Pseudohalide tectons within the coordination sphere of the uranyl ion: An experimental and theoretical study of C—H···O, C—H···S and chalcogenide non-covalent interactions.

Stefano Nuzzo,¹ Brendan Twamley,¹ James A. Platts,² and Robert J. Baker¹*

1 School of Chemistry, University of Dublin, Trinity College, Dublin 2, Ireland

2 School of Chemistry, Main Building, Cardiff University, Park Place, Cardiff, CF10 3AT, U.K.
ABSTRACT. A series of uranyl thiocyanate and selenocyanate of the type \([R_4N]_3[UO_2(NCS)_5]\) \((R_4 = \text{^n}Bu_4, \text{Me}_3\text{Bz}, \text{Et}_3\text{Bz})\), \([\text{Ph}_4\text{P}][UO_2(NCS)_3(NO_3)]\) and \([R_4N]_3[UO_2(NCSe)_5]\) \((R_4 = \text{Me}_4, \text{^n}\text{Pr}_4, \text{Et}_3\text{Bz})\) have been prepared and structurally characterized. The resulting non-covalent interactions have been examined and compared to other examples in the literature. The nature of these interactions is determined by the cation so that when the alkyl groups are small, chalcogenide···chalcogenide interactions are present, but this ‘switches off’ when \(R = \text{^n}\text{Pr}\) and charge assisted \(U=O\cdots H—C\) and \(S(e)\cdots H—C\) hydrogen bonding remain the dominant interaction. Increasing the size of the chain to \(^n\text{Bu}\) results in only \(S\cdots H—C\) interactions. Spectroscopic implications of these chalcogenide interactions have been explored in the vibrational and photophysical properties of the series \([R_4N]_3[UO_2(NCS)_5]\) \((R_4 = \text{Me}_4, \text{Et}_4, \text{^n}\text{Pr}_4, \text{^n}\text{Bu}_4, \text{Me}_3\text{Bz}, \text{Et}_3\text{Bz})\), \([R_4N]_3[UO_2(NCSe)_5]\) \((R_4 = \text{Me}_4, \text{^n}\text{Pr}_4, \text{Et}_3\text{Bz})\) and \([\text{Et}_4N]_4[UO_2(NCSe)_5][\text{NCSe}]\). The data suggest that \(U=O\cdots H—C\) interactions are weak and do not perturb the uranyl moiety. While the chalcogenide interactions do not influence the photophysical properties, a coupling of the \(U=O\) and \(\delta(\text{NCS})\) or \(\delta(\text{NCSe})\) vibrational modes is observed in the 77K solid state emission spectra. A theoretical examination of representative examples of \(\text{Se}^\cdots\text{Se}, \text{C}—\text{H}^\cdots\text{Se}\) and \(\text{C}—\text{H}^\cdots\text{O}=\text{U}\) by molecular electrostatic potentials, NBO and AIM methodologies give a deeper understanding of these weak interactions. \(\text{C}—\text{H}^\cdots\text{Se}\) are individually weak but \(\text{C}—\text{H}^\cdots\text{O}=\text{U}\) interactions are even weaker supporting the idea that the –yl oxo’s are weak Lewis bases. An Atoms in Molecules study suggest that the chalcogenide interaction is similar to lone pair…\(\pi\) or fluorine…fluorine interactions. An oxidation of the NCS ligands to form \([(UO_2)(\text{SO}_4)_2(\text{H}_2\text{O})_4].3\text{H}_2\text{O}\) was also noted.

KEYWORDS: Non-covalent interactions; uranyl chemistry; photoluminescence; AIM
**Introduction**

An understanding of the plethora of non-covalent interactions is of importance, not just in structural chemistry but also materials and biological chemistry. While the hydrogen bond is the most well studied, over the past few years other non-conventional noncovalent interactions have garnered significant amount of attention, such as halogen bonding,\(^1\) pnictogen bonding,\(^2\) and even aerogen bonding.\(^3\) Chalcogenide bonding\(^4\) is a relatively recent addition to the toolbox of crystal engineering and follows the same type of bonding scheme as halogen bonding, namely a $\sigma$-hole.\(^5\)

This bonding scheme can be viewed comparably to hydrogen bonding as $X - D \cdots A$, where $X$ is any atom, $D$ is the donor atom and $A$ is the acceptor atom. The $\sigma$-hole is visualized as a region of positive electrostatic potential found on an empty $\sigma^*$ orbital and is dependent upon two parameters: the $\sigma$-hole becomes more positive (and hence stronger interactions) when (1) $D$ is more polarizable and (2) when the $X$ atom is more electron withdrawing. The majority of chalcogenide bonding features attractive interactions between $S$ or $Se$ and an electronegative $N$ or $O$ donor. Computational studies on $Se\cdots O$ or $S\cdots N$ interactions have recently been conducted and shown them to be dependent on the substituents on the selenium atom and the interactions described as a charge transfer from the $N$ lone pair to the $\sigma^*$ $Se\cdots X$ orbital (Chart 1).\(^{4j,6}\) Synthetic molecular balances based on formamide and thioformamide units have confirmed this, and shown that chalcogenide...$H\cdots C$ interactions can also be described in this manner.\(^{4b}\) The atomic polarizability increases down the group so that $S = 19.3$ a.u. and $Se 25.4$ a.u. (a.u. = atomic units); thus a stronger interaction should occur when descending the group.\(^7\) Moreover, as the atomic radius increases steric effects become less pronounced as the group is descended. These interactions can be comparable in strength to hydrogen and halogen bonding. For example, the computed interaction energy of $FHS\cdots NH_3$ is -8.4 kcal mol$^{-1}$ compared to -11.3 kcal mol$^{-1}$ for $FHSe\cdots NH_3$ and -10.3 kcal
mol$^{-1}$ for the halogen bonded F—Cl···NH$_3$.$^5$d Perhaps more illuminating is the low temperature structure of the F$_4$S···NEt$_4$ adduct which displays an S—N bond length of 2.384(2) Å, typical for a dative covalent bond;$^8$ theory supports this hypothesis$^9$ and reports a binding energy of -14.4 kcal mol$^{-1}$.

Halogen interactions have been further delineated by the geometric parameters shown in Chart 1. 0°≤|θ$_1$-θ$_2$|≤15° are classified as type I; 15°≤|θ$_1$-θ$_2$|≤30° are quasi-type I/type II and 30°≤|θ$_1$-θ$_2$| are type II.$^{10}$ Type I are generally thought to be due to crystal packing, while Type II are stabilizing interactions. A recent examination of the Cambridge database showed that the majority of S···S interactions in sp$^2$ hybridized sulfur are Type I ($θ_1$ ≈ $θ_2$).$^{11}$ It is also apparent that S···O interactions are prevalent in molecular recognition and host-guest interactions in biological systems.$^{12}$ However few examples of chalcogenide interactions exist where a metal ion is included.$^{13}$ Indeed, even the more mature field of halogen bonding features limited examples, notably iodoform adducts of trans-[PtX$_2$(NCNR)$_2$]$_2$,$^{14}$ or in a uranyl organic framework based on halogen substituted isonicotinic acids.$^{15}$
**Chart 1.** Types of halogen and chalcogenide bonding.

