



Reactivity and Thermal Stability of Hexafluorides

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Reactivity and Thermal Stability of Hexafluorides

N.P.Galkin and Yu.N.Tumanov

The review deals with the possibility of using various physicochemical characteristics of the hexafluorides of p, d, and f elements for a comparative estimation of their thermal stability and reactivity. The force constants of EF₆ molecules, the average bond energy and length, and also the thermodynamic stability constants of the octahedral EF₆ molecules have been used as characteristic properties.

It has been shown that the force constants of the E-F bond are not effective characteristics of the thermal stability and reactivity of all hexafluorides in reactions where more than one E-F bond is dissociated.

Only the thermodynamic stability constants can serve as an objective criterion for the assessment when account is taken of definite dissociation mechanisms of EF₆ molecules (complete or incomplete dissociation).

Within the limits of individual groups of 4d, 5d, and 5f elements and also sulphur, selenium, and tellurium, the variation of the force constants and thermodynamic stability in relation to the incomplete dissociation of the hexafluorides corresponds to the experimental sequences of their thermal stabilities.

The bibliography includes 85 references.

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I. INTRODUCTION

Hexafluorides are a class of fluoride compounds formed by certain elements having not less than $\sin p$, d, or f valence electrons. Up to the present time, the hexafluorides of 18 elements have been identified: sulphur, chromium, selenium, molybdenum, technetium, ruthenium, rhodium, tellurium, xenon, tungsten, rhenium, osmium, iridium, platinum, polonium, uranium, neptunium, and plutonium. In addition one may expect, by analogy, the formation of the hexafluorides of palladium, iodine, astatine, radon, and americium.

The common properties of hexafluorides are low melting and boiling points and comparatively high volatility under ordinary conditions (Table 1). With the exception of XeF₆, the molecules of hexafluorides EF₆ have the configuration of a regular octahedron belonging to the $O_{\rm h}$ point group with the element E in the centre. The reactivities of hexafluorides are not the same. Some hexafluorides, for example CrF₆, are so unstable that they can be synthesised and investigated only at very low temperatures, while others (for example sulphur hexafluoride) resemble the noble gases in their chemical inertness.

Hexafluorides have attracted much interest, since those of many elements are obtained and used in individual stages of the technology of the manufacture of nuclear fuel ¹⁻³. In particular, this accounts for the publication of a large number of studies on the physicochemical properties of both individual hexafluorides and individual groups. Reviews have also appeared on hexafluorides, among which mention should be made in the first place of those by Weinstock ²⁻⁴ and O'Donnel and coworkers ⁵⁻⁸.

In Weinstock's work attention was concentrated on the molecular constants of hexafluorides. However, there

Table 1. Physicochemical properties of hexafluorides 3,4,8.

		,					
Hexa- fluoride	Type of charac- teristic elec- trons of element E	No. of free electrons	Symmetry of ground state of EF ₆	M.p., °C	B.p. (sublimation temp.) at p = 1 atm., °C	Critical temp., °C	Vapour pressure at 15°C, mmHg
	1				1		
SF_6	3 p	0	A _{1g}	-50.4*	-63.7	45.5	
SeF_6	4 p		A ₁₀	-34.7*	-45.9	72.35	
TeF ₈	5 p	-	A_{1g}	-37.0*	-38.9	83.25	_
PoF_6	6 p		A_{1g}^{-3}	_	_		
CrF_6	3 d	0		_	_	_	_
MoF ₆	ļ	0	A _{1g}	17.4	35	_	361
TcF,	4 d	1	G _{3/2g}	37.4	55.3		135
RuF _e			F _{2g}	50.4	45.9		58
RhF ₆		2 3	G _{3/2g}	70	_	_	38
WF ₆		0	A _{1g}	2.0	17.5	_	698
ReF ₆	5 d	1	Gulag	18.5	33.7	_	358
OsF ₆		2	$G_{3/2g}$ E_g, F_{2g}	32.1	45.9	_	312
IrF ₆		2 3	G _{3/2g} 2g	43.8	53.6	177	227
PtF ₆		4	$A_g^{3/2g}$	61.3	69.14	_	66
UF,		0	A_{1g}	64.1**	56.5	245	56
NpF ₆	5 f	1	$E_{5/2}^{1g} u$	55.1**	55.2	-	64
PuF ₆	,	2	A _{1g}	51.6**	62,2	_	53
XeF ₆		2		26		_	12

^{*} Under the pressure of its own vapour.

are certain data on their thermal stabilities 3 , 4 . O'Donnel and coworkers attempted to make a systematic arrangement of the hexafluorides of certain elements in terms of their reactivities. In particular, they showed that the reactivities of hexafluorides fall on moving down the Groups of the Periodic System, while the thermal stabilities increase on going from elements with 3d electrons to elements with 5d electrons $^{5-7}$.

^{**} Triple point.

In the solution of certain practical problems associated with the presence in a chemical system of the hexafluorides of several elements or their lower fluorides, it is necessary to employ some qualitative criterion in order to predict the changes in the system which are potentially possible. Thermodynamic analysis is in many cases impossible, since the thermochemical and thermodynamic constants of many fluorides are unknown. On the other hand, an empirical systematic arrangement may prove to be unreliable, since the rate of a chemical reaction depends on the experimental conditions. For example, in the system "uranium hexafluoride-hydrogen" the products may be uranium pentafluoride, tetrafluoride, trifluoride, or the so called "intermediate" fluorides U2F9, U4F17, etc., depending on the temperature, the reactant ratio, and the effect of light. Until recently it was also believed that molybdenum and tungsten hexafluorides are vigorous fluorinating agents 9,10, but later studies 5,11 showed that these two hexafluorides are inert.

In the present review the possibility of using various physicochemical characteristics of hexafluorides for a comparative estimation of their reactivities and thermal stabilities is discussed. Such characteristics may be the force constants of the EF_6 molecules, the $\mathrm{E-F}$ bond energies and lengths, and the equilibrium dissociation constants of hexafluorides. An attempt has been made to discover regularities in thermal stability both for the entire class of hexafluorides and for individual series.

