THE FO RADICAL AND HIGH PRESSURE REACTIONS OF N\textsubscript{2}F\textsubscript{2}

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THE FO RADICAL

AND

PRESSURE REACTIONS OF N₂F₂

Eleventh Quarter Report

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FOREWORD

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This is a progress report and the results presented should be regarded as tentative.

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ABSTRACT

Efforts to generate gaseous OF centered upon flash photolysis of CF₃COOF at low temperature. Two transient phenomena were found: at 77°K violet phosphorescence from the solid and at higher temperature two diffuse absorption bands from the gas. Because the phosphorescence is observed from the solid of several fluorocarbonyl compounds, the phosphor is likely an impurity excited by energy transfer from the host lattice. The transient absorption spectrum is not produced from the other carbonyl compounds nor from products of CF₃COOF. Therefore, the transient is related to CF₃COOF photolysis. Attempts to identify the transient are in progress.

Mass spectrometric search for OF from OF₂ and other hypofluorites was impeded because of poor mass resolution by the MS-10 mass spectrometer. This condition was corrected and decomposition studies are in progress.

For electrolytic generation of OF₂ from wet HF, ohmic overvoltage at the nickel anode was determined by both potential decay and superimposed square wave. The ohmic overvoltage at normal electrolysis conditions was surprisingly low (0.25 to 0.40 v. at 4.8 v. anode). Also, changes in product yield did not give rise to changes in the ohmic overvoltage. The origin of product yield changes at the anode will be sought by other electrochemical methods.

Work continued on the structure of N₂F₂ complexes. Solutions of N₂F₂ in SbF₅, blue when fresh but rapidly fading to yellow, were examined by electron paramagnetic resonance to determine if the color arises from free radicals. An EPR spectrum of the yellow solution could be resolved.
into two separate spectra. The sixteen-line spectrum is consistent with the SbF$_5^-$ radical while the five-line spectrum arises from an N$_2$ containing radical of as yet unknown composition.
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I. INTRODUCTION

This study involves two areas of inorganic fluorine chemistry—the OF radical and the chemistry of difluorodiazine (N₂F₂). Both substances are potential intermediates for synthesis of new fluorine compounds.

Program objectives are:

1. Study the existence of the OF radical produced by photochemical and electrochemical techniques.
2. Investigate the behavior of N₂F₂, both alone and in the presence of other inorganic NF reagents, with and without catalysts, at 10⁴ atm and higher.

The work carried out during this quarter is presented in three parts: fluid-phase formation and detection of OF, electrolysis of wet hydrogen fluoride, and high-pressure reactions of N₂F₂.
II. THE OF RADICAL: FORMATION AND DETECTION IN FLUID PHASES

Introduction

Although OF has now been identified by elegant infrared methods in low-temperature matrix experiments, the radical has so far proven too unstable or too reactive to be detected spectroscopically in the gas phase. As part of our program on flash photolysis of hypofluorites\textsuperscript{2,3,4}, we have now examined trifluoroacetyl hypofluorite, CF\textsubscript{3}COF, both gaseous and in solution at low temperature. Thermolytic studies\textsuperscript{5} have indicated that the CF\textsubscript{3}O–F is the weakest bond to thermal dissociation, but flash photoexcitation can perhaps result in CF\textsubscript{3}C=OF bond cleavage. A high photolytic rate of formation coupled with reduction of unimolecular decay rates by low-temperature operation should favor accumulation of OF for optical spectroscopic detection. This report describes photolytic intermediates and products found in our survey of CF\textsubscript{3}COF photolysis.

Mass spectrometric detection is more sensitive than optical spectroscopy, and we are preparing to use this method to seek OF from flow thermolyses and photolyses.\textsuperscript{2} However, instrument difficulties have delayed its use to date.

Preparation of CF\textsubscript{3}COF

The hypofluorite was prepared by fluorination of CF\textsubscript{3}COOH\textsuperscript{6,7}; fresh batches were made for each photolytic experiment.

\[
\text{CF}_3\text{COOH} + \text{F}_2 \xrightarrow{\text{HeO}} \text{CF}_3\text{COF} + \text{HF}
\]
The product, purified by fractionation and stripping, decomposed (spark or thermal) to give roughly a twofold pressure rise, corresponding to \( CO_2 + CF_4 \) formation. Infrared analysis showed that the final product contained small amounts of \( CF_3COOH, SiF_4, \) and \( CF_2O \), in addition to \( CF_3COOF \) (1860 cm\(^{-1}\), 1070 cm\(^{-1}\)) and the \( CO_2 + CF_4 \) decomposition products. Infrared frequencies for the hypofluorite were assigned because they disappeared as the hypofluorite decomposed, and because the hypsochromic shift in stretching frequencies is about that predicted for carboxylic double and single bonds when H is replaced by F. Stewart and Cady have reported\(^5\) thermal decomposition of \( CF_3COOF \) at room temperature by a chain reaction second order in hypofluorite. We found that the hypofluorite, at an initial pressure of about 10 torr, disappeared with an \( \tau_0 \)-sec. half-life, about three times that estimated from the published specific rate. The appearance of \( CF_2O \) as a primary decomposition product suggests oxygen contamination in our reaction vessel and infrared cell--both inhibition of the decomposition and \( CF_2O \) formation due to oxygen addition were noted by Stewart and Cady. However, our photolysis experiments were carried out in vacuum-tight cells and handling equipment and therefore could not have been contaminated with oxygen.

