See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/255776540

Formation and characterization of uranium(VI) chloride fluorides, UFnCl6-n(n=1-5)

ARTICLE in JOURNAL OF THE CHEMICAL SOCIETY DALTON TRANSACTIONS · JANUARY 1984

Impact Factor: 4.1 · DOI: 10.1039/DT9840002127

CITATIONS	READS
10	26

2 AUTHORS, INCLUDING:



Anthony J Downs
University of Oxford

258 PUBLICATIONS 5,184 CITATIONS

SEE PROFILE

Formation and Characterization of Uranium(VI) Chloride Fluorides, UF_nCl_{6-n} (n = 1-5)

Anthony J. Downs * and Christopher J. Gardner
Inorganic Chemistry Laboratory, University of Oxford, South Parks Road, Oxford OX1 3QR

Fluorine-19 n.m.r. measurements show that UF₆ undergoes facile halide exchange with either SiMe₃Cl or UCl₆ in CFCl₃ or CF₂Cl₂ solution at temperatures below -60 °C. Hence all the uranium(v_1) chloride fluorides in the series UF_nCl_{6-n} (n = 1—5) have been characterized virtually definitively. The visible and mass spectra of the mixed halides have been investigated also. Unlike UF₆ and UCl₆, the mixed halides decompose at temperatures above -60 °C liberating chlorine gas and giving halides of uranium(v_1) or uranium(v_2).

Mixed uranium(vi) chloride fluorides, UF_nCl_{6-n} (n = 1--5), have been postulated recently as intermediates in the reaction of uranium hexafluoride with chlorotrimethylsilane, SiMe₃Cl.¹ Linear interpolation of enthalpies of formation and entropies implies that UF₅Cl and UF₄Cl₂ are thermodynamically unstable at room temperature with respect to the formation of β-UF₅ and UF₄ respectively together with gaseous chlorine. However, the only direct evidence of such species reported to date is provided by the i.r. spectra of mixtures of UF6 with TiCl4, HCl, or BCl3 in liquid xenon held at temperatures below -60 °C.2 Unfortunately there is no help to be gained from isotopic effects and the assignment of i.r. absorptions to particular UF_nCl_{6-n} species rests perforce on (i) the order of appearance of new bands and the dependence of their intensities on the relative proportions of the reagents, and (ii) comparisons with the wavenumbers calculated for these species.

We have prepared mixtures of the chloride fluorides UF_nCl_{6-n} (n=1-5) by the exchange reactions of UF_6 with either $SiMe_3Cl$ or UCl_6 in a Freon solvent at temperatures below -60 °C and studied their spectroscopic properties and chemical stability. Here we report the ¹⁹F n.m.r. spectra of the mixtures, the multiplet patterns and parameters of which provide an unambiguous basis for the identification of each of the species UF_5Cl , cis- and trans- UF_4Cl_2 , mer- and fac- UF_3Cl_3 , cis- and trans- UF_2Cl_4 , and $UFCl_5$. We also describe the visible absorption spectra exhibited by such mixtures and the mass spectrum of the most volatile species UF_5Cl . The mixed halides decompose at temperatures much above ca. -60 °C with the release of elemental chlorine and the formation of tetra- or penta-halides of uranium.

Experimental

Apparatus.—All the samples including uranium hexafluoride were prepared using either (i) a metal vacuum-line constructed in $\frac{3}{8}$ -in copper tubing with brass 'Swagelok' unions and brass-bodied Hoke valves and equipped with a handling section made from Teflon-FEP tubing with Teflon unions and needle valves (Production Techniques Ltd.), or (ii) under an atmosphere of dry argon ($H_2O \le 20$ p.p.m.) in a glove-box. Reactions were carried out in Teflon-FEP tubes (Production Techniques Ltd.) with thermally moulded ends, each fitted with a Teflon needle valve. Before use, the tubes were conditioned by admitting fluorine gas to a pressure of 1 atm (ca. 10^5 Pa), typically for 16 h.

