will be interesting to observe how well the chemistry of the N-H and P-H bonds in difluoramine and difluorophosphine parallel one another.

Acknowledgment.---Partial support of this work by a grant from the Petroleum Research Fund of the American Chemical Society is gratefully acknowledged.

CONTRIBUTIOS FROM THE DEPARTMENT OF CHEMISTRY, UNIVERSITY OR CALIFORSIA, BERKELEY, CALIFORNIA 94720

Pentafluorosulf ur and Trifluorome thy1 Oxydifluoramines. Preparations and Properties'

BY WILLIAM H. HALE, JR., AND STANLEY M. WILLIAMSON

Received April 14, 1965

Tetrafluorohydrazine reacts with pentafluorosulfur hypofluorite and trifluoromethyl hypofluorite to form SOF₄, SF₆, FNO, NF_3 , and SF_6ONF_2 and COF_2 , CF_4 , FNO , NF_3 , and CF_3ONF_2 , respectively. The new difluoramines are colorless, do not react with glass or mercury at room temperature, and have normal boiling points of -10.0 and -60° , respectively, for the pentafluorosulfur and trifluorometkyl oxydifluoramines. The molecular compositions have been proved by physical methods and the reaction schemes for the formation and thermal decomposition of the compounds are given.

Tetrafluorohydrazine is known to exist in equilibrium with NF_2 radicals at room temperature.² Many studies, with and without ultraviolet light, have shown that N_2F_4 is an efficient reagent for the introduction of the NF₂ group into organic and inorganic substrates.³⁻¹¹

Since $SF₅OF¹²$ and $CF₃OF¹³$ have been reported to give small amounts of $F₅SO$ and $F₃CO$ radicals upon irradiation with ultraviolet light, it was thought that N_2F_4 should react with SF₅OF and CF₃OF to give SF₅- ONF_2 and CF_3ONF_2 , respectively. The experiments described below confirm their discovery and characterization.

Experimental

Materials.--Pentafluorosulfur hypofluorite and trifluoromethyl hypofluorite were prepared by the methods of Dudley'l and Kellogg,¹⁵ respectively. The crude SF₅OF was purified by fractional codistillation¹⁶ and pure $SF₅OF$ was identified by its molecular weight and infrared spectrum. The crude $CF₃OF$ was placed over water for 3 days to remove the COF₂. The gaseous mixture was then led through a trap at -95° and into a

- *(3)* K. C. Petry and J. P. Freeman, *J. Am. Chenr.* SOC., **83, 3912** (l9til).
- (6) *C..* B. Colhum and F. **A.** Johnson, *Inorg. Chem.,* **1,** 715 (1962).
- (7) G. H. Cady, D. F. Eggers, and B. Tittle, *Proc. Chem. Soc.*, 65 (1963). *(8)* M. Lustig and G. H. Cady, *Inorg. Chein.,* **2,** *888* (1963).
- (9) E. C. Stump, Jr., C. D. Padgett, and W. S. Brey, Jr., $ibid.,$ 2, 648 (1963).
	- (10) M. Lustig, C. L. Bumgardner, and J. K. Ruff, *ibid.,* **3,** 917 (1964).
	- (11) L. C. Duncan and G. H. Cady, *ibid.,* **3, 1045** (1964).

trap at -183° where a white solid was collected. The infrared spectrum of this material was identical with the literature spectrum of $CF_3OF,$ ¹⁷ although it is possible that some CO_2 was present. A previous report states that $CF₃OF$ is a liquid at - 183'.13 Tetrafluorohydrazine from E. I. du Pont de Nemours and Co. was sufficiently pure to use without additional purification.

Reaction of SF_5OF with N_2F_4 . --The gases for this and all other reactions were handled in a vacuum line system with its stopcocks lubricated by No. 90 Kel-F grease. The appropriate quantities of the two reactant gases were condensed in zones in the cold finger of a 1-1. Pyrex glass vessel and then were allowed to warm to room temperature.

The reaction products were separated by means of fractional codistillation¹⁶ and were identified by their molecular weights and characteristic infrared spectra.