II. PRINCIPAL GROUPS OF HEXAFLUORIDES

Depending on the nature of the valence shell of the element forming the hexafluoride or the type of its valence electrons, all hexafluorides can be arranged systematically in groups according to the position of the elements in the Periodic System. The positions of the hexafluorides of various elements in these groups are consistent with the positions of the elements themselves, either in the Periods (horizontal series) or in the Group (vertical series) of the Periodic System. The horizontal series of hexafluorides comprise the groups of hexafluorides of 3d (CrF₆), 4d (MoF₆, TcF₆, RuF₆, RhF₆), 5d (WF₆, ReF₆, OsF_6 , IrF_6 , PtF_6), and 5f elements (UF_6, NpF_6, PuF_6) . The vertical series of hexafluorides comprise (1) the hexafluorides of the non-metals in Group VI of the Periodic System (SF₆, SeF₆, TeF₆, PoF₆), (2) the hexafluorides of the transition elements of Group VI (CrF6, MoF_6 , WF_6 , UF_6), (3) the hexafluorides of elements with one unshared electron (TcF₆, ReF₆, NpF₆), (4) the hexafluorides of elements with two unshared electrons (RuF6, OsF_6 , PuF_6), (5) the hexafluorides of elements with three unshared electrons (RhF₆, IrF₆), and (6) platinum hexafluoride (PtF₆). The platinum atom in this molecule has four unshared electrons which, judging from the type of the ground state of the PtF₆ molecule (Table 1), are paired. Xenon hexafluoride, the molecule of which does not belong to the symmetry point group O_h , occupies a special place.

The electronic ground states of eleven hexafluorides of sulphur, selenium, tellurium, molybdenum, tungsten, uranium, neptunium, plutonium, rhodium, iridium, and platinum are either non-degenerate or the degeneracy is predominantly of the spin type. The normal vibrational spectra of these molecules correspond to O_h symmetry. The vibrational spectra of the hexafluorides of technetium, ruthenium, rhenium, and osmium have appreciable anomalies (compared with the preceding hexafluorides),

which are attributed to the dynamic Jahn-Teller effect 4,12-15. According to the Jahn-Teller theorem 16, the orbital degeneracy of the electronic ground state of octahedral molecules is reduced by distortion of their configuration, i.e. by a decrease of molecular symmetry. This is the so called static Jahn-Teller effect. However, in the present case the static Jahn-Teller effect is not observed, but there is an anomaly in the infrared absorption spectra owing to a vibronic interaction.

Table 2. Width of the half-absorption bands of certain hexafluorides (cm⁻¹) 4.

	ν,	/2			ν ₁	/2	Δν _{1/3}	
Molecule	v _t +v _s	v _z +v ₃	Δν _{1/2}	Molecule	ν ₁ +ν ₃	v ₂ +v ₃		
MoF ₆ TcF ₆ RuF ₆ RhF ₆	20 23 21 19	31 113 — 27	0 84 - 0	WF6 ReF6 OsF6 IrF6 PtF6	17 36 23 18 20	24 68 84 26 26	0 43 59 0	
Mean*	21	29		Mean*	20	25		

^{*} The band widths for "anomalous" hexafluorides were disregarded in the averages.

The infrared spectra of the "normal" hexafluorides usually have two characteristic bands corresponding to the compound frequencies $\nu_1+\nu_3$ and $\nu_2+\nu_3$ in the region of 1140-1710 cm $^{-1}$ and having almost identical intensities and outlines (Table 2). For each hexafluoride, the long wavelength band corresponds to the compound frequency $\nu_2+\nu_3$ and the short wavelength band to the frequency $\nu_1+\nu_3$. However, whereas the outlines of the $\nu_1+\nu_3$ bands are virtually identical for both "normal" and "anomalous" hexafluorides, those of the $\nu_2+\nu_3$ bands for different types of hexafluorides are not the same, which is a qualitative criterion of the "anomaly" of the hexafluorides.

The relations between the observed frequencies and the number of unshared valence electrons in the molecules of the "normal" hexafluorides, derived by Weinstock and Goodman 3 ,4, make it possible to estimate the vibration frequencies ν_2 and ν_5 which the molecules of the hexafluorides of technetium, ruthenium, rhenium, and osmium would have in the absence of the dynamic Jahn-Teller effect. The points corresponding to the ν_2 and ν_5 frequencies of the "anomalous" hexafluorides deviate from the graphical relations. It is interesting to note that the anomalies in the vibrational spectra of the hexafluorides of 4d and 5d elements are not repeated in the series of the hexafluorides of 5f elements.

III. RELATION BETWEEN THE VARIATION OF THE FORCE CONSTANTS OF HEXAFLUORIDES AND THEIR THERMAL STABILITIES AND REACTIVITIES

In certain reviews (for example those of O'Donnel and Stewart 5,6) attempts are made to relate the regularities in the variation of the reactivities of hexafluorides in the first vertical series (CrF₆, MoF₆, WF₆, UF₆) to the valence force constants ($f_{\rm d}$ in Wilson's terminology 17) and average energies of the E-F bonds. Several comments of

a general nature must be made concerning the estimation of the reactivities of hexafluorides with the aid of the force constants. The force constant† $f_{\rm d}=(\partial^2 V/\partial q^2)_{\rm q}\to_0$, where V is the potential energy of the molecule and q the vibrational coordinate, characterises the resistance of the molecule to the deformation of the equilibrium configuration. However, even for diatomic molecules there is no unambiguous relation between the force constant and the dissociation energy¹⁸. The quantity $(\partial^2 V/\partial q^2)_{\rm q}\to_0$ characterises merely the curvature of the potential well in the vicinity of the equilibrium position but does not necessarily indicate its depth ¹⁹.

The multiplicity of models proposed for the description of the force field of a polyatomic molecule is due to the existence of a set of elastic structural elements the number of which is greater than the number of chemical bonds and independent nuclear coordinates. However, regardless of the choice of model, none of the different types of coefficients can provide a simple individual characteristic of a bond in a polyatomic molecule 20 . The extension of one bond involves a whole group of structural elements, since the valence force constant $f_{\rm d}$ depends not only on the interaction between two atoms but also on their interaction with other structural elements of the molecule. Only a set of force constants characterises an individual bond in a polyatomic molecule 21 .

We shall consider to what extent various force constants of hexafluoride molecules can characterise their thermal stability and reactivity. In the present paper we shall employ the terminology adopted by Claassen ²²:

- (1) fd-valence force constant;
- (2) f_{α} —deformation force constant for the bond angle;

Interaction Constants

- (3) f_{dd} —for the interaction of bonds at right angles to one another;
- (4) $f_{\alpha\alpha}$ —for the interaction of adjacent angles in the same plane;
- (5) $f_{d\alpha}$ —for the interaction of an angle with the bonds forming its sides;
- (6) δ—for the interaction of two opposite bonds;
- (7) γ —for the interaction of an angle with a bond perpendicular to its plane;
- (8) k—for the interaction between two angles in perpendicular planes having a bond in common;
- (9) ϵ —for the interaction between two angles in perpendicular planes without a bond in common.

