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CORROSION ASSOCIATED WITH FLUORINATION IN THE OAK RIDGE NATIONAL LABORATORY FLUORIDE VOLATILITY PROCESS

> A. P. Litmon A. E. Goldmon

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METALLURGY DIVISION

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A. P. Litman and A. E. Goldman

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### CONTENTS

Summ	ary-	
[ <b>.</b> ]	Mark	I Volatility Pilot Plant L Nickel Fluorinator
	Α.	Material Selection and Fabrication
	Β.	Operational History
	С.	Reaction to Environment
		1. Chemistry
		2. Dimensional Analysis
		3. Metallographic Study
		4. Summary of Corrosive Attack
	D.	Discussion of Results
		1. Individual Actions of F <sub>2</sub> , UF <sub>6</sub> , and Fused Fluoride Salts
		2. Collective Attack During Volatility Process Fluorination
		a. Interior Bulk Losses
		b. Interior Intergranular Attack
		c. Exterior Intergranular Attack
		d. Grain-Size Variations
. 1	Mark	II Volatility Pilot Plant L Nickel Fluorinator
	Α.	Material Selection and Fabrication-Design Changes
	Β.	Operational History
	C.	Reaction to Environment
		1. Visual and Vidigage Inspections
		2. Chemistry
		3. Dimensional Analysis
		4. Metallographic Study
		5. Summary of Corrosive Attack
	D.	Discussion of Results
		1. Interior Bulk Losses
		2. Interior Intergramular Penetration
		3. Exterior Intergranular Attack
		4. Grain-Size Variations

r

,

8

•

	Ε.	Corrosion of Internal Components from the Mark II VPP Fluorinator	71
III.	Benc	h-Scale Fluorination Corrosion Studies	77
	Α.	Inconel Fluorinator	79
		1. Test Method	79
		2. Discussion of Results	80
	Β.	A Nickel Miniature Fluorinators	84
		1. Test Method	84
		2. Discussion of Results	89
	С.	INOR-8 Fluorinators	95
		1. Test Method	95
		2. Discussion of Results	96
IV.	Vola	tility Pilot Plant Scouting Corrosion Tests	108
	Α.	Material Selection	108
	Β.	Test Method	113
	С.	Reactions to Environments	113
	D.	Discussion of Results	119
	Ε.	Future Studies	123
V.	Argo	onne National Laboratory Fluorination Corrosion Studies	125
	Α.	Test Method	125
	Β.	Discussion of Results	126
VI.	Supp	elementary Volatility Pilot Plant Equipment	129
Ack	nowle	dgment	129
Bib	liogr	aphy	130
App Sp	endix ecime	A - Photomicrographs of VPP Scouting Corrosion Test	131
App	endix Com Was Abs Val Flu Pro	B - Supplementary VPP Equipment	153 155 159 166 173 174 184

#### CORROSION ASSOCIATED WITH FLUORINATION IN THE OAK RIDGE NATIONAL LABORATORY FLUORIDE VOLATILITY PROCESS

A. P. Litman and A. E. Goldman

#### SUMMARY

This report evaluates chemical corrosion on reaction vessels and equipment used during the fluorination of fused-salt fuels and subsequent associated operations in the Oak Ridge National Laboratory (ORNL) Fluoride Volatility Process and is a continuation and expansion of the Metallurgy Division assistance to the Chemical Technology Division in this regard. The fluorination phase consists of converting uranium tetrafluoride to volatile uranium hexafluoride by fluorine sparging of molten fluoride salts and subsequent decontamination and recovery of the uranium hexafluoride.

For convenience in reporting, this document is divided into six sections. Sections I and II describe the corrosion behavior of full-size fluorination vessels fabricated from L nickel and used during Volatility Pilot Plant (VPP) operations. Section III covers corrosion evaluations of bench-scale fluorinators made of A nickel, Inconel, and INOR-8, which were operated by the Volatility Studies Group, Chemical Development Section A, of the Chemical Technology Division. Section IV describes scouting tests of many proprietary and nonproprietary materials exposed to the pilot plant fluorinator environments and the reactions of the various materials to those service conditions. Appendix A shows selected photomicrographs of the corrosion specimens described in Section IV. For comparison, results of some of the corrosion tests performed by the Argonne National Laboratory on metal coupons under simulated fluorination conditions are reported in Section V. Section VI and Appendix B deal with results of examinations of supplementary VPP equipment including a radioactive-products trap, a waste-salt line, the absorbers, valves and fittings, the fluorine-disposal system, and process-gas lines.

In this report, corrosive attack is reported as mils per month based on molten salt residence time or mils per hour based on fluorine exposure time. These rates are included specifically for comparison purposes, are not exact, and should not be extrapolated into longer time periods for design work or other applications.

