Synthesis, structure and bonding of actinide disulphide dications in the gas phase†

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Actinide disulphide dications, AnS$_2^{2+}$, were produced in the gas phase for An = Th and Np by reaction of An$^{3+}$ cations with the sulfur-atom donor COS, in a sequential abstraction process of two sulfur atoms, as examined by FTICR mass spectrometry. For An = Pu and Am, An$^{2+}$ ions were unreactive with COS and did not yield any sulphide species. High level multiconfigurational (CASPT2) calculations were performed to assess the structures and bonding of the new AnS$_2^{2+}$ species obtained for An = Th, Np, as well as for An = Pu to examine trends along the An series, and for An = U to compare with a previous experimental study and DFT computational scrutiny of US$_2^{2+}$. The CASPT2 results showed that, like in the case of uranium, the new AnS$_2^{2+}$ ions have ground states with triangular geometries, corresponding to the presence of a persulphide in the case of thorium that formally leads to a stable ThIVS$_2^{2+}$ species, while a supersulphide appears to be present in the case of U, Np and Pu, formally leading to an AnIVS$_2^{2+}$ species. The computations also revealed that linear thioactinyl structures are higher in energy, with a difference that increases fourfold upon moving from U to Pu, apparently indicating that it will be even more pronounced for Am.

Introduction

The actinyl dications, An$^{IV}$O$_2^{2+}$, are especially significant in the chemistry of U, Np, Pu, and Am.$^1$ Uranyl in particular has received widespread consideration from both the experimental and theoretical perspectives.$^2$–$^{10}$ The synthesis of analogues of uranyl, where oxygen is replaced by another main-group element or moiety, such as carbene, imido, and terminal chalcogenide, has multiplied in recent years and is still an ongoing endeavour.$^{11}$–$^{15}$ For the less-studied transuranium actinides, only recently has a first actinyl V or VI analogue been successfully prepared, specifically a trans-bisimido complex of Np(V).$^{16}$

A particular goal has been to synthesize species that integrate bonds between uranium and sulphur, the closest chalcogenide homologue of oxygen, namely thiouranyl, US$_2^{2+}$. We recently reported a detailed study of the gas-phase reactions of monopositive and dipositive uranium cations with COS, with focus on the formation of US$_2^{2+}$, potentially the thiouranyl ion.$^{18}$ Remarkably, density functional theory (DFT) computations indicated that the ground-state structure for this species is triangular US$_2^{2+}$, with a side-on $\eta^5$-S$_2$ ligand, while the linear thiouranyl isomer, [S=U=S]$_2^{2+}$, was computed to be 40.9 kcal mol$^{-1}$ higher in energy.$^{18}$

In actinide chemistry, the issue of the extent of covalency in metal–ligand interactions has been pervasive and long lasting.$^{19}$–$^{22}$ Molecules involving metal–ligand interactions with multiple-bond character are strong indicators of the extent of
covalency, and changes along the actinide series can be particularly illuminating. Therefore, in the present work, gas-phase studies of the formation of actinide disulphide dications were performed through reactions of An\(^{2+}\) ions with COS, for An = Th, Np, Pu, and Am, to complement previous studies with uranium and evaluate potential trends along the actinide series. Thorium was included in this study as reference, because it can only achieve a maximum oxidation state of +4 and cannot form actinyl or analogue species. Gas-phase ion-chemistry studies are generally conducted in combination with computational studies so that experimentally observed species can be elucidated. Here, we have opted for using high-level multiconfigurational (CASPT2) calculations to assess the structure and bonding of observed and non-observed AnS\(_2^{2+}\) species, and try to illuminate any trends for the early actinides.

There are very few reported studies on elementary molecular actinide disulphides. Andrews and co-workers accomplished the production of the neutral ThS\(_2\) and US\(_2\) by the reaction of sulphur vapour with laser-ablated metal atoms in cryogenic matrices.\(^{23,25}\) In a previous study of An sulphides, we were able to produce AnS\(_2^{2+}\) ions for An = Th, Pa, U, and Np by sequential reactions of An\(^+\) ions with OCS.\(^{26}\) Earlier experiments by Gibson had shown that AnS\(_2^{2+}\) could be obtained using thiols as reagents for An = U, Np, Pu, and Am.\(^{27}\)