### Halogen Bonding

**Type I**

\[
\theta_1 - \theta_2 = 0 - 15^\circ
\]

**Type II**

\[
\theta_1 - \theta_2 < 30^\circ
\]

\[
\theta_1 \sim \theta_2
\]

### Chalcogenide Bonding

\[
\theta_1 \sim 180^\circ
\]

\[
\theta_2 \sim 90^\circ
\]

Lone pair

**Chart 1.** Types of halogen and chalcogenide bonding.
The uranyl ion has a number of features that make it attractive for a study on non-covalent interactions. The coordination geometry is generally limited by the linear O=U=O (–yl) fragment so all coordination will be directed to the equatorial plane. Changing the equatorial ligands can influence the properties of the uranyl ion in catalytic applications or the formation of uranyl coordination polymers with great effect. Over the last few years the –yl moiety has been shown to act as a secondary directing influence in supramolecular assembly to form 3-dimensional structures. However, as the oxo groups are generally considered as poor Lewis bases these interactions have been somewhat neglected. Interactions with metal cations can elongate the U=O bond length to a degree that is noticeable crystallographically. Some evidence suggests that this weakening of the U=O bond is due to the increase in electron density around the uranyl cation and increase electrostatic repulsions within the U=O group. For weaker interactions, hydrogen bonding acceptor via C—H···O=U interactions, sometimes via charge assisted hydrogen bonding, can be used to recognize or separate the uranyl ion selectively from complex mixtures. More recently, the uranyl ion as a halogen bond acceptor has been observed. Finally, additional supramolecular interactions utilizing the equatorial ligands as tectons have been utilized. For example, the elegant studies using uranyl thiocyanate to explore non-covalent bonding in halogenated pyridinium cations showed some evidence for S···S close contacts. In relation to this we recently reported close contacts between two selenium atoms in [Et₄N]₄[UO₂(NCSe)₅][NCSe] (dₗS…Se = 3.427(1) Å). A series of [R₄N]₃[UO₃(NCS)₃] (R = Me, Et, nPr) have been structurally characterized as examples of species present in spent nuclear fuel reprocessing. Moreover, the uranyl thiocyanate ion has shown to have unusual thermochromic behavior in ionic liquids.

In this contribution we have extended our study of the coordination chemistry of the [NCSe]⁻ pseudohalide as a tecton in uranyl coordination chemistry, of which there is only one other
We also compare the uranyl selenocyanate chemistry to that of uranyl thiocyanate compounds which we and others have reported, and explore the other non-covalent interactions, namely charge assisted $\text{C—H} \cdots \text{O} = \text{U}$ and $\text{C—H} \cdots \text{S(e)}$ hydrogen bonding interactions. To do this we have utilized X-ray crystallography, vibrational and photoluminescence spectroscopy and theoretical methods to further our understanding of these weak interactions.

**Results and Discussion**

*Synthesis and Structural studies.*

The syntheses of all the compounds described herein are simple (Equations 1 and 2) and X-ray quality crystals grown for all. We will use the nomenclature $\text{RE}$, where $\text{R}$ is the alkyl group of the ammonium cation and $\text{E}$ is $\text{S}$ or $\text{Se}$ to distinguish these compounds. Thus we have a small library of compounds that we can compare structural parameters and packing motifs (i) within the NCS series, (ii) within the NCSe series and (iii) with the same cations. In one case we also obtained a small amount of a decomposition product presumably from $[\text{NCS}]^-$ oxidation and this has been structurally characterized. When the reaction of cyanates were examined, we consistently observed hydrolysis of the $[\text{NCO}]^-$ fragment and $\text{K}_4[\text{UO}_2(\text{CO}_3)_3]$ the only product obtained. In contrast, the reaction of $[\text{NCTe}]^-$ with uranyl nitrate afforded immediate decomposition to a grey precipitate of presumably tellurium metal and, from the acetonitrile soluble fraction, single crystals of $[\text{Et}_4\text{N}]_2[\text{UO}_2\text{Cl}_4]$ were deposited. $[\text{NCTe}]^-$ chemistry is dominated by this decomposition.

$$[\text{UO}_2(\text{NO}_3)_2].6\text{H}_2\text{O} + 5 \text{NaNCS} + 3\text{R}_4\text{NCl} \rightarrow [\text{R}_4\text{N}]_3[\text{UO}_2(\text{NCS})_5] \quad (1)$$

$$[\text{UO}_2(\text{NO}_3)_2].6\text{H}_2\text{O} + 5 \text{KNCS} + 3\text{R}_4\text{NCl} \rightarrow [\text{R}_4\text{N}]_3[\text{UO}_2(\text{NCSe})_5] \quad (2)$$
(i) $[R_4N]_3[O_2(NCS)_5]$ series ($R = Me, Et, "Pr, "Bu, Me_3NBz, Et_3NBz$)

The solid-state structures of MeS, EtS and $"PrS$ have been previously described, but the focus of that work was not on the supramolecular structures, whereas $"BuS, Me_3NBzS$ and $Et_3NBzS$ are new. An example of the structure of $Me_3NBzS$ is shown in Figure 1, while selected metric parameters for the RS series are reported in Table 1; other structural data are collated in Figure S1-S3 and Tables S1-S8. In all compounds the uranyl geometry is, as expected, pentagonal bipyramidal with a linear O=U=O moiety, and the U—N, N=C and C=S bond lengths are statistically identical. The only differences are in the orientation of how the $[NCS]$ ion coordinates to the uranium; in particular the U—N=C angle and the deviation of the S atoms out of the UN$_5$ plane, which diverges significantly.
Figure 1. Asymmetric unit of Me$_3$NBzS with atomic displacement parameters shown at 50% probability. Hydrogen atoms omitted for clarity.
Table 1. Selected average bond lengths (Å) and angles (°) for the RS series.

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<tr>
<th>Compound</th>
<th>U=O</th>
<th>U—N</th>
<th>N=C</th>
<th>C=S</th>
<th>O=U=O</th>
<th>U—N=C</th>
<th>S o.o.p (Å)</th>
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<td>1.770(4)</td>
<td>2.45(2)</td>
<td>1.15(2)</td>
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<td>179.7(1)</td>
<td>154.1(4) to 176.2(4)</td>
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<td>1.14(1)</td>
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<td>2.45(1)</td>
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<td>1.62(1)</td>
<td>179.21(13)</td>
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<td>S1 -0.797</td>
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Apparent in the structures of MeS and EtS are S···S close contacts within the van der Waals radii for two S atoms (3.78 Å). In MeS these are Type I interactions (d_{S···S} = 3.587 Å, \(\theta_2-\theta_1 = 0^\circ\)). In contrast, EtS forms chains along the crystallographic \(b\)-axis and the angles \(\theta_1\) and \(\theta_2\) suggest these are not crystal packing (Figure 2a), in accord with the conventional description for halogen bonding (Chart 1); we believe that this is applicable here as the interactions are both of a \(\sigma\)-hole type. The S···S distances and angles are: d_{S···S} = 3.470 Å, \(\theta_2-\theta_1 = 38.3^\circ\) and d_{S···S} = 3.591 Å, \(\theta_2-\theta_1 = 20.8^\circ\). There are also S···H—C weak hydrogen bonds (d_{S···C} = 3.540(8) to 3.895(7) Å) where the alkylammonium cation links two chains. It appears that the S···H—C interaction is responsible for the movement of the sulfur atom out of the UN5 plane. Also present are longer C—H···O=U hydrogen bonds (d_{O···C} = 3.21 and 3.47 Å). As the size of the R group increases the close contacts between two S centers are no longer present, and in the \(^{\text{n}}\)Pr case, S···H—C interactions are prevalent (Figure 2b; d_{S···C} = 3.74 – 3.84 Å) along with bifurcated hydrogen bonding to the uranyl ion (d_{O···C} = 3.40 and 3.49 Å). We have prepared and structurally characterized \(^{\text{n}}\)BuS and there are no S···S close contacts. Thus it appears that there is a steric effect of the cation in the nature of these non-covalent interactions, whereby the conformationally non-rigid arms of the n-propyl chain are enough to "turn-off" the S···S close contacts, while the S···H—C interactions dominate all
structures. Support for this hypothesis comes from the structures of \([\text{Me}_3\text{NBz}]_3[\text{UO}_2(\text{NCS})_2]\) and
[\([\text{Et}_3\text{NBz}]_3[\text{UO}_2(\text{NCS})_2]\)] where there are short contacts between two sulfur atoms, but the angles
suggest a type I interaction, \textit{i.e.} due simply to packing (Me\textsubscript{3}NBzS: \(d_{S\cdots S} = 3.350\) Å, \(\theta_2-\theta_1 = 0^\circ\);
Et\textsubscript{3}NBzS: \(d_{S\cdots S} = 3.540\) Å, \(\theta_2-\theta_1 = 14.0^\circ\)). However, these systems enable us to explore the C—
H\cdots S interactions in more detail (Me\textsubscript{3}NBzS: \(d_{S\cdots C} = 3.690(3)\) to \(3.824(3)\) Å; Et\textsubscript{3}NBzS: \(d_{S\cdots C} =
3.711(4)\) to \(3.849(5)\) Å). It has been reported from computational studies that the strength of this
weak hydrogen bond is dependent upon the hybridization of the C—H donor and this decreases as
hybridization increases\textsuperscript{34} but in the structures of Me\textsubscript{3}NBzS and Et\textsubscript{3}NBzS there is no obvious
correlation. From this description it is clear that the supramolecular structures are influenced subtly
by the steric effects of the cation.
Figure 2. Supramolecular structure of (a) EtS highlighting the non-covalent S⋯S interactions (black lines) and (b) the S⋯H-C hydrogen bonding in nPrS (blue lines). Color code: U – pink; O – red; N- blue; C – grey; S – yellow; H – white.