**Flash Photolysis of CF\(_3\)COF**

Because low-temperature solutions afford a relatively easy experimental approach to low-temperature flash photolysis, we attempted to prepare solutions of \( CF_3COF \) in liquid \( NF_3 \) (b.p. 149\(^\circ\)K). As with other hypofluorites (excepting \( FOF \) and \( FOOF \)\(^3,4\)), the compound was too insoluble to allow examination by absorption spectroscopy.
Low-temperature gas phase flash photolyses were carried out in vacuum-jacketed gas cells containing an internal coil which could be cooled by liquid nitrogen or other coolants. For survey experiments, we used this experimental pattern:

A) A known amount of CF₃COOF, purified and checked by infrared on an aliquot, was condensed (77°K) as a white solid on the cooling coil of the gas cell. An absorption spectroflash of this cell taken during a photolytic flash was used as a reference for further absorption spectra since no gas was present in the cell (p < 0.1 torr).

B) The coolant was flushed from the coil and the contents were allowed to warm. Coil temperature was followed by a thermocouple—the internal gas temperature was always slightly less than that of the coil. The solid melted and vaporized as it warmed. At intervals, the cell contents were photoflashed; flash absorption spectra were taken during the photoflash to reveal any transient absorptions accompanying photolysis, and after the photoflash, to give the absorption spectra of photoproducts. The latter spectrogram includes thermal products, but we expect thermal reactions to be slow below 250°K.

C) After the cell had warmed completely (1-2 hours), ultimate products were determined by infrared.

From flash photolysis of CF₃COF we found two transient phenomena: first, a bright violet phosphorescence in the solid on the
cooling coil; and second, at higher temperatures, two diffuse absorption bands of a transient in the gas phase.

The violet phosphorescence (about 3-sec. life time) was present only in solid CF$_3$COOF and disappeared at the melting point (about 160°K). Following total volatilization of the sample, the phosphorescence could be obtained again if the sample or its decomposition products were recondensed and photoflashed. Other solid fluoro-carbonyl compounds when photoflashed gave qualitatively the same phosphorescence. Phosphorescence intensity increased in the order CF$_2$O $<$ CF$_3$COOH $<$ CF$_3$COOF $<$ CF$_3$COCF$_3$. Because the absorption edges for these compounds move to the visible in the same order, the fraction of photo-excitation from the photoflash ("white" light from 2000-7000 Å tailing off below 2000 Å) increases in this same pattern. Since the same phosphorescence is produced from these sources, the phosphor most likely is an impurity which is excited by energy transfer from the host lattice. The phosphorescence does not seem to be associated with primary photolysis of the hypofluorite.

The transient absorption spectrum, on the other hand, does appear to be associated with CF$_3$COOF photolysis. Flash photolysis with the cell cooled below 200°K shows no short-lived absorption. However, the partial pressure of CF$_3$COOF (b.p. 251°K) is low in this region, so little photolytic light is absorbed. (We have not yet obtained a good absorption spectrum of CF$_3$COOF; however, its strong bands, like those of CF$_3$COCH, appear to lie in the far ultraviolet with only tailing into
the quartz ultraviolet.) Between 200°K and 250°K, the photoflash is attended by the clear appearance of two band maxima at 258 μ and 255 μ. Further bands may appear at 248 μ and in the 260-270 μ interval, but these regions are overlaid by other absorptions and are quite tentative. The new bands are not generated from any of the other carbonyl derivatives and cannot be produced from products present after CF₃COOF has been completely decomposed. Therefore, the transient seems clearly related to CF₃COOF photolysis; however, assignment has not been made. Carriers with known spectra in this region, as CF₂, need to be ruled out. Better defined spectra, kinetic spectroscopic measurements of the transient lifetime, and further correlation experiments with potential transient sources are planned to aid assignment.

**Absorption Spectra of Flash Photolysis Products from CF₃COF**

At low temperature (77°K), the gas cell was transparent in the ultraviolet. As the cell warmed, flash spectra following flash photolyses of the cell contents showed new diffuse absorption bands due to decomposition products from CF₃COOF. The results are summarized as follows:
Approximate Temperature, °K | State | Gaseous Absorption Spectrum, max in mμ (absorbance)
--- | --- | ---
77 | White solid | Nil
150 | White solid | Onset of absorption at short wavelength edge.
200 | Liquid | a) 245, 270 (0.04)
b) 280 (0.003)
c) 296 (0.04), 314
d) 368 (0.16)
   (c)
250 | Trace of liquid | a) 245, 270 (0.05)
b) 280 (0.10)
c) 296 (0.10), 314
d) 368 (0.30)
   (c)

a) Temperature of internal coil-cell temperature slightly lower.
b) Condensate on coil.
c) Transient appears during photoflash—see preceding section.

Increased absorption below 250 mμ is expected as the cell warms and the CF₃COOF vaporizes. By the time the cell temperature reaches 200°C the contents have been exposed to 3 to 4 photoflashes. New bands, shown in the table, reflect products from photolysis of the gas and, to some extent, of the solid remaining on the cooling coil. New bands fall into a pattern of four groups which seem to change in intensity together. The table lists absorbances for a band in each group determined from densitometry of flash spectra. Bands in the near u.v. appear much like carbonyl bands—the group "C" in particular resemble CF₃COCF₃. It is then tempting to consider a diketone such as CF₃COCOCF₃ for the higher band at 368 mμ, but no identification has been made yet; polyoxygen difluorides are also possible absorbers. The
sharply rising band near 280 μ falls where F₂ absorbs and may represent
a primary or secondary photochemical formation of the element. When
cells are recooled following a sequence of flash photolysis with warm-
ing, these absorbances are largely eliminated. The 280 μ band decreases
more slowly, as would be expected if it were due to F₂, which will not
condense at 77°K but must disappear by reaction.

After warming to room temperature and standing for 1-2 hours,
the products (IR) are largely CO₂, CF₄, CF₂O, HF; SiF₄ from reaction
with the cell is also found. No hypofluorite remains.

These survey experiments indicate products consistent with
photolysis of CF₃COOF at the carbonyl group are formed. We plan now to
conduct flash photolysis at 200°K with provision for both IR and mass
spectrometric analysis of products. We also seek to assign the product
ultraviolet spectra by correlation with known compounds.