**Experimental details**

Gas-phase experiments were performed by Fourier transform ion-cyclotron resonance mass spectrometry (FTICR-MS), using procedures described in detail in our previous studies of actinide sulphides,\(^{18,26}\) as well as in the ESL.\(^\dagger\) The An\(^{2+}\) ions were produced by direct laser desorption/ionization of small pieces of alloys that contained \(\sim 20\%\) of Th or \(\sim 5\%\) of the remaining actinide metals (Np, Pu, Am) in a Pt matrix. The COS reagent, a commercial product with \(>99\%\) purity as confirmed by electron ionization mass spectra, was introduced into the spectrometer through a leak valve to constant pressures of \((1–3)\) Torr.\(^{12}\)\(^{13}\) The reactant An\(^{2+}\) ions or sequential product ions were isolated through a leak valve to constant pressures of \((1–3)\) Torr. The reactant An\(^{2+}\) ions or sequential product ions were isolated and product ion intensities were monitored as a function of the reaction time. The reactant ions were thermalized by collisions with argon, maintained at constant pressures in the range of \((1–5) \times 10^{-6}\) Torr. Rate constants, \(k\), were determined from the pseudo-first-order decay of the relative signals of the reactant ions as a function of time at constant neutral pressures. Reaction efficiencies are described as \(k/k_{\text{COL}}\), where the \(k_{\text{COL}}\) is the collisional rate constant (see the ESL\(^\dagger\)).

**Computational details**

Complete active space self-consistent field (CASSCF)\(^{28,29}\) calculations were performed using the MOLCAS 8.0 package\(^{30}\) to generate a reference space (CAS\([n,m]\)), which consists of \(n\) electrons in the \(m\) molecular orbitals that describe the essential static correlation of the zeroth order wavefunction. All atoms were described with Atomic Natural Orbital\(^{31}\) basis sets of the relativistic core contracted (ANO-RCC\(^{32–34}\)) type. These calculations involved a default atomic mean field integral\(^{35}\) (AMFI) computation with a second-order Douglas-Kroll–Hess Hamiltonian (DKH2).\(^{36,37}\) The following contraction schemes were used: for Th, \(27s24p18d14f6g3h \rightarrow 9s8p6d4f2,\) for U, Np and Pu, \(26s23p17d13f5g3h \rightarrow 9s8p6d4f2g,\) and \(17s12p5d \rightarrow 4s3p1d\) for S. CASPT2 numerical gradients were performed without symmetry constraints to obtain stationary points on the lowest energy root within each multiplicity starting from a previously B3LYP optimised guess of the highest spin state of every system. Convergence thresholds were set to \(10^{-3}\) and \(10^{-6}\) a.u. for coordinate and gradient change respectively. The Cholesky\(^{38}\) integral decomposition technique was chosen to speed up the optimisations. The default IPEA value of 0.25 a.u. was used for the zeroth order wavefunction in perturbative calculations. An imaginary level shift of 0.1 a.u. was imposed in the CASPT2 stage to alleviate the intruder state problem.

The triangular \(\eta^5\)-AnS\(_2^{2+}\) species underwent a CASPT2(4,8) (An = U), CASPT2(5,8) (An = Np) and CASPT2(6,8) (An = Pu) state specific geometry optimisation for every spin state. The linear ThS\(_2^{2+}\) and US\(_2^{2+}\) species are closed shell with a significant HOMO-LUMO gap so an optimisation at the CASPT2 level becomes unnecessary. Therefore, for computational efficiency, an MP2 optimisation was performed on these diamagnetic species and the CASPT2 calculations run on top of the optimised geometry. As is the case with active spaces with many doubly occupied or empty MOs (NOOs > 1.98, NOOs < 0.06) some inactive or external MOs outside of our model space intruded the CAS. To keep the correlation treatment consistent a rotation constraint was imposed on these intruder orbitals to maintain them outside the CAS via the Supsym keyword. This produced only a slight change in the final PT2 energy and a very small decrease in the weight of the zeroth order wavefunction of the perturbed solution. The An = U, Np, Pu linear cations underwent geometry optimisations that included all the bonding and anti-bonding combinations of the 3p orbitals. A single point evaluation with the same \(n,13\) active space run on the remainder of all \(\eta^5\)-AnS\(_2^{2+}\) isomers to assess the linearization energies.

Symmetrised models of each stationary point underwent a single point CASPT2(\(n,13\)) run with \(C_{2v}\) symmetry in order to obtain the ground state term symbol of each model in this point group (see ESL\(^\dagger\)).

**Results and discussion**

**Gas-phase synthesis of actinide disulphide dications**

Gas-phase reactions of dipositive actinide cations, An\(^{2+}\) (An = Th, Np, Pu, Am), with COS were examined by FTICR-MS, with a focus on the formation of disulphide dications, AnS\(_2^{2+}\) and for comparison with the previously studied U\(^{2+}\).\(^{18}\) As in the case of uranium, carbonyl sulfide was selected because it is a thermodynamically favorable sulfur-atom donor (\(D(\text{OC-S}) = 73.6 \pm 0.2\) kcal mol\(^{-1}\) and \(D(\text{O-CS}) = 160.6 \pm 0.5\) kcal mol\(^{-1}\))\(^{39}\) and also because it has a rather high ionization energy (IE(COS) = 11.18 eV)\(^{39}\) that could disfavor electron-transfer pathways in the reactions with An\(^{2+}\) ions.
AnS$_2^{2+}$ ions were produced for An = Th and Np in a sequential abstraction process of two sulfur atoms by the An$_2^{2+}$ ions, similarly to what was previously observed for U$_2^{2+}$. For An = Pu and Am, An$_2^{2+}$ ions were reactive with COS (within the detection limit, $k_{k_{\text{COL}}} \leq 0.005$) and did not yield any sulphide species. Fig. 1 and 2 show representative mass spectra obtained in the reactions of Th$_2^{2+}$ and Np$_2^{2+}$, respectively.