A further secondary interaction occurs between the C—H of the cation and the uranyl group. The –yl oxo is well known to act as a hydrogen bond acceptor.\textsuperscript{21,22} In all the examples in this series, except for nBuS, there are numerous short contacts with d_C⋯O between 3.0-3.5 Å; Figure 3 shows a specific example. These hydrogen bonding interactions do not perturb the U=O bond lengths, as
all examples are essentially identical to that of BuS which does not feature any U=O—H—C close contacts, indicating that the hydrogen bonding interactions are very weak (vide infra).

Figure 3. Supramolecular interactions in Me₃NBzS: dashed red line U=O—H—C and dashed blue line S···H—C.

During one attempt at the synthesis of [Ph₄P]₃[UO₂(NCS)₅] we obtained a few single crystals that were shown to be [Ph₄P]₂[UO₂(NCS)₃(NO₃)] by single crystal X-ray diffraction. We were able to increase the yield using the correct stoichiometry, albeit as a polymorph, but have been unable to
crystallize the homoleptic thiocyanate complex with this cation. The structure of this compound is shown in Figure 4. The bond lengths are similar to the homoleptic series described above, with the exception that the U-N bond lengths are slightly shorter (2.412(8) and 2.402(11) Å). The U—O bond length (2.503(7) Å) is typical for a nitrate ion coordinated in a bidentate fashion. There are no S⋯S close contacts in this case and the dominant interactions are C—H⋯O hydrogen bonding to the nitrate ion (d_C⋯O = 3.237(10) Å) and longer C⋯H⋯S (d_C⋯S = 3.736(9) Å); the [NCS]− ion involved in this interaction is bent out of the UN₅ plane (0.307 Å; C(1)-N(1)-U(1) = 166.2(7)°) in comparison to that [NCS]− ion that does not have close contacts which is in the plane (C(2)-N(2)-U(1) = 180.0°), adding weight to our thesis that it is the weak hydrogen bonding that is responsible for the disruption in the U—NCS coordination environment.
**Figure 4.** Molecular structure of [Ph₄P]₂[UO₂(NCS)₃(NO₃)] with atomic displacement parameters shown at 50% probability. Hydrogen atoms omitted for clarity. Symmetry unique atoms labelled only.

In an attempted reaction of uranyl nitrate, sodium thiocyanate and Me₃SnCl, we did not isolate the expected species, but instead structurally characterized [(UO₂)₂(SO₄)₂(H₂O)₄].3H₂O. The mechanism of oxidation of the [NCS]⁻ ion is not immediately apparent, but there is some literature precedent for oxidation of coordinated thiocyanate to sulfate. Nevertheless the structure is α-uranyl sulfate determined in 1978, and similar to the mineral Shumwayite, [(UO₂)₂(SO₄)₂].5H₂O. Our refinement is significantly better so we have included it herein but will not comment further.

(ii) [R₄N]₃[UO₂(NCSe)₃] series (R = Me, Et, "Pr, Et₃NBz)

The number of structurally characterized uranyl selenocyanate compounds numbers only two: we reported the structural characterization of [Et₄N]₄[UO₂(NCSe)₃][NCSe], and Walensky and co-workers reported a tridentate salicylaldiminate uranyl complex of [NCSe]⁻. We have extended our initial study to include [Me₄N]₃[UO₂(NCSe)₃].H₂O (MeSe), "PrSe and Et₃NBzSe, all of which have been structurally characterized. The gross solid-state structures are the same as for the RS series, and an exemplar, "PrSe, is shown in Figure 5, while the other structures are in Figures S4-S6. The pertinent metric parameters are collated in Table 2 and Tables S9-S14, and there are no significant differences in the series.
Table 2. Selected average bond lengths (Å) and angles (°) for the RSe series.

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<th>Compound</th>
<th>U=O</th>
<th>U—N</th>
<th>N=C</th>
<th>C=Se</th>
<th>O=U=O</th>
<th>U—N=C</th>
<th>Se o.o.pa</th>
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<tr>
<td>MeSe</td>
<td>1.769(5)</td>
<td>2.46(2)</td>
<td>1.15(3)</td>
<td>1.80(3)</td>
<td>179.0(3) to 179.0(7)</td>
<td>162.0(6)</td>
<td>Se1 +0.609&lt;br&gt;Se2 +0.710&lt;br&gt;Se3 +0.114&lt;br&gt;Se4 +0.227&lt;br&gt;Se5A +1.552&lt;br&gt;Se5B +1.879&lt;br&gt;Se6 -1.145&lt;br&gt;Se7 +0.135&lt;br&gt;Se8 +0.249&lt;br&gt;Se9 -0.073&lt;br&gt;Se10 +0.018</td>
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<td>179.57(9) to 177.6(3)</td>
<td>171.0(2)</td>
<td>Se1 -0.164&lt;br&gt;Se2 +0.482&lt;br&gt;Se3 -0.258&lt;br&gt;Se4 +0.496&lt;br&gt;Se5 -0.481</td>
</tr>
<tr>
<td>&quot;PrSe</td>
<td>1.7632(19)</td>
<td>2.46(1)</td>
<td>1.15(1)</td>
<td>1.78(1)</td>
<td>179.48(10) to 173.8(2)</td>
<td>157.1(2)</td>
<td>Se1 +0.329&lt;br&gt;Se2 -0.673&lt;br&gt;Se3 -0.035&lt;br&gt;Se4 -0.638&lt;br&gt;Se5 -0.498</td>
</tr>
<tr>
<td>Et₃NBzSe</td>
<td>1.765(5)</td>
<td>2.45(2)</td>
<td>1.14(3)</td>
<td>1.79(3)</td>
<td>167.3(7) to 174.9(7)</td>
<td>Se1 +0.232</td>
<td></td>
</tr>
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<td>---------</td>
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<tr>
<td></td>
<td>1.756(5)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Se2 +0.499</td>
<td></td>
</tr>
<tr>
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<td>Se3 +0.203</td>
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<td></td>
<td></td>
<td></td>
<td>Se5 -0.267</td>
<td></td>
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</tbody>
</table>

a distance out of the UN₅ plane.