Seven independent force constants may be isolated for hexafluoride molecules 4:

$$(1) A = f_d; (9)$$

(1)
$$A = I_d;$$
 (5) $E = f_a - \gamma;$ (2) $B = \delta;$ (6) $F = 2 (I_{pq} - \gamma)$

(2)
$$B = \delta;$$
 (6) $F = 2(f_{\alpha\alpha} - \gamma);$ (3) $C = 2f_{dd};$ (7) $G = 2(k - \epsilon).$

(4)
$$D=2(f_{da}-\delta);$$

Many investigators 4,22-36 analysed the normal vibrations of octahedral hexafluoride molecules and calculated the force constants. The force constants depend on the choice of model for the force field. In the present study only the force constants quoted in the review of Weinstock and Goodman 4 are considered. They are listed in Table 3.

Figs. 1 and 2 present a graphical interpretation of the relation between the force constants of EF_6 molecules and the charge number of the element E (see p. 157).

Analysis of these relations permits the following conclusions: (1) the valence force constant exceeds approximately by two powers of ten all the remaining force constants; this is a general rule 37; (2) the independent force constants A, B, E, and partly G of hexafluoride molecules increase on going from 4d to 5d elements and sharply decrease on going to 5f elements; (3) the force constants C and F also increase on going from 4d to 5delements; on going to 5f elements, the constant Cincreases and the constant F changes in different ways for each of the vertical series; (4) the constant D decreases on moving downwards along the vertical series; (5) the constants A, B, C, and partly G decrease sharply on moving along the series of 4d, 5d, and 5f elements from left to right, which is consistent with the variation of the thermal stabilities of the hexafluorides; (6) the constants D, E, and F increase along the series of 4a, 5a, and 5felements (from left to right).

Table 3. Force constants of octahedral hexafluoride molecules 4 [cm⁻¹/(0.01 Å)²].

		-		-				
Force constant	SF.	SeF ₆	MoF ₆	TcF _t	RuF,	RhF ₆	TeF.	WF.
A B C D E F G	28.163 —1.125 3.580 5.634 4.346 0.443 0.848	25.663 0.666 1,232 2,160 2.648 0.316 0.676	24.673 1.673 2.597 0.713 0.843 -0.555 -0.415	24,483 0,860 1,888 0,720 1,083 0,183 0,214	23.304 0.331 1.268 0.767 1.199 0.050 0.205	21.770 -0.654 0.986 0.806 1.260 0.221 0.234	26.032 0.773 0.711 0.499 1.090 -0.317 -0.002	26.480 2.246 0.709 0.488 0.970 -0.455 0.434
	ReF.	Os F ₆	IrF.	PtF,	UF.	NpF.	PuF _¢	
A B C D E F	26.353 1.802 2.293 0.480 1.243 -0.006 0.173	26.185 1.190 1.744 0.531 1,400 0.307 0.194	25.181 0.209 1.418 0.543 1.432 0.476 0.213	23.033 -1.033 1.322 0.526 1.446 0.605 0.168	19.412 0.066 3.038 0.210 0.663 0.083 0.100	19.029 -0.314 2.702 0.244 0.832 0.229 0.050	18,451 -0,425 2,314 0,255 0,899 0,255 0,035	

One may conclude that the variation of the majority of force constants along the horizontal and vertical series of hexafluorides is of the same type as that of the valence force constant $f_{\bf d}(A)$. The force constants for sulphur, selenium, and tellurium hexafluorides are quoted only for comparison.

Analysis of the feasible configurations of the xenon hexafluoride molecule 38 showed that it cannot have the structure of a regular octahedron. A model close to the structure of iodine heptafluoride, in which the missing bond is replaced by the electron pair of xenon, is more likely. This is confirmed both by the analysis of the infrared spectrum of xenon hexafluoride 39 and by the study of its structure by electron diffraction 40,41 . Therefore the force constants of XeF₆ cannot be compared with those of other hexafluorides.

[†] For diatomic molecules.

On the hypothesis of a relation between the force constants, thermal stabilities, and reactivities of EF₈ molecules, the latter can be arranged in the following series in terms of their thermal stability:

Vertical series:

$$\begin{split} &SF_6 > TeF_6 > SeF_6; \\ &WF_6 > MoF_6 > UF_6; \\ &ReF_6 > TcF_6 > NpF_6; \\ &OsF_6 > RuF_6 > PuF_6; \\ &IrF_6 > RhF_6. \end{split}$$

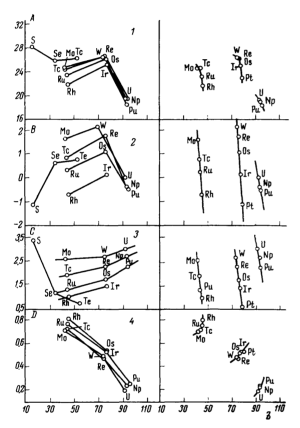


Figure 1. Variation of the force constants A, B, C, and D of EF_6 molecules with the charge number of the central atom of the element E in the vertical and horizontal series of hexafluorides.

Horizontal series:

$$\begin{split} MoF > TcF_6 > RuF_6 > RhF_6; \\ WF_6 > ReF_6 > OsF_6 > IrF_6 > PtF_6; \\ UF_6 > NpF_6 > PuF_6. \end{split}$$

When all the elements are arranged in a single series, the following sequence is obtained:

$$\begin{split} &SF_6>WF_6>ReF_6>OsF_6>TeF_6>SeF_6>lrF_6>MoF_6>TcF_6>\\ &>RuF_6>PtF_6>RhF_6>UF_6>NpF_6>PuF_6. \end{split}$$

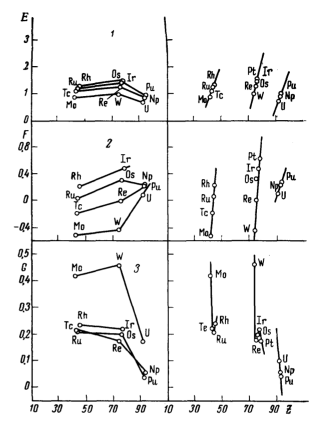


Figure 2. Variation of the force constants E, F, and G of EF_6 molecules with the charge number of the element E in the vertical and horizontal series of hexafluorides.

The last series probably does not fully reflect the actual relation between the thermal stabilities and reactivities of the hexafluorides. Therefore, before dealing with a comparative analysis of the experimental data, it is necessary to analyse the above relations by other methods.