Chemicals.—Uranium hexafluoride (British Nuclear Fuels Ltd.) was freshly vacuum-sublimed into a conditioned Kel-F trap held at -45 °C and its vapour pressure measured to check its purity.³ Uranium hexachloride was prepared by

the reaction of BCl₃ with UF₆ following the procedure of O'Donnell and Wilson.⁴ The Freons CFCl₃ and CF₂Cl₂ (B.D.H. Ltd.) were initially dried over 4A molecular sieves, purified by trap-to-trap distillation *in vacuo* and stored over fresh P₂O₅. Chlorotrimethylsilane (Hopkins and Williams) was purified and stored in a similar manner. Nitrogen monoxide was used as supplied by Matheson; nitrosyl chloride was prepared by the reaction between KCl and NO₂⁵ and purified by trap-to-trap distillation *in vacuo*.

Procedures and Spectrometers.—Samples for n.m.r. investigation kept at low temperature were sealed in a section of FEP tubing (length 5—10 cm, external diameter 4 mm).⁶ The ends of the plastic were trimmed and the cold sample was inserted into a standard precision n.m.r. tube which also included a little CD₂Cl₂ as a 'lock'. Fluorine-19 n.m.r. spectra were recorded with the aid of a Bruker FT AM 250 or WM 250 spectrometer operating at 235.7 MHz.

Visible absorption spectra were recorded for solutions containing UF₆ and either SiMe₃Cl or UCl₆ kept at low temperatures. For this purpose, the solution was contained in an FEP tube with specially flattened parallel sides which was supported in an unsilvered Dewar vessel equipped with inner and outer Pyrex windows to give optical access to the sample. The sample was cooled, typically to ca.-83 °C, by a stream of cold nitrogen gas produced from a liquid nitrogen boiler and the temperature was monitored using a copper-constantan thermocouple. Spectra were measured on a Perkin-Elmer Lambda 3 spectrophotometer.

Mass spectroscopic measurements were carried out with a V.G. Micromass 12B mass spectrometer operating with an electron beam of 70 eV and an accelerating voltage of 3.0 kV; the vapour sample was admitted via a nozzle at room temperature. The spectra were calibrated with reference to peaks due to air or 'L'-grade perfluorokerosene. Infrared spectra of vapours and solids were measured with a Pye-Unicam model SP 2000 and Raman spectra of solids with a Spex Ramalog 5 spectrophotometer.

In a typical experiment, approximately 70 mg (0.2 mmol) of UF₆, measured tensimetrically, was condensed in a conditioned FEP reaction vessel at -196 °C. About 5 cm³ of liquid CFCl₃ or CF₂Cl₂ was co-condensed with the UF₆, followed by the required amount of SiMe₃Cl, also measured tensimetrically. The reaction mixture was then warmed up to -63 °C by supporting the vessel in a chloroform 'slush' bath and shaken periodically. As soon as the Freon melted, the mixture took on a deep red or brown colouration. With molar proportions SiMe₃Cl: UF₆ \leq 6:1, one mol of SiMe₃F, identified by its i.r.⁷ and ¹H and ¹⁹F n.m.r.⁸ spectra, was produced for every mol of SiMe₃Cl taken, as demonstrated for example by

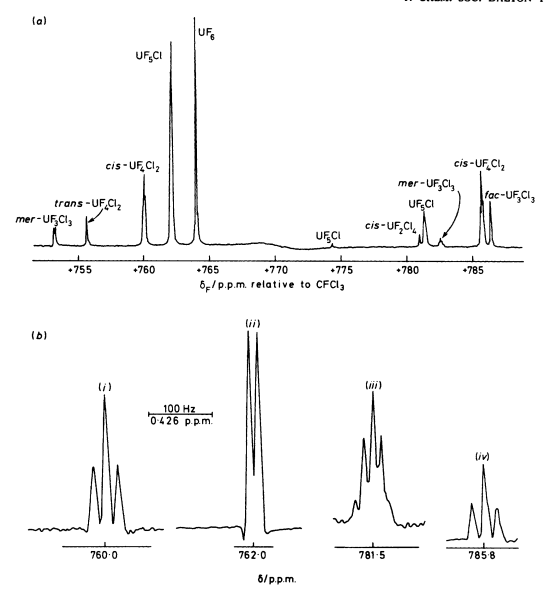


Figure 1. (a) Part of a ¹⁹F n.m.r. spectrum (235 MHz) of a CFCl₃ solution containing an approximately equimolar mixture of UF₆ and SiMe₃Cl at -73 °C. (b) Some details of the ¹⁹F n.m.r. spectrum shown in (a): (i) lower frequency triplet from cis-UF₄Cl₂; (ii) doublet, and (iii) quintet from UF₅Cl; (iv) higher frequency triplet from cis-UF₄Cl₂

fractionation of the volatile components of the reaction mixture contained in CF_2Cl_2 .

