

Interaction between Uranium Tetrafluoride Oxide and the Pentafluorides of Arsenic, Niobium, Tantalum, and Bismuth †

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No reaction occurs between UF_4O and AsF_5 but $\text{UF}_4\text{O} \cdot 3\text{MF}_5$ ($\text{M} = \text{Nb}$ and Ta) and $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$ have been obtained as yellow or orange solids by warming mixtures of UF_4O with excess of the appropriate pentafluoride in anhydrous HF or by combination of the oxide tetrafluoride with excess of the pentafluoride as a melt. The solid adducts have been characterized by their reaction stoichiometries, chemical analyses, and vibrational spectra. Like $\text{UF}_4\text{O} \cdot 2\text{SbF}_5$, the adducts are fluorine bridged with some ionic character. The thermal decomposition of the adducts results in the production of uranyl species.

Although the adducts $\text{UF}_4\text{O} \cdot n\text{SbF}_5$ ($n = 2$ or 3) are rather easily obtained by dissolution of UF_4O in excess of SbF_5 and subsequent removal of SbF_5 by pumping,^{1,2} related UF_4O -Lewis acid adducts have not been previously obtained. In the present study an estimate of the extent of the donor behaviour of UF_4O has been made by examination of its reactions with a series of Lewis-acid pentafluorides in addition to SbF_5 .

Arsenic pentafluoride was found to be unreactive with UF_4O under the conditions used but the new adducts, $\text{UF}_4\text{O} \cdot 3\text{MF}_5$ ($\text{M} = \text{Nb}$ and Ta) and $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$, were readily prepared and characterized. Vibrational spectroscopic examination has shown that, like $\text{UF}_4\text{O} \cdot 2\text{SbF}_5$, the adducts are essentially fluorine bridged but possess some ionic character. The thermal decompositions of the new adducts do not yield lower adducts but, instead, uranyl derivatives are formed.

The preparation of $\text{UF}_4\text{O} \cdot \text{SbF}_5$,² which necessitates the warming of stoichiometric quantities of UF_4O and SbF_5 in anhydrous HF and removal of the solvent, has three problems associated with it. The first is that instability of $\text{UF}_4\text{O} \cdot \text{SbF}_5$ -HF solutions gives rise to UF_2O_2 compounds.³ This can be minimized by keeping the reaction time to a minimum. The second is the facility with which $\text{UF}_4\text{O} \cdot 2\text{SbF}_5$ is formed and the fact that the small excess of SbF_5 , used to ensure complete reaction of the UF_4O ,² results in contamination of the product with 1 : 2 adduct. The third is that, because of the low solubility of UF_4O in HF, traces of unreacted UF_4O always remain. The present study includes an improved method of preparation of $\text{UF}_4\text{O} \cdot \text{SbF}_5$.

Results

Gaseous arsenic pentafluoride does not react with solid UF_4O at temperatures up to 60 °C, nor is there evidence for formation of $\text{UF}_4\text{O} \cdot \text{AsF}_5$ adducts when large excesses of AsF_5 are added to UF_4O in anhydrous HF between room temperature and 40 °C for periods of up to 12 h. On prolonged exposure, evidence for the formation of uranyl species is observed.

The $\text{UF}_4\text{O} \cdot 3\text{MF}_5$ ($\text{M} = \text{Nb}$ and Ta) adducts were prepared by shaking UF_4O with a greater than five-fold excess of the appropriate pentafluoride in anhydrous hydrogen fluoride for 1 h at 50–60 °C followed by removal of solvent at room temperature and excess of pentafluoride by sublimation at 40–50 °C.

Owing to the low solubility of the UF_4O in HF, it is difficult to achieve complete reaction and some UF_4O is always present in the final solid residue. This is the case even

when $\text{UF}_4\text{O} \cdot \text{MF}_5$ mixtures of up to a 1 : 12 molar ratio are employed. Secondly, the adducts formed, like their SbF_5 analogues, are unstable in anhydrous HF. On standing for periods longer than 1 h the orange colour in the solvent begins to fade and colourless crystals of uranium hexafluoride appear. After 2 h, in addition to UF_6 , pale yellow solids containing mainly uranyl species are deposited and the solutions assume a green tinge.