**Figure 5.** Molecular structure of ⁹PrSe with atomic displacement parameters shown at 50% probability. Hydrogen atoms omitted for clarity.
The supramolecular interactions in the RSe series are greater and more complex than the RS series; in particular Se···Se interactions closer than the sum of the van der Waals radii (3.64 Å)\textsuperscript{32} are preponderant, as expected if the bonding is via a σ-hole type mechanism. Se···H—C interactions\textsuperscript{38} and U=O···H—C weak hydrogen bonds are also prevalent.

The packing of MeSe is shown in Figure 6. There are a number of Se···Se distances which are all slightly longer than the van der Waals radii (d_{Se···Se} = 3.671(2) - 3.755(3) Å), but theory shows that these are stabilizing interactions (vide infra). Interestingly Se(1)···Se(7), Se(4)···Se(8) and Se(3)···Se(9) form 1D sheets along the c-direction (Figure 6b), with C—H···Se weak hydrogen bonds (d_{C···Se} = 3.75 – 4.07 Å) linking the layers. C—H···O=U hydrogen bonding (d_{C···O} = 3.35 and 3.57 Å) interactions are also present, along with hydrogen bonding from the free water to U=O (d_{O···O} = 3.13 Å) and Se (d_{O···Se} = 3.79 Å). The water presumably comes adventitiously from the recrystallization process; notably all other compounds were prepared and recrystallized in air so it may be that the small cation allows for the inclusion of water in this case. Irrespective of the source of the water, it clearly engages in hydrogen bonding to the preference of C—H···O=U hydrogen bonding and possibly chalcogenide bonding.
Figure 6. Packing of MeSe along the crystallographic a-axis. (a) A perspective view and (b) a view highlighting the Se···Se interactions (black lines).

PrSe shows no Se···Se interactions, probably due to the steric effect of the longer n-propyl arms that completely shield the uranyl complex, as illustrated in the space-filling structure (Figure 7). There are a plethora of Se···H—C (d_{Se···C} = 3.64 to 4.00 Å) and C—H···O=U (d_{C···U} = 3.175(3), 3.271(3) and 3.387(3) Å) close contacts that link the uranyl ions in layers. As in the previous series, it appears that Se···H—C interactions bend Se out of the UN₅ plane.
Figure 7. Space-filling view of PrSe (orange – Se; blue – N, dark Grey – C; light gray – H; red – O; uranyl axis is parallel to the page).

In Et₃NBzSe (Figure 8) there are a number of Se···Se close contacts (d_{Se···Se} = 3.565(2) Å, θ₂-θ₁ = 20.7°; d_{Se···Se} = 3.665(2) Å, θ₂-θ₁ = 66.1°). There are also multiple Se···H—C (d_{C···Se} = 3.790(8) - 3.913(8) Å) and U=O···H—C (d_{C···U} = 3.310(9) - 3.59(1) Å) interactions present.
Figure 8. Supramolecular interactions in Et$_3$NBzSe: (a) a view of the C—H…O and C—H…Se; (b) the packing along the crystallographic $a$-axis and (c) along the crystallographic $b$-axis highlighting the chalcogenide interactions (dashed black line Se⋯Se; dashed red line U=O⋯H—C and dashed blue line Se⋯H—C).

(iii) Structural comparisons within the RS and RSe series — $^n$PrS/$^n$PrSe and Et$_3$NBzS/Et$_3$NBzSe.

A summary of the types of interactions is shown in Table 3. We can directly compare structural parameters between two series with the same cation. For $^n$Pr, the obvious conclusion is that the
chalcogenide···chalcogenide interactions are not present. Perhaps this is due to the non-rigid nature of the propyl arms which preferentially engage in S(e)···H—C and U=O···H—C interactions, implying that these are more favored over S(e)···S(e) interactions. The charge assisted hydrogen bonds to the uranyl ion are about the same length. Interestingly, the Et₃NBz series shows that as the polarizability of the chalcogenide atom increases then Type II interactions become preferred, even though these are isostructural pairs. The charge assisted hydrogen bond distances do not significantly change, indicating that perhaps these weak interactions are purely electrostatic in origin. A similar trend is observed for the S(e)···H—C interactions.

Table 3. Structural comparison of [UO₂(NCE)₃]³⁻ compounds.

<table>
<thead>
<tr>
<th>Compound</th>
<th>S···S (vvdv)</th>
<th>S···H—C</th>
<th>O···H—C</th>
<th>Compound</th>
<th>Se···Se (vvdv)</th>
<th>Se···H—C</th>
<th>O···H—C</th>
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<tr>
<td>MeS</td>
<td>Type I (95)</td>
<td>multiple</td>
<td>multiple</td>
<td>MeSe</td>
<td>Type I (101)</td>
<td>multiple</td>
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<td></td>
<td></td>
<td></td>
<td>Type II (100)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>EtS</td>
<td>Type II (92)</td>
<td>multiple</td>
<td>multiple</td>
<td>EtSe</td>
<td>Type II (94)</td>
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</tbody>
</table>
A useful method for comparing solid state structures is via the Hirschfeld surfaces.\textsuperscript{39} We have employed this method to examine the non-covalent interactions present in the RS and RSe series, along with selected other homoleptic uranyl thiocyanate complexes.\textsuperscript{28,40} A quantitative analysis is shown in Figure 9. The chalcogenide···chalcogenide interactions decrease as the R group increases in size, but those with the R\textsubscript{3}NBz group are type I for S and type II for Se. However it is clear that the S(e)···H—C interactions are the largest non-covalent interaction present but also decrease in the same order.

<table>
<thead>
<tr>
<th></th>
<th>&quot;PrS</th>
<th>no</th>
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<th>&quot;PrSe</th>
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<tbody>
<tr>
<td>&quot;BuS</td>
<td>no</td>
<td>one</td>
<td>no</td>
<td>-</td>
<td>Me\textsubscript{3}BzNS</td>
<td>Type I</td>
<td>multiple</td>
<td>multiple</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td>(88)</td>
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<tr>
<td>Et\textsubscript{3}BzNS</td>
<td>Type I</td>
<td>multiple</td>
<td>multiple</td>
<td>Et\textsubscript{3}BzNSe</td>
<td>Type II</td>
<td>multiple</td>
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</tr>
<tr>
<td></td>
<td>(94)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>(98)</td>
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</tr>
</tbody>
</table>
Spectroscopic characterization.

All the compounds discussed above have been characterized spectroscopically with the aim of observing if the structural perturbations are detectable. NMR spectroscopy is uninformative, while solid-state $^{77}$Se NMR spectroscopy does not show a clear trend (Figure S25).