Results

Fluorine-19 N.M.R. Spectra.—The ¹⁹F n.m.r. spectrum of a CFCl₃ solution containing a mixture of UF₆ and SiMe₃Cl held at ca. -75 °C showed resonances attributable to UF₆ 9 and SiMe₃F 8 at δ +764 and -159 p.p.m. respectively relative to CFCl₃. In addition there were numerous resonances, some with distinctive multiplet structures, between δ +745 and +790 p.p.m. the intensities of which varied markedly with the molar proportions SiMe₃Cl: UF₆.

These new resonances are illustrated in Figure 1(a) which shows the relevant part of the spectrum exhibited by a solution approximately 6 mmol dm⁻³ in UF₆ and having SiMe₃Cl: UF₆ = 1:1; the multiplet structures of some of the resonances are illustrated in the expanded traces of Figure 1(b). A solution

with a similar concentration of UF₆ but with SiMe₃Cl: UF₆ = 1:3 gave a spectrum consisting essentially of the singlets at $\delta + 764$ and -159 p.p.m. due to UF₆ and SiMe₃F respectively, together with a quintet centred at $\delta + 781.5$ p.p.m. and a doublet at δ +762.0 p.p.m. The quintet and doublet had relative intensities of ca. 1:4 and neither this nor the positions of the resonances varied with the reacting proportions or concentrations of UF₆ and SiMe₃Cl (even with 100-fold dilution of the solution). These results give unequivocal evidence for the formation of the pseudo-octahedral molecule UF₅Cl. By contrast, a solution with the proportions SiMe₃Cl: UF₆ = 5: 1 gave a strong singlet at δ +774.3 p.p.m. corresponding presumably to UFCl₅ and two much weaker singlets at $\delta + 781.0$ and + 746.1 p.p.m. corresponding to cisand trans-UF₂Cl₄. Systematic variations of the proportions SiMe₃Cl: UF₆, combined with observations of the frequencies, multiplet patterns, and relative intensities of the new resonances, allow us to identify all eight of the uranium(vi) chloride fluorides: UF₅Cl, cis- and trans-UF₄Cl₂, mer- and fac-UF₃Cl₃,

Table 1. Details of the 19 F n.m.r. spectra of the compounds UF_nCl_{6-n} (n = 1—6) in CFCl₃ solution at -73 °C

Compound	N.m.r. system	$\delta_F^*/p.p.m.$	$^{2}J(^{19}F^{-19}F)/Hz$
UF ₆	A_6	+764.0	
UF₅Cl	A ₄ X	$A_4 + 762.0$ X + 781.5	13.3 \pm 0.6
cis-UF ₄ Cl ₂	A_2X_2	$A_2 + 760.0$ $X_2 + 785.8$	
trans-UF4Cl2	A_4	+755.5	
mer-UF ₃ Cl ₃	A_2X	$A_2 + 753.0$ X + 782.6	24.0 \pm 2.5
fac-UF3Cl3	A_3	+ 786.4	_
cis-UF2Cl4	A_2	+781.0	_
trans-UF2Cl4	A_2	+746.1	
UFCI ₅	Α	+774.3	

* 19 F Chemical shifts: frequency measured with reference to the resonance due to the CFCl₃ solvent; estimated uncertainty ± 0.1 p.p.m.

cis- and trans-UF₂Cl₄, and UFCl₅. Details of the relevant n.m.r. parameters are given in Table 1.

Analogous measurements carried out with a CCl₃F solution containing a mixture of UF₆ and UCl₆, again held at ca. -75 °C, yielded a ¹⁹F n.m.r. spectrum identical with that derived from a mixture of UF₆ and SiMe₃Cl having the same proportions of fluoride and chloride, apart from the absence of a resonance due to SiMe₃F. Changing the proportions of UF₆ and UCl₆ had the same effect as changing the proportions of UF₆ and SiMe₃Cl.