The NbF_5 and TaF_5 adducts seem less stable in anhydrous HF than those of SbF_5 . The reaction conditions outlined are optimum ones, giving a minimum of unreacted UF_4O and a minimum of uranyl derivative. Attempts to attain complete reaction by employing longer reaction times or higher temperatures resulted only in the production of higher proportions of uranyl impurity. Attempts to induce complete reaction by the addition of trace quantities of fluorine or antimony pentafluoride were also unsuccessful.

Attempts to prepare lower $\text{UF}_4\text{O} \cdot n\text{MF}_5$ ($\text{M} = \text{Nb}$ or Ta ; $n = 1$ or 2) adducts by the reaction of the appropriate stoichiometric ratios of UF_4O and MF_5 in anhydrous HF at temperatures up to 80 °C resulted in the production of only those adducts described previously together with unreacted UF_4O and uranyl impurity.

Reaction of UF_4O with a greater than five-fold excess of BiF_5 with shaking in anhydrous HF at 40–50 °C and removal of solvent at room temperature and excess of pentafluoride at 50–60 °C, as in the niobium and tantalum case, yielded $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$.

In contrast to the NbF_5 and TaF_5 cases, the greater reactivity of BiF_5 and the higher solubility of the BiF_5 adduct in anhydrous HF ensured completeness of the reaction, and the greater stability of the adduct is evidenced by the absence of significant uranyl impurity.

Efforts to produce stable $\text{UF}_4\text{O} \cdot 3\text{BiF}_5$ were unsuccessful. Fused mixtures of UF_4O and BiF_5 in a 1 : 3 ratio exhibited vibrational spectra and X-ray powder diffraction patterns which could be matched with those of $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$ but displayed additional features which were not coincident with those associated with BiF_5 . On heating, however, BiF_5 was slowly lost until $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$ remained. In view of the fact that $\text{UF}_4\text{O} \cdot 3\text{SbF}_5$ decomposes on gentle heating to give the more stable 1 : 2 adduct,² it seems likely that, in the bismuth case, only the 1 : 2 compound is stable. The fact that the 1 : 3 mixture showed no X-ray lines coincident with those of BiF_5 may be because of changes due to the BiF_5 molecules being weakly associated.

Gentle warming of $\text{UF}_4\text{O} \cdot \text{MF}_5$ ($\text{M} = \text{Nb}$, Ta , and Bi) mixtures at 50–60 °C under a pressure of argon (*ca.* 200 mmHg) and subsequent removal of argon and unreacted

† Non-S.I. unit employed: mmHg $\approx$ 134 Pa.

Table 1. Vibrational data (1 000—400 cm^{-1}) of the adducts of UF_4O with NbF_5 and TaF_5 compared with those of UF_4O , NbF_5 , and TaF_5

UF_4O		TaF_5 Raman ^c	NbF_5		$\text{UF}_4\text{O} \cdot 3\text{TaF}_5$		$\text{UF}_4\text{O} \cdot 3\text{NbF}_5$	
I.r. ^a	Raman ^b		I.r. ^c	Raman ^c	I.r.	Raman	I.r.	Raman
					(982w) ^d 910ms 903 (sh)	910m	(980w) ^d 907ms 900 (sh)	906w
890vs	895vs 889vs 882vs	757vvs		766vvs 752w	753ms 748 (sh)	756ms 742m	744 (sh) 736 (sh)	767vs 755mw 741w 719s
		727m	734vs	716vs	725 (sh) 705 (sh)	726w 702m	708s	
		699ms	688s			686mw 677w	692s	698vw
660vs	665s	671w 646mw	661m	668w 656m	663vs,br 642vs 585m 571 (sh)	661w 633w 581w,br	672 (sh) 662vs 610mw 580mw 568m 552mw	670w 659mw
550s	550m							
466m,br			514ms 479w		506m,br 470m,br	480w,br	488m,br 432w,br	

^a This work. ^b Ref. 2. ^c Ref. 17. ^d Band due to uranyl impurity.

pentafluoride under vacuum also yields the adducts $\text{UF}_4\text{O} \cdot 3\text{MF}_5$ ($\text{M} = \text{Nb}$ and Ta) and $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$.