Reaction 1.-Pyrex glass vessel at room temperature with initial pressures of 80 mm. for each reactant. The vessel was allowed to stand 17 hr., 8 of which were in the dark. Products found were NF_3 , NO, SIF_4 , SF_6 , SO_2F_2 , SOF_4 , SF_6ONF_2 , and NO_2 . The yield of SF_9ONF_2 was 40 mole $\%$ with respect to the reactant, SF_6OF , and all of the SF_6OF and N_2F_4 was consumed.

Reaction 2.---Pyrex glass vessel at room temperature with initial pressures of 41 mm. for each reactant. The vessel was allowed to stand for 23 hr., 12 of which were in the dark. All the $SF₅OF$ and $N₂F₄$ was consumed, but no $SF₅ONF₂$ was produced. Products observed were NF₃, NO, SiF₄, SF₆, SO₂F₂, SOF₄, and $NO₂$.

Reaction 3.^{-The} vessel used was a prefluorinated 0.25-in. 0.d. closed-end nickel tube fitted with a Xo. 327 Hoke valve (Teflon seal on stem). The initial pressure of each reactant in the vessel was 4 attn. After 24 hr. at room temperature, no SFaONF2 was observed; *80%* of each reactant was recovered unreacted. The remainder of the gas did not contain $NO₂$ until the gas had contacted the glass vacuum system. This behavior is indicative of the production of FXO as a primary reaction product. Gases present other than SF_6OF , N_2F_4 , and NO_2 were NF_3 , SiF_4 , SF_6 , SO_2F_2 , and SOF_4 .

Reaction 4.--Pyrex glass vessel at room temperature with initial pressures of 140 mm. for each reactant. Ultraviolet irradiation for 21 hr. by a General Electric Type H100-A4/T lamp produced a 60 mole $\%$ yield of SF₅ONF₂ (based on the reactant,

^{(1) (}a) This report is taken from the Ph.D. thesis of W. H. Hale and was presented at the 150th Xational Meeting of the American Chemical Society, Atlantic City, N. J., Sept. 1965. (h) After this paper had been submitted for publication, the authors learned that G. H. Cady and L. C. Duncan had prepared CF₃ONF₂ several years ago, but because of governmental classification they did not report it or continue its characterization.

⁽²⁾ (a) F. **A.** Johnson and *C.* **B.** Colburn, *J.* **Ain.** *Chem. SOC.,* **82,** 2400 (1960) ; (b) L. H. Piette, F. **A.** Johnson, K. **A.** Booman, and C. B. Colburn, *J. Chem. Phys.,* **36,** 1481 (1961).

⁽³⁾ K. C. Petry, *J. Am. Chein. SOC.,* **32,** 2400 (1960).

⁽⁴⁾ J. W. Frazer, *J. Inoi,g. h-tacl. Chein.,* **16,** 6.1 (1960).

⁽¹²⁾ C. I. Merrill and G. H. Cady, *J. Am. Chem.* SOC., **83,** 298 (1961). (13) J. **A.** C. Allison and G. H. Cady, *ibid.,* **81,** 1089 (1959).

⁽¹⁴⁾ F. B. Dudley, *G.* **H.** Cady, and D. **F.** Eggers, Jr., *ibid.,* **78, 1553** (1956).

^(1.5) K. **R.** Kellogg **and** *C,.* **Ti. Cntly.** *ihid., 70,* 8986 (lW8).

⁽¹⁶⁾ G. H. Cady and D. P. Siegwarth, *Anal. Chem.*, 31, 618 (1959).

⁽¹⁷⁾ K. J. T,agemann, E. **A. Jones,** and **P.** J. H. Woltz, *J. Chrm. 7'hy3., 20,* 1768 (1952).

 $SF₅OF$). Some unreacted $N₂F₄$ and $SF₅OF$ were present and minor amounts of NO, SiF_4 , SF_6 , SO_2F_2 , and NO_2 were formed.

Reaction of $CF₃OF$ with $N₂F₄$. --Pyrex glass vessel at room temperature with initial pressures of 300 mm. for each reactant. Ultraviolet irradiation for 24 hr. produced a 40 mole $\%$ yield of $CF₃ONF₂$ (based on the reactant, $CF₃OF$). Other products observed were NF₃, CF₄, and COF₂. A white scaly solid which slowly evolved NO, $NO₂$, and SiF₄ remained in the flask. About *20y0* of each reactant was recovered unreacted.