Table 1 summarizes the product distributions, rate constants and efficiencies for primary (An$_2^{2+}$) and secondary (AnS$_2^{2+}$) reactions with COS; the results previously obtained for the case of uranium are included for comparison. The reaction efficiencies of the primary An$_2^{2+}$ ions followed the order: Th$_2^{2+}$ > U$_2^{2+}$ > Pu$_2^{2+}$ ≈ Am$_2^{2+}$ ≈ 0, a reactivity trend that has been observed before in the reactions of An$_2^{2+}$ cations with other neutral molecules, namely oxidants and hydrocarbons.40,41 Besides An$_2^{2+}$, primary formation of AnO$_2^{2+}$ was observed for Th (and U) but not for Np; these results are in accord with $D(\text{An}_2^{2+} - \text{O}) > D(\text{O-CS})$ for Th (and U) but not for Np.42 Interestingly, the reaction efficiencies of the primary AnS$_2^{2+}$ ions followed the order: Np$_2^{2+}$ > U$_2^{2+}$ > Th$_2^{2+}$.

As observed previously for uranium,18 the new AnS$_2^{2+}$ ions (An = Th, Np) reacted further with COS to form AnS$_3^{2+}$ species. The reaction of ThS$_2^{2+}$ could also be examined and, like uranium, yielded the tetrasulphide, ThS$_4^{2+}$, while the reaction of NpS$_3^{2+}$ could not be studied due to poor ion signal. Details on these reactions (product distributions and kinetics) are presented in the ESI† (Table S1).

To summarize the experimental results, new actinide disulphide dications, AnS$_2^{2+}$, could be effectively prepared for An = Th and Np. However, gas-phase synthesis of a species with a specific composition and charge does not directly reveal structure and bonding. In our previous study of uranium sulphide cations,18 DFT computations indicated that the ground state of the US$_2^{2+}$ ion has a triangular structure, with the linear thiouranyl isomer 40.9 kcal mol$^{-1}$ higher in energy. For the produced ThS$_2^{2+}$ ion, the scenario should be different, as Th can only achieve a maximum oxidation state of +4, and, therefore, the formation of a stable ThV persulphide can be anticipated. For the case of NpS$_2^{2+}$, it is possible that a linear thioneptunyl is produced or, like in the case of US$_2^{2+}$, a triangular species with a side-on η$_1$-S$_2$ ligand. To assess the structure and bonding of the new AnS$_2^{2+}$ species obtained for An = Th, Np, as well as for An = Pu to uncover trends along the An series, and for An = U to complement the previous study by DFT,18 we performed CASSCF computations that are presented in the following section.

### Structure and bonding of actinide disulphide dications

Since the crystal field of the 5f orbital environment in the actinide dications – herein they shall be named 5f$\alpha$, 5f$\beta$, 5f$\gamma$, 5f$\delta_1$, 5f$\delta_2$, 5f$\delta_1$, 5f$\delta_2$ (see Fig. 3 below and ESIF for their representation) – produces very small splittings, the degree of static correlation will be significant and the high symmetry of these species will render it difficult or unreliable the use of single determinant methods such as DFT to reproduce the electronic structure of these species. Moreover, the small size of the systems is an advantage from a computational perspective and as such we performed a perturbative wavefunction geometry optimisation and analysis of actinide disulphide dications, AnS$_2^{2+}$. Pre-trials were conducted to assess the most
adequate active space for the triangular and linear species. For the An = U, Np, Pu linear species, optimisations were performed at the CASPT2(12,13) and CASPT2(13,13) levels. The triangular isomers required a smaller active space (CASPT2(4,8), CASPT2(5,8) and CASPT2(6,8)) due to their lower symmetry which breaks the degeneracy of the \( \pi \) orbitals (see Computational details).