**Mass Spectrometric Detection**

For some time the MS-10 mass spectrometer has been giving poor
mass resolution and irreproducible peak heights. The ratio of peaks is
necessary in our study to help show the molecular species from which
the mass peak is derived. Instability in peak heights introduces in-
tolerable errors in peak ratio measurements.

Origin for the difficulty was first sought in the associated
electronics and then in the mass analyzer element. No sources of diffi-
culty were found in extensive searches for possible electronic contribu-
tions. Repeated cleaning of the working parts in the mass spectrometer
gave no permanent improvement. The offending part was finally identified as the supporting frame piece bearing the 90° ion beam-defining slit (α-slit). This was done by substituting parts from a new MS-10 analyzer. Normal MS-10 operation was restored after the defective piece had been etched strongly with hydrochloric acid and polished.

A likely cause of the instability is a semiconducting layer on the front metal around the α-slit. Slower than normal dissipation of charges (resulting from impingement of ion beams on the surface) could cause spreading of ion beams, which would in turn cause loss in mass resolution and ion beam intensity. Whether attack by OF₂ was responsible for the difficulty may be revealed when we resume studies with OF₂. We can now return to the thermolysis study of OF₂ and examine OF/OF₂ (35/54) m/e ratio with changing temperature and the ionization efficiency curves for the two ions.

Experimental

CF₂O, CF₃COCF₃ (Penninsular), N₂F₃ (Air Products and Chemicals) and CF₃COOH (Allied Chemical) were chemically pure compounds used as received.

CF₃COOF was prepared according to the recipe of Cady and Kellog⁴--CF₃COOH, F₂, and H₂O, each in an N₂ carrier gas stream, were allowed to react in polyethylene vessels. The effluent stream was passed through successively colder traps to remove unconverted reactants. Finally, products were condensed (77°K) in a trap attached to a vacuum line and pumped to remove F₂ and N₂.
Products were transferred by vacuum line methods either to infrared or to photolysis cells. Infrared cells (5 cm path) were made of Teflon with AgCl windows.

Vacuum-jacketed silica cells for flash photolysis of low-temperature solutions and gases were described in previous reports. Low-temperature solutions were prepared by condensing (77°K) first CF₃COOF and then NF₃ solvent into the cell to make up the desired liquid volume. With the gas cell, samples were condensed on an internal cooling coil filled with liquid nitrogen. Flash photolytic and flash spectroscopic techniques were described earlier.

**Program**

Since product spectra from these survey experiments are consistent with compounds produced by photocleavage of bonds adjacent to the carbonyl group in CF₃COF, our detection of a photo-transient spectrum from this compound is particularly interesting. We therefore plan further experiments to seek to identify the photo-transient. These include: efforts to brighten the transient spectrum; kinetic spectroscopic measurements of the lifetime and of quenching of the transient; and correlation experiments with other compounds which might give fragments in common with those from CF₃COF. Using mass spectrometry, we shall resume study of OF₂ thermolysis and photolysis. Hypofluorites will also be examined as possible sources for OF. These experiments will complete our projected work on optical detection of OF for this year.
III. THE OF RADICAL: ELECTROLYSIS IN WET HF

The OF radical may be an intermediate in the formation of OF₂ during the electrolysis of wet HF. Our study of this electrolysis has a two-fold objective: (1) formulate a reaction mechanism on the basis of operating conditions, electrolyte composition, and product distribution, and (2) use electrochemical measurements to test and supplement this mechanism and to attempt to detect reaction intermediates.

During the electrolysis, gaseous products are swept out by helium and are sampled for gas chromatography. Changing product yields are readily seen by a rapid gas chromatographic analysis of the gaseous products (O₂, O₃, OF₂, H₂).

Water concentration was found to be important to OF₂ yield, but even careful control of water concentration was not sufficient to maintain constant OF₂ yield. However, periodic interruption of the electrolysis gave constant OF₂ yields over several hours. We believe interrupted operation gives constant yield by maintaining a steady-state composition of the anode surface film.

Studies at constant cell voltage showed the effects of H₂O and KF concentrations, interruption rate, temperature, and mercury impurity on the OF₂ yields. Later, when the Hg/Hg₂F₂ reference electrode was sufficiently developed for use in wet HF, the effects of anodic voltage on OF₂ yields were studied.

During this quarter changing the anode voltage-drop during interruption was found to have a small effect on yield. Components of
the anodic potential were determined by a decay method and a superimposed square wave method for resistance measurement seeking clues to electrode mechanisms. The ohmic overvoltage (mostly due to anodic film resistance) is surprisingly small, while activation overvoltage is at least as great as the limiting potential (i.e., $E_0 + \text{Nernst terms}$) as determined by decay measurements.

The action of gaseous oxygen atoms on NiF$_2$ was examined for clues to OF$_2$ formation. No OF$_2$ was observed.

**Experimental**

The main electrolysis cell$^{2a}$ (now with an anode area of 2.5 cm$^2$ and a cathode area of 10 cm$^2$), analytical techniques$^{2b}$, and the reference electrode$^{4a}$ have been described. Ohmic overvoltage was determined by two methods: potential decay and superimposed square wave. Potential decay was also used to evaluate activation overvoltage and the limiting potential.