The relative (integrated) intensities of the ¹⁹F resonances associated with the different members of the series UF, Clowere estimated for mixtures with different proportions of fluoride and chloride in an attempt to assess the distribution of the products. It is of course difficult to relate the peak areas to the concentrations of the resonating nuclei without some knowledge of the relaxation times. However, we found that the integrated intensities of the quintet and doublet due to UF₅Cl were in the ratio 3.9:1 (as opposed to the expected ratio of 4:1). These two resonances span about one-half of the total range of chemical shifts exhibited by the UF_rCl_{6-r} molecules and we have therefore been encouraged to assume that there is a constant proportionality between the peak area and concentration of the resonating nucleus. For a mixture with $SiMe_3Cl: UF_6 = ca. 1.3: 1$, the relative concentrations are thus estimated to be as follows: UF₆: UF₅Cl: UF_4Cl_2 (both isomers): UF_3Cl_3 (both isomers) = 1:2.95: 2.78: 1.06 (with estimated errors in the order of 5%). For a random distribution of products in such a mixture we expect $UF_6: UF_5Cl: UF_4Cl_2: UF_3Cl_3 = 1:1.66:1.15:0.42.$ cis-UF₄Cl₂: trans-UF₄Cl₂ and fac-UF₃Cl₃: mer-UF₃Cl₃ were found for this and other mixtures to be 12:1 and 1.5: 1 respectively, whereas the spectra of mixtures richer in chloride indicated that cis-UF₂Cl₄: trans-UF₂Cl₄ = 7:1. Random statistics lead one to expect cis and trans isomers of pseudo-octahedral MX₄Y₂ molecules to be formed in the proportions 4: 1 and fac and mer isomers of MX₃Y₃ molecules in the proportions 1.5:1.

Visible Absorption Spectra.—The effect of adding successive aliquots of SiMe₃Cl to a solution of UF₆ in CFCl₃ was also monitored by reference to the visible absorption spectrum of the solution maintained at ca. -80 °C; such experiments were carried out with solutions typically 60 mmol dm⁻³ in UF₆.

Solutions rich in UF_6 exhibited a new band centred at 510 nm in addition to the strong absorption which appears at wavelengths shorter than 450 nm (see Figure 2). The position

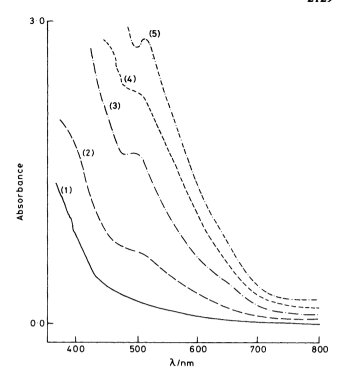


Figure 2. Visible absorption spectra of solutions containing varying amounts of SiMe₃Cl added to a solution of UF₆ (30 mmol dm⁻³) in CFCl₃ held at -83 °C. The proportions UF₆: SiMe₃Cl corresponding to each trace are as follows: (1) 10:1, (2) 5:1, (3) 3.3:1, (4) 2.5:1, (5) 2:1

of the new band and the general appearance of the spectrum were independent of both the initial UF₆ concentration and the proportions SiMe₃Cl: UF₆ within the range 1:20 to 1:2.5. Under these conditions and with due allowance for the loss of a small amount of material through conditioning of the cell (leading to a lower absorbance than expected at the lowest concentrations of SiMe₃Cl), the absorbance of the band at 510 nm increased linearly as a function of the concentration of SiMe₃Cl initially present. This indicates that the band arises from a single product, UF₅Cl, formed in accordance with equation (1). The absorption coefficient of the band, $\varepsilon_{\rm max}$, was estimated to be 290 \pm 30 cm² mol $^{-1}$.

$$UF_6 + SiMe_3Cl UF_5Cl + SiMe_3F$$
 (1)

As the proportion of SiMe₃Cl in the solution increased, the appearance of the absorption spectrum between 400 and 800 nm changed in the manner illustrated in Figure 2. When the ratio SiMe₃Cl: UF₆ reached or exceeded 6:1, the spectrum became virtually invariant with respect to the initial concentration of SiMe₃Cl and was identical with that of a solution prepared from an authentic sample of UCl₆.