**ThS\(_2\)\(^{2+}\) isomer pair**

We begin with the simplest system, \( \eta^2-\text{ThS}_2\)\(^{2+}\), where the most plausible bonding scenarios are the binding of two \( \text{Th}^{2+} + (\times)\text{S}^2\)\(^-\) fragments resulting in a triplet state, or alternatively a closed shell molecule partitioned as \( \text{Th}^{4+} + \text{S}_2^{2-} \) (persulphide), in view of the significant stability of the +4 oxidation state of Th.\(^3\) A CASPT2(2,2) optimization for the ground-state singlet and triplet were performed and the resulting energy differences unequivocally favour a closed shell singlet. The triplet is 29.7 kcal mol\(^{-1}\) higher in energy and consists of a singly occupied \( \pi^* \) MO from the di-sulfide unit plus one \( 6\delta_{2\pi} \) orbital holding the remaining electron. A re-optimised structure with the MP2 method provides a geometry which is very close to the one obtained with CASPT2(2,2) and with identical reference weights (0.8) in the perturbed wavefunction confirming the lack of multi-reference character of the ground-state singlet (which is 98.7% mono-determinantal). The calculated CASPT2(2,2) equilibrium distances of the ground state are \( d(\text{Th-S}) = 2.477 \) Å and \( d(\text{S-S}) = 2.173 \) Å and those with MP2 \( d(\text{Th-S}) = 2.480 \) Å and \( d(\text{S-S}) = 2.169 \) Å. For comparative purposes a single point evaluation using the CASPT2(10,13) method was used and the linearization energy evaluated to be +59.4 kcal mol\(^{-1}\).

The analogous MP2 calculation as expected provided a very similar energy (56.2 kcal mol\(^{-1}\)).

**US\(_2\)\(^{2+}\) isomer pair**

For the \( \eta^2-\text{US}_2\)\(^{2+}\) system the most suitable active space was found to be the seven 5f orbitals plus the out of plane \( \pi_1^- (2p_x \pm 2p_y) \) linear combination set of the \( \text{S}_2 \) fragment (Fig. 3). The lowest singlet, triplet and quintet states were optimised at the CAS(4,8)PT2 level. The results are summarised in Table 2. One of the more interesting aspects is the low energy value of the quintet state with respect to the ground-state triplet which had already been an important conclusion in a preceding paper.\(^18\) This quintet state can be described as a \( \text{U}^{3+}(5f^3) + \text{S}_2^{2-} \) ferromagnetic interaction as evidenced by the natural orbital occupancies (NOOs). The 5f\(_{6\delta}\) and the \( \pi^- \) MOs do not mix as in the lower spin states as these NOOs remain localised.

In the triplet ground state there is some mixing between the 5f\(_{6\delta}\) and the \( \pi^- \) orbital from sulphur. The CI coefficients of the CASPT2(4,8) first order wavefunction show a strong multi-reference character: triplet state has the leading (30%) configuration \( \lambda_1 = \{ (5f_1)^1 (5f_2)^0 \} \{ (5f_6)^0 (5f_6^- + \pi_1^-)^2 \} \{ (5f_8)^0 \} \{ (5f_{10})^0 \} \{ (5f_{12})^0 \} \{ (5f_{14} - \pi_1^-)^0 \} \) followed by the configuration state function (CSF) with an additional open shell 23% \( \lambda_2 = \{ (5f_1)^1 \} \{ (5f_6)^0 \} \{ (5f_6^- + \pi_1^-)^1 \} \{ (5f_8)^0 \} \{ (5f_{10} + \pi_1^-)^0 \} \{ (5f_{12})^0 \} \{ (5f_{14} - \pi_1^-)^0 \} \). The makeup of the \( \lambda_3 \) CSF is two determinantal: \( 1/2 \{ (5f_6^- + \pi_1^-)^1 \} - 1/2 \{ (5f_6^- + \pi_1^-)^0 \} \) where the minus sign shows that this is the \( M_e = +1 \) component of the \( |S = 1; S_1 = 3/2; S_2 = 1/2 \rangle \) coupled state rather than from \( |S = 2; S_1 = 3/2; S_2 = 1/2 \rangle \). The natural spin density of the unperturbed CAS wavefunction, shown in Fig. 4, highlights the spin coupling at the different An(5f) and \( \pi^- \) sites.

The singlet state is 16.2 kcal mol\(^{-1}\) higher in energy and presents a complex electronic structure. Its leading configuration (34%) is \( \lambda_1 = \{ (5f_6)^1 \} \{ (5f_8)^0 \} \{ (5f_6^- + \pi_1^-)^2 \} \{ (5f_{10})^0 \} \{ (5f_{12})^0 \} \{ (5f_{14} - \pi_1^-)^0 \} \) followed by \( \lambda_2 = \{ (5f_1)^1 \} \{ (5f_6)^0 \} \{ (5f_{10} + \pi_1^-)^0 \} \{ (5f_{12})^0 \} \{ (5f_{14} + \pi_1^-)^0 \} \) (14%) admixed with several other configurations each below 10% in overall weight.