Thermal Decomposition of the Difluoramines.--A 200-ml. Pyrex glass vessel containing 56.5 mm. pressure of SF_5ONF_2 was immersed in a water bath at 85° and within 1 hr. a brown gas appeared. After the flask had been at *85"* for 14 hr., analysis of the gas mixture showed that all the reactant had decomposed to give SF₆, NO, NO₂, and SiF₄ in molar ratios of $4.0 \pm 0.2:2.5 \pm 0.2:2.5$ $0.2: 1.9 \pm 0.2: 1$, respectively. Under similar conditions at 190° for 16 hr., CF_3ONF_2 decomposed to give CF₄, NO, NO₂, and SiF₄ in molar ratios of $4.1 \pm 0.2:2.0 \pm 0.2:2.1 \pm 0.2:1$, respectively. Although SF₅ONF₂ decomposed slowly at 75°, decomposition of CF_3ONF_2 did not set in until about 140°. A small amount of a white solid that was decomposed by water remained in the flask in each case.

Properties of SF_5ONF_2 and CF_3ONF_2

Molecular Weights.--From vapor density measurements assuming perfect gas behavior, the average weight per molar volume of SF_5ONF_2 and CF_3ONF_2 was 195.9 and 137.8, respectively. The theoretical values are 195.1 and 137.0, respectively. Neither compound reacted with glass, mercury, Kel-F grease, or sodium chloride.

Melting Points.—Both compounds are white solids at -183° and melt to colorless liquids. A melting point of $-139 \pm 1^\circ$ was obtained for SF₅ONF₂ by visual observation of a sample under autogenous pressure as it warmed in an isopentane cold bath. The melting point of CF_3ONF_2 was not determined. It was a liquid at -160° .

Vapor Pressures and Boiling Points.---For SF_6ONF_2 , equilibrium boiling points at a series of fixed pressures were obtained using an apparatus similar to that of Kellogg and Cady.16 Some representative data are *(T,* OK., *P,* mm.): 210.2, 48.5; 219.7, 89.0; 225.4, 123.8; 232.9, 184.5; 238.4, 247.7; 242.8, 312.0; 246.0, 357.7; 250.4, 439.2; 255.8, 558.2; 261.2, 713.6; 264.2, 794.1; 264.9, 822.4. These data can be represented by the equation

$$
\log P \text{ (mm.)} = 7.6009 - \frac{1.242 \times 10^3}{T}
$$

The maximum deviation of experimentally observed vapor pressures from calculated values is 2.2% (average deviation 0.9%) in the range 48 to 822 mm. pressure. The normal boiling point of SF_5ONF_2 as calculated from this equation is -10.0° . Using the Clausius-Clapeyron equation and assuming ideal gas behavior, the molar heat of vaporization is 5700 cal., corresponding to a Trouton constant of 21.6 cal. mole⁻¹ $deg. -1$.

Because of the high volatility of $CF₃ONF₂$, its vapor pressure curve was determined only approximately by measuring equilibrium vapor pressures at various temperatures as shown by the following data $(T, \ ^{\circ}K, P,)$ mm.): 139.7, 2.3; 162.5, 20.2; 176.5, 77.4; 194.7,

252.0. These data give a normal boiling point of $-60 \pm 1^{\circ}$.

Mass Spectra.-The mass spectra were obtained from a Consolidated Electrodynamics Corp. Model 103-C mass spectrometer with an ionization potential of 70 volts. As is generally the case in highly fluorinated molecules, the parent ion was not observed for either compound.