Two fluorinators were used in the VPP to carry out the fluorination reactions. These vessels, Mark I and Mark II, were fabricated into right cylinders, approx 4-1/2 ft in height, from the same heat of L (low carbon) nickel. The first vessel contained equimolar NaF-ZrF<sub>4</sub> or NaF-ZrF<sub>4</sub>-UF<sub>4</sub> (48-46-4 mole %) for approx 1250 hr at 600-725°C. Over a period of 61 hr, 57 500 standard liters of  $F_2$  were sparged into the salts. This constituted a  $F_2$ :U mole ratio of 3:1 beyond theoretical requirements. The Mark II fluorinator contained fluoride salts of approximately the same compositions plus small additions of  $PuF_{4_1}$  during three runs. The salts were kept molten at 540-730°C for approx 1950 hr and about 60 500 standard liters of  $F_2$  were sparged into the Mark II melts in 92 hr.

Both fluorinators sustained large corrosion losses consisting of extensive wall thinning, severe interior intergranular attack, and a moderate exterior oxidation attack. Maximum deterioration on the Mark I vessel occurred in the middle vapor region at a calculated rate of 1.2 mils/hr, based on fluorine sparge time, or 46 mils/month, based on time of exposure to molten salts. The second vessel showed maximum attack in the salt-containing region at similarly calculated rates of 1.1 mils/hr and 60 mils/month. Some evidence was found to indicate that the intergranular attack may have resulted from sulfur in the systems. Bulk metal losses from the vessel's walls were believed to be the result of cyclic losses of  $\text{NiF}_{\mathcal{O}}$  "protective" films. 'Fne films were formed on the interior walls of the fluorinators during conditioning and fluorination treatments and lost as the result of rupturing, spalling, fluxing, washing actions, and/or dissolution in highly corrosive condensates formed during operations. The shift in maximum corrosion attack geometry in the two fluorinators is believed to have resulted from differences in operating conditions. The Mark II vessel experienced higher temperatures, longer fluorine exposure times, and extended uranium residence times in its salt baths.

Specimens removed from the wall of the first fluorinator showed a variation in average ASTM grain-size number of 5-6 to > 1, the largest grains being found in the middle vapor region. The second vessel had a more uniform grain-size pattern, average ASTM grain-size numbers varying from 3-5 to 2-4.

The variations in grain sizes are believed to have resulted from variable heating rates during initial usage. Low rates permit more complete internal stress recovery prior to the start of recrystallization which results in fewer nucleation sites and therefore larger grains during recrystallization. Metallographic examinations did not provide evidence of a causal relationship between grain size and fluorinator wall corrosion.

Examinations of bench-scale reactors, where simulated fluorination environments were provided to study process variables and corrosion, showed that A nickel had the highest degree of corrosion resistance as a fluorinator material of construction when compared with Inconel and INOR-8. Intergranular penetration and subsequent sloughing of whole grains seemed to be the predominant mode of corrosive attack on the Inconel vessel. At the higher test temperatures, 600°C, INOR-8 miniature fluorinators showed large bulk metal losses plus selective losses of chromium, molybdenum, and iron from the exposed alloy surfaces. Evidence of a marked reduction in attack on nickel and INOR-8 was found during lower temperature studies at 450-525°C. These lower temperature operations were made possible by adding lithium fluoride to the sodium fluoride-zirconium tetrafluoride salt mixtures.

Scouting corrosion tests were performed in the VPP's fluorinators using rod, sheet, or wire specimens of commercial and developmental alloys. These tests were subjected to serious limitations due to the lack of control over operating conditions and thus considerable variation in the corrosion of L nickel control specimens resulted. Those nickel-rich alloys containing iron and cobalt showed some superiority in corrosion resistance when compared with L nickel specimens. This was probably because of the low volatility of iron and cobalt fluorides. Nickel-rich alloys containing molybdenum additions showed variable behavior in the fluorination environment. Some of the data suggested improved resistance over L nickel while other tests showed the reverse.

- 3 -

Since both of the known molybdenum fluorides that could be formed during fluorination have very high volatility, one would not expect improved resistance from molybdenum additions. The experiments emphasize that the present method of selection of test materials based on the low volatility of metal fluorides that may form during fluorination continues to have merit. Additional experimental nickel-base alloy corrosion specimens, containing magnesium, aluminum, iron, cobalt, or manganese, have been fabricated and will be used in future screening tests in a subsequent pilot plant fluorinator.

A review of one fluorination test series conducted by the Argonne National Laboratory gave general agreement with ORNL scouting corrosion test specimen results, although comparisons were hampered by different test conditions. The Argonne National Laboratory has suggested that the corrosion problem be attacked by further studies on the use of cold wall vessels, spray towers, or low-melting salts for volatility processes.

Visual and metallographic examinations plus ultrasonic measurements of other VPP vessels and equipment fabricated generally from Monel or Inconel showed a wide variation in resistance to those various local service conditions. The studies suggest that Inconel can continue to be used as a material of construction for some components but frequent inspections are indicated. Monel appears generally satisfactory for the applications to date.

From a corrosion standpoint, the fluorination vessel in the VPP continues to be the most vulnerable to attack due to the nature of the contained environment and the high temperature necessary for fluorination. The continued use of L nickel for the fluorination vessel does not appear prohibitive for batch operations only due to the present high value of the pilot plant's product. At present, the only guarantee for improved service life for nickel fluorinators seems to be utilization of the lowest practical temperature. Although not conclusively proven for the fluorination vessels, reduction of sulfur contamination and the ensuring of a uniform, small-grain size in the vessels may improve vessel performance. For long-time fluorinator integrity, selection or development of a new material of construction, the use of salts with lower melting points, or the use of a cold wall vessel seems necessary.