**Potential Decay**—The components are those described previously$^{4b}$ for chronopotentiometry, but they have been rearranged to give decay curves. This is shown in Figure 1, which also shows the timer for interrupted operation.$^{2a}$

In normal operation, switch S$_1$ is set to the automatic timer, while switch S$_3$ is set to position 2 to include relay R$_1$ but not R$_2$. During interrupted operation the potentiostatic control voltage is switched by the timer from CV No. 1 to CV No. 2 for 1 sec every 120 sec. The potential drop during interruption is controlled by CV No. 2.
FIGURE 1

POTENTIAL DECAY

110 V AC

AUTOMATIC TIMER

S1

S2

67 V

P & B Hg WETTED
RELAYS J.M. 2.115.22

1 2 3

R1

R2

0.10 V

C.V. NO 1

C.V. NO 2

7 V

1K

1M

680 mmf

REFERENCE COUNTERTOP WORKING CONTROL VOLTAGE INPUT

POTENTIOSTAT

COUNTER ELECTRODE

REFERENCE ELECTRODE

WORKING ELECTRODE

OSCILLOSCOPE

WORKING GROUND

A

B

OSCILOSCOPE

REFERENCE ELECTRODE

WORKING ELECTRODE
With switch $S_1$ still set for normal operation, the oscilloscope is triggered during the interruption, but the AC from the timer produces noise on the oscilloscope. Therefore, a D.C.-operated trigger for decay was incorporated. When a decay curve is desired, switch $S_1$ is shifted to the 67 v. battery position, and $S_3$ is set to position 1 to activate both relay $R_1$ and relay $R_2$. Then switch $S_2$ is closed, disconnecting the potentiostat from the electrolysis cell and triggering the oscilloscope.

Potential decay was also used to measure ohmic overvoltage during continuous electrolysis. The automatic timer was not used and $S_3$ was at position 2. To obtain a decay curve, the above procedure is repeated; namely, $S_1$ is set to the 67 v. battery position, $S_3$ is set to position 1, and $S_2$ is closed.

The value of the ohmic overvoltage ($\eta_R$) can be determined from the break in the first 0.1 millisecond of the decay curve (Figure 2). The limiting potential ($E_{\infty}$) is determined when the decay curve levels off. The optimum time for this measurement has not yet been determined, but it is on the order of minutes. Combined activation and concentration overvoltage ($\eta_A + \eta_C$) is calculated by difference.

Superimposed Square Wave--A Hewlett-Packard square-wave generator (211A) at 500 pulses per second was used. However when it was superimposed on the potentiostatically controlled cell, the potentiostat responded and distorted the square wave. This was avoided by switching from potentiostatic to galvanostatic operation for the measurement. The components and arrangement are shown in Figure 3.
FIGURE 2
SAMPLE DECAY CURVE

Volts

Hg/Hg₂F₂

Anode voltage \( \eta_R \)

\( \eta_A (+\eta_C) \)

E∞

Time

0.1 M SEC.  2 MIN.
FIGURE 3
SUPERIMPOSED SQUARE WAVE

AUTOMATIC TIMER 110 V AC

P & B Hg WETTED RELAYS JM-2-115-2

WENKING POTENTIOSTAT-GALVANOSTAT

COUNTER ELEC.RODE
REFERENCE ELEC.RODE
WORKING ELEC.RODE

SQUARE WAVE GENERATOR

CONTROL VOLTAGE INPUT

REF. COUNTER WORK

REFERENCES

OSCILLOSCOPE

WORK-GROUND
In normal operation (interrupted-potentiostatic control), switch S₁ is at position a (Figure 3). To permit a measurement, switch S₁ is reversed to position b. This joins the counter and reference circuits from the Wenking to the counter electrode via a variable resistance (R₁) to give galvanostatic operation. By proper adjustment of R₁ and the control voltage (CV No. 1 and 2), the cell voltage and current can be kept approximately the same as during potentiostatic operation. This switching and adjusting takes 15-20 seconds and therefore is a longer interruption than normal (1 sec).

The square-wave output and R std. are adjusted to give a known current at 500 cycles/sec, when superimposed on the electrolysis cell. The IR drop of this signal can be determined from its oscilloscope pattern. Because stray 60-cycle AC distorts the 500-cycle signal, the signal cannot be displayed continuously. Instead, the scope is triggered to give a single display by closing S₂ which activates relay R₃. From the voltage drop found on this display and the known square-wave current, an electrode resistance can be calculated. This resistance and the known DC current give the ohmic overvoltage η_R.

Results

Interrupted Voltage Drop--The effects of impressed voltage during interruption are shown in Figure 4. In the range of 1.5 to 3.5 anode voltage during interruption there is no effect on yields. The voltage drop is in this range when no voltage is imposed during interruption and confirms an assumption made in our last report when studying
FIGURE 4
EFFECT OF VOLTAGE DROP DURING INTERRUPTION

CONDITIONS:
H2O ≈ 0.67 MOLE %
KF ≈ 1.0 MOLE %
TEMP. = 1.2°C
INTERUPTION RATE = 1 SEC./120 SEC.
SOLID POINTS = DUPLICATE RUN

ANODE TO CATHODE VOLTAGE DURING INTERRUPTION

ANODE TO REFERENCE (Hg2F2) VOLTAGE DURING INTERRUPTION

3 TO 4 HR. YIELD, % OF CURRENT

CURRENT DENSITY, AMPS/CM²

0.15
0.10
0.05
effect of cell voltage. However, at large voltage drops the OF$_2$ yield appears to be increasing slightly.

**Ohmic Overvoltage**—The ohmic overvoltage ($\eta_R$) of both electrodes determined by the two methods are shown in Figure 5 as a function of current density. The cathode show values of less than 0.1 v. by both methods, and is thus a negligible factor with cell voltages that generally are 6.0 v. or more.

The anode values are also quite small. At normal cell conditions (0.1 amp/cm$^2$; cell voltage = 6.2; anode voltage = 4.8) the potential decay method showed 0.25 v. while the square wave method gave ~0.40 v. The discrepancies between the two methods are possibly due to effects of the alternating square wave on the electrode surface; as both measurements show low $\eta_R$, no attempt was made at this time to resolve the difference.

The $\eta_R$ of continuous operation was also found to be quite low, in fact even lower than interrupted operation. Since determination of $\eta_R$ by potential decay requires interrupting the current, repeated determinations approach the interruption values. However, this is due to changes in the current density with interruption rather than any great change in $\eta_R$.