X-Ray powder diffraction patterns of $\text{UF}_4\text{O} \cdot 3\text{NbF}_5$ and $\text{UF}_4\text{O} \cdot 3\text{TaF}_5$ resemble each other as do those of $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$ and $\text{UF}_4\text{O} \cdot 2\text{SbF}_5$, however, neither pair is isostructural.

Infrared and Raman spectral data of powdered $\text{UF}_4\text{O} \cdot 3\text{NbF}_5$ and $\text{UF}_4\text{O} \cdot 3\text{TaF}_5$ are recorded in Table 1. The similarity of the vibrational spectra of the new adducts to those of $\text{UF}_4\text{O} \cdot 3\text{SbF}_5$,² is not surprising. Since the pentafluorides themselves are all tetramers in the solid state⁴ their adducts with UF_4O might reasonably be expected to have related structures. In all three cases the U—O vibrational stretching frequency indicates that the oxygen is non-bridging and the shift to higher frequency, relative to that of UF_4O itself, is indicative of some ionic contribution to the bonding (*cf.* $\text{UF}_4\text{O} \cdot 2\text{SbF}_5$,²). It is clear, however, that the adducts are not salts since the spectra do not contain bands which match those of $[\text{MF}_6]^-$ in $\text{Cs}[\text{MF}_6]$ ^{5,6} or $[\text{M}_2\text{F}_{11}]^-$ in $\text{Cs}[\text{M}_2\text{F}_{11}]$ ⁷ ($\text{M} = \text{Nb}$ and Ta). In fact, the spectra are too complex to be definitively assigned but bands in the 760—600 cm^{-1} region can be attributed to terminal M—F stretching, in the 600—500 cm^{-1} region to terminal M—F and U—F stretching, and in the 500—400 cm^{-1} region to $\text{M} \cdots \text{F} \cdots \text{U}$ and $\text{M} \cdots \text{F} \cdots \text{M}$ bridge stretching. It seems likely, therefore, that the structures of these adducts will be related to that of $\text{UF}_2\text{O}_2 \cdot 3\text{SbF}_5$ which contains SbF_6 units and Sb_2F_{11} side-chains fluorine bridged to the uranium atoms.⁸ The ionic contribution to the bonding from $[\text{UF}_2\text{O}][\text{MF}_6][\text{M}_2\text{F}_{11}]$ will be only small.

The vibrational data for $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$ are compared with those of UF_4O and BiF_5 in Table 2. In common with $\text{UF}_4\text{O} \cdot n\text{SbF}_5$ ($n = 1-3$)² and $\text{UF}_4\text{O} \cdot 3\text{MF}_5$ ($\text{M} = \text{Nb}$ and Ta), $\nu(\text{U}=\text{O})$ in $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$ is shifted to a higher frequency than that of UF_4O . Comparison of the spectra with those of $\text{Cs}[\text{BiF}_6]$ ⁹ and other $[\text{BiF}_6]^-$ salts involving the cations O_2^+ ,^{10,11} NO^+ ,¹² NF_4^+ ,¹³ H_3O^+ ,¹⁴ and $[\text{ClOF}_2]^+$ ¹⁵ shows that frequencies characteristic of the hexafluorobismuthate ion are not observed. The possible occurrence of $[\text{Bi}_2\text{F}_{11}]^-$ in the adduct cannot be so readily discounted. Previous reports of Raman spectra of the $[\text{Bi}_2\text{F}_{11}]^-$ ion have been published,^{11,16}