Mass Spectra.—A two-fold excess of UF₆ was co-condensed with a frozen CFCl₃ solution of SiMe₃Cl in an FEP tube and the mixture maintained at -63 °C for 4 h. After all the material volatile at this temperature had been evaporated under high vacuum, the tube was connected directly to the stainless-steel gas inlet system of the mass spectrometer. With the sample at -63 °C, no mass peaks attributable to a volatile uranium compound could be detected but with the sample at -45 °C the spectrum of the vapour showed peaks originating not only in the molecular ions UF₆+ but also in UF₅Cl+ as well as fragments of these species. The relevant peaks are

Table 2. Mass spectrum of the vapour over the solid reaction mixture formed by UF₆ and SiMe₃Cl (UF₆: SiMe₃Cl = 2:1) and held at -45 °C

m/e	Relative intensity	Ion 4
370	1.0	UF ₅ 37Cl+
368	3.0	UF ₅ 35Cl+
352	b	UF ₆ +
351	3.5	UF ₄ ³⁷ Cl ⁺
349	10.5	UF ₄ 35Cl+
333	b	UF ₅ +
330	7.2	UF ₃ 35Cl+
314	b	UF ₄ +
311	2.9	UF ₂ 35Cl+
295	b	UF ₃ +
292	2.0	UF35Cl+
276	b	UF ₂ +
273	1.6	U35Cl+
257	b	UF+
238	b	U+
157	Ь	UF ₄ 2+
147.5	b	UF_3^2
138	b	UF ₂ ²⁺
128.5	b	UF2+
119	b	U ²⁺

^a U corresponds to ²³⁸U, F to ¹⁹F. ^b Mass peaks corresponding to UF_n^+ and UF_n^{2+} (n=0—6) were at least one order of magnitude more intense than those due to UF_nCl^+ (n=0—5).

listed in Table 2. The identity of each of the ions UF_5Cl^+ and UF_4Cl^+ was confirmed by the appearance of a doublet with components having relative intensities of 1:3 corresponding to the abundances of the naturally occurring chlorine isotopes ^{37}Cl and ^{35}Cl respectively. However, mass peaks associated with the ions $UF_n^{37}Cl^+$ (n=0-3) could not be resolved because of the high intensity of the adjacent peaks associated with the ions UF_{2+n}^+ .

Chemical Studies.—CFCl₃ solutions containing SiMe₃Cl and UF₆ in various proportions were allowed slowly to warm up from -63 °C to room temperature, with constant stirring. All the volatile components were then evaporated under vacuum and the products investigated by spectroscopic and chemical analysis.

Mixtures with SiMe₃Cl: UF₆ \leq 1:1 gave a pale green involatile solid which was soluble in dry acetonitrile and identified as β -UF₅ on the evidence of its i.r. and Raman spectra. By contrast, a mixture with SiMe₃Cl: UF₆ = 2:1 gave a darker green solid which was insoluble in acetonitrile; elemental analysis indicated that this was UF₄, a conclusion supported by its i.r. spectrum. Hence our results agree with those reported recently by Brown *et al.* When the reaction mixture included an excess of SiMe₃Cl, with SiMe₃Cl: UF₆ \geq 6:1, vacuum evaporation of all the material volatile at -63 °C left what appeared to be a black solid stable at room temperature. This was shown by elemental analysis and by its i.r., ¹² u.v.-visible, ¹³ and mass spectra to be UCl₆.

The volatile material derived from reaction mixtures which had been allowed to warm up to room temperature included, in addition to the solvent, SiMe₃F, UF₆ (from mixtures rich in fluoride), and elemental chlorine (identified by the Raman spectrum of the solid condensate formed on a copper block at –196 °C ¹⁴). Experiments were also carried out with a more volatile solvent to facilitate vaporisation at temperatures low enough to avoid significant decomposition of the uranium(v1) chloride fluorides. For example, a CF₂Cl₂ solution containing an equimolar mixture of UF₆ and SiMe₃Cl was kept at

-63 °C while being subjected to continuous pumping followed by trap-to-trap fractionation of the volatile material. Under these circumstances the vapour species included only the solvent, SiMe₃F, UF₆, and a brown product; little or no chlorine was detected. The brown product collected with UF₆ in a trap held at -83 °C; attempts to separate it from UF₆ have as yet been unsuccessful. Analysis of a typical condensate formed at -83 °C indicated that it was a mixture containing UF₆ and UF₅Cl in roughly equal proportions. When such a condensate was allowed to warm much above -60 °C, it decomposed with the formation of pale green β-UF₅ and gaseous chlorine.