The same values of $\eta_R$ were obtained regardless of whether current density was progressively increased or decreased (Figure 5). However, the yield of OF$_2$ was not constant and Figure 6 shows a drop in OF$_2$ yield as current density was decreased. The effect was observed
FIGURE 5
OHMIC OVERTONAGE

\[ \eta, \text{ VOLTS} \]

\[ \text{CURRENT DENSITY, AMPS/CM}^2 \]

Interrupted Operation
- --- SQUARE WAVE METHOD
○ VOLTAGE INCREASED
● VOLTAGE DECREASED

Potential Decay
- --- POTENTIAL DECAY
△ VOLTAGE INCREASED
▲ VOLTAGE DECREASED

Continuous Operation
- --- POTENTIAL DECAY
FIGURE 6
OF₂ YIELD VARIATION WHILE MEASURING OHMIC OVERVOLTAGE

CELL CONDITIONS:
\( \text{H}_2\text{O} = 0.68 \text{ MOLE \%} \)
\( \text{KF} = 1.0 \text{ MOLE \%} \)
\( \text{TEMP.} = 1.2^\circ \text{C} \)
\( \text{INTERRUPTION RATE} = 1 \text{ SEC./120 SEC.} \)

WITH POTENTIAL DECAY METHOD

WITH SQUARE WAVE METHOD

CURRENT DENSITY, AMPS/CM²
in both methods but was most pronounced in the square-wave method (lower half of Figure 6). This drop is not short-lived because measurements made a day later gave low OF$_2$ yields even at the higher current densities. But again the $\eta_R$'s were unchanged. The anode $\eta_R$ increases during the induction period at approximately the same rate as the current (Figure 7); hence, since $\eta_R = IR$, we find the resistance ($R$) of the anode film is constant during the induction period.

**Limiting Potential ($E^\infty$), Activation Overvoltage ($\eta_A$) and Concentration Overvoltage ($\eta_C$)**—The optimum conditions for these determinations are still being studied. Thus far, $E^\infty$ values vary over a 0.5 v. range (1.7 to 2.2 v.) with 2.0 v. average. Within this spread no correlation with OF$_2$ yields could be noted.

The $\eta_A + \eta_C$ had a wider range (2.5-3.5 v.) and appear to be the largest and probably the most important. Because the solution is well stirred by sparging and evolving gases, $\eta_C$ should be small but not zero. Using the Hg/H$_2$F$_2$ electrode as a reference, Figure 8 shows an estimate of the components of cell voltage at a normal operating condition.

**Oxygen Atom Reactions**—Standard vacuum-line techniques with a Raytheon Microtherm unit were used to produce oxygen atoms. The behavior of nickel fluoride to $\cdot$* atoms is shown by the following equation:

$$\text{NiF}_2\cdot4\text{H}_2\text{O (green) + }\cdot* \rightarrow \text{yellow amorphous powder (NiF}_2\text{) with a black crust}$$
FIGURE 7
INDUCTION PERIOD

CELL CONDITIONS:

H$_2$O $\approx$ 0.72 MOLE %
KF $\approx$ 1.0 MOLE %
TEMP. = 1.2°C
VOLTAGE - CONTROL = 5.0 V
CELL = 6.2 - 6.5
INTERUPTION RATE = 1 SEC./120 SEC.

![Graphs showing current density and voltage over time for anode and cathode.](image-url)
**FIGURE 8**

**COMPONENTS OF CELL VOLTAGE**

\[ E_{\text{CELL}} (6.2 \text{ V}) \]

\[ E_{\text{CATH}} (1.4) \]

\[ E_{\text{ANODE}} (4.8) \]

\[ \eta_R (0.3) \]

\[ \eta_A + \eta_C (2.5) \]

\[ E_\infty (2.0) \]

\[ \eta_A + \eta_C (1.0) \]

\[ \eta_R (0.1) \]

\[ E_\infty (0.3) \]

**VOLTAGE vs Hg/Hg_2F_2**

\[ E_\infty = E_0 + \frac{RT}{ZF} \ln \left( \frac{\text{REACTANT}}{\text{PRODUCTS}} \right) \]
The yellow powder is amorphous as it did not give an X-ray pattern. However, when NiF₂·4H₂O (B & A, Technical grade) was dried in an oven at 140°C, the resulting yellow powder was crystalline enough to show NiF₂ patterns. The black crust gave weak X-ray patterns that may be iron or chromium oxides, but no nickel oxide was detected. Possibly the Ni oxides are amorphous just as the yellow NiF₂.

Nickelic oxide (Baker C.P. analyzed) was reacted with 48% aqueous HF and with wet HF in the presence of KF as represented by the following equation:

\[
\text{Nickel oxide} + \text{HF} \rightarrow \text{NiF}_2 + \text{H}_2\text{O} + \text{gas}
\]

The gas from this reaction gave a positive KI test. However, when this gas was analyzed with the G.C. no OF₂ was found, and at most a trace of O₃, but there is an unidentified component.

None of these reactions give any definite support for our proposal that decomposition of H₂O gives oxygen atoms and these attack the NiF₂·HF system to form OF₂.

**Discussion**

Voltage drop during interruption has no effect on OF₂ yield for intermediate values, but large drops seem to be beneficial. The present study did not examine very small voltage drops to determine how closely the actual operating potential can be approached before interruption becomes ineffective. Previous results indicate that suitable selection of interruption schedule can always raise OF₂ yield. Control of voltage drop during interruption may have a similar effect in permitting maximum OF₂ yields.
We expected the ohmic overvoltage at the anode to comprise a large portion of the anodic potential for two reasons: (1) nickel in HF is not attacked, presumably because of an impervious film; and (2) the large ohmic overvoltage observed during electrolysis of dry HF with nickel anodes has been attributed to anode film resistance. However, we did observe that anode films in wet HF appeared thin and quite different from those in dry HF. Furthermore, the current density was higher in wet HF. Apparently, the impervious anode film obtained during electrolysis of wet HF is a reasonably good conductor, or perhaps a semiconductor.