but the spectra in one¹¹ are so dominated by the presence of bands due to $[\text{BiF}_6]^-$ that they are not useful for comparison. Those of $[\text{XeF}_3][\text{Bi}_2\text{F}_{11}]$, $[\text{XeF}][\text{Bi}_2\text{F}_{11}]$, and $\text{Cs}[\text{Bi}_2\text{F}_{11}]$ ¹⁶ are more complex than that of $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$ and exhibit additional bands in the 560—530 cm^{-1} region which may be the frequency range of the $\text{Bi} \cdots \text{F} \cdots \text{Bi}$ bridge in the $[\text{Bi}_2\text{F}_{11}]^-$ ion. However, the complex $\text{NF}_4 \cdot \text{BiF}_5 \cdot n\text{BiF}_5$ ($n = 0.6-1.5$)¹³ exhibits a Raman band at 452 cm^{-1} and the i.r. spectrum of pure solid BiF_5 contains a weak broad band at 450 cm^{-1} ,¹⁷ both of which have been correlated with fluorine bridging or the presence of polyanionic species.^{13,17} Since $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$ exhibits broad bands at 477 and 430 cm^{-1} , which might therefore be reasonably attributed to bridging fluorine, and a number of bands in the terminal Bi—F stretching region, the adduct probably contains pentafluoride-like polymeric units. Because of the close chemical similarity between SbF_5 and BiF_5 it seems likely that the structure will be closely related to that of the fluorine-bridged adduct $\text{UF}_4\text{O} \cdot 2\text{SbF}_5$.²

Although complete reaction of $\text{UF}_4\text{O} \cdot \text{BiF}_5$ mixtures of stoichiometries between 1 : 1 and 1 : 2 in anhydrous HF could not be achieved, the i.r. spectra of the products exhibited bands which might be attributed to $\text{UF}_4\text{O} \cdot \text{BiF}_5$. A typical spectrum showed peaks at 904m, (896mw), (892mw), 673mw (661m), 616s, 585 (sh), 568s, (552 sh), 510vw,br, and 485w,br cm^{-1} (peaks in parentheses are close to those associated with UF_4O). The position of $\nu(\text{U}=\text{O})$ at 904 cm^{-1} is different from that observed for $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$ but is at approximately the frequency expected for $\text{UF}_4\text{O} \cdot \text{BiF}_5$ and correlates well with the value of 907 cm^{-1} found for $\text{UF}_4\text{O} \cdot \text{SbF}_5$ ² (SbF_5 being more acidic than BiF_5).

The niobium and tantalum adducts are almost insoluble in HF, SO_2ClF , $\text{CF}_2\text{ClCCl}_2\text{F}$, CF_2Cl_2 , and WF_6 , and react with CH_3CN . As a result, ^{19}F n.m.r. spectra were not obtained. However, at 90 °C in molten TaF_5 the $\text{UF}_4\text{O} \cdot 3\text{TaF}_5$ adduct exhibited a broad band at *ca.* 90 p.p.m. upfield of CFCl_3 due to exchange.

The heating of $\text{UF}_4\text{O} \cdot 3\text{MF}_5$ ($\text{M} = \text{Nb}$ and Ta) to 100 °C, under dynamic vacuum in glass sublimation tubes, results in their complete decomposition. The $\text{UF}_4\text{O} \cdot 2\text{BiF}_5$ adduct, like $\text{UF}_4\text{O} \cdot 2\text{SbF}_5$, is thermally much more stable. Decomposition

Table 2. Vibrational data (1 000–200 cm⁻¹) of the adduct UF₄O·2BiF₃, compared with those of pure UF₄O and BiF₃

UF ₄ O		BiF ₃		UF ₄ O·2BiF ₃	
I.r. ^a	Raman ^b	I.r. ^c	Raman ^c	I.r.	Raman
				(998vww) ^d	
				911s	911m
				906s	
890vs	895vs 889vs 882vs			678ms	678m
660vs	665s	627s		672 (sh)	
				632 (sh)	
				620vs	
				610 (sh)	611s
550s	550m		595	587s	590vs
			570	568s	664 (sh)
466m,br				ca. 500vw (sh)	
		450m,br		477m,br	489w,br
	345vw 276m			430m,br	
				370w,br	
				272w,br	247mw (sh)
		220m	255	235w (sh)	230mw
	201m			214mw,br	216mw (sh)
	148s		167		175w,br
	117m		101		153w,br

^a This work. ^b Ref. 2. ^c Ref. 17. ^d Band arising from uranyl impurity.

at 100 °C is slow and no loss of BiF₃ is observed. In all cases vibrational spectroscopic and X-ray diffraction studies show that lower adducts, such as UF₄O·MF₃, are not produced. However, i.r. spectra of the solid residues after heating show bands characteristic of uranyl species and fewer terminal metal–fluorine stretching bands are observed. It seems likely, therefore, that adducts related to UF₂O₂·nSbF₃ (*n* = 2–4)^{3,8} are produced.

