D. Sommer, *Eur. J. Inorg. Chem.*, 2001, 693.
- 44 W. Kläui, N. Mocigemba, A. Weber-Schuster, R. Bell, W. Frank, D. Mootz, W. Poll and H. Wunderlich, *Chem.–Eur. J.*, 2002, **8**, 2335.
- 45 W. Kläui and P. C. Kunz, Inorg. Synth., 2010, 35, 125.
- 46 W. Kläui, Z. Naturforsch., 1979, 34B, 1403.
- 47 S. K. Cary, K. S. Bowland, J. N. Cross, S. A. Kozimor and B. L. Scott, *Polyhedron*, 2017, **126**, 220.
- 48 G.-C. Wang, Y.-M. So, K.-L. Wong, K.-C. Au-Yeung, H. H.-Y. Sung, I. D. Williams and W.-H. Leung, *Chem.-Eur. J.*, 2015, **21**, 16126.
- 49 M. Wedler, J. W. Gilje, M. Noltemeyer and F. T. Edelmann, *J. Organomet. Chem.*, 1991, **411**, 271.
- 50 D. Baudry, M. Ephritikhine, W. Kläui, M. Lance, M. Nierlich and J. Vigner, *Inorg. Chem.*, 1991, 30, 2333.
- 51 T. C. H. Lam, E. Y. Y. Chan, W.-L. Mak, S. M. F. Lo, I. D. Williams, W.-T. Wong and W.-H. Leung, *Inorg. Chem.*, 2003, 42, 1842.
- 52 A. C. Grunwald, C. Scholtysik, A. Hagenbach and U. Abram, *Inorg. Chem.*, 2020, **59**, 9396.
- 53 J. W. Napoline, S. J. Kraft, E. M. Matson, P. E. Fanwick, S. C. Bart and C. M. Thomas, *Inorg. Chem.*, 2013, 52, 12170.
- 54 W. Kläui, D. Matt, F. Balegroune and D. Grandjean, *Acta. Cryst.*, 1991, **C47**, 1614.

- 55 A. M. Mounce, H. Yasuoka, G. Koutroulakis, J. A. Lee, H. Cho, F. Gendron, E. Zurek, B. L. Scott, J. A. Trujillo, A. K. Slemmons, J. N. Cross, J. D. Thompson, S. A. Kozimor, E. D. Bauer, J. Autschback and D. L. Clark, *Inorg. Chem.*, 2016, 55, 8371.
- 56 R. E. Wilson, Y.-J. Hu and H. Nitsche, *Radiochim. Acta*, 2005, 93, 203.
- 57 M. R. Antonio, C. W. Williams, J. A. Sullivan, S. Skanthakumar, Y.-J. Hu and L. Soderholm, *Inorg. Chem.*, 2012, **51**, 5274.
- 58 C. J. Windorff, C. A. P. Goodwin, J. M. Sperling, T. E. Albrecht-Schönzart, Z. Bai, W. J. Evans, Z. K. Huffman, R. Jeannin, B. N. Long, D. P. Mills, T. N. Poe and J. W. Ziller, *Inorg. Chem.*, 2023, 62, 18136.
- 59 J. Su, T. Cheisson, A. McSkimming, C. A. P. Goodwin, I. M. DiMucci, T. Albrecht-Schönzart, B. L. Scott, E. R. Batista, A. J. Gaunt, S. A. Kozimor, P. Yang and E. J. Schelter, *Chem. Sci.*, 2021, 12, 13343.
- 60 C. Berger, C. Marie, D. Guillaumont, C. Tamain, T. Dumas, T. Dirks, N. Boubals, E. Acher, M. Laszczyk and L. Berthon, *Inorg. Chem.*, 2020, 59, 1823.
- 61 D. Lemire, T. Dumas, C. Marie, F. Giusti, G. Arrachart, S. Dourdain and S. Pellet-Rostaing, *Eur. J. Inorg. Chem.*, 2023, **26**, e202300461.
- 62 L. Yao, W. Junli, L. Yuhang, W. Hui, W. Wentao, L. Baole and Y. Taihong, *RSC Adv.*, 2024, **14**, 560.
- 63 C. Berger, E. Moreau, C. Marie, D. Guillaumont, A. Beillard and L. Berthon, *Solvent Extr. Ion Exch.*, 2021, **40**, 290.
- 64 J. A. Macor, J. L. Brown, J. N. Cross, S. R. Daly, A. J. Gaunt, G. S. Girolami, M. T. Janicke, S. A. Kozimor, M. P. Neu, A. C. Olson, S. D. Reilly and B. L. Scott, *Dalton Trans.*, 2015, 44, 18923.
- 65 X.-B. Li, Q.-Y. Wu, C.-Z. Wang, J.-H. Lan, M. Zhang, Z.-F. Chai and W.-Q. Shi, *J. Phys. Chem. A*, 2023, **127**, 7479.
- 66 V. S. Koltunov, R. J. Taylor, S. M. Baranov, E. A. Mezhov, V. G. Pastushchak and I. May, *Radiochim. Acta*, 2000, 88, 65.
- 67 V. S. Koltunov, V. I. Marchenko, G. I. Zhuravleva and O. A. Savilova, *Radiochemistry*, 2001, **43**, 334.
- 68 V. S. Koltunov, S. M. Baranov and V. G. Pastushchak, *Radiochemistry*, 2001, **43**, 346.
- 69 P. Tkac, M. Precek and A. Pulenova, Proc. Global, 2009, 6.
- 70 P. Tkac, M. Precek and A. Paulenova, *Inorg. Chem.*, 2009, 48, 11935.
- 71 R. J. Taylor and I. May, Czech J. Phys., 1999, 49, 617.
- 72 R. J. Taylor, C. Mason, R. Cooke and C. Boxall, *J. Nucl. Sci. Technol.*, 2002, **39**, 278.
- 73 V. I. Marchenko, K. N. Dvoeglazov, O. A. Savilova and V. I. Volk, *Radiochemistry*, 2012, 54, 459.
- 74 S. L. Yarbro, S. B. Schreiber, E. M. Ortiz and R. L. Ames, *J. Radioanal. Nucl. Chem.*, 1998, 235, 21.
- 75 National Nuclear Data Center and Brookhaven National Laboratory, 2021, accessed April 21, 2021. https:// www.nndc.bnl.gov/nudat2/.
- 76 B. W. Stein, S. A. Kozimor and V. Mocko, *The Plutonium Handbook*, ed. D. L Clark, D. A. Geeson and R. J. Hanrahan Jr, American Nuclear Society, La Grange Park, Illinois, 2nd Edition, 2019, p. 2951.

77 APEX3, SADABS, SAINT, SHELXTL, XCIF, and XPREP, Bruker 80 K. Momma and F. Izumi, J. Appl. Crystallogr., 2011, 44, AXS, Inc., Madison, WI, 2016.

78 G. M. Sheldrick, Acta Crystallogr., 2015, C71, 3.

Chemical Science

- 79 C. B. Hübschle, G. M. Sheldrick and B. Dittrich, J. Appl. Crystallogr., 2011, 44, 1281.
- 1272.
- 81 O. Dolomanov, L. Bourhis, R. Gildea, J. Howard and H. Puschmann, J. Appl. Crystallogr., 2009, 42, 339.