The much lower ohmic overvoltage in wet HF than in dry HF shows that water causes a large decrease in the anode film resistance. We do not know how the water affects the NiF₂ film. Formation of a hydrate (NiF₂·4H₂O) is unlikely as it is dehydrated by anhydrous HF. Our infrared reflectance spectra showed OH bonds and adsorbed water on an anode after use, but no special precautions were taken to prevent adsorption of moisture from the air after the anode was rinsed with HF and evacuated. We cannot be certain that absorptions in the infrared are due to material that survived contact with the electrolyte.

The unchanging resistance during the induction period and when OF₂ yields are different is surprising because we had assumed a marked change in the anode film. The nature of these changes will be investigated by other means.
Program

We plan to determine if products in the electrolyte contribute to the $E_\infty$ of the potential decays and to an arrest in chronopotentiograms. Using coulometry and chronopotentiometry, we will also study anode films during the induction period and after steady $\text{OF}_2$ yields have been attained at several levels.
IV. ICH PRESSURE REACTIONS OF DIFLUORODIAZINE

During the last quarter we have made additional attempts to prepare $\text{N}_2\text{F}^+\text{ClO}_4^-$, have attempted to add $\text{F}_2$ to cis-$\text{N}_2\text{F}_2$, and have improved the procedure for isomerizing trans-$\text{N}_2\text{F}_2$ to cis-$\text{N}_2\text{F}_2$. Development of a capsule for use at very high pressures also was continued.

Further infrared studies of $\text{N}_2\text{F}_2:$SbF$_5$ complexes in SbF$_5$ solutions have failed to detect N-F stretches. However, an electron paramagnetic resonance examination of these solutions has revealed free radical species.

**Preparation of $\text{N}_2\text{F}^+\text{ClO}_4^-$**

Further attempts were made to form $\text{N}_2\text{F}^+\text{ClO}_4^-$ by the reactions of various combinations of NF$_3$, $\text{N}_2\text{F}_2$, and Mg(ClO$_4$)$_2$. At conditions of $-78^\circ\text{C}$ and 35,000 psig for 4 hours, no reaction took place with either a mixture of NF$_3$, cis-$\text{N}_2\text{F}_2$, and Mg(ClO$_4$)$_2$ in a mole ratio 7.5 to 1. In several attempts, mixtures of NF$_3$, cis-$\text{N}_2\text{F}_2$, and Mg(ClO$_4$)$_2$ in a mole ratio of approximately 10 to 4 to 1 ignited at about 0$^\circ$C, and 50 to 83% of the cis-$\text{N}_2\text{F}_2$ decomposed. In contrast, mixtures of NF$_3$ and cis-$\text{N}_2\text{F}_2$ in a mole ratio of about 10 to 4 show only 20% decomposition of cis-$\text{N}_2\text{F}_2$ up to temperatures of $82^\circ$C. Apparently, the Mg(ClO$_4$)$_2$ either acts as a catalyst for the decomposition of cis-$\text{N}_2\text{F}_2$ or reacts first with NF$_3$.

**Addition of $\text{F}_2$ to cis-$\text{N}_2\text{F}_2$**

All previous attempts to add to the $\text{N}_2\text{F}_2$ molecule to form liquids or solids have been unsuccessful. The addition of $\text{F}_2$ to $\text{N}_2\text{F}_2$
was not tried previously because the product would be a gas, N$_2$F$_4$. However, it was of interest to try this reaction in order to see if such an addition is possible. A 3 to 1 mole ratio of F$_2$ and cis-N$_2$F$_2$ was maintained at 87,000 psig and 69°C for 4 hours. No N$_2$F$_4$ was formed, but 8% of the cis-N$_2$F$_2$ decomposed. Apparently, F$_2$ is not as effective a diluent to prevent decomposition as NF$_3$ where only 20% decomposition occurred at 5 to 2 NF$_3$ to N$_2$F$_2$ at 82°C.

**Isomerization of trans-N$_2$F$_2$**

Isomerization of trans-N$_2$F$_2$ to cis-N$_2$F$_2$ was carried out in a stainless steel vessel for one hour at 75°C by a variation of Hurst's procedure. Hurst activated bombs for the isomerization reaction by decomposing a small amount of trans-N$_2$F$_2$ in them. We have been unsuccessful in activating bombs in this way. Instead, we have found that isomerization will take place if a small amount of F$_2$ is mixed with the trans-N$_2$F$_2$. For example, when 150 mg of F$_2$ was added to 4 g of trans-N$_2$F$_2$, the yield of the cis isomer was about 94%, with a total recovery of 97%.

**Very-High-Pressure Apparatus**

We have continued evaluating capsules for use at very high pressures. Figure 9 shows a nickel capsule, fabricated in our shops, of semi-bellows design complete with valve. Figure 10 presents drawings of two electroformed nickel bellows with .002 and .010 wall thickness (Servomecoder Corporation, Clifton, N. J.). These bellows were joined to the internal valve and lower closure by silver soldering.
FIGURE 9
SEMI-BELLOWS TYPE NICKEL CAPSULE

- NICHROME CONTROL WIRE
- 1/8" O.D. x 0.038" I.D.
- 304 SS TUBE

- 0.200"
- 0.7750"
- 0.2450"
- 0.030"
- 0.0615"
- 0.1340"
- 0.10"
- 9/64"
- 0.030"
Figure 10
Electro-Formed Nickel Bellows

C'EN END

OPEN END

WALL

4 FINS

OPEN END

WALL

5 FINS
The capsule with semi-bellows was tested in the very-high-pressure apparatus with piston-cylinder device using ethane in the capsule. The capsule held until about 150,000 psi when the ethane was suddenly released. The internal valve was found to be deformed possibly due to squeezing by the cylinder walls of the device.