IV. VARIATION OF THE AVERAGE BOND ENERGIES OF HEXAFLUORIDES IN HORIZONTAL AND VERTICAL SERIES

The average bond energy $E_{\rm o}$ of the EF $_{\rm n}$ molecule is defined by the formula

$$E_0 = \frac{D_0^0}{n},$$
 (1)

where D_0^0 is the heat of atomisation of the molecule and n is the number of bonds. The values of D_0^0 are not known for all hexafluorides and the accuracy of the calculation or estimation of some of them is low. For example, one may mention the discrepancy between the data of different workers for the heat of formation of gaseous plutonium hexafluoride: -460 kcal mole $^{-1}$, 42 -473 kcal mole $^{-1}$, 43 -411.2 kcal mole $^{-1}$, 44 and -418 ± 8 kcal mole $^{-1}$ 45. The last value was recommended by Rand 45 as the most accurate.

The bond energies and heats of atomisation of certain hexafluorides are listed in Table 4. For the majority of the known hexafluorides, the heats of formation are known only under standard conditions, i.e. only the values of $\Delta H_{\rm f,298}^{\circ}$ are available, and in order to find D_0^0 $(-\Delta H_0^0)$ a correction must be calculated from the equation

$$\left[\frac{\partial \left(\Delta H_T^{\circ}\right)}{\partial T}\right]_p = \Delta C_p,\tag{2}$$

where ΔC_p is the heat capacity change from 0° to 298°K.

Table 4. Heats of formation, heats of atomisation, and bond lengths and energies in hexafluoride molecules.

Hexafluoride	Heat of formation of hexafluoride from elements under standard conditions ΔH_{2}^{R} , 298, kcal mole ⁻¹	References	Heat of atomisation of hexafluoride under standard conditions -D298, keal mole-1	References	Heat of atomisation of hexafluoride at absolute zero D_0^0 , keal mole ^{-1*}	References	Bond dissociation energy E_0^0 , kcal	References	E-F bond length, Å	References
SF ₆ SeF ₆ TeF ₆ MoF ₆ TcF ₆ RuF ₆ RuF ₆ ReF ₆ VF ₆	-291.7 -266.9 -327.2 -320 -300 -411.5 -278 -130 -510.5 -461 -418 -82.9 kcal at 393° K	46 46 46 47 42 — 48 42 42 42 42 42 45 50		51 51 51 51.52 51.52 	474.6 434.9 483.2 649.2 565 — 725 573 522 399.5 — 735.96 675.6 619.5	55 56 55 55 55 55 55	79.0 72.5 80.6 108.2 94.2 — 120.9 ~95.5 ~86.9 66.55 — 112.8 103.21 32.3	50	$\begin{array}{c} 1.564 \pm 0.01 \\ 1.688 \pm 0.010 \\ 1.688 \pm 0.03 \\ 1.83 \\ 1.8512 \\ 1.8775 \\ 1.8738 \\ 1.833 \\ \dots \\ 1.83 \\ 1.833 \\ 1.833 \\ 1.934 \\ 1.994 \\ 1.990 \\ 1.972 \\ 1.90 \end{array}$	57 57 58 60 56 60 60 59 22 22 22 22 22 38

^{*} Estimated from Eqn. (2).

The accuracy of the determination of ΔH_{298}° is usually not sufficiently high to make this correction, but nevertheless in the present study the correction was introduced.

Analysis of the results in Table 4 shows that the bond energies decrease in the horizontal series from left to right like the valence force constants (A) and the constant B, C, and partly G (see curves 1-3 in Fig. 1 and curve 3 in Fig. 2). In the vertical series the bond energy increases in the series of hexafluorides of 4d, 5d, and 5f elements. In the series of hexafluorides of sulphur, selenium, and tellurium, the E - F bond energy is 79.0, 72.5, and 80.6 kcal mole-1 respectively. The Xe-F bond has the lowest dissociation energy (only 32.3 kcal mole⁻¹).

Thus if it is supposed that the average bond energy characterises the thermal stability and reactivity of hexafluoride molecules (as was done by O'Donnel and Stewart5), then hexafluorides can be arranged in the following series in terms of decreasing thermal stability:

$$\begin{aligned} UF_6 > WF_6 > NpF_6 > MoF_6 > PuF_6 > ReF_6 > TcF_6 > OsF_6 > \\ > TeF_6 > SF_6 > SeF_6 > IrF_6 > XeF_6 \; . \end{aligned}$$

The discrepancy between this series and the experimental stabilities and activities of hexafluorides will be discussed below.

INTERATOMIC DISTANCES AND BOND STRENGTH

The use of interatomic E-F distances to characterise the chemical properties of hexafluorides (Table 4) appears to be of doubtful value. In the series of hexafluorides of uranium, neptunium, and plutonium, the interatomic E-F distances decrease but in the series of hexafluorides of 4d and 5d elements it is difficult to establish a similar sequence. Probably Badger's equation 61, whereby the E-F bond lengths were found (see, for example, Ref. 56 and 60), does not ensure the necessary accuracy owing to the dependence of the calculated force constants on the force field model.

ANALYSIS OF THE THERMODYNAMIC STABILITIES OF HEXAFLUORIDES

A rigorous characteristic of the dissociation of any chemical compound is its thermodynamic stability in this reaction. The thermodynamic stability is characterised by the reciprocal of the equilibrium constants K_e for the corresponding dissociation process:

$$\alpha = \frac{1}{K_e} = \exp\left(\frac{\Delta H_o^0}{RT} - \frac{\Delta \Phi_T^*}{R}\right), \tag{3}$$

where $\Delta\Phi_{\mathbf{T}}^{\boldsymbol{*}}$ is the reduced free energy of the dissociation reaction, R the gas constant, and T the absolute temperature. The values of $K_e(\alpha)$ may be related to the corresponding yields of the dissociation products of hexafluorides. For example, for the reaction

$$(EF_6)_{gas} \to (E)_{gas} + 6(F)_{gas}, \tag{4}$$

$$(EF_6)_{gas} \rightarrow (E)_{gas} + 6(F)_{gas}, \qquad (4)$$

$$\alpha = \frac{(1-x)(1+6x)}{4.6656 \cdot 10^4 x^7 p^6}, \qquad (5)$$

where p is the pressure and x the number of moles of hexafluoride which have reacted.

The values of α depend both on the enthalpy change (ΔH_0^0) and on the change in the reduced free energy $(\Delta\Phi_{\mathbf{T}}^{*})$, which are calculated from energy and geometrical molecular constants.