Since NO reacts with UF₆ to form the nitrosonium salt NO⁺UF₆⁻,¹⁵ we investigated the interaction of nitrogen monoxide with a UF₆-SiMe₃Cl reaction mixture in the hope of fixing UF₅Cl or UF₄Cl₂ in the form of the corresponding hexahalogenouranate(v) anion. In fact, admission of NO to a CFCl₃ or CF₂Cl₂ solution having the initial composition SiMe₃Cl: UF₆ = 1:1 or 2:1 and held at -70 °C gave only NOCl (identified by its i.r. spectrum ¹⁶) and either β -UF₅ or UF₄ formed in accordance with an equation such as (2).

$$UF_5Cl + NO \longrightarrow UF_5 + NOCl$$
 (2)

Evidently UF₅ and NOCl do not interact even at -70 °C.

Discussion

Our results demonstrate that facile halide exchange occurs between UF₆ and either SiMe₃Cl or UCl₆ in Freon solutions at temperatures below $-60\,^{\circ}$ C. The solutions vary in colour from red to almost black according to the concentrations and the relative proportions of chloride and fluoride. With the aid of the ¹⁹F n.m.r. spectra it has been possible to identify all the members of the series UF_nCl_{6-n} (n=1-5). The facility of halide exchange tends to frustrate any attempt to separate individual species, although mixtures rich in UF₆ and containing essentially only UF₆ and UF₅Cl may yet prove amenable to low-temperature fractionation.

The assignment of some of the 19F resonances summarised in Table 1 and illustrated in Figure 1 follows directly from the multiplet patterns. This is the case with (i) UF₅Cl characterized by a doublet and a quintet with relative intensities of 4:1. (ii) cis-UF₄Cl₂ characterized by two triplets of equal intensity, and (iii) mer-UF₃Cl₃ characterized by a doublet and a triplet with relative intensities of 2:1. However, the other products each contain fluorine in a single magnetic environment and exhibit only a single 19F resonance. We have identified these resonances by reference to the way in which the intensity varies with the relative proportions of chloride and fluoride. Thus the singlet due to trans-UF₄Cl₂ grows in step with the distinctive multiplets due to the cis isomer and likewise the singlet due to fac-UF₃Cl₃ grows in step with the distinctive multiplets due to the mer isomer. UFCl₅ is identified as the source of the only 19F resonance to persist in solutions containing a large excess of chloride. To distinguish the resonances arising from cis- and trans-UF₂Cl₄, we have constructed a Dean-Evans plot 17 showing how the chemical shift varies with the number of fluorine substituents cis to the resonant fluorine atom (Figure 3). Smooth curves are obtained only if the resonance at lower frequency is assigned to trans-UF₂Cl₄.

The Dean-Evans plot brings to light three features of interest in relation to molecules of the type UF_nCl_{6-n} . (i) We note that fluorine atoms *trans* to chlorine resonate at consistently higher frequencies than fluorine atoms *trans* to a second fluorine atom. Similar behaviour is found for fluorine *trans* to OMe in individual molecules of the type UF_{n-1} of UF_{n-1} but does not necessarily extend from one member

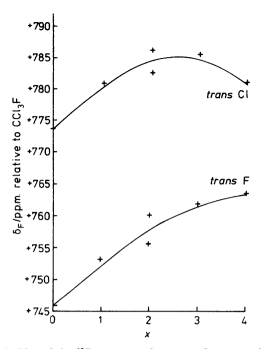


Figure 3. Plot of the ¹⁹F resonance frequency, δ_F , vs. number of fluorine atoms (x) cis to the resonant fluorine in the series UF_n-Cl_{6-n} (n=1—6). The upper curve relates to a resonant fluorine which is *trans* to a chlorine and the lower curve to a resonant fluorine which is *trans* to another fluorine

of the series to another. By contrast, the reverse frequency pattern applies, mutatis mutandis, to molecules in the series $UF_n(OTeF_5)_{6-n}^{19}$ and $WF_nCl_{6-n}^{20}$ (ii) It appears that the resonance of a linear F^-U^-F moiety moves to higher frequency as the number of cis-fluorine atoms increases. Again this parallels the behaviour of molecules in the series $UF_n^-(OMe)_{6-n}^{18}$ but is opposed to the trend exhibited by the series $UF_n(OTeF_5)_{6-n}^{19}$ and $WF_nCl_{6-n}^{20}$ (iii) The resonance frequency of a linear F^-U^-Cl moiety first increases, reaches a maximum and then decreases as the number of cis-fluorine atoms increases. Such a trend is intermediate between those displayed by the series $UF_n(OMe)_{6-n}^{18}$ and $UF_n(OTeF_5)_{6-n}^{19}$.