The mass spectrum of $SF₅ONF₂$ beginning at 19 mass units consists of the following *(M/Q,* species, relative intensity): 19, F+, 6.3; 30, NO+, 100.0; 33, NF+, 0.7; 48, SO+, 3.4; 49, FNO+, 1.2; 51, SF+, 3.8; 52, NF2+, 3.3; 67, FSO+, 12.4; 68, F_2NO^+ , 0.2; 70, SF_2^+ , 4.2; 86, F_2SO^+ , 7.9; 89, SF_3^+ , 11.4; 105, F_3SO^+ , 8.4; 108, SF_4^+ , 2.6; 127, $SF₅$ ⁺, 28.6. Peaks that corresponded to fragments from $SF₅ONF₂ containing S³³ and S³⁴ were observed$ in the proper abundances along with the corresponding **Ss2** peaks.

The mass spectrum of $CF₃ONF₂$ beginning at 19 mass units consists of the following *(M/Q,* species, relative intensity: 19, F⁺, 2.0; 23.5, FCO²⁺, 0.4; 28, $CO⁺$, 23.5; 30, NO⁺, 61.3; 31, CF⁺, 5.4; 33, NF⁺, 5.1; 47, FCO⁺, 33.8; 49, FNO⁺, 0.9; 50, CF₂⁺, 0.8; 52, NF_2^+ , 85.8; 66, F_2CO^+ , 16.0; 69, F_3C^+ , 100.0; 85, F_3CO^+ , 14.8; 118, F_3COMF^+ , 0.1.

Although intense peaks for F_8CO^+ and F_2CO^+ are observed in $CF₃ONF₂$, the corresponding $F₅SO⁺$ and $F₄SO⁺$ ions are not formed. This is not unexpected since neither F_5 SOOS F_5^{12} nor F_5 SOC F_3^{18} forms the $F₅SO⁺$ or $F₄SO⁺$ ions.

Nuclear Magnetic Resonance Spectra.—The n.m.r. spectra were obtained using a Varian Associates HR-60 n.m.r. spectrometer with a 56.4-Mc. radiofrequency probe for use in F19 resonance work. A reproduction of the spectrum of SF_5ONF_2 is given in Figure 1, in which the NF_2 and SF_6 resonances are shown on different ordinate and abscissa scales. The zero value in Figure 1 corresponds to the center of the SF_4 resonance. The sulfur-fluorine nuclear resonance is typical of the $SF₅$ group in that there is considerable second-order spin-spin interaction between the apical and the four basal fluorine atoms. Preliminary determination of line positions and chemical shift between the two types of sulfur fluorine atoms was done using the method of Merrill, Williamson, Cady, and Eggers.¹⁹ The chemical shift obtained in this manner was used in combination with *JAB* values of 150.0, 152.0, and 154.0 C.P.S. as suggested by Merrill, et al , 19 for calculations of the **AB4** spectrum on an I.B.M. 7094 computer using the method of McConnell, McLean, and Reilly.²⁰ Excellent agreement between the experimental and calculated spectra was obtained for the selected value of J_{AB} = 154.0 c.p.s. and the experimentally determined δ = 4.45 p.p.m. The apical fluorine nuclear resonance is upfield from that of the basal fluorines as is usually the

⁽¹⁸⁾ L. C. Duncan and G. H. Cady, *Inovg. Chem.,* **3,850 (1964).**

⁽¹⁹⁾ C. **I.** Merrill, *S.* M. Williamson, G. H. Cady, and D. F. Eggers, Jr.. *ibid.*, **1,** 215 (1962).

⁽²⁰⁾ H. M. McConnell, A. D. McLean, and C. A. Reilly, *J. Chem. Phys.,* **23, 1152 (1955).**

case for SF_6O groups.¹⁹ Peak areas of the three different types of fluorine atoms were close to the ratio of $1:4:2$ as would be expected for FSF_4ONF_2 . Experimental peak areas were $1:4.2 \pm 0.2:2.1 \pm 0.2$.

The n.m.r. spectrum of CF_3ONF_2 shows two resonances with an upfield to downfield peak area ratio of 1.6 ± 0.2 . The upfield resonance, due to the CF₃ fluorines, is split into a sharp 1:2:1 triplet by the NF_2 fluorine atoms. The downfield resonance is a very broad triplet caused by splitting of the NF_2 fluorine atom resonance by the N^{14} quadrupole moment.

A summary of the experimental n.m.r. data for $SF_{5}^ ONF₂$ and $CF₃ONF₂$ is given in Table I.