### Table 2 Energies, bond distances and NOOs of the optimised \( \eta^2-\text{US}_2\)\(^{2+}\) species in several spin states

<table>
<thead>
<tr>
<th>Total spin</th>
<th>0</th>
<th>1</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \Delta E / \text{kcal mol}^{-1} )</td>
<td>+16.2</td>
<td>0</td>
<td>+2.8</td>
</tr>
<tr>
<td>( d(U-S)/\text{Å} )</td>
<td>2.577</td>
<td>2.599</td>
<td>2.630</td>
</tr>
<tr>
<td>( d(S-S)/\text{Å} )</td>
<td>2.060</td>
<td>2.055</td>
<td>2.049</td>
</tr>
</tbody>
</table>

**Natural orbital occupations [NOOs]**

<table>
<thead>
<tr>
<th>(5f(_{6\delta})(^{1+})) ( \pi_1^- )</th>
<th>1.160</th>
<th>1.110</th>
<th>0.988(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5f(_{6\delta})</td>
<td>0.676</td>
<td>0.910</td>
<td>0.030</td>
</tr>
<tr>
<td>5f(_{6\delta}^-)</td>
<td>0.645</td>
<td>0.788</td>
<td>0.223</td>
</tr>
<tr>
<td>5f(_{8\delta})</td>
<td>0.173</td>
<td>0.084</td>
<td>0.888</td>
</tr>
<tr>
<td>5f(_{10\delta})</td>
<td>0.196</td>
<td>0.170</td>
<td>0.767</td>
</tr>
<tr>
<td>5f(_{12\delta})</td>
<td>0.275</td>
<td>0.205</td>
<td>0.822</td>
</tr>
<tr>
<td>5f(_{14\delta}-\pi_1^-)</td>
<td>0.543</td>
<td>0.692</td>
<td>0.171(^a)</td>
</tr>
<tr>
<td>5f(_{14\delta})</td>
<td>0.279</td>
<td>0.016</td>
<td>0.085</td>
</tr>
</tbody>
</table>

\(^a\) Fully localised orbital, orbital in brackets not present.
The non-integer orbital occupations are a consequence of the near degeneracy of one or more states that is present in these highly symmetric systems, specifically the 5f orbitals that are degenerate. A multi-state analysis would resolve all these states but a spectroscopic analysis is beyond the scope of this work.

Both the triplet and singlet states have a 5fδl + π1− MO composition that is approximately 11% 5fδl + 89% π1− (S) whereas its anti-bonding counterpart 5fδl − π1− is 87% 5f + 14% π− (S) (see ESI†). Performing a 2 × 2 rotation of the 5fδl ± π1− orbital pair yields the orthogonal localised 5fδl and π1− MO pair, which have the occupation numbers of 0.08 and 1.12 respectively. Both these spin states show evidence of exhibiting a side-on S2− radical fragment which gradually donates some electronic density as the spin state gets higher. In the quintet state, as seen from Table 2, the (localised) π1− MO will lower its NOO to 0.988 involving an approximate back-donation increase of 0.13 e− from the triplet to the quintet.

From an orbital interaction perspective, the most important interactions in η2-US22+− are between the antibonding π1 and π2 set of di-sulphur and their linear combination with the 5f and 6d functions of uranium (Scheme 1).

Given that the 6d + π2− MO is in the inactive group of orbitals its NOO is precisely 2.0. The Mayer–Mulliken bond order is a quantitative measure of a bond strength and will be used in this work to assess these magnitudes and serve to compare the compounds under analysis. The U–S bond order calculated from the CASPT2(4,8) density in the 3X state is 0.997 while the S–S bond has a value of 1.078.

Some notable aspects emerge when comparing the bonding scenario with the analogous η2-UO22+ species to help understand the preference of the uranyl cation for linearity and the thiolated uranium cations for the chelated form. To this aim we also optimised the triplet η2-UO22+ geometry, which lies 5 kcal mol−1 below the quintet state, and performed a comparative orbital analysis. Owing to the shorter O–O distance, the 6d/π2− overlap and back-donation is not as effective in η2-UO22+.

This becomes apparent from the analysis of the relevant MO compositions in both systems. Whereas in η2-US22+ the natural orbital is composed of the following atomic orbitals:

\[ \Psi_{trp}(6d + \pi_2^-) = 0.135\phi_U(6p_x) + 0.366\phi_U(6d_d) + 0.599\phi_{S_1}(3p) - 0.598\phi_{S_2}(3p) + \ldots \]

in η2-UO22+ the 6d contribution decreases while the oxygen 2p orbitals have a slightly higher contribution:

\[ \Psi_{trp}(6d + \pi_2^-) = 0.270\phi_U(6p_x) + 0.269\phi_U(6d_d) + 0.661\phi_{O_1}(2p) - 0.691\phi_{O_2}(2p) + \ldots \]

where the 6pπ orbital in uranium though filled is more accessible to overlap with the π2− group MO.

Two important points from this analysis of the η2-US22+ and η2-UO22+ cations become evident: (1) a π type of overlap between the 6d orbital with the π2− MOs (as shown in Scheme 1) is energetically more stabilising than it would be in a linear [SUS]2+ species since the corresponding 6d/3p overlap would then only have π symmetry; (2) U–O bonds are not stable in a side-on arrangement since the 6d orbitals can never have a significant overlap with the 2p set because of the electronegativity of the oxygen atoms.