The thermodynamic functions for gases consisting of polyatomic molecules (including EF, molecules) are calculated in many cases in terms of the rigid rotator-harmonic oscillator approximation. The components of the thermodynamic functions due to translational, rotational, and vibrational motion are calculated. This is because the electronic ground states of polyatomic molecules are as a rule of the singlet type and therefore the partition function with respect to the intramolecular motion of the polyatomic molecule is equal to the sum of the vibrational and rotational partition functions of the electronic ground state 17. The thermodynamic functions obtained are called virtual, since they do not include the components due to nuclear spins and the mixing of isotopes.

The multiplicity of the electronic ground state is taken into account by introducing into the thermodynamic functions the term $R \ln p_{\mathbf{M}}$, where $p_{\mathbf{M}}$ is the statistical weight

It is assumed 17 that the electronic ground state of polyatomic molecules with an even number of electrons is of the singlet type (the statistical weight of the state is unity); electronic states with one free electron have a statistical weight of two. Some workers have calculated the thermodynamic properties of many hexafluorides in

terms of the approximation of the rigid rotator-harmonic oscillator model. The entropies, reduced free energies, heat capacities, and enthalpy changes have been calculated in the range $50-1600^{\circ}$ K at p=1 atm. for the hexafluorides of uranium ⁵⁵, plutonium ^{55,52}, molybdenum, tungsten, iridium, and neptunium ⁵⁶ and in the range $50-2000^{\circ}$ K for the hexafluorides of ruthenium, rhodium ⁸⁰, and technetium ⁵⁶. It should be noted that some of the molecular constants employed by Nagarajan to calculate the thermodynamic functions of the above hexafluorides were subsequently revised by Weinstock and Goodman ⁴ (Table 5).

Table 5. Fundamental vibration frequencies of hexafluoride molecules.

Molecule *	v ₁ (1)	ν ₂ (2)	v _s (3)	v. (3)	v. (3)	ν. (3)
SF ₆	770	640	639	614	522	349
SeF ₆	708	(661)	780	437	(403)	(262)
TeF ₆	701	674	752	325	313	(195)
PoF ₆	[700]	[682]	[741]	[253]	[243]	[144]
MoF ₆	741	643	741	262	(312)	(122)
TcF ₆	(712)	(639)**	748	265	(257)**	[174]
RuF ₆	(675)	(624)**	735	275	(283)**	[186]
RhF ₆	(634)	(592)	724	283	269	(189)
PdF ₆	[590]	[525]	[711]	[280]	[258]	[191]
WF ₆	(771)	(673)	711	258	(315)	(134)
ReF ₆	755	(671)**	715	257	(295)**	(193)
OsF ₆	(733)	(668)**	720	272	(276)**	[205]
IrF ₆	(701)	(646)**	719	276	(258)	(206)
PtF ₆	655	(600)	705	273	(242)	[211]
UF ₈ NpF ₆ PuF ₆ AmF ₈ CrF ₆	667	535	624	(184)	(201)	(140)
	(648)	(528)	624	(198)	(205)	(165)
	(628)	(523)	616	(203)	(211)	(173)
	[609]	[500]	[603]	[205]	[216]	[178]
	(720)	(650)	790	[266]	[309]	[110]

*The degree of degeneracy of the given vibration is indicated in brackets after ν_1 , ν_2 , etc.

** The estimate was made from the variation of the vibration frequencies with the number of free electrons in the hexafluoride molecules 4; these are postulated frequencies in the absence of vibronic interaction. The frequencies obtained from the above relation and virtually identical with the experimental values are indicated in round brackets; the frequencies calculated from the Raman frequencies are indicated in square brackets.

The ν_6 frequencies for molybdenum and tungsten hexafluorides, the ν_2 frequency for technetium hexafluoride, etc. have been revised. The Mo-F bond length in the MoF_6 molecule, assumed by Nagarajan to be 1.56 Å, $^{\rm 83}$ is inconsistent with a later value of 1.83 Å. $^{\rm 59}$ Moreover, the statistical weight of the electronic ground state in the case of its degeneracy was not taken into account in any of the calculations. This applies to the hexafluorides of technetium, ruthenium, rhodium, rhenium, osmium, iridium, and neptunium.

According to Moffit et al. ⁶⁴ the lower electronic states of the hexafluorides of rhenium and osmium are orbitally degenerate and the Jahn-Teller theorem predicts instability of the octahedral configuration. As already mentioned above, such instability does not lead to a static distortion of the symmetrical configuration in the gas phase. This is shown by the similarity of the electronic spectra of the hexafluorides of rhenium and osmium on the one hand and those of iridium and platinum on the other. For the last two hexafluorides, the Jahn-Teller theorem does not predict instability of the octahedral configuration. The number of electronic bands in the spectra of all four

hexafluorides corresponds to the number predicted by the ligand field theory for the configuration with the $O_{\rm h}$ symmetry; the absence of the static distortion of the EF₆ molecules was demonstrated by electron diffraction and by the analysis of the infrared spectra of the "anomalous" hexafluorides. In the case of static distortion, one might expect that the force constants for the longer bonds (or larger angles) would differ from the force constants for the shorter bonds (or smaller angles); moreover, one might observe splitting of the infrared absorption bands. The analogies in the vibrational spectra have been interpreted on the basis of a vibronic interaction 12 ,65. Quantitatively this interaction was investigated by Weinstock and coworkers 4 ,14,15,66.

Table 6. Effect of vibronic interaction on the thermodynamic properties of hexafluorides 4.

Hexafluoride	T, °K	ΔΦ [*] T, cal mole deg l	$\Delta (H_{\rm T}^0 - H_0^0),$ cal mole ⁻¹	ΔST, cal mole-1 deg-1
ReF ₆	200	0.103	18.5	0.196
	298.15	0.136	20.2	0.203
	900	0.175	12.4	0.189
TcF ₆	200	0.087	17.1	0.172
	298 15	0,117	19.1	0.181
	900	0.154	11.1	0.166

The empirical relations between the ground vibration frequencies and the number of unbound electrons 4 enabled Weinstock and Goodman to estimate the frequencies of the hitherto not investigated (CrF $_6$, PoF $_6$) and not even synthesised (AmF $_6$) hexafluorides. The effect of the vibronic interaction on the thermodynamic properties was also estimated by Weinstock and Goodman 4 (Table 6). The corrections obtained ($\Delta\Phi_{T}^*$ and ΔS_{T}^0) are virtually negligible for TcF $_6$ and ReF $_6$ in the range 200–900°K. Such an error in the calculations of the thermodynamic functions is possible as a result of an error of 1–2 cm $^{-1}$ in the estimation of the vibration frequencies.