The piston-cylinder device was eliminated from the apparatus and tests were carried out hydrostatically. A capsule made from 0.002 inch thick bellows containing ethane was subjected to 50,000 psi. After depressuring the internal valve was opened and the ethane partially recovered. However, cracks were found in the recovered capsule; these probably occurred during ethane recovery and indicate the walls were too brittle.

A capsule made from a bellows with .010 inch wall was annealed before use. It was filled with liquid ethane and pressured hydrostatically to 125,000 psi. When pressure was dropped to atmospheric the internal valve could not be opened. However, when the capsule was removed from the pressure vessel and cooled in liquid nitrogen, the valve opened easily and the ethane was recovered. The walls of this capsule were intact.

In future experiments, attempts will be made to reach 200,000 psi and to pressure N2F2.

**Infrared Study of N2F2:SbF5**

In preparing solutions of N2F2:SbF5 in SbF5, we observed a royal blue solution when N2F2 contacted the surface of the SbF5. We are
attempting to obtain infrared spectra of this blue solution and the colorless or yellow solution from the $\text{N}_2\text{F}_2$:SbF$_5$ complex to determine whether a nitrogen-containing species such as $\text{N}_2\text{F}^+$ is present.

Our initial failure to detect N-F stretches at 1057 cm$^{-1}$, as found in solid state spectra, was attributed to inadequate mixing and sampling of the solution.$^4$ Therefore, the apparatus was modified so that the flow of SbF$_5$ and $\text{N}_2\text{F}_2$ into the Kel-F reaction tube could be alternated through a three-way valve$^{12}$ connected to Teflon tubing that extended to the bottom of the reaction tube. With this arrangement, a uniform blue solution was produced, and the color persisted so long as the $\text{N}_2\text{F}_2$ bubbled through the SbF$_5$. However, the solid complex eventually plugged the tubing and interrupted the flow. Nitrogen pressure was used to force the SbF$_5$ through the blockage and to force some of the solution into the infrared cell.$^{48}$ The blue color disappeared before an infrared spectrum could be obtained. The resulting yellow solution failed to show N-F stretches.

To eliminate the plugging problem, the Teflon tubing was replaced with a glass bubbler. With this arrangement, the SbF$_5$ was introduced through a Kel-F valve at the bottom of the bubbler, and the $\text{N}_2\text{F}_2$ was introduced through a tube at the top. Again, a uniform blue solution was produced, but it turned yellow a few seconds after bubbling of $\text{N}_2\text{F}_2$ was stopped, and no evidence for a N-F compound was found in its infrared spectrum.
EPR Study of \( \text{N}_2\text{F}_2: \text{SbF}_5 \)

On the possibility that the blue color may indicate the presence of free radicals, the \( \text{N}_2\text{F}_2: \text{SbF}_5 \) solutions were examined by electron paramagnetic resonance. The solutions were prepared directly in the EPR cells in an apparatus similar to one that had been used for infrared studies. The cells were of two types. The first was 1/4-in. O.D. Kel-F tubing heat-sealed at the bottom. It was not satisfactory because it would not hold SbF\(_5\) for more than one hour before a hole developed. The second, shown in Figure 11, was pyrex glass. Spectra were obtained on a modified Varian Model V4500 spectrometer operated at x-band frequencies of about 9500 mc/sec. For increased sensitivity, a Varian Model V4560 100-kc Modulating Unit was used. The rectangular cavity operated in the \( \text{TE}_{012} \) mode. The cavity is actually a low-loss tuneable short. Such an adjustable wave-guide short is of the non-contracting type for minimum r-f loss.

Figure 12 shows blank EPR spectra for: (a) the empty Kel-F cell, (b) the empty glass cell, and (c) freshly distilled SbF\(_5\) in the glass cell. No signal is apparent in the Kel-F spectra, but a weak signal is evident in the other two.

Figures 13 and 14 show the spectra for the yellow \( \text{N}_2\text{F}_2: \text{SbF}_5 \) solution in Kel-F and in glass. The sample in Kel-F shows two distinct species—at \( g = 2.002 \) and 2.027—whereas the sample in glass shows only one—at \( g = 2.027 \). To test whether Kel-F was contributing to the spectrum, Kel-F particles were added to a glass tube before the \( \text{N}_2\text{F}_2: \text{SbF}_5 \) was added. The EPR spectrum of this sample is shown in Figure 15. Two species are
FIGURE 11
GLASS CELL FOR EPR STUDIES

FEMALE SYRINGE

CA 1'' OD

INNER TUBE 3-4 mm OD

LARGE TUBE 9 mm OD

CA ¼''
FIGURE 12
BLANK EXPERIMENTS

a. KEL F TUBING
S. 8x100
M. 4x100
RES 2 SC 2/4

b. PYREX GLASS TUBE
S. 2x1000
M. 4x100
RES 3 SC 2/6 HI

c. DISTILLED SbF₅ IN GLASS TUBE
S. 2x1000
M. 4x100
RES 3 SC 2/6 HI

GAUSS
FIGURE 13
EPR SPECTRUM OF CIS-N$_2$F$_2$-SbF$_5$ SOLUTION IN KEL-F

~120 GAUSS
~30 GAUSS
8 GAUSS

FIGURE 14
EPR SPECTRUM OF CIS-N$_2$F$_2$-SbF$_5$ SOLUTION IN GLASS

$g=2.027$

S-1×1K
M-5×1
RES 2 SC 2/2

$\sim$120 GAUSS
FIGURE 15
EPR SPECTRUM OF CIS-N₂F₂-SbF₅ SOLUTION IN GLASS CONTAINING KEL-F

S-1×1000
M-4×100
RES 3 SC 2/4 Hi

- 38 -
g=2.093  g=2.027

3340  3420  3560
GAUSS
evident, but only the one at $g = 2.027$ corresponds to the spectrum of $\text{N}_2\text{F}_2:\text{SbF}_5$ in glass; the other represents a new species at $g = 2.093$.

