[UF₆]²⁻: A Molecular Hexafluorido Actinide(IV) Complex with Compensating Spin and Orbital Magnetic Moments

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Abstract: The first structurally characterized hexafluorido complex of a tetravalent actinide ion, the [UF₆]²⁻ anion, is reported in the (NEt₄)[UF₆]·2H₂O salt (1). The weak magnetic response of 1 results from both U⁴⁺ spin and orbital contributions, as established by combining X-ray magnetic circular dichroism (XMCD) spectroscopy and bulk magnetization measurements. The spin and orbital moments are virtually identical in magnitude, but opposite in sign, resulting in an almost perfect cancellation, which is corroborated by ab initio calculations. This work constitutes the first experimental demonstration of a seemingly non-magnetic molecular actinide complex carrying sizable spin and orbital magnetic moments.

The chemistry of uranium compounds and the diversity of potential uses for uranium in novel materials are currently experiencing a renaissance.[1] Molecular complexes with uranium in either + III or + V oxidation states have been shown to generally exhibit slow relaxation of magnetization,[2] together with a single U⁴⁺ complex that appears as an exception.[3] Such complexes are also interesting building blocks for magnetic polynuclear complexes and one-dimensional coordination polymers.[4] Recent spectroscopic and theoretical investigations have aimed at a detailed understanding of the electronic structure of actinide ions, which is significantly more complicated than for lanthanides.[5] The large spin–orbit coupling of actinide ions and the less shielded nature of their 5f orbitals relative to the 4f orbitals, result in much stronger interactions of the f electrons with the surrounding atoms. Hence, the concomitant stronger ligand field cannot be considered as a perturbation, particularly for the early members of the 5f series. Furthermore, spectroscopic data of actinide systems are typically very rich and the determination of the electronic energy level-splitting and -composition remains quite challenging. For the magnetic characterization of actinide-based materials, powder and thermodynamically averaged magnetization data bring only limited information about the underlying physics. However, since the magnetic properties are predominantly defined by the energy levels that are thermally populated at room temperature and below, any additional experimental information on these low-lying energy levels would be of great relevance. For this task, X-ray magnetic circular dichroism spectroscopy is a powerful tool able to deconvolute the macroscopically measured magnetic moment into its spin and orbital contributions.[6] Despite its routine applications in magnetism, this technique has never been applied to molecular actinide systems.[7] Inspired by the early works of Ryan et al. and Brown et al.,[8a,b] we herein report the synthesis and crystal structure of (NEt₄)[UF₆]·2H₂O (1) featuring the first structurally characterized example of an octahedral fluoride complex in the tetravalent actinide family.[9] Although the octahedral [UF₆] and [UF₆]²⁻ complexes have been well-described and structurally characterized,[9] the existence of an octahedral [UF₆]²⁻ has only been inferred from vibrational spectroscopy, but not confirmed by X-ray crystallography.[8a,b] Indeed, more generally, despite the vast number of known fluoride complexes and solids of the actinide ions, no...
hexafluoride complex of the tetravalent actinides has yet been structurally characterized.

The reaction of UF₄·xH₂O (x ≈ 1.5) with NEt₄F·H₂O in propylene carbonate under a dry N₂ atmosphere yields a pale green solution. Addition of acetone induces the crystallization of a pale-green material suitable for single-crystal X-ray diffraction. The crystal structure analysis of (NEt₄)₂[UF₄]·2H₂O (1) reveals the presence of slightly compressed octahedral [UF₄]²⁻ complexes (Figure 1), with U–F bond lengths being longer for fluoride ligands engaged in hydrogen bonding (2.177(2) and 2.181(2) Å versus 2.124(2) Å; Supporting Information, Figure S1).[14] Selected bond lengths [Å] and angles [°]: U–F 2.124(2), U–F2 2.177(2), U–F3 2.181(2), cis–F–U–F 89.23(7)–90.90(7). U purple, F green, O red, C black, H gray.

Figure 1. Molecular structure of the [UF₄]²⁻ complex in 1 (ellipsoids set at 50% probability) and the hydrogen bonding pattern linking [UF₄]²⁻ complexes into supramolecular chains running along the crystallographic c direction (Supporting Information, Figure S1).[14] The temperature dependence of the magnetic susceptibility–temperature product, χT, of 1 is shown in Figure 2a. From 0.35 cm³Kmol⁻¹ at 300 K, it nearly vanishes to 5 × 10⁻³ cm³Kmol⁻¹ at 1.8 K, which is far from the 1.60 cm³Kmol⁻¹ value expected for the Russell–Saunders atomic ground state term of the 5f² configuration (5H₄, gₛ = 4/5).[12] The electronic ground state of U(IV) in octahedral symmetry is a singlet state composed of a linear combination of the |mₛ = 0⟩ and |mₛ = ±4⟩ states.[19] Thus, the magnetic susceptibility is expected to be dominated by temperature-independent paramagnetism (TIP). The simulated χT product, using the model developed in reference [13] with our ab initio results (see below), is shown in Figure 2a. The experimental departure from the idealized TIP dependence differs from the behavior of [UC₅H₄]²⁻ and [UB₅H₅]²⁻ analogues,[14] but is nevertheless commonly observed for U(IV) systems.[20] To obtain experimental information on the nature of the electronic ground state of uranium in 1, X-ray absorption spectroscopy (XAS) and X-ray magnetic circular dichroism (XMCD) were employed.