For the hexafluorides of osmium and ruthenium, the corrections are of the same order of magnitude as those for ReF_6 and TcF_6 , which are listed in Table 6. This implies that the ν_2 and ν_5 frequencies (Table 5) may be employed without additional correction to calculate the thermodynamic properties of rhenium, technetium, ruthenium, and osmium hexafluorides.

Table 7 lists the reduced free energies of certain hexafluorides required for the calculation of their thermodynamic stabilities. They were calculated in terms of the rigid rotator—harmonic oscillator approximation taking into account the corrections for frequencies in the review of Weinstock and Goodman⁴ and for the degeneracy of the electronic ground state.

The reduced free energies of the elements forming hexafluorides were taken from Kulikov's monograph ⁵² and the paper of Gurvitsh and Jungman ⁶⁷. The thermodynamic stability constants of the hexafluorides of certain elements were calculated by Eqn. (3) (Table 8).

Fig. 3 presents the logarithms of the thermodynamic stability constants of the hexafluorides in the vertical series at 298°, 1200°, and 2400°K. Analysis of these relations permits the following conclusions:

1. The thermodynamic stabilities of hexafluorides in relation to atomisation reactions decrease along the horizontal series (4d, 5d, and 5f elements).

<i>T</i> , °K	SF.	SeF₄	TeF₄	MoF _€	TcF₄	WF.	ReF ₆	OsF.	IrF.	UF.	NpF.	PuF₄
298 400 800 1200 1600 2000 2400 2800 3200 3600 4000	56.64 62.01 74.88 85.21 93.38 100.11 105.84 110.82 115.21 119.16 122.72	59,06 64.21 80.07 91.46 100.27 107.43 113.46 118.67 123.24 127.33 131.01	62.42 68.13 85.06 96.88 105.91 113.22 119.34 124.61 129.24 133.36 137.08	64.46 70.52 88.10 100.17 109.32 116.72 122.85 128.19 132.84 136.99 140.73	65.90 71.83 89.30 101.82 110.47 117.84 124.01 129.32 133.37 138.10 141.85	65. 11 71. 16 88. 67 100. 71 109. 86 117. 23 123. 40 128. 71 133. 36 137. 51 141. 25	66.63 72.56 89.89 101.87 110.99 118.34 124.50 129.80 134.45 138.15 141.90	66.93 72.84 90.13 102.10 111.22 118.57 124.73 130.03 134.67 138.75 142.43	66.68 72.65 90.06 102.08 111.23 118.60 124.77 130.07 134.73 138.85 142.60	68.74 75.44 94.26 106.85 116.30 123.86 130.15 135.55 140.28 144.37	69.48 76.08 94.73 107.26 116.67 124.22 130.50 135.89 140.61 144.82 148.58	67.36 73.93 92.54 105.06 114.48 122.02 128.30 133.69 138.41 142.62 146.38

Table 7. Reduced free energies of certain hexafluorides (cal mole⁻¹ deg⁻¹) at p = 1 atm. in the ideal gas state.

Table 8. Thermodynamic stability constants of hexafluorides

<i>т</i> °, қ	SF ₆	SeF₄	TeF.	MoF _€	TcF.	WF.	ReF.	OsF.	IrF.	UF.	NpF ₆	PuF.
1600 2000 2400 2800 3200	1.8·10 ⁻² 1.48·10 ⁻⁸ 3.02·10 ⁻¹³ 7.07·10 ⁻¹⁷	4.47 · 10 ³⁴ 5.13 · 10 ³ 6.45 · 10 ⁻⁵ 1.48 · 10 ⁻¹⁰ 4.67 · 10 ⁻¹⁵ 5.49 · 10 ⁻¹⁸	5.61 · 10 ²⁸ . 5	7,88 · 10 ²⁹ 9,13 · 10 ¹⁷ 4,21 · 10 ⁹ 1,63 · 10 ⁸ 1,91 · 10 ⁻²	1.45.10 ⁸⁰ 5.10 ⁸⁴ 3.17.10 ¹⁹ 1.66.10 ⁹ 9.6.10 ¹ 2.35.10 ⁻⁶ 1.48.10 ⁻⁸ 7.1.10 ⁻¹⁸	4·10 ⁹¹ 1.78·10 ⁹⁵ 2·10 ⁸⁸ 1.6·10 ²⁷ 1.74·10 ¹⁷ 1.34·10 ⁸ 4.46·10 ² 1.94·10 ⁻²	5.62.10 ⁵¹ 7.93.10 ³⁵ 1.26.10 ²⁰ 10 ¹⁰ 3.39.10 ² 1.13.10 ⁻³ 4.22.10 ⁻⁸ 1.29.10 ⁻¹¹	5.62·10 ⁵¹ 1.26·10 ²⁸ 1.15·10 ¹⁴ 4.17·10 ⁴ 8.84·10 ⁻³ 6.17·10 ⁻⁸ 8.5·10 ⁻¹¹ 7.24·10 ⁻¹⁵	1.91.1080 1.59.1012 1.74.101 1.23.10-6 1.2.10-12 1.82.10-15 3.98.10-19 8.9.10-21	4,47.1098 6,31.1059 7.95.1039 1,59.1036 2,82.1014 2,37.109 6,02.103 2,56.10-1	1.59.1083 2.1082 7.93.1033 2.3.1021 3.8.1012 5.94.105 2.82 1.44.10-5	1.7.10 ⁷³ 7.94.10 ⁴² 7.94.10 ²⁷ 1.5.10 ¹⁵ 9.5.10 ⁸ 9.10 ¹ 1.55.10 ⁻³ 6.68.10 ⁻⁸

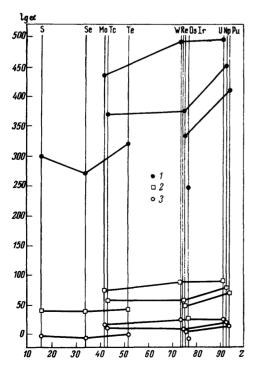


Figure 3. Variation of the logarithms of the thermal stability constants of hexafluorides with the charge number of the central atom; temperature (°K): 1) 298; 2) 1200; 3) 2400.

2. The sequence of hexafluorides in terms of decreasing thermodynamic stability depends little on temperature. In the vertical series the hexafluorides of the actinide elements are the most stable:

$$\label{eq:uf_s} \text{UF}_6\!>\!\text{WF}_6\!>\!\text{MoF}_6; \; \text{NpF}_6\!>\!\text{ReF}_6\!>\!\text{TcF}_6; \; \text{PuF}_6\!>\!\text{OsF}_6.$$

However, the difference between the values of $\lg \alpha$ diminishes as the temperature is raised, owing to an increase of the contribution by the entropy factor to the equation $\Delta G = \Delta H - T \Delta S$ compared with the heat of reaction.