The adduct UF₄O·2SbF₃ was obtained by gentle warming of stoichiometric quantities of SbF₃ with UF₄O in anhydrous HF for less than 1 h and removal of the hydrogen fluoride solvent by decantation. Repeated washing with and decantation of anhydrous HF from the product resulted in the removal of UF₄O and UF₄O·2SbF₃ impurities, and pure UF₄O·SbF₃ was obtained.

Discussion

X-Ray single-crystal structures of UF₄O·2SbF₃² and UO₂F₂·3SbF₃⁸ have shown that both are fluorine-bridged Lewis acid–base complexes. The adducts UF₄O·3MF₃ (*M* = Nb and Ta) and UF₄O·2BiF₃ have vibrational spectra which are related to the antimony pentafluoride compounds and imply related structures. In all of the compounds some small degree of ionicity occurs with the extent of the ionic character implicit in the values of the frequencies of the U=O stretching vibrations. The shifts to higher frequency than those of UF₄O itself are a result of a withdrawal of electron density from the UF₄O towards the more acidic pentafluorides. The magnitude of these shifts is generally small, and in some cases less than likely experimental error, but agrees with the expected relative Lewis acidities of the pentafluorides, SbF₃ ≥ BiF₃ > TaF₃ > NbF₃ (Table 3).

The ready derivation of uranyl species and UF₆ from UF₄O·SbF₃ adducts³ and from the adducts described in this paper is interesting in view of recent work¹⁸ which suggests that from a thermochemical point of view UF₄O behaves as a very loosely bound 1 : 1 complex of UF₂O₂ and UF₆.

Experimental

Starting Materials.—Uranium tetrafluoride oxide was prepared as described by Wilson.¹⁹ Bismuth pentafluoride (Ozark-Mahoning Co.) was purified by vacuum sublimation at 120 °C. Tantalum, niobium, and arsenic pentafluorides were prepared from the elements as described previously.²⁰ The purification of anhydrous HF has also been described elsewhere.⁸

Syntheses.—All manipulations were carried out under anhydrous conditions using procedures outlined previously.²

The uranium tetrafluoride oxide adducts were prepared by two methods. In the first, weighed amounts of UF₄O and MF₃ (*M* = Nb, Ta, or Bi) in molar ratios of 1 : ≥5 were introduced into weighed prefluorinated ¼-in FEP or ¾-in Kel-F reactors in a dry-box. Typically, 0.5–1.0 mmol quantities of UF₄O were employed. The reactors were pumped to high vacuum before the introduction of anhydrous HF solvent. After allowing the mixtures to warm to room temperature they were agitated gently for 1 h with further warming [50–60 °C (Nb and Ta), 40–50 °C (Bi)] to give orange-yellow solutions over orange-yellow solids. In no case was complete solution achieved. The adducts formed were all less soluble than their SbF₃ counterparts, a qualitative order of solubilities suggested Bi > Ta > Nb. After 1 h the solvent was removed, initially under static vacuum and finally by pumping on the solid adduct and excess of pentafluoride. In the second method of preparation weighed amounts of UF₄O and excess of MF₃ (*M* = Nb, Ta, or Bi) were warmed gently to 50–60 °C until the pentafluoride became molten and the UF₄O dissolved to give an orange solution. In order to prevent sublimation of the pentafluoride from the reaction zone, a pressure of argon (ca. 200 mmHg) was introduced into the reaction tube. In both reaction methods, after completion of reaction and, in the reactions using HF, the removal of solvent, the excess of pentafluoride was moved to the upper part of the reaction tube by vacuum sublimation at 40–50 °C (Nb, Ta) and 50–60 °C (Bi). When ¼-in FEP reaction tubes were used the weight of the reaction product, and hence the