Vibrational Spectroscopy. The vibrational spectra of crushed single crystals clearly show the uranyl stretch and the N=C and C=S(e) bond stretches. In the series the bands attributable to these stretches do not change significantly (Table S15); the Raman active $v_1$(U=O) symmetric stretching mode lies in a narrow range between 842 and 850 cm$^{-1}$ while the IR active $v_3$(U=O) asymmetric
stretching mode comes between 910 and 940 cm\(^{-1}\) (Figures S7-S8). For comparison, the 
\([\text{UO}_2(\text{H}_2\text{O})_3]^{2+}\) ion shows these bands at ca. 860-880 cm\(^{-1}\) and 930-960 cm\(^{-1}\) for the \(v_1\) and \(v_3\) modes respectively;\(^{41}\) the presence of the donor atoms in the equatorial plane cause a redshift, as expected. From the IR and Raman spectra the stretching force constant \((k_1)\) and the interaction force constant \((k_{12})\) can be calculated, as \(k_1\) gives an indication of bond strength while \(k_{12}\) describes the interaction between the two \(-\text{yl}\) oxygen atoms.\(^{42}\) In our case any changes must be due to the difference in the hydrogen bonding, and we can benchmark our data to the unperturbed \(U=O\) bond of "BuS. The force constants do not change significantly from this value ("BuS, \(k_1 = 6.92\) mdyn/Å; range = 6.84 – 7.06 mdyn/Å) and there is no correlation to the donor···acceptor bond distances (Table S15) so the hydrogen bonding does not influence these metrics, as observed by others.\(^{22}\) There is however a larger \(k_1\) and \(k_{12}\) associated with the NCSe ligands compared to the NCS analogues as shown for the "PrS//PrSe (S: \(k_1 = 6.91\) mdyn/Å, \(k_{12} = -0.14\) mdyn/Å; Se: \(k_1 = 6.98\) mdyn/Å, \(k_{12} = -0.14\) mdyn/Å) and Et\(_3\)NBzS/Et\(_3\)NBzSe (S: \(k_1 = 6.78\) mdyn/Å, \(k_{12} = -0.10\) mdyn/Å; Se: \(k_1 = 6.86\) mdyn/Å, \(k_{12} = -0.17\) mdyn/Å) families. The structural information showed no significant changes in donor···acceptor bond lengths, so this could be a manifestation of the differing donor abilities of the coordinating ligands.\(^{20a}\)

**Photoluminescence Spectroscopy.** The photophysical properties of the uranyl ion have been well studied and the optical properties are due to a ligand-to-metal charge transfer (LMCT) transition involving promotion of an electron from bonding \(-\text{yl}\) oxygen orbitals \((\sigma_u, \sigma_g, \pi_u\) and \(\pi_g)\) to non-bonding uranium 5f\(_5\) and 5f\(_9\) orbitals, centered at ca. 420 nm. The characteristic green emission at ca. 520 nm arises from de-excitation of the \(^3\Pi_u\) triplet excited state. Superimposed on the absorption and emission bands are bands arising from strong coupling of the ground state Raman active symmetric vibrational \(O=U=O\) \((v_1)\) mode with the \(^3\Pi_u\) electronic triplet excited state.\(^{43}\)
There are some general trends that are apparent in the luminescence measurements, such as a correlation between the electron donating ability of the ligand and the $E_{0-0}$ values; i.e. the more electron donating the ligand the lower the energy value of the emission.\textsuperscript{44} We recently showed for a series of $[\text{UO}_2X_4]^{2-}$ ions ($X = \text{Cl, Br, I}$) that the emission maxima and lifetimes of the $\text{O}_\text{yl} \rightarrow \text{U}$ LMCT band are not particularly sensitive to variation of the halide.\textsuperscript{45} Recently the influence of the tecton in $[\text{UO}_2\text{Cl}_4]^{2-}$ has been interrogated using luminescence spectroscopy\textsuperscript{46} and it may be that those ligands that have vibrational modes that can couple with the $\text{U}=\text{O}$ vibrations to directly influence the luminescence properties. The emission and excitation spectra of RS and RSe have been recorded both in the solid state and in solution, where all non-covalent interactions break up and mononuclear species dominate.\textsuperscript{47} In the solution emission spectra (Figure S22 and Table S16) there are clear differences between the RS and RSe family and the $[\text{NCS}]^-$ ion is a more electron donating ligand than the $[\text{NCSe}]^-$ ion in this series and on par with the nitrate ion. The differences within each family are small and mainly due to the changes in the vibronic coupling, and is also observed in the room temperature solid state emission spectra (Figure S23). Given the poor resolution of these spectra, we investigated the low temperature (77 K) photoluminescent properties of all compounds and selected examples are shown in Figure 10 (Figure S24 shows the remainder, except for MeSe which did not show any resolution enhancement). Immediately apparent is the increase in resolution, to the point that additional weaker bands appear, and are labelled as set A and B. The weak bands of set B are separated from each other by a vibrational progression of ca. 850 cm$^{-1}$ which is due to coupling to the $v_1(\text{U}=\text{O})$ vibration, but the differences between set A and B are ca 490 cm$^{-1}$. It is noteworthy that these additional bands were observed in the room temperature emission spectra of $[\text{UO}_2(\text{NCS})_3]^3-$ in ionic liquids and ascribed to changes in the coordination environment. However a close examination of the Raman spectra of the RS
compounds show a number of rather weak bands at 500 and 544 cm\(^{-1}\) due to the \(\delta\)(NCS) mode\(^{48}\), so it appears that this can couple with the uranyl vibrational mode; the same pattern is observed in \([\text{Ph}_4\text{P}]_2[\text{UO}_2\text{(NCS)}_3\text{(NO}_3\text{)}]\) (Figure S24). Supporting this is the smaller and weaker coupling of ca. 450 cm\(^{-1}\) in the RSe family which corresponds to the very weak \(\delta\)(NCSe) vibrational bend at ca. 430 cm\(^{-1}\) in the Raman spectra (Figues S9). The only other mode that could couple is the U—N bend but in An(IV) thiocyanates this comes at ca 200 cm\(^{-1}\).\(^{49}\) The differences between the RS and RSe families are not marked at low temperature. Finally we have measured the lifetimes of the emissions both in the solid state and solution (Table S16), and they vary substantially across the series so no definitive conclusions can be drawn.

**Figure 10.** Emission spectra of selected examples of RS and RSe at 77 K in the solid state \((\lambda_{ex} = 340 \text{ nm})\).
In summary, the spectroscopic data show some small differences between the [NCS] and [NCSe] coordinated ions, but suggest that the C–H···O interactions are not sufficiently strong to perturb the electronic structure of the uranyl ion, while the consequences for chalcogenide interactions are not observable.

*DFT calculations.*

To deepen our understanding of the non-covalent interactions in these species, we turned to DFT calculations. Supramolecular calculations of energy would be dominated by charge-charge interactions, so instead we have used molecular electrostatic potential (MEP), Atoms-in-Molecules (AIM) and natural bond orbital (NBO) derived properties to characterize U=O···H—C, Se···H—C and chalcogenide interactions in molecules and dimers selected from the crystal structures reported herein.

(i) *Chalcogenide interactions.* Se···Se interactions in [Et₄N][UO₂(NCSe)₅][NCSe], where the Se···Se distances are substantially shorter than the van der Waals radii, lead us to hypothesize the presence of a σ-hole on Se. Figure 11 shows the molecular electrostatic potential projected onto the 0.001 au isodensity surface for [UO₂(NCS)₅]³⁻ and [UO₂(NCSe)₅]³⁻. The overall negative charge of each complex means that the MEP is negative across the entire surface, but the plots nevertheless show regions of relative electron depletion (*i.e.* a σ-hole) on both S and Se, with a more pronounced hole on Se (-0.231 au) than on S (-0.246 au). There are also small differences in the –yl oxygens and corroborates the differing electron donating ability of the ligands from the photoluminescence spectroscopy and force constants.
Figure 11. Molecular electrostatic potential projected onto the 0.001 au isodensity surface for [UO$_2$(NCS)$_3$]$^{3-}$ (top) and [UO$_2$(NCSe)$_3$]$^{3-}$ (bottom).
We elected to do DFT calculations for dimers of $[\text{UO}_2(\text{NCSe})_5]^3^-$ from MeSe and the complex of $[\text{UO}_2(\text{NCSe})_5]^3^-$ with NCSe$^-$ extracted from EtSe, as these gave representative examples of the weak interactions present in the series. NBO analysis indicates that the major interactions between species are donation from a Se lone pair of the coordinated [NCSe]$^-$ to a $\sigma^*$ orbital on another [NCSe]$^-$ moiety (1.94 kcal/mol and 0.58 kcal/mol) in the dimers extracted from MeSe. In the dimer extracted from EtSe, donation from a Se lone pair of the coordinated [NCSe]$^-$ to the C=Se $\sigma^*$ orbital of free [NCSe]$^-$ (2.94 kcal/mol), along with donation from the free Se lone pair to the coordinated C=Se $\sigma^*$ orbital (2.20 kcal/mol). Figure 12 shows the orbitals involved in EtSe. The energies of the MeSe dimers are lower than that of EtS and in keeping with the longer Se…Se distances in the structures. Finally, the interaction energies of model compounds $\text{Me}_2\text{Se}^\cdot\cdot\cdot\text{SeMe}_2$ (-2.82 kcal/mol) and $\text{Me}_2\text{Se}^\cdot\cdot\cdot\text{Se(Me)CN}$ (-4.62 kcal/mol)$^{50}$ are comparable to our systems.