The NOOs in the 5fδl ± π− set do not change to a significant extent between η2-UO22+ and η2-US22+ and neither does their composition (5fδl ± π1− = 25% 5fδl + 75% π1−; 5fδl ± π1− = 65% 5fδl + 35% π1−). The Mayer–MullikenCASPT2 bond orders are 0.67 for U–O and 0.96 for O–O. The U–O bond is indubitably weaker than that of U–S in these chelates.

The closed shell [SUS]2+ species affords no special difficulty in its computational treatment. The calculated U–S equilibrium bond length is 2.193 Å (Table 3) and the (S–U–S) angle is precisely 180°. The CASPT2(12,13) electronic energy difference of linear [SUS]2+ with respect to the dihapto species

<table>
<thead>
<tr>
<th>ΔE(kcal mol−1)</th>
<th>d(U–S)/Å</th>
</tr>
</thead>
<tbody>
<tr>
<td>8.4</td>
<td>2.193</td>
</tr>
</tbody>
</table>

Table 3 NOOs and single point energy of the closed shell linear [SUS]2+ with respect to the 3X ground state of η2-US22+.
is +8.4 kcal mol$^{-1}$, a comparatively smaller figure than the previous$^{18}$ DFT value of +40.9 kcal mol$^{-1}$.

### NpS$_2$$_2^+$$^-$$^+$$^-$ isomer pair

With the neptunium system an analogous chemical bonding picture emerges (Table 4), the only difference being the electron count. The ground state is a quadruplet ($^4$X), the sextet state is located slightly higher in energy (3.4 kcal mol$^{-1}$) with respect to the triplet-quintet gap (2.8 kcal mol$^{-1}$) in $\eta^2$-US$_2$$^+$. In the sextet state the 5f$_{3/2}$ orbital also does not mix with the $\pi_2^-$ group MO which leads to a ferromagnetic Np$^3+$[5f$^6$] + S$_2$ coupling analogously to the $\eta^2$-US$_2$$^-$ quintet state.

Upon orbital localisation of the $^4$X CASPT2(5,8) wavefunction the occupation numbers of the two 5f$_{3/2}$ and $\pi_2^-$ orbitals are 0.120 and 1.12 respectively, values similar to the ones previously obtained for the $\eta^2$-US$_2$$^-$ cation. Indeed, the only significant change with respect to the $\eta^2$-US$_2$$^-$ species is that there is one additional electron in the 5f$_{3/2}$ atomic orbital:

- 34% $^4$w$_{1}$ = ((5f$_{3/2}$)$^1$ (5f$_{3/2}$)$^9$ (5f$_{3/2}$ + $\pi_2^-$)$^2$ (5f$_{5/2}$)$^1$ (5f$_{5/2}$)$^0$ (5f$_{5/2}$ + $\pi_2^-$) + 18% $^4$w$_{1}$ = ((5f$_{5/2}$)$^1$ (5f$_{5/2}$)$^9$ (5f$_{5/2}$ + $\pi_2^-$)$^2$ (5f$_{3/2}$)$^1$ (5f$_{3/2}$)$^0$ (5f$_{5/2}$ + $\pi_2^-$) which is entirely analogous to the $^2$A$_2$$^+$ CSF in $\eta^2$-US$_2$$^+$. This spin coupling can be demonstrated by the CASSCF natural spin density shown in Fig. 5.

Since nothing is known about the linear [SNpS]$^2^+$ structure a survey of the most chemically intuitive spin states was undertaken (Table 5). The lowest energy micro-state is also a quartet but situated some +31.8 kcal mol$^{-1}$ above the ground state of $\eta^2$-NpS$_2$$^+$. This linear structure has an equilibrium distance of 2.256 Å and the (S-Np-S) angle is very close to 180°.

![Fig. 5 CASSCF(5,8) spin density plot obtained from the natural spin orbitals of the $^4$X state of $\eta^2$-NpS$_2$$^+$. Positive spin density is blue, negative spin density is yellow.](image)

### PuS$_2$$_2^+$$^+$ isomer pair

We focus now on the analysis of the isomers of the PuS$_2$$^+$. This state comes as a natural extension of the electronic structure seen in $\eta^2$-US$_2$$^+$ and $\eta^2$-NpS$_2$$^-$ (triplet and quartet, respectively). The difference now is that little di-radical coupling is present between the $\pi$ antibonding MOs in di-sulphur and the remainder of the 5f manifold. This ground state is more complex in its configurational make-up than has been described heretofore. The leading CASPT2 CSF is $^5$w$_{1}$ = ((5f$_{5/2}$)$^1$ (5f$_{5/2}$)$^9$ (5f$_{5/2}$ + $\pi_1$)$^2$ (5f$_{5/2}$)$^1$ (5f$_{5/2}$)$^0$ (5f$_{5/2}$ + $\pi_1$) + 16% $^5$w$_{1}$ = ((5f$_{5/2}$)$^1$ (5f$_{5/2}$)$^9$ (5f$_{5/2}$ + $\pi_1$)$^2$ (5f$_{5/2}$)$^1$ (5f$_{5/2}$)$^0$ (5f$_{5/2}$ + $\pi_1$) which is entirely analogous to the $^2$A$_2$$^+$ CSF in $\eta^2$-US$_2$$^+$. This spin coupling can be demonstrated by the CASSCF natural spin density shown in Fig. 5.