Table 3. A comparison of the trend observed for $\nu(\text{U}=\text{O})$ in the vibrational spectra (cm^{-1}) of the $\text{UF}_4\text{O}\cdot\text{MF}_3$ adducts ($\text{M} = \text{Bi}, \text{Nb}, \text{Sb}$, or Ta) with the relative Lewis acidities of the pentafluorides

(i) $\text{UF}_4\text{O}\cdot 3\text{SbF}_5^a$

I.r.	Raman
920	921
914	

$\text{UF}_4\text{O}\cdot 3\text{TaF}_5$

I.r.	Raman
910	910
903	

$\text{UF}_4\text{O}\cdot 3\text{NbF}_5$

I.r.	Raman
907	906
900	

UF_4O

I.r. ^b	Raman ^a
890	895
	889
	882

SbF_5

TaF_5

NbF_5

← increasing $\nu(\text{U}=\text{O})$

← increasing e^- withdrawal from UF_4O

← increasing Lewis-acid strength

(ii) $\text{UF}_4\text{O}\cdot 2\text{SbF}_5^a$

I.r.	Raman
912	912

$\text{UF}_4\text{O}\cdot 2\text{BiF}_5$

I.r.	Raman
911	911
906	

UF_4O

I.r. ^a	Raman ^b
890	895
	889
	882

SbF_5

BiF_5

← increasing $\nu(\text{U}=\text{O})$

← increasing e^- withdrawal from UF_4O

← slight increase in Lewis-acid strength

^a This work. ^b Ref. 2.

stoichiometry, was determined by removing the section of the tube containing the sublimed excess of pentafluoride with a knife in the dry-box, refitting the valve to the remainder of the tube containing the adduct, washing out and drying the detached piece of tube, closing one end by moulding, and weighing when evacuated. After evacuating and weighing the, now shortened, reaction tube the weight of solid product was calculated. When $\frac{3}{4}$ -in Kel-F reaction vessels were employed the excess of pentafluoride was removed from the upper part of the reactor in the dry-box using a micro-spatula. In all cases the weight of product was in close agreement with the formulations $\text{UF}_4\text{O}\cdot 3\text{MF}_3$ ($\text{M} = \text{Nb}$ and Ta) or $\text{UF}_4\text{O}\cdot 2\text{BiF}_5$. However in both reaction methods i.r. spectra indicate that the niobium and tantalum compounds were contaminated by a trace amount of uranyl impurity.

The washing of $\text{UF}_4\text{O}\cdot\text{SbF}_5$ free of $\text{UF}_4\text{O}\cdot 2\text{SbF}_5$ and UF_4O in the improved preparation of $\text{UF}_4\text{O}\cdot\text{SbF}_5$ was achieved by carrying out the reaction in a $\frac{1}{4}$ -in h-shaped FEP reactor. After the reaction was complete and the solid had settled, the HF solvent was decanted into the side arm. Repeated distillation of the HF onto the product followed by decantation into the side arm gradually removed $\text{UF}_4\text{O}\cdot 2\text{SbF}_5$ contaminant.

Characterizations.—Reaction stoichiometries were monitored and X-ray diffraction, vibrational spectroscopic, and n.m.r. data were obtained as outlined previously.^{2,8} Thermal-decomposition studies were carried out under dynamic vacuum in Pyrex tubes. Chemical analyses were not attempted for $\text{UF}_4\text{O}\cdot 3\text{MF}_3$ ($\text{M} = \text{Nb}$ and Ta). Analytical results for the bismuth compound confirmed the formulae determined from the weight of product obtained [Found: Bi, 43.75; F, 28.2; O, 1.90; U, 25.15 (by difference). Calc. for $\text{UF}_4\text{O}\cdot 2\text{BiF}_5$: Bi, 44.55; F, 28.35; O, 1.70; U, 25.4%].

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