The spectrum in Figure 13 can be resolved into two separate spectra, as shown by the hand drawings in Figure 16. The five-line spectrum is consistent with an N$_2$ containing radical in which the $^{14}\text{N}$ atoms, with nuclear spin (I) of one, give rise to the splitting:

$$2I_{\text{Total}} + 1 = 2 \times 2 + 1 = 5.$$

The splitting and the expected intensities are diagrammed in Figure 17. The five-line spectrum exhibits the same intensity pattern. Furthermore, the $g$-value of 2.002 is consistent with a nitrogen-containing radical, and the overall band width of about 30 gauss is compatible with $^{14}\text{N}$ splittings. However, the number of bonded fluorines cannot be ascertained from these spectra but may be revealed if fine structure EPR spectra can be obtained, showing the $^{19}\text{F}$ splittings. Absence of the N$_2$ radical signal in glass (Figure 14) may be due to adsorption.

The 16-line spectrum in Figure 16 was extracted from Figure 13 with the aid of the spectrum in glass in Figure 14. The $g$-value is $2.077$ and the band width is about 120 gauss. Evidence shows it arises from an antimony-containing radical. The $g$-value coincides with that of the six-line spectrum ($I_{121}\text{Sb} = 5/2$) of the paramagnetic sites in powdered Sb$_2$Se$_5$, as shown in Figure 18. The band width is similar to that found in aqueous NO$^+\text{SbF}_6^-$ (Figure 19) and SbCl$_3$ in acetone (Figure 20).

Spin-spin coupling constants for antimony are chosen to be 16, 24 or 32 gauss to fit the observed band width, while that for $^{19}\text{F}$ is
FIGURE 16

RESOLUTION OF EPR SPECTRUM IN KEL-F
FIGURE 17
SPLITTING DIAGRAM FOR N₂ RADICAL SPECIES

GROUND LEVEL

(1=1)

SPLIT BY FIRST ¹⁴N

SPLIT BY SECOND ¹⁴N

RELATIVE INTENSITIES 1 2 3 2 1
Figure 18

EPR Spectrum of Solid Antimony Pentasulfide

S: 2x10^4
M: 4x10^5
Res 3 x 2/6 H}

Gauss
FIGURE 19
EPR SPECTRUM OF NO\textsuperscript{+}SbF\textsubscript{6}\textsuperscript{-} IN WATER
1 DAY LATER
ALSO ADDED
MORE H\textsubscript{2}O
S - 2 \times 1K
M - 4 \times 100
RES 3 SC 2/6 Hi

FIGURE 20
EPR SPECTRUM OF SbCl\textsubscript{3} IN ACETONE
S - 2 \times 1K
M - 4 \times 100
RES 3 SC 2/5 Hi
about 8 gauss. Several antimony fluoride species are possible, such as SbF₄, SbF₅⁻, SbF₆, or SbF₆⁻ free radicals. The combinations of antimony and fluorine spin-spin couplings for each structure as shown in Figure 21. The constructed spectrum for SbF₅⁻, using the Sb coupling constant of approximately 16, fits the observed spectrum very closely in predicted splittings (16 lines total) and intensities (Figure 22). In Figure 22, and even more so in Figure 14, the lack of symmetry with respect to the center of the pattern is evident. This phenomenon apparently arises from anisotropy of the radical species interacting with the nuclear orientations possible in the external field, and is consistent with the trigonal bipyramide structure for SbF₅. The lack of fit of the spectrum with SbF₆ appears to exclude the location of the electron on an Sb atom of the linear chain structure attributed to liquid SbF₅. Perhaps addition of an electron to the Sb in such a chain frees that SbF₅ unit.

The presence of SbF₅⁻ as the counter species to the N₂ radical species suggests a single electron transfer:

\[ \text{N}_2\text{F}_2 + \text{SbF}_5(\text{soln}) \rightarrow \text{N}_2\text{F}_2^{+}\text{SbF}_5^- (\text{soln}) \]

However, that the N₂ species arises from N₂F₂ rather than from an impurity, such as N₂O or N₂O₂, has to be verified.

**Program**

We will continue attempts to add to N₂F₂ at high pressure and will attempt to pressure N₂F₂ to near 200,000 psi. Infrared and EPR studies of N₂F₂ solutions in SbF₅ will be continued.
Figure 21
SPLITTING DIAGRAMS FOR ANTIMONY FLUORIDE RADICAL SPECIES
(SHOWN FOR HALF OF SPECTRUM)

SbF₆⁻

144 GAUSS
17 LINE SPECTRUM

196 GAUSS
25 LINE SPECTRUM

SbF₅⁻

120 GAUSS
16 LINE SPECTRUM

160 GAUSS
21 LINE SPECTRUM

SbF₄⁻

96 GAUSS
13 LINE SPECTRUM

128 GAUSS
17 LINE SPECTRUM

160 GAUSS
21 LINE SPECTRUM

RELATIVE INTENSITIES OF A LINE SPECTRUM
Figure 22: Comparison of Calculated and Observed Spectrum for $\text{SF}_5^-$
V. REFERENCES

2. Second Annual Report, this contract: (a) pp. 44 and A-9; (b) pp. A-10 and A-14.
3. Ninth Quarter Report, this contract.
4. Tenth Quarter Report, this contract: (a) p. 5; (b) p. 6; (c) p. 17; (d) p. 25; (e) Figure 13; (f) Figure 10; (g) Figure 11.