The uranium M₄₅ XAS spectra are dominated by strong resonance lines, so-called white lines, that are due to dipole-allowed transitions from the spin–orbit split 3d₃/2,5/2 levels into the magnetic 5f states along with much weaker 3d—6p and 3d—continuum transitions. According to the dipole selection rule (|ΔJ| = 1), the larger Mₛ white line intensity primarily reflects the population of the 5f₃,5 states and the weaker M₄ intensity only the 5f₄,4 states.[15] In Figure 3a, the XAS M₄₅ spectra are shown on the same relative energy scale that is obtained by shifting the original spectra (Figure 3b, gray trace) to perfectly overlap the first EXAFS oscillation (inset Figure 3a). It is apparent, that the absorption maxima of the M₄ and M₅ white lines are at different photon energies separated by about 0.8 eV, with the M₅ peak being at lower photon energies. This energy splitting is due to spin–orbit coupling of the 5f states. In a spherically symmetric ligand field, the magnitude of this gap can be approximated by ΔE = 7ζᵣᵣ, where ζᵣᵣ is the spin–orbit parameter for the 5f electrons. The angular part of the spin–orbit interaction for the 5f states, ⟨Σ₃/2,₅/2⟩, can be obtained using the so-called spin–orbit sum rule, which relates its ground-state expectation value to the branching ratio (BR) of the isotropic X-ray absorption intensities.[16] The experimental BR value, 0.727 ± 0.005, of 1 results in ⟨Σ₃/2,₅/2⟩/⟨Σ₃/2,₅/2⟩ ≈ 5.7, which is significantly larger than the values obtained by atomic calculations on the 5f² configuration of –3–, –4–, and –3.88 in the Russell–Saunders, j–j, and intermediate coupling scheme, respectively.[11] It should be noted that the spin–orbit sum rule has been derived neglecting the core–valence interaction. The observed deviation from theory is thus not unexpected because of the highly localized nature of the 5f states in 1, where core-valence interaction could be very strong. Indeed, such discrepancies with theoretical models have already been

Figure 2. a) Temperature dependence of the χT product (χ being the magnetic susceptibility defined as M(μ₀H)/H) over the 1.8–100 K range (0.3 K min⁻¹) of a polycrystalline sample of 1 obtained with μ₀H = 1.0 T. The gray dashed line is the calculated χT product using the model of Ref. [13] with a T₁–T₂ energy splitting of 1300 cm⁻¹. b) Field-dependence of the magnetization, M(μ₀H), obtained at 4 K in the 0–7 T range and the scaled field dependence of the XMCD signal measured in the 0–17 T range at a photon energy of 3727.3 eV corresponding to the maximum of the XMCD signal at the uranium M₄ edge.
observed in electron energy loss spectroscopy results obtained on various U$^{IV}$ oxide minerals.$^{[20]}$

The normalized XMCD spectra at the uranium M$_5$ edges are shown on Figure 3b. Sizeable dichroism signals are observed at both M$_5$ and M$_6$ edges. The spectral shape of the XMCD at the M$_5$ edge is in qualitative agreement with the one estimated from multiplet calculations for the 5f$^2$ configuration with an octahedral ligand field (Figure 3b, green trace), and with experimental results on U$^{IV}$-containing metals.$^{[21]}$ This favorable comparison of the uranium M$_5$ XMCD spectral shape, which is extremely sensitive to the 5f occupation, confirms also the 5f$^2$ configuration of uranium in 1. To deduce the magnitudes of the magnetic moments carried by the uranium 5f states, the so-called magneto-optical sum rules can be used, which relate the integrals of the XMCD spectra to the orbital (orbital sum rule)$^{[22]}$ and spin (spin sum rule)$^{[23]}$ magnetic moments. Unfortunately, the spin sum rule is based on the same approximation as the spin–orbit sum rule, and can therefore hardly be applied to our XMCD spectra. Nevertheless, the orbital sum rule remains valid irrespective of this approximation and numerical integration of the XMCD spectra shown in Figure 3b affords $M_{\text{orbital}} = 0.47 \mu_B$. The spin moment, $M_{\text{spin}}$, may hereafter be determined by scaling the magnetization curve measured by monitoring the M$_4$ XMCD signal as a function of the applied magnetic field (shown in Figure 2b) to the macroscopic magnetization data. This procedure allows the estimation of the absolute magnetization at 17T, which amounts to only $M_{\text{total}} = 0.060(4) \mu_B$, thereby yielding $M_{\text{spin}} = -0.41 \mu_B$ through the $M_{\text{total}} = M_{\text{orbital}} + M_{\text{spin}}$ relation. Thus, the analysis of the XMCD spectra reveals, unambiguously, the existence of sizable orbital and spin magnetic moments in 1 despite a low bulk magnetic moment. Similar cancellation of spin and orbital magnetizations for uranium has already been observed in itinerant 5f systems like UFe$_2$, using neutron scattering and confirmed by XMCD.$^{[21]}$ Here, we demonstrate that the spin–orbital cancellation is not limited to metallic systems with strongly delocalized 5f states but is also present in a molecular system with much higher degree of 5f localization.