The energy of the lower spin states is much higher, 44 and 66 kcal mol$^{-1}$ above the ground state for the triplet and singlet (Table 6), respectively.

Analogously to the $\eta^2$-US$_2$$^+$ cation, a low lying higher spin state exists in the form of a septet which is pratically...
The majority contribution stems from the \((5f_{s}^1 m_1)(5f_{p}^1 m_2)(5f_{d}^1 m_2)(5f_{f}^1 m_2)(5f_{d}^2 m_2)(5f_{f}^2 m_2)(5f_{p}^1 m_2)(5f_{p}^2 m_2)(5f_{s}^1 m_2)\) configuration state function and trailing below it in predominance is \((5f_{s}^1 m_2)(5f_{p}^1 m_2)(5f_{d}^1 m_2)(5f_{f}^1 m_2)(5f_{d}^2 m_2)(5f_{f}^2 m_2)(5f_{p}^1 m_2)(5f_{p}^2 m_2)(5f_{s}^1 m_2)\) (37%) where now the 5f\(_{d}\) orbitals do not mix as seen in the highest spin states of the previous disulphides.

The linear \([\text{SPuS}]^{2+}\) dication has also never been addressed computationally so three spin states \(S = 0, 1\) and \(2\) were optimised to ascertain the nature of its ground state and compare it to the triangular species in the extended active space. Although the linear species tend to have lower spin states due to the An–R_S triple bonds, in the case of plutonium this is no longer the case since the 5f\(_{s} + \text{s}^0_{C0}\) MO becomes partially occupied just as it was seen in \([\text{SNpS}]^{2+}\) and three other 5f orbitals are singly occupied (Table 7). The triplet state is doubly degenerate under \(C_{2v}\) symmetry (3A\(_{2}\) + 3B\(_{2}\), see ESI†) and is 1.8 kcal mol\(^{-1}\) higher in energy and quasi-degenerate with the quintet state (5A\(_{2}\)). The latter resolution into a single state explains why the latter NOOs are close to either 0, 1 or 2.

### Global view

The most significant conclusion of these calculations is that the thermodynamic energy for the linearization process increases fourfold upon changing from U to Pu (Table 8). Presumably this trend will be more pronounced for the later actinides since orbital mixing is expected to take on a greater role.\(^{19}\) Additionally, the depopulation of an An–S\(_{\text{s}}\) σ bond in the linear species is already present in the Np and Pu sulfides. This finding

### Table 6

<table>
<thead>
<tr>
<th>Total spin</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\Delta E/(\text{kcal mol}^{-1}))</td>
<td>+66.1</td>
<td>+44.3</td>
<td>0</td>
<td>+0.7</td>
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<tr>
<td>(d(\text{Pu–S})/\text{Å})</td>
<td>2.552</td>
<td>2.539</td>
<td>2.579</td>
<td>2.581</td>
</tr>
<tr>
<td>(d(\text{S–S})/\text{Å})</td>
<td>2.050</td>
<td>2.047</td>
<td>2.050</td>
<td>2.041</td>
</tr>
</tbody>
</table>

### Table 7

<table>
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<th>Total spin</th>
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<th>1</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\Delta E/(\text{kcal mol}^{-1}))</td>
<td>+32.8</td>
<td>+1.8</td>
<td>0</td>
</tr>
<tr>
<td>(d(\text{Pu–S})/\text{Å})</td>
<td>2.301</td>
<td>2.279</td>
<td>2.279</td>
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</table>

### Table 6

<table>
<thead>
<tr>
<th>Energies, NOOs and bond distances of the optimised di-haptic plutonium disulphide species [CASPT2(6,8)] in several spin states</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total spin</td>
</tr>
<tr>
<td>(\Delta E/(\text{kcal mol}^{-1}))</td>
</tr>
<tr>
<td>(d(\text{Pu–S})/\text{Å})</td>
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<tr>
<td>(d(\text{S–S})/\text{Å})</td>
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### Table 7

<table>
<thead>
<tr>
<th>NOOs, bond distances and optimised energies [CASPT2(14,13)] of linear PuS(_{2}^{2+}) with respect to the (\tilde{X}) ground state</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total spin</td>
</tr>
<tr>
<td>(\Delta E/(\text{kcal mol}^{-1}))</td>
</tr>
<tr>
<td>(d(\text{Pu–S})/\text{Å})</td>
</tr>
</tbody>
</table>

### Notes

- Localised orbital, orbital in brackets not participating.
defies conventional chemical intuition such that in the linear structures the π bonds are resilient and σ bonds are weakened. Since σ overlap is no longer as efficient as in the early actinides, a side-on coordination will become energetically more rewarding. From an empirical viewpoint, the decreasing stability of high oxidation states beyond Pu supports this analysis.