Infrared Spectra.-The infrared spectra reproduced in Figures 2 and **3** were obtained on a Perkin-Elmer Model 421 grating spectrophotometer using a Pyrex

TABLE I1

			TABLE II				
		INFRARED DATA OF SF_5ONF_2 and CF_3ONF_2					
		-SF&ONF2-					
	Rel.		Rel.		Rel.		Rel.
$Cm. -1$	int.	$\rm Cm.$ ⁻¹	int.	$\rm Cm.$ $^{-1}$	int.	$\rm Cm.$ $^{-1}$	int.
1966	VW	1130	vw	2598	VW	1296	VS.
1908	vw	1091	VW	2525	vw	1243	vs
1843	v w	1029	s	2420	vw	1215	vs
1706	vw	928	vvs	2233	vw	1066	m
1639	W	858	vs	2150	m	1027	s
1620	w	844	vs	1921	vw	933	s
1575	w	786	$_{\rm VS}$	1797	vw	892	s
1304	VW	706	s	1745	vw	865	vs
1226	w	\sim 600	vs.	1560	v w	712	S
1154	vw			1494	vw	628	m
				1411	VW		

Figure 2.-Infrared spectrum of $SF₅ONF₂$.

Figure 3.—Infrared spectrum of $CF₃ONF₂$.

cell with 9-cm. path length and NaCl windows. Polystyrene calibration indicates that the reported values are accurate within 3 cm^{-1} . The data obtained are summarized in Table 11.

The assignments for the intense bands along with the data on which these assignments are based are given in Table 111.

The frequencies of the asymmetric in-plane SF_{4-} square stretch and the asymmetric N-F stretch lie very close together. For SF_6ONF_2 , we feel that these two bands overlap and could explain the unusual intensity of the band at 928 cm ⁻¹. Also lying close together are the N-0 and S-F stretching frequencies. For SF₅ONF₂ both bands are observed (at 858 and 844 $cm. -1$), but their assignment is uncertain. The symmetric out-of-plane SF_4 -square deformation around 600 cm.-l was not determined exactly because of the NaCl cutoff at this frequency.

The PQR branching observed in the bands at 712 and 706 cm.⁻¹ affords additional evidence for assign-

TABLE I11 INFRARED FREQUENCY ASSIGNMENTS FOR CF_3ONF_2 and SF_6ONF_2 $(\nu \text{ IN CM.}^{-1})$

$Com-$									
pound	ν A	ν B	νC	$\nu_{\rm F}$	ν _G	νĦ	νŢ	νŢ	
CF ₃ ONF ₂	1296	1243	1215	1027	933	712	892	865	
CF ₃ OF ^a	1282	1259	1220				880		
$CF8OSF8b$	1275	1238	1198						
NF ₃				1032	905	647			
$SfsNF_{2}^{d}$				1029	946	679			
$NF2OSO2Fe$				1032	913				
CH_3ONH_2'								858	
Com-									
pound	νD.	νE	ν F	иG	vн	νJ	vк	νL	
SF ₅ ONF ₂	\sim 928	~100	1029	\sim 928	706	858?	844?	786	
$SF_6OCF_3^b$							854		
NFs^c			1032	905.	647				
$S F_5 N F_2^d$	912	605	1029	946	679				
$\mathrm{NF_{2}OSO_{2}F^{\sigma}}$			1032	913				778	
SF ₆	940	615							
SFeCl ^h	908	599							
${\rm SF}_{{\rm s}}{\rm OF}^i$	935	637							
FSO ₂ OF ^j							852	789	
$\mathrm{SO}_2\mathrm{F}_2{}^k$							852		
${\rm SF}_b {\rm OOSFs}^l$							848		
$CH3OSO2Fj$							857	787	
$\nu_A \equiv CFs$ asym. str.		$vG \equiv N-F$ asym, str.							
$\nu_B \equiv C F_3$ asym. str.		$\nu_{\rm H} \equiv N_{\rm F2} X$ sym. def.							
$\nu c \equiv C F_3$ sym. str.		$v_I \equiv C-O$ str.							
ν n \equiv asym. in-plane SF4-square str.		$v_I \equiv N-O str.$							
$\nu_{\rm E} \equiv$ sym, out-of-plane SF4-square def.		ν K \equiv S-F str.							
$v_L \equiv S$ -0 str. $\nu_F \equiv N-F$ sym. str.									