- 3. In the series of non-metal hexafluorides the stability falls from sulphur hexafluoride to selenium hexafluoride, but increases sharply on going to tellurium hexafluoride, so that $\text{TeF}_6 > \text{SF}_6 > \text{SeF}_6$.
- 4. The sequence of hexafluorides in terms of decreasing thermodynamic stability in relation to the atomisation reaction is as follows:

$$\text{UF}_6 > \text{WF}_6 > \text{NpF}_6 > \text{MoF}_6 > \text{PuF}_6 > \text{TcF}_6 > \text{ReF}_6 > \text{OsF}_6 > \text{TeF}_6 > \text{SF}_6 > \text{SeF}_6 > \text{IrF}_6.$$

The positions of the hexafluorides in this series depend to a considerable degree on the accuracy of the determination of ΔH_0^0 (Table 9). Xenon hexafluoride should occupy the last place in this series.

We shall examine the changes in the thermodynamic stability of hexafluorides in partial dissociation reactions in the first vertical and third horizontal series. The decomposition to the tetrafluoride may be employed as a reaction of this kind:

$$(EF_6)_{gas} \rightarrow (EF_4)_{gas} + 2(F)_{gas}. \tag{6}$$

The reduced free energies of the tetrafluorides required for the calculation of the stability constants were evaluated

in terms of the rigid rotator—harmonic oscillator model using an empirical method for the estimation of the equivalent geometrical and energy constants ⁶⁸. The results of the calculation are presented in Table 10. The calculation was made for only six hexafluorides, since only in these cases was it possible to calculate the heats of reaction at the standard temperature (298.16°K) and the absolute zero (Table 9).

Table 9. Enthalpies of dissociation of hexafluorides by reaction (6) under standard conditions (298.16°K) and at the absolute zero (kcal mole⁻¹).

Hexa- fluoride	ΔH ² 298	ΔĦ°	References to literature employed in calculations	Hexa- fluoride	ΔH ² 298	ΔH°	References to literature employed in calculations
UF	184.4	181.4	49,69,70	WF ₆	234	232	42.48
NpF	131	129	42,71	MoF ₆	194	191	42.47
PuF	102.5	101.3	45	SF ₆	—	152,261	17

Table 10. Thermodynamic stability constants of hexafluorides in reaction (6).

<i>т</i> °, К	UF.	NpF.	PuF,	WF.	MoF,	SFa
800	103,	1,32 · 1021	2.24.1013	1.78-1048	7.93·10°7	4.36.1028
1200	1.12.1017	1,12-109	4.8.103	5.38.1028	1.13.1020	2.1.1014
1600	1,2.109	1.23 · 10 ³	1.04.10-1	1.27.1016	1.76.1011	1.48.104
2000	2.76:104	1.34.10-1	1.78-10-4	5.49.109	7.24.105	8.9.10-
2400	1.66 · 101	1.37.10-3	2.29.10-6	3.03.105	$2.21 \cdot 10^{2}$	1.23.10
2800	8.92.10-1	2.81.1(\-5	1.1.10-7	2.95.102	6.88-100	2.2.10
3200	1,74.10-3	1.62.10-6	1.15.10-8	1.61.100	9.10-3	3.81.10-

The logarithms of the thermodynamic stability constants of the hexafluorides decrease from left to right along the series of 5f elements. When the hexafluorides are arranged in a common series in terms of decreasing thermal stability with respect to reaction (6), the following sequence is observed up to 2300°K: WF $_6 > {\rm MoF}_6 > {\rm UF}_6 > {\rm SF}_6 > {\rm NpF}_6 > {\rm PuF}_6$. Above 2300°K, SF $_6$ and NpF $_6$ change places.

VII. ANALYSIS OF EXPERIMENTAL DATA FOR THE THERMAL STABILITIES AND REACTIVITIES OF HEXAFLUORIDES

The chemical reactivities; of non-metal hexafluorides differ markedly from those of metal hexafluorides. With the exception of polonium hexafluoride, non-metal hexafluorides exhibit unusual chemical inertness. In the reviews by Weinstock already mentioned^{2,3}, it is reported that the thermal stabilities in this series decrease and the chemical reactivities increase from sulphur hexafluoride to tellurium hexafluoride, the hydrolysis of tellurium hexafluoride, which is the most reactive of the three hexafluorides, requiring not less than 24 h for its completion at

room temperature³. The information available at present about the thermal stability of sulphur hexafluoride is fairly adequate. The compound is stable in an electric arc 72 , which can be accounted for 3 by its high thermal stability. At $2300-2500^{\circ} \rm K$ sulphur hexafluoride molecules dissociate into atoms 46 . A wide variety of fragments of SF $_6$ molecules, obtained in an electric arc, have been isolated $^{73},^{74}$. The reason for the pronounced insulating properties of sulphur hexafluoride is probably the high electron affinity not only of fluorine atoms and molecules, but also of SF $_6$ molecules and their fragments 75 .

We shall consider the experimental regularities in the variation of the thermal stability and reactivity in the vertical and horizontal series of metal hexafluorides. The first vertical series includes the hexafluorides of chromium (3d), molybdenum (4d), tungsten (5d), and uranium (5f). The molecules of all these hexafluorides do not have free electrons. Their chemical reactivity may be characterised by the reactions involving the fluorination of the lower fluorides of various elements (including the lower fluorides of the elements forming the hexafluorides), the sensitivity to hydrolysis, substitution and addition reactions, etc.

The feasibility of reaction (3) is a direct indication that tungsten hexafluoride is chemically less reactive than molybdenum hexafluoride:

$$(WF_4)_{solid} + 2(MoF_6)_{gas} \rightarrow 2(MoF_5)_{solid} + (WF_6)_{gas}. \tag{7}$$

Chromium hexafluoride is thermally unstable at -100° C. ⁷⁶ Even chromium pentafluoride is more reactive than the hexafluorides of molybdenum, tungsten ¹¹, and uranium ⁶. This was demonstrated by studies of the reactions of the above fluorides with the trifluorides of phosphorus, arsenic, bismuth, etc. The reaction with carbon disulphide is particularly instructive:

$$CS_2 + 8CrF_5 \rightarrow 8CrF_3 + CF_4 + 2SF_6$$
, (8)
 $2CS_2 + 6MoF_6 \rightarrow (CF_3)_2 S_2 + 2S + 6MoF_5$. (9)

Reaction (8) takes place explosively¹¹, reaction (9) is also vigorous¹¹, but tungsten hexafluoride does not react with carbon disulphide¹¹. Uranium hexafluoride fluorinates carbon disulphide, various fluorination products being obtained, depending on temperature.⁶,⁷⁷

On the basis of studies of the behaviour of these four hexafluorides, O'Donnel⁶,⁷,¹¹ believes that the reactivity of the hexafluorides in the first vertical series diminishes in the following sequence: $\text{CrF}_6 > \text{UF}_6 > \text{MoF}_6 > \text{WF}_6$. Thus the hexafluoride of uranium (a 5f element) is more reactive than the hexafluorides of molybdenum (4d) and tungsten (5d).