The ¹⁹F resonance frequencies of octahedral or pseudooctahedral species of the type $[SnF_nX_{6-n}]^{2-}$ (e.g. X = Cl, Br, I, OR, NCO, NCS, or N₃) led Dean and Evans ¹⁷ to the empirical expression (3) for the resonance frequency of nucleus Z, δ_{Z_1} , where p is the number of fluorine atoms cis and

$$\delta_{\mathbf{z}} = pC + qT \tag{3}$$

q the number trans to Z, and C and T are constants for a particular system. This simple linear relation gives a tolerable account of δ_F for the members of the series $UF_n(OMe)_{\delta-n}$, ¹⁸ $UF_n(OTeF_5)_{\delta-n}$, ¹⁹ and $WF_nCl_{\delta-n}$ but Figure 3 shows it to be wide of the mark for the series $UF_nCl_{\delta-n}$.

The relative intensities of the ¹⁹F resonances due to the molecules UF_nCl_{6-n} suggest non-random scrambling, with the equilibria favouring somewhat the mixed halides UF₅Cl, UF₄Cl₂, and UF₃Cl₃. Hence there is probably a small but significant enthalpy change associated with reactions of type (4). It is noteworthy also that although the isomers fac- and

$$\frac{n}{6}UF_6 + \frac{6 - n}{6}UCl_6 \longrightarrow UF_nCl_{6-n}$$
 (4)

mer-UF₃Cl₃ are formed in proportions consistent with random statistics, this is not the case with UF₄Cl₂ and UF₂Cl₄, the cis

isomers of which are 2—3 times more abundant than random statistics would lead us to expect. This possibly reflects the additional dipolar contributions made by interactions between the cis isomers and the CCl₂F solvent molecules.

The visible absorption spectrum associated with UF₅Cl in CFCl₃ solution (see Figure 2) displays two significant features: (i) the distinctive new band centred at ca. 510 nm, and (ii) the marked increase in absorption at shorter wavelengths (≤450 nm). In view of the ambiguity about the correct ordering of the molecular orbitals in the symmetrical UCl₆ molecule,²¹ it is difficult to give an unequivocal interpretation of this spectrum. However, comparisons with the absorption spectra of gaseous UF₆ ($\lambda_{\text{max}} = 369 \text{ nm}$) ²² and UCl₆ in either the gas phase $(\lambda_{\text{max}} = ca. 485 \text{ nm})^{13}$ or in CFCl₃ solution $(\lambda_{\text{max}} = 495 \text{ nm})^{23}$ suggest that the band at 510 nm has significant Cl \rightarrow U chargetransfer character. Presumably the absorption at shorter wavelengths involves mainly F-U charge transfer and the increased intensity reflects the reduction in symmetry of UF₅Cl compared with UF₆ although some admixture of Cl-U charge transfer cannot be discounted. The weak band at 369 nm exhibited by gaseous UF₆ has been ascribed to forbidden $a_{1g} \rightarrow t_{2u}$ and $t_{1g} \rightarrow a_{2u}$ transitions; ²⁴ under C_{4v} symmetry, the corresponding transitions of UF₅Cl are allowed $(a_1 \rightarrow e \text{ and } e \rightarrow b_2)$. This is consistent with the observed absorption coefficients, with $\varepsilon_{\rm max}=ca$. 54 cm² mol⁻¹ for UF₆ at $\lambda=369$ nm ²¹ and $\varepsilon\geqslant460$ cm² mol⁻¹ for UF₅Cl at $\lambda \leq 400$ nm. Unfortunately the absorption of the Teffon-FEP cell precluded measurements at wavelengths shorter than 400 nm.