*^a*See ref. 17. * See ref. 18. M. K. Wilson and *S.* R. Polo, *J. Chem. Phys.,* 19, 534 (1951). *d* See ref. **7. e** See ref. 8. M. Davies and N. **A.** Spiers, *J. Chem. Soc.,* 3971 (1959). **0** R. T. Lagemann and E. **A.** Jones, *J. Chem. Phys.,* 19, 534 (1951). * L. H. Cross, H. L. Roberts, P. Goggin, and L. **A.** Woodward, *Trans. Faraday Soc., 56,* 945 (1960). *i* See ref. **14.** *i* F. B. Dudley, G. H. Cady, and D. F. Eggers, Jr.. *J. Am. Chem. Soc.,* **78,** 290 (1956). *k* W. D. Perkins and M. K. Wilson, *J. Chem. Phys.*, 20, 1791 (1952). ^{*l*} See ref. 12.

ment of these bands to the NFzX symmetric deformation since this would be a parallel vibration of a near symmetric top.

Discussion

In addition to the formation of $SF₅ONF₂$ and $CF₃$ - ONF_2 , the respective hypofluorites act both as fluorinating and oxidizing agents to N_2F_4 . Fluorination reactions of SF_5 OF with $CO,^{21}$ SOF₂,¹² and SO_2^{22} produce SOF_4 and COF_2 , SOF_4 , and SO_2F_2 , respectively. The corresponding reactions of CF30F with *SOz,* **23** sulfur, 24 and various organic compounds¹³ produce $COF₂$ and SOzFz, SF4, and fluorinated hydrocarbons, respectively.

The fluorination reactions that are occurring in our work are SF₈OF + N₂F₄ \longrightarrow SOF₄ + 2NF₈ (1)

$$
SF8OF + N2F4 \longrightarrow SOF4 + 2NF8
$$
 (1)

$$
CF8OF + N2F4 \longrightarrow SOF4 + 2NF3
$$

\n
$$
CF8OF + N2F4 \longrightarrow COF2 + 2NF3
$$
 (2)

Pentafluorosulfur hypofluorite can oxidize *SO2* to SO_8^{22} and CCl₄ to COF₂,²¹ whereas CF₃OF oxidizes SO_2 to a mixture of several compounds such as CF_8OSO_2F , $(CF_8)_2SO_4$, etc.²³

 Beach^{25} has reported the oxidation of N₂F₄ by air.
 $\text{N}_2\text{F}_4 + \frac{1}{202} \longrightarrow \text{FNO} + \text{NF}_3$ (3)

$$
F_2F_4 + \frac{1}{2}O_2 \longrightarrow \text{FNO} + \text{NF}_3 \tag{3}
$$

(21) S. M. Williamson and G. H. Cady, *Inorg. Chem.*,
$$
1, 673
$$
 (1962).

(22) G. Pass and H. L. Roberts, *ibid.,* **8,** 1016 (1963).

(24) R. L. Porter and G. H. Cady, *ibid.,* **79,** 5625 (1957).

In a moist glass system, Beach noticed SiF₄ and $NO₂$ and no FNO. The products from our reactions suggest that the hypofluorites oxidize N_2F_4 in the following manner.

$$
SFsOF + N2F4 \longrightarrow FNO + NF3 + SF6
$$
 (4)
CF_sOF + N₂F₄ \longrightarrow FNO + NF₃ + CF₄ (5)

$$
CF3OF + N2F4 \longrightarrow FNO + NF3 + CF4
$$
 (5)

Reactions 4 and 5 are analogous to reaction **3.** Ruff, Menzel, and Neumann²⁶ have studied the reaction of FNO and glass.

$$
4\text{FNO} + \text{SiO}_2 \longrightarrow \text{SiF}_4 + 2\text{NO} + 2\text{NO}_2 \tag{6}
$$