The value of the electron density at the critical point, $\rho_{\text{bcp}}$, is often taken as a proxy for bond strength. Here the Se…Se contact exhibits a bond critical point with $\rho_{\text{bcp}} = 0.009$ au for both dimers, corroborating the NBO analysis that a weak but a stabilizing interaction is present. This is noteworthy as in MeSe the Se…Se contacts were slightly greater than the sum of the van der Waals radii. There are few reports of Se…E examples characterized via this methodology to compare, but XRSe…OH$_2$ and XRSe…NH$_3$ (R = H; Me; X = H, F, Me, CF$_3$, Cl, OH, OMe, NH$_2$, NHMe, CN) $\rho_{\text{bcp}}$ range from 0.053-0.270 au for the Se…O/N interaction, or in F$_2$C=Se…NX (X = CH; H$_3$; HCH$_2$; Cl; Me; $\rho_{\text{bcp}} = 0.0089-0.0197$) and F$_2$C=Se…XY (XY = BrCl, ClF and BrF; $\rho_{\text{bcp}} = 0.0276-0.0440$).$^{51}$ The optimized dimers of (X$_2$C=S)$_2$ range from $\rho_{\text{bcp}} = 0.039$ to 0.050 with the sum of the charge transfer 1.52 to 3.12 kcal/mol (X = H, NH$_2$, OH, F, Cl).$^{11}$ This data can be put into a broader context by comparing to other non-conventional bonding that has been reported. Whilst it is clear that substitution effects are important, these chalcogenide interactions are on par with the
weak fluorine bonding (e.g. $\rho_{\text{bcp}} = 0.0073$ for the interaction in H—F…F—F$^{52}$), or halogen lone pair…$\pi$ interactions,$^{53}$ and weaker than, for example, F—Br..I—F ($\rho_{\text{bcp}} = 0.0320$)$^{54}$ as anticipated from the $\sigma$-hole mechanism of bonding which depends on the polarizability of the atom.

![Figure 12](image.png)

**Figure 12.** Theoretical analysis of $[\text{Et}_4\text{N}]_4[\text{UO}_2(\text{NCSe})_2][\text{NCSe}]$ (a) NBO orbitals involved in the coordinated to free, (b) free to bound lone pair to $\sigma^*$ orbitals.

**(ii) weak hydrogen bonding.** We have characterized dimers bound by Se···H—C and U=O···H—C hydrogen bonding, taken from MeSe and "PrSe, an AIM approach as this features interactions. A dimer taken from MeSe contains three C—H···O=U bond critical points with $\rho_{\text{bcp}} = 0.007, 0.006$ & 0.005, while the H-bonds from "PrSe: 6 x C—H···Se = 0.008, 0.006, 0.005, 0.005, 0.004, 0.003. Thus, individual C—H···Se are individually weak but rather numerous, lending support to the hypothesis that these dominate in cases such as "PrSe with many such donors. C—H···O=U interactions seem even weaker on this basis, again supporting the observation noted above that the terminal O in uranyl are relatively weak Lewis bases. *Intramolecular* C—H···O=U hydrogen
bonding has recently been theoretically analyzed and in this case $\rho_{bcp} = 0.002-0.0243$ a.u., indicating that intramolecular hydrogen bonding is perhaps stronger than intermolecular hydrogen bonding, although geometric constraints may influence this.

**Conclusions**

A series of uranyl thiocyanate and selenocyanate compounds have been prepared and structurally characterized. The nature of the alkyl ammonium group dictates the non-covalent interactions observed in the solid state and chalcogenide, $C-H\cdots O=U$ and $C-H\cdots S(e)$ are observed. Chalcogenide interactions are observed in some of the structures and are more prevalent in the [NCSe] series than the [NCS], and in line with the increased polarizability of the Se which renders a more pronounced $\sigma$-hole on Se (-0.231 au from MESP) than on S (-0.246 au from MESP). The molecular orbitals responsible for this were identified and correspond with the known ideas of $n$ to $\sigma^*$ charge transfer in these non-classical interactions, albeit in this example the non-bonded pair of electrons resides in a $\sigma$ orbital. Comparison with other weak $\sigma$-hole type bonds puts these chalcogenide interactions similar to interactions involving fluorine or, more commonly used in synthetic chemistry, lone pair-$\pi$ interactions.\(^{55}\) $C-H\cdots S(e)$ interactions are individually weak from an AIM analysis but numerous and accounts for their preponderance in the solid state structures. The vibrational data suggest that the charge assisted hydrogen bonding to the $\cdots$yl oxygens does not influence the spectroscopic properties and DFT studies show that these interactions are the weakest of the three studied. Unusually, the coupling the $\delta$(NCS) and $\delta$(NCSe) vibrational modes to the uranyl mode manifests itself in the photophysical properties of these species in the solid state. Taken together these results show that collectively non-classical
interactions can have an important influence of the solid state structures, but are in these cases not observable by conventional spectroscopic techniques. Nonetheless, these add to the growing field of chalcogenide interactions in the coordination sphere of metals.

**Experimental Section**

*Caution!* Natural uranium was used during the course of the experimental work. As well as the radiological hazards, uranium is a toxic metal and care should be taken with all manipulations. Experiments using uranium materials were carried out using pre–set radiological safety precautions in accordance with the local rules of Trinity College Dublin.

$^1$H, $^{13}$C{$^1$H}, $^{31}$P{$^1$H} and $^{77}$Se{$^1$H} spectra were recorded on a Bruker AV400 spectrometer operating at 400.23 MHz, 155.54 MHz, 161.98 MHz and 76.33 MHz respectively, and were referenced to the residual $^1$H and $^{13}$C resonances of the solvent used or external H$_3$PO$_4$ or Me$_2$Se. Solid state $^{77}$Se NMR spectra were recorded on a Bruker 400 HD at 76.3 MHz. IR spectra were recorded on a Perkin Elmer Spectrum One spectrometer with attenuated total reflectance (ATR) accessory. Raman spectra were obtained using 785-nm excitation on a Renishaw 1000 micro-Raman system. UV-vis measurements were made on a Perkin Elmer Lambda 1050 spectrophotometer, using fused silica cells with a path length of 1 cm. Steady-state photoluminescence spectra were recorded on a Horiba-Jobin-Yvon Fluorolog-3 spectrofluorimeter. Luminescence lifetime data were recorded following 372-nm excitation, using time-correlated single-photon counting (a PCS900 plug-in PC card for fast photon counting). Lifetimes were obtained by tail fit on the data obtained, and the quality of fit was judged by
minimization of reduced chi-squared and residuals squared. X-ray crystallography data were measured on a Bruker Apex diffractometer at 100 K. The structure was solved by direct methods and refined by least squares method on $F^2$ using the SHELXTL program package. Crystal data, details of data collections and refinement are given in Table S17. CCDC 1586895-1586903 contains the supplementary crystallographic data for this paper. This data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif. Structural figures were prepared using VESTA or mercury. All other chemicals and solvents were obtained from commercial sources and used as received.