Ab initio CASSCF$^{[22]}$ and NEVPT2$^{[23]}$ calculations on the [UF$_6$]$^-$ complex predict significantly lower orbital and spin magnetic moments of the ground state ($M_{\text{orbital}} = -0.051 \mu_B$ and $M_{\text{spin}} = 0.018 \mu_B$, NEVPT2). Considering the idealized high-symmetry environment of the [UF$_6$]$^-$ moiety, it is reasonable to explore the influence of vibronic effects on the magnetic properties of the complex. The calculations indicate that the $A_{1g}$ ground state is separated from the excited states by more than 1000 cm$^{-1}$ for both the crystal structure model and an idealized octahedral geometry. Thus, there is no ground state degeneracy and vibronic effects must be related to a pseudo-Jahn–Teller mechanism.$^{[24]}$ To evaluate if vibronic coupling impacts the magnetic moments, a Hamiltonian was built with diagonal elements corresponding to state energies including spin–orbit effects and a Zeeman term. The state interaction between the ground state, 0, and an excited state, $a$, is represented by the following expression [Eq. (1)]:

$$\mathcal{H}_{ia} = -\mu_B \mu_0 (g_{ia} S_{ia} + L_{ia}) \cdot \mathbf{H} + F_{ia} \lambda$$  \hspace{1cm} (1)$$

where $S_{ia} = \langle \Psi_i | \hat{S}_z | \Psi_0 \rangle$ and $L_{ia} = \langle \Psi_i | \hat{J}_z | \Psi_0 \rangle$ for $i = x, y,$ and $z$. The first term corresponds to the Zeeman interaction and the second is the linear vibronic contribution with coupling constants [Eq. (2)]:

$$F_{ia} = \langle \Psi_0 | \frac{d \mathcal{H}_{ia}}{d \lambda} | \Psi_i \rangle$$  \hspace{1cm} (2)$$

where $\mathcal{H}_{0i}$ is evaluated considering the crystallographically determined geometry, and $\Psi_0$ and $\Psi_i$ are ground and excited state wave functions, respectively (see Supporting Information for further details). Vibronic mixing quickly increases the magnetic moment and then stabilizes for low displacement values (ca. $\lambda = 0.03$ Å; Supporting Information, Figure S5). If we consider 0.03 Å as a representative displacement, we obtain $M_{\text{orbital}} = 0.48 \mu_B$ and $M_{\text{spin}} = -0.26 \mu_B$ which are in reasonable agreement with the experiment. Thus, the influence of the vibronic couplings on the magnetic properties is remarkably important for the [UF$_6$]$^-$ complex, but there are no reasons to believe that this effect is restricted to this system.
particular system. Indeed, the magnitude of the magnetic moment in UF₄ could only be explained by invoking dynamic Jahn–Teller distortions and the inelastic neutron scattering spectra of the lowest-lying crystal field states of UF₄ and UCl₄ have revealed the signatures of strong couplings between electronic states and vibrations.

In conclusion, this work reports on the UF₄⁻ complex, which is the first example of a structurally characterized hexafluorido actinide(IV) complex. Despite the very weak total magnetic moment of UF₄⁻, the detailed study and analysis of its magnetic properties combining state-of-the-art X-ray absorption spectroscopy, magnetometry, and quantum chemical calculations have demonstrated the existence of relatively large spin and orbital magnetic moments. These moments are roughly one hundred times larger than those found in related 5d systems with electronic singlet ground states. These significant spin and orbital components should be considered in the future analyses of the magnetic exchange interactions in uranium(IV)-based materials and fuel the curiosity for a fundamental understanding of the electronic structure and magnetism of other actinide-containing molecules and materials.

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Conflict of interest

The authors declare no conflict of interest.

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