Therefore, all of the by-products found in our reactions are accounted for by the appropriate reactions above, except for the traces of SO_2F_2 which most likely are produced by some decomposition of the $SOF₄$ or transient F₅SOOSF₅.¹² 2SOF₄ \rightarrow SO₂F₂ + SF₆ (7)

$$
2\text{SOF}_4 \longrightarrow \text{SO}_2\text{F}_2 + \text{SF}_6 \tag{7}
$$

In these reactions, FNO is absorbed by $SiO₂$ to form a white, scaly compound which slowly evolves NO, $NO₂$, and $SiF₄$ *in vacuo*. Evolution of these gases occurs in all-glass systems as well as systems in contact with Kel-F stopcock grease. It is felt that the $NO₂$ observed must come from $FNO + glass$, because when $SF₅OF$ and $N₂F₄$ are allowed to react in a nickel vessel, no brown gas was present until the product gases had been exposed to the glass vacuum system for a minute or so.

As an additional check on the role of FNO in the reaction, SF_6OF and NO gave only SOF₄, SiF₄, NO, and $NO₂$ in a Pyrex glass reaction vessel, but from a nickel vessel, rapid infrared analysis of the product gases showed FNO and SOF4 to be present, probably from the following fluorination reaction.
 $SF_6OF + 2NO \longrightarrow SOF_4 + 2FNO$ (8)

$$
SF5OF + 2NO \longrightarrow SOF4 + 2FNO
$$
 (8)

For maximum yield of the addition compounds, N_2F_4 hypofluorite reactions should be run at high pressures in the presence of ultraviolet light. The extreme pressure dependence of the N_2F_4 -SF₅OF reaction is shown by the 40% yield of SF₅ONF₂ starting with 80 mm. of each of the reactants and the total lack of S_{F_6} ONF₂ formation starting with 41 mm. of each reactant. However, light is necessary since reactions in the dark with 4 atm. each of N_2F_4 and SF₅OF produced no SF₅- $ONF₂$.

The relatively low rates of the photolytic addition reactions could probably be increased by using quartz apparatus since Pyrex glass is a poor transmitter of the higher-energy mercury lamp radiation. The relative yields of the addition compounds might be increased since the side reactions are probably not photolytic.

The thermal decomposition products and relative amounts are consistent with the reactions

$$
F_{\delta}S\text{ONF}_{2} \xrightarrow{\Delta} F_{\delta}S_{\cdot} + \cdot\text{ONF}_{2} \longrightarrow SF_{\theta} + \text{FNO} \qquad (9)
$$

$$
F_3CONF_2 \xrightarrow{\Delta} F_3C \cdot + \cdot ONF_2 \longrightarrow CF_4 + FNO \quad (10)
$$

(26) 0. Ruff, **W.** Menzel, and W. Neumann, *2. anovg. allgem. Chem.,* **108,** 293 (1932).

⁽²³⁾ W. **E'.** van Meter and G. H. Cady, *J. Am. Chem. Soc.,* **81,** 6005 (1960).

⁽²⁵⁾ 1,. K. Beach, *J. Inovg. Nucl.* Chem., **26,** 2033 (1964).

followed by eq. 6. Approximate calculations of ΔH show that eq. 9 and 10 are the favored reactions despite the weakness of the O-N bond (\sim 35 kcal.)²⁷ relafive to the *S*-O $({\sim}65$ kcal.)²⁷ and *C*-O $({\sim}81$ kcal.) bonds. Rupture of the O-N bond would lead to $SOF₄$ (or COF_2) and NF₃. None of the latter compounds was obtained. *B* higher temperature is required for

(27) G. S. Paulett and M. Lustig, *J. Am. Chem. Soc.*, 87, 1020 (1965).

the decomposition of $CF₃ONF₂$ than for $SF₅ONF₂$ because of the greater strength of the C-0 bond relative to the *S-0* bond.

Acknowledgments.—We wish to thank Mr. James A. Ferretti for his assistance with the calculated and experimental n.m.r. spectra. W. H. H. appreciates the financial assistance given by the Standard Oil Company of California.