DFT calculations were carried out using the meta-hybrid M06-2X DFT method Gaussian09, without symmetry constraints. Monomer or dimer coordinates were extracted from relevant crystal structures, and where necessary C—H bond lengths normalized to values from neutron diffraction (1.082 Å), with all heavy atom positions frozen at crystalline values. Scalar relativistic effects in uranium were included through use of effective core potentials as defined in Preuss et al’s ECP/basis set on U, while for lighter atoms 6-31+G(d,p) was used. Natural bond orbital (NBO) analysis was performed using Gaussian09; Atoms-in-Molecules (AIM) analysis used AIMAll. Topological analysis of the electronic density ($\rho$) is based upon those points where the gradient of the density, $\nabla \rho$, vanishes. In this work we consider points where one curvature (in the inter-nuclear direction) is positive and two (perpendicular to the bond direction) are negative, termed (3, -1) or bond critical points. Properties evaluated at such points characterize the bonding interactions present.
All solvents and reagents were obtained commercially and used as received. \([R_4N]_3\text{[UO}_2\text{(NCS)}_5]\) (R = Me, Et, "Pr) were synthesized via a slight modification of literature procedure, whereby the reactants were mixed in MeCN rather than H\(_2\)O.

**Synthesis of uranyl thiocyanates**

To a solution of \(\text{UO}_2\text{(NO}_3\text{)}_2 \cdot 6\text{H}_2\text{O}\) (0.30 g) in acetonitrile (30 cm\(^3\)) were added \(\text{Na}[\text{NCS}]\) (5 equivalents) and the appropriate cation (3 equivalents). The solution mixture was stirred at room temperature for 60 minutes. The resulting yellow solution was filtered and the solvent was reduced in volume. Crystals suitable for X-ray diffraction were obtained by vapor diffusion of \(^4\text{Pr}_2\text{O}\) into MeCN solutions of the products and yellow crystals obtained after ca. 1 week.

\([\text{"Bu}_4\text{N}]_3\text{[UO}_2\text{(NCS)}_5]\). Yield: 57\%, 0.44 g; \(^1\text{H} \text{NMR (400 MHz, CD}_3\text{CN): } \delta 1.00 \text{ ppm (t, 36H, J}_\text{H-H} = 7.36 \text{ Hz, CH}_3), \delta 1.38 \text{ ppm (m, 24H, J}_\text{H-H} = 7.43 \text{ Hz, CH}_2), \delta 1.63 \text{ ppm (m, 24H, J}_\text{H-H} = 8.55 \text{ Hz, CH}_2), \delta 3.11 \text{ ppm (m, 24H, J}_\text{H-H} = 8.62 \text{ Hz, CH}_2); \text{}^{13}\text{C}\{\text{}^{1}\text{H}\} \text{ NMR (155 MHz, CD}_3\text{CN): } \delta 12.89 \text{ ppm (CH}_3), \delta 19.39 \text{ ppm (CH}_2), \delta 23.36 \text{ ppm (CH}_2), \delta 58.33 \text{ ppm (CH}_2); \text{} \text{IR (ATR, v/cm}^{-1}\text{): 2960 (m), 2872 (w), 2052 (s, C=\text{N}), 1467 (m), 1380 (w), 1152 (w), 1108 (w), 1067 (w), 1026 (w), 919 (s, U=O), 882 (m), 736 (m, C=S); \text{} \text{Raman (v/cm}^{-1}\text{): 2099 and 2059 (C=\text{N}), 1456, 1324, 909, 850 (U=O), 736 (C=S); \text{} \text{UV-vis (}\varepsilon, \text{ dm}^3 \text{ mol}^{-1} \text{ cm}^{-1}\text{): (298 K, MeCN, } \sim 10^{-5} \text{ M): 330 nm (4868), 284 nm (6240.5), 232 nm (17618).}

\([\text{Me}_3\text{NBz}]_3\text{[UO}_2\text{(NCS)}_5]\). Yield: 55\%, 0.29 g; \(^1\text{H} \text{NMR (400 MHz, CD}_3\text{CN): } \delta 3.05 \text{ ppm (s, 27H, CH}_3), \delta 4.43 \text{ ppm (s, 6H, CH}_2), \delta 7.52 \text{ ppm (m, 6H, o-CH), } \delta 7.53 \text{ ppm (m, 3H, p-CH), } \delta 7.56 \text{ ppm (m, 6H, m-CH); } \text{}^{13}\text{C}\{\text{}^{1}\text{H}\} \text{ NMR (155 MHz, CD}_3\text{CN): } \delta 52.4 \text{ ppm (CH}_3), \delta 69.3 \text{ ppm (CH}_2), \delta 127.6
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ppm (ipso-C), δ 129.2 ppm (m-CH), δ 130.7 ppm (p-CH), δ 132.9 ppm (o-CH); IR (ATR, v/cm⁻¹): 3420 (m), 3025 (w), 2030 (s, C=N), 1634 (w), 1581 (w), 1475 (m), 1408 (m), 1375 (m), 1211 (m), 1076 (w), 1036 (w), 970 (m), 916 (s, U=O), 885 (m), 779 (m, C=S), 725 (s), 700 (s); Raman (v/cm⁻¹): 2090 and 2062 and 2048 and 2041 (C=N), 1065, 1584, 1471, 1446, 1216, 1184, 1157, 1033, 1003, 845 (U=O), 823, 808 (C=S), 725, 621; UV-vis (ε, dm³ mol⁻¹ cm⁻¹), (298 K, MeCN, ~10⁻⁵ M): 342 nm (14461), 300 nm (17597).

[Et₃NBz]₃[UO₂(NCS)₅]. Yield: 62%, 0.41 g; ¹H NMR (400 MHz, CD₃CN): δ 1.37 ppm (t, 27H, J_H-H = 7.11 Hz, CH₃), δ 3.18 ppm (q, 18H, J_H-H = 7.18 Hz, CH₂), δ 4.35 ppm (s, 6H, CH₂), δ 7.51 ppm (m, 3H, p-CH), δ 7.53 ppm (m, 6H, m-CH), δ 7.55 ppm (m, 6H, o-CH); ¹³C{¹H} NMR (155 MHz, CD₃CN): δ 8.13 ppm (CH₃), δ 53.5 ppm (CH₂), δ 61.1 ppm (CH₂), δ 128.1 ppm (ipso-C), δ 130.2 ppm (m-CH), δ 131.5 ppm (p-CH), δ 133.4 ppm (o-CH); IR (ATR, v/cm⁻¹): 3437 (w), 2994 (w), 2086 (m), 2042 (s, C=N), 1455 (m), 1399 (m), 1330 (w), 1214 (w), 1182 (w), 1150 (m), 1086 (m), 1033 (m), 910 (s, U=O), 788 (m) 752 (s, C=S), 693 (s); Raman (v/cm⁻¹): 2089 and 2042 (C=N), 1005, 842 (U=O), 816 and 812 and 804 (C=S), 679; UV-vis (ε, dm³ mol⁻¹ cm⁻¹), (298 K, MeCN, ~10⁻⁵ M): 333 nm (4268 nm), 286 nm (5610), 230 nm (8503).