Conclusions

Uranium hexafluoride reacts rapidly with either chlorotrimethylsilane or uranium hexachloride in Freon solution at temperatures below $-60\,^{\circ}\mathrm{C}$ to produce uranium(vi) chloride fluorides of the type $\mathrm{UF_nCl_{6-n}}$ (n=1-5) which have been characterized by their ¹⁹F n.m.r. and visible absorption spectra. Unlike the hexafluoride and hexachloride, the chloride fluorides decompose at room temperature with the evolution of gaseous chlorine and the formation of uranium(v) or uranium(v) halides. They react with nitrogen monoxide to form not the appropriate nitrosonium hexahalogenouranate(v), $\mathrm{NO^+UF_nCl_{6-n}^-}$ $(cf.\ \mathrm{UF_6^-})$, but nitrosyl chloride and a lower-valent uranium halide.

Acknowledgements

We thank the S.E.R.C. and British Nuclear Fuels, Springfields Works, for support of this research, including the funding of a studentship (to C. J. G.), Drs. E. H. Appelman and J. H. D. Eland for practical help with the measurement of the mass spectra, and Drs. A. E. Derome and B. Hunter and Miss K. Nicholls for similar help with the ¹⁹F n.m.r. spectra.

References

- D. Brown, J. A. Berry, J. H. Holloway, and G. M. Staunton, J. Less-Common Met., 1983, 92, 149.
- 2 W. B. Maier II, W. H. Beattie, and R. F. Holland, J. Chem. Soc., Chem. Commun., 1983, 598; see also W. H. Beattie and W. B. Maier II, Polyhedron, 1983, 2, 1371.
- 3 G. D. Oliver, H. T. Milton, and J. W. Grisard, J. Am. Chem. Soc., 1953, 75, 2827.
- 4 T. A. O'Donnell and P. W. Wilson, in 'Inorganic Syntheses,' ed. F. Basolo, McGraw-Hill, 1976, vol. 16, p. 143.
- 5 C. T. Ratcliffe and J. M. Shreeve, in 'Inorganic Syntheses,' ed. W. L. Jolly, McGraw-Hill, 1968, vol. 11, p. 199.
- 6 R. R. Reinhard, Rev. Sci. Instrum., 1965, 36, 549.

- 7 H. Kriegsmann, Z. Anorg. Allg. Chem., 1958, 294, 113.
- 8 E. Schnell and E. G. Rochow, J. Am. Chem. Soc., 1956, 78, 4178; S. G. Frankiss, J. Phys. Chem., 1967, 71, 3418.
- 9 K. Seppelt and N. Bartlett, Z. Anorg. Allg. Chem., 1977, 436, 122.
- 10 E. Jacob, Z. Anorg. Allg. Chem., 1973, 400, 45; D. Brown, B. Whittaker, J. A. Berry, and J. H. Holloway, J. Less-Common Met., 1982, 86, 75.
- 11 W. Krasser and H. W. Nürnberg, Spectrochim. Acta, Part A, 1970, 26, 1059.
- 12 W. Kolitsch and U. Müller, Z. Anorg. Allg. Chem., 1975, 418, 235.
- 13 W. B. Maies II, R. F. Holland, and W. H. Beattie, Spectrosc. Lett., 1983, 16, 233.
- 14 M. Suzuki, T. Yokoyama, and M. Ito, J. Chem. Phys., 1969, 50, 3392.
- 15 J. R. Geichman, E. A. Smith, S. S. Trond, and P. R. Ogle, *Inorg. Chem.*, 1962, 1, 661.

- 16 L. H. Jones, R. R. Ryan, and L. B. Asprey, J. Chem. Phys., 1968, 49, 581.
- 17 P. A. W. Dean and D. F. Evans, J. Chem. Soc. A, 1968, 1154.
- 18 E. A. Cuellar and T. J. Marks, *Inorg. Chem.*, 1981, 20, 2129.
- 19 K. Seppelt, Chem. Ber., 1976, 109, 1046.
- 20 G. W. Fraser, C. J. W. Gibbs, and R. D. Peacock, J. Chem. Soc. A, 1970, 1708.
- 21 G. Thornton, N. Edelstein, N. Rösch, R. G. Egdell, and D. R. Woodwark, J. Chem. Phys., 1979, 70, 5218.
- 22 W. B. Lewis, L. B. Asprey, L. H. Jones, R. S. McDowell, S. W. Rabideau, A. H. Zeltmann, and R. T. Paine, J. Chem. Phys., 1976, 65, 2707.
- 23 C. J. Gardner, unpublished work.
- 24 M. Boring and J. W. Moskowitz, Chem. Phys. Lett., 1976, 38, 185.

Received 29th December 1983; Paper 3/2287