> CONTRIBUTION FROM THE DEPARTMENT OF CHEMISTRY. IOWA STATE UNIVERSITY, AMES, IOWA

Molecular Structure and Bonding of $N_2(CF_3)_4$. **An Electron Diffraction Study1,'**

BY L. S. BARTELL AND H. K. HIGGINBOTHAM

Received March 15, 1965

The molecule $(CF_3)_2NN(CF_3)_2$ represents an interesting example in which, paradoxically, a bond appears to be strengthened and shortened by virtue of strong nonbonded repulsions across it. The substance has been reported to be exceptionally stable to thermal decomposition, and its central N-N bond was observed in this study to be perhaps 0.05 Å. shorter than its counterpart in N_2H_4 . The crowding of trifluoromethyl groups is found to flatten the two $>N-$ pyramids almost to planarity and to set the dihedral angle between opposite $-N(CF_3)_2$ groups at a value of about 90° . A molecular orbital description suggests that the short N-N bond may be understood in terms of the enhancement of π bonding ensuing from the nearly D_{2d} symmetry imposed by steric forces. Structural parameters and amplitudes of vibration of the free (gaseous) molecules were determined by electron diffraction. All bond lengths except for N-N were found to be normal. The large deformations from normal bond angles lead to a rather efficient solution of the molecule's steric problems. Sonbonded repulsions are not borne by a few highly compressed atoms but are distributed among two dozen atom pairs in normal van der Waals contact.

Introduction

The compound tetrakis(trifluoromethy1)hydrazine poses several interesting questions. It has been reported by Young and Dresdner³ to be enormously more stable to the thermal rupture of its N-N bond than a number of other hydrazine derivatives. Its stability is even more remarkable when it is considered that it is among the most crowded of the derivatives studied and must suffer substantial repulsions between its bulky CF_3 groups. If the molecule were assumed to have normal bond lengths and to have bond angles similar to those of hydrazine itself, the implied nonbonded $F \cdots F$ distances would be almost 1 Å. shorter than any nongeminal $F \cdots F$ distances ever observed. Since such short distances are highly improbable it seemed worthwhile to investigate the deformations in structure introduced by steric stress. It was hoped that a structural analysis might also shed light on the surprising stability of the substance.

Experimental

A sample of $N_2(CF_3)_4$ was donated by Professor John A. Young. Gas phase chromatography indicated that the compound was approximately 99% pure. Diffraction patterns were taken with the Iowa State University sector electron diffraction apparatus? using a sample pressure of **15** torr. Absorbancies of diffraction patterns were converted to intensities in a manner described elsewhere.⁴⁻⁶

Structure Analysis

Leveled experimental intensities $I_0(q)$ were converted to reduced molecular intensities by the procedure outlined in ref. 6. The leveled intensities and associated background functions are shown in Figure 1. Experimental radial distribution functions, incorporating a modification function $exp(-0.00166s^2)$, were calculated using theoretical data for the range $q = 0$ to $q = 16$ and experimental data for $q = 17$ to $q = 120$.

The experimental distribution curves bore scant resemblance to a variety of initial synthetic curves calculated for plausible molecular parameters. Therefore it was decided to make some rather severe simplifying assumptions about the structure and to optimize. parameters before relaxing the assumptions.

It is reasonable to expect the present molecule to have C₂ symmetry and, accordingly, 25 independent structural parameters. If all C-F bonds are assumed to have the same length, all C-N bond lengths are taken

⁽¹⁾ This research was supported **by** a grant from the **Natiuval** Science Foundation.

⁽²⁾ Rased in part on a dissertation by H. K. Higginbotham to the Graduate School, Iowa State University, in partial fulfillment of the requirements for the degree of Doctor of Philosophy, 1964.

⁽³⁾ J. A. Young and R. D. Dresdner, *J. Org. Chem.*, **28**, 833 (1963).

⁽⁴⁾ L. S. Bartell, K. Kuchitsu, and R. J. deKeui, *J. Chem. Phys.,* **36,** 1211 (1961).

^{(&}gt;, I,. *S.* Bartell and I,. 0. Rrockway, *J. Appl. Phys.,* **24,** fii6 **(1953).**

⁽⁶⁾ R. A. Bonham and L. S. Bartell, *J. Chem. Phys.*, **31,** 702 (1959).