To explain the observed trend in the linearization energy two factors must be taken into account: one is the strength of the S–S bonds in the triangular structure versus the strength of the An–S bonds in the linear structures. If the former is constant, the latter will dictate the trend in isomer conversion energy. Indeed, a linear correlation can be established between the difference in bond order which is indeed significant due to the change in bond order. This energy change is incomparably high. This may not just be because the S–S bond is stronger than a single O–O bond. On this basis there is an additional incentive for the uranyl species to remain linear.

As reported above, the polysulphides ThS₄⁺, NpS₃⁺ and ThS₄⁺ were also synthesized by sequential abstractions of S atoms from COS. Although these species were not calculated, it is reasonable to surmise regarding possible structures. The computed structures of US₄⁺ and US₂⁺ were reported in ref. 18. Given that these previous DFT calculations predicted the same triangular triplet structure for US₂⁺ as was computed at the higher level of theory employed here, it is expected that the previously computed ground state structures for US₂⁺ and US₄⁺ are also valid; both are triplet, with the former having one η²-coordinated S₂ moiety, and the latter having two η²-coordinated S₃ moieties. It is surmised that the structures of ThS₃⁺ and NpS₂⁺ similarly have an S₃ moiety that is η²-coordinated to the metal center. The proposed ground state structures for ThS₃⁺ and NpS₂⁺ might be accessible from the ground state structures of the disulphides via insertion of an S atom into an An–S or S–S bond; both insertions likely present substantial barriers such that the experimentally produced trisulphides may not exhibit the lowest energy structures. It should be remarked that formation of triangular AnS₂⁺ from AnS²⁺ by addition of an S atom does not require An–S or S–S bond cleavage and should thus present a substantially lower, perhaps negligible, kinetic barrier. The structure of ThS₄⁺ may be analogous to the [[(η⁵S₂)U(η⁵S₂)]²⁺ structure reported in ref. 18, although the alternative [(η⁵S₃)Th]²⁺ structure cannot be excluded. The triplet spin state of US₂⁺ suggests a formally U⁴⁺ oxidation state but the U–S bond distances indicate four U–S single bonds and a potentially higher oxidation state, which is inaccessible in the case of Th for which the tetravalent oxidation state is realistically the highest. As with the trisulphides, formation of tetrasulphides by S atom addition may present sufficiently high barriers that the structure obtained under low-energy experimental conditions is not necessarily the computed ground state structure.

Conclusions

The gas-phase synthesis of new actinide disulphide dications, AnS₂⁺ (An = Th, Np), was accomplished by sequential reactions of An²⁺ ions with COS; for An = Pu and Am, An²⁺ ions were unreactive. CASPT2 computations were performed to assess the structures and bonding of the new AnS₂⁺ ions [An = Th, Np], of PuS₂⁺ to uncover trends along the An series, and of US₂⁺, previously studied experimentally and examined by DFT. These computations revealed that, like in the case of uranium, the AnS₂⁺ ions have ground-state structures of the metallacycle type, with the presence of a persulphide ligand in η²-ThS₂⁺, formally corresponding to a Th(Ⅳ) species, and of supersulphide ligands in US₂⁺, NpS₂⁺, and PuS₂⁺, formally corresponding to An(Ⅲ) species. Molecular orbital and energetic analysis of the dicationic species reveal that the existence of linear actinide di-chalcogenides are likely confined to the uranyl ([UO₂⁺)]

![Fig. 8 Difference in An–S Mayer–Mulliken bond order between the linear and the triangular structures taken as a function of the isomer conversion energy.](image-url)
species due to three factors: the destabilising effect that the poor 2p → 6d donating ability exerts upon the system with side-on coordination ([n^2-UO_2]^{2-}), the stronger σ bond present in supersulphide (S_2^{2-}) stabilising the triangular species with respect to the linear isomer and finally the depopulation of the 5f^2 + σ^* bonding orbitals in the ground states of the [SAnS]^{2+} species which weaken the strength of this interaction and de-stabilise the linear isomers.

The An(n+1) reduction potentials clearly indicate a significant stability of Th(n) species and an increasing relative stability of the An(n) when moving from U to Pu, culminating in a clear predominance of Am(n). Linear thiaocenyl structures are higher in energy, with a difference that increases fourfold upon changing from U to Pu. An inverse correlation was found concerning the difference in the An–S bond order between linear and triangular structures and the linearization energy, showing that the di-sulphide side-on coordination gains strength thus stabilising the triangular isomers with heavier actinides.

Acknowledgements

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