In the series of hexafluorides of 5f elements (uranium, neptunium, and plutonium), the reactivity increases from UF₆ to PuF_6 . $^{2-4}$, 6 , 71 . Uranium hexafluoride may be obtained by the fluorination of the tetrafluoride with plutonium hexafluoride 88 . Neptunium and plutonium hexafluorides are readily decomposed photochemically, so that it is impossible to record their Raman spectra. Americium hexafluoride, the existence of which may be postulated by analogy with platinum hexafluoride, is probably even more unstable 79 .

The transition elements of the second intercalary decade (4d elements) form four hexafluorides: MoF_6 , TcF_6 , RuF_6 , and RhF_6 . Ruthenium and rhodium hexafluorides oxidise xenon even at room temperature, like platinum and plutonium hexafluoride 50 , in contrast to UF_6 , NpF_6 , IrF_6 , and OsF_6 . Technetium hexafluoride is reduced more readily than rhenium hexafluoride 80 . The hexafluorides of technetium and ruthenium are thermally

[‡] As in the work of Canterford and O'Donnel⁷, it is assumed that the increase of the thermal stability of the molecule is accompanied by a decrease of its chemical reactivity.

and photochemically unstable, so that it is impossible to obtain their Raman spectra. Molybdenum hexafluoride is the most stable in this series, the other hexafluorides being probably thermally unstable. The experimental evidence for the increase of the reactivity of the hexafluorides from left to right in this series is inadequate, but it is postulated that rhodium hexafluoride is thermally less stable than ruthenium hexafluoride⁸¹.

The series of hexafluorides of 5d elements contains WF₆, ReF₆, OsF₆, IrF₆, and PtF₆. Platinum hexafluoride fluorinates the fluorides of neptunium and plutonium to the corresponding hexafluorides 72 . Selenium tetrafluoride reduces platinum hexafluoride to the tetrafluoride, but the hexafluorides of osmium and iridium pass to the pentavalent state under the same conditions 82 . Tungsten hexafluoride does not react with nitric oxide 83 , osmium hexafluoride forms NO 4 OsF₆, 83 and iridium and platinum form (NO)₂IrF₆ 85 and (NO)₂PtF₆ 84 respectively. Canterford and O'Donnel arranged the hexafluorides of 5d elements in the following sequence in terms of decreasing thermal stability: WF₆ > OsF₆ > IrF₆ > PtF₆. In addition 7 , OsF₆ > RuF₆, IrF₆ > RhF₆.

Among the least thermally stable metal hexafluorides of various series, plutonium hexafluoride has a higher thermal stability than platinum hexafluoride; neptunium and uranium hexafluorides are more stable than ruthenium, rhodium, and platinum hexafluorides. The least stable hexafluoride is probably that of chromium§. There is evidence that the thermal stability of rhodium hexafluoride is lower than that of platinum hexafluoride 85 . From this standpoint, the least stable metal hexafluorides can probably be arranged in the following series in terms of decreasing thermal stability: $PuF_6 > PtF_6 > RhF_6 > CrF_6$.

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Analysis of the results presented in this review leads to the conclusion that the thermal stability of a particular hexafluoride, as a characteristic of its reactivity, depends on the type of the dissociation reaction considered. Depending on whether the hexafluoride dissociates completely into atoms or only partially to the penta- or tetrafluoride, a particular sequence of hexafluorides in the vertical and horizontal series in terms of varying thermal stability is obtained.

The experimental thermal stability and reactivity series characterising the incomplete conversion of metal hexafluorides, mainly to the penta- or tetra-fluorides within the limits of individual groups, correspond to changes in the majority of the force constants.

However, if all the hexafluorides are arranged in a single sequence in terms of decreasing valence force constants, the latter is not consistent with the experimental thermal stability series.

The experimental data for non-metal hexafluorides do not agree fully with the variation of the force constants. Probably a higher force constant for the bond is a necessary but not sufficient condition for a higher thermal stability of the hexafluoride. The shifts occurring in the dissociation of the E-F bond are so large, even on partial dissociation, that they no longer depend on the force constants. Moreover, if stepwise dissociation of the E-F bond is postulated, then for reactions (6) one

must consider not only the fd constant of the EF₆ molecule, but also the fd constant of the EF₅ molecule, since on transition from the hexafluoride to the pentafluoride both the geometrical configuration and the average bond energy of the molecule change. For example, the average energy of the U-F bond in the uranium hexafluoride molecule is 122.24 kcal and in the pentafluoride $137 \text{ kcal mole}^{-1}$.

The equilibrium constants for the dissociation of hexafluorides to the corresponding tetrafluorides [reaction (6)] probably constitute a more objective criterion of the stability of the hexafluoride in relation to incomplete dissociation (Table 10). In this case the thermal stability of the hexafluoride decreases in the following sequence:

$$WF_6>MoF_6>UF_6>SF_6>NpF_6>PuF_6>PtF_6>RhF_6>CrF_6$$
 .

From this point of view, the great inertness of sulphur hexafluoride is to be accounted for not by thermodynamics but by kinetic characteristics. The reasons for the position of the hexafluorides SeF_6 , TeF_6 , TcF_6 , OsF_6 , IrF_6 , and RuF_6 in this series are still not quite clear.

Probably the most objective characteristic of the stability of hexafluorides in relation to total dissociation (atomisation) is provided by the thermodynamic stability The estimates of thermal stability from the average energies of the E-F bonds in hexafluoride molecules agree fully, except for rhenium hexafluoride, with the thermodynamic estimates. The valence force constants are altogether unsuitable to characterise the stability of hexafluorides in relation to the atomisation reaction. The discrepancy between the thermal stability and force constant series constitutes indirect evidence that the decomposition of hexafluorides to atoms takes place via stepwise abstraction of fluorine atoms and, instead of a single force constant of the bond, one must take into account all the force constants of the E-F bonds in the molecules of the hexa-, penta-, tetra-, tri-, di-, and mono-fluorides of the elements.

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^{\$} Only the hexafluorides having the octahedral configuration with the O_h symmetry are considered.

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