[Ph₄P]₂[UO₂(NCS)₃(NO₃)]. Yield: 51%, 0.36 g; ¹H NMR (400 MHz, CD₃CN): δ 7.71 ppm (m, 8H, J_H-H = 8.43 Hz, m-CH), δ 7.78 ppm (m, 8H, J_H-H = 7.93 Hz, o-CH), δ 7.95 ppm (m, 4H, J_H-H = 7.53 Hz, p-CH); ¹³C{¹H} NMR (155 MHz, CD₃CN): δ 118.4 ppm (d, J_P-C = 96.60 Hz ipso-C), δ 130.3 ppm (d, J_P-C = 12.8 Hz, m-CH), δ 134.7 ppm (d, J_P-C = 10.3 Hz, o-CH), δ 135.4 ppm (d, J_P-C = 2.94 Hz, p-CH); ³¹P{¹H} NMR (162 MHz, CD₃CN): δ 22.7 ppm; IR (ATR, v/cm⁻¹): 3054 (w), 2029 (s, N=C), 1584 (w), 1481 (w), 1433 (m), 1341 (w), 1312 (w), 1269 (w), 1189 (w), 1163
(w), 1104 (s), 1025 (w), 996 (m), 928 (s, U=O), 754 (m, C=S), 720 (s), 678 (s), 616 (w), 523 (s); Raman (v/cm\(^{-1}\)): 2079 and 2034 (C=N), 1586, 1575, 1187, 1165, 1111, 1095, 1037, 1001, 860 (U=O), 826 and 819 (C=S), 677, 615, 278, 257, 244; UV-vis (\(\epsilon, \text{dm}^3 \text{mol}^{-1} \text{cm}^{-1}\)), (298 K, MeCN, \(\sim 10^{-5} \text{M}\)): 330 nm (4727), 275 nm (11131), 232 (49865).

**Synthesis of uranyl selenocyanates**

To a solution of UO\(_2\)(NO\(_3\))\(_2\) \cdot 6H\(_2\)O (0.30 g) in acetonitrile (30 cm\(^3\)) were added KNCSe (5 equivalents) and the appropriate cation (3 equivalents). The solution mixture was stirred at room temperature for 60 minutes in the dark. The resulting orange solution was filtered and the solvent was reduced in volume. Crystals suitable for X-ray diffraction were obtained by vapor diffusion of \(\text{t}^\text{Pr}_2\text{O}\) into MeCN solutions of the products and yellow crystals obtained after ca. 1 week.

\([\text{Me}_4\text{N}]_3[\text{UO}_2(\text{NCSe})_5]\) \cdot H\(_2\)O. Yield: 39%, 0.20 g; \(^1\text{H NMR} (400 \text{MHz, CD}_3\text{CN}): \delta \ 3.14 \text{ ppm (s, 12H, CH}_3\)), \delta \ 2.25 \text{ ppm (s, 2H, H}_2\text{O); }^{13}\text{C}\{^1\text{H}\} \text{ NMR} (155 \text{MHz, CD}_3\text{CN}): \delta \ 54.9 \text{ ppm (CH}_3\)); \(^{77}\text{Se}\{^1\text{H}\} \text{ NMR} (76 \text{MHz, CD}_3\text{CN}): \delta \ -340.5 \text{ ppm (NCSe); } \text{IR (ATR, v/cm}^{-1}\)): 3027 (w), 2956 (w), 2060 and 2042 (s, C=N), 1738 (w), 1480(m), 1410 (w), 1365 (w), 944 (s), 920 (s, U=O), 632 (m, C=Se); Raman (v/cm\(^{-1}\)): 2071 and 2091 (C=N), 1586, 1575, 1187, 1165, 1111, 1095, 1037, 1001, 860 (U=O), 826 and 819 (C=S), 677, 615, 278, 257, 244; UV-vis (\(\epsilon, \text{dm}^3 \text{mol}^{-1} \text{cm}^{-1}\)), (298 k, MeCN, \(\sim 10^{-5} \text{M}\)): 329 nm (7311), 250 nm (25068).

\([\text{t}^\text{Pr}_4\text{N}]_3[\text{UO}_2(\text{NCSe})_5]\). Yield: 45%, 0.36 g; \(^1\text{H NMR} (400 \text{MHz, CD}_3\text{CN}): \delta \ 0.977 \text{ ppm (t, 36H, J}_{\text{H-H}} = 7.27 \text{ Hz, CH}_3\)), \delta \ 1.69 \text{ ppm (m, 18H, J}_{\text{H-H}} = 9.27 \text{ Hz, CH}_2\)) \delta \ 3.10 \text{ ppm (m, 18H, J}_{\text{H-H}} = 5.20 \text{ Hz, CH}_2\)); \(^{13}\text{C}\{^1\text{H}\} \text{ NMR} (155 \text{MHz, CD}_3\text{CN}): \delta \ 10.8 \text{ ppm (CH}_3\)), \delta \ 16.0 \text{ ppm (CH}_2\)), \delta \ 61.0 \text{ ppm
(CH₂); $^{77}\text{Se}^1\text{H}$ NMR (76 MHz, D-Acetone): δ -305 ppm (NCSe); IR (ATR, ν/cm$^{-1}$): 2968 (m), 2937 (w), 2877 (w), 2094 (w), 2060 and 2046 (s, C=N), 1482 (m), 1455 (m), 1385 (m), 968 (m), 926 (s, U=O), 842 (w), 756 (m, C=Se), 627 (m); Raman (ν/cm$^{-1}$): 2100 and 2079 and 2065 (C=N), 1459, 1316, 1140, 1110, 1037, 852 (U=O), 783, 757, 640 (C=Se), 557, 333, 313, 210; UV-vis (ε, dm$^3$/mol$^{-1}$/cm$^{-1}$), (298 k, MeCN, $\sim$10$^{-5}$ M): 330 nm (9696).

$[\text{Et}_3\text{NBz}]_3[\text{UO}_2(\text{NCSe})_5]$. Yield: 54%, 0.44 g; $^1\text{H}$ NMR (400 MHz, D-Acetone): δ 1.55 ppm (t, 27H, J$_{H-H}$ = 7.31 Hz, CH$_3$), δ 3.47 ppm (q, 18H, J$_{H-H}$ = 7.33 Hz, CH$_2$), δ 4.70 ppm (s, 6H, CH$_2$), δ 7.55 ppm (m, 3H, p-CH), δ 7.57 ppm (m, 6H, m-CH), δ 7.66 ppm (m, 6H, o-CH); $^{13}\text{C}^1\text{H}$ NMR (155 MHz, D-Acetone): δ 7.35 ppm (CH$_3$), δ 52.7 ppm (CH$_2$), δ 60.3 ppm (CH$_2$), δ 128 ppm (ipso-C), δ 129 ppm (m-CH), δ 131 ppm (p-CH), δ 133 ppm (o-CH); $^{77}\text{Se}^1\text{H}$ NMR (76 MHz, D-Acetone): δ -342 ppm (NCSe); IR (ATR, ν/cm$^{-1}$): 2984 (w), 2086 (w), 2047 (s, C=N), 1451 (m), 1395 (m) 1210 (w), 1186 (w), 1154 (m), 1078 (w), 1030 (m), 969 (w), 920 (s, U=O), 831 (w), 806 (w), 791 (m), 751 (s, C=Se), 701 (s); Raman (ν/cm$^{-1}$): 2087 and 2056 and 2043 (C=N), 1604, 1460, 1212, 1051, 1033, 1003, 842 (U=O), 677, 643, 624; UV-vis (ε, dm$^3$/mol$^{-1}$/cm$^{-1}$), (298 k, MeCN, $\sim$10$^{-5}$ M): 329 nm (5383), 264 nm (8167), 248 nm (18259).

ASSOCIATED CONTENT

AUTHOR INFORMATION

Corresponding Author
* Tel: +353-1-8963501. Fax: +353-1-6712826. E-mail: bakerrj@tcd.ie

Author Contributions
The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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**Supporting Information.** Full crystallographic descriptions (CIF), spectroscopic and computational details. This material is available free of charge via the Internet at http://pubs.acs.org.

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Chalcogenide interactions

C-H⋯S(e)/C-H⋯O=U bonds