- 4 -

The evaluation of process corrosion that occurred during the development studies of hydrogen fluoride dissolution of uranium-bearing fuel elements, the head-end cycle of the volatility process, will be covered in a separate report.<sup>1</sup>

#### I. Mark I Volatility Pilot Plant L Nickel Fluorinator

#### A. Material Selection and Fabrication

The selection of material for the first pilot plant fluorinator was made by members of the Chemical Technology Division after a study of the available corrosion literature and the ASME Boiler and Pressure Vessel Code.<sup>2,3,4</sup> Nickel seemed to be the most likely candidate material of construction, although at 600-700°C, the anticipated operating temperature range of the fluorinator, Myers and DeLong reported penetration rates of fluorine on nickel of 16-34 mils/month. The ASME Code allowable design stress data above approx 315°C were not available for commercial purity A nickel (0.05-0.15 wt % C). This was because of the known effects of embrittlement through intercrystalline precipitation of graphite in nickel containing carbon after long-time exposure to high temperatures.<sup>5</sup> However, satisfactory design data were available for low-carbon L nickel at approx 650°C, so this material was selected for the first pilot plant fluorinator.

The Mark I fluorinator was fabricated at ORNL from L nickel using a heat with the vendor's analysis of 99.36% Ni-0.02% C-0.23% Fe-0.06% Cu-0.26% Mn-0.04% Si-0.005% S. Annealed plate stock of 1/4-in. thickness was rolled into

<sup>1</sup>A. E. Goldman and A. P. Litman, Corrosion Associated with Hydrogen Fluoride Dissolution in the Fluoride Volatility Process, ORNL-2833 (to be published).

<sup>2</sup>W. R. Myers and W. B. DeLong, "Fluorine Corrosion," <u>Chem. Engr. Prog.</u> (May, 1948).

<sup>3</sup>"Engineering Properties of Nickel," <u>Tech. Bull. T-15</u>, The International Nickel Company, Inc., New York, Revised, p. 21, July, 1949.

<sup>4</sup>Rules for Construction of Unfired Pressure Vessels, ASME Boiler and Pressure Vessel Code, Section VIII, <u>Am. Soc. Mech. Eng.</u>, 1956 Edition.

<sup>5</sup>W. A. Mudge, "Nickel and Nickel-Copper, Nickel-Manganese, and Related High-Nickel Alloys," <u>The Corrosion Handbook</u> (ed. by H. H. Uhlig), p. 683, John Wiley and Sons, Inc., New York, 1948. a 14-in.-diam cylinder, 54 in. in height, for the vessel shell and longitudinally seam welded using an inert-gas metal-arc (nonconsumable) process. The filler material used was INCO-61 welding wire and ORNL Reactor Material Specification RMWS-5 was used as the basis of the joining procedure.<sup>6</sup> A nominal 3/8-in.-thick L nickel flanged and dished head was welded to the shell to form the bottom of the vessel (Fig. 1).

#### B. Operational History

The Mark I fluorinator was used by the Unit Operations Section of the Chemical Technology Division during preliminary fluorination equipment studies for a period of about three months. During that time, no fluorine or uranium-containing molten salts were in contact with the vessel. Table I cites the process conditions in detail for those studies and for the more extensive "M" equipment shakedown and "C" process demonstration runs performed later in the VPP.

Figure 2 shows the position of the Mark T fluorinator during the VPP runs while Fig. 3 shows the interior piping, gas dispersion assembly, and the placement of an early group of corrosion test specimens. The lower half of the fluorination vessel was surrounded by a vertical tube-type electric-resistance furnace of 30-kw rating to provide the necessary heat (600-725°C) for operations. During the pilot plant runs, rod-type electric resistance heating elements with a total rating of 9 kw were installed on the upper exterior walls of the fluorinator.

Prior to exposing the fluorinator to elemental fluorine during actual fluorination of the fused salts, a "conditioning" cycle was performed wherein fluorine was introduced into the vessel which was heated to 20-150°C to induce the formation of NiF<sub>2</sub> "protective" films. Fluorine used in the VPP was obtained in steel tank trailers from the Oak Ridge Gaseous Diffusion Plant (ORGDP) fluorine generating station. A flowing stream sample analyzed by ORGDP personnel indicated the analysis of the fluorine was 95%  $F_2$ , < 5% HF,

R. M. Evans (ed.) Oak Ridge National Laboratory, Reactor Materials Specifications, TID-7017, pp. 117-128 (October 29, 1958).

- 6 -



Fig. 1. Mark I Volatility Pilot Plant Fluorinator.

- 7 -

	Phase I Unit Operations Runs "N	Phase II (" Runs (1-48) "C	Phase III " Runs (1-15)	Total
Temperature; max (°C)	600-700	600-725	600-725	600-725
Thermal cycles (room temperature to 600-725°C)	\$ 50	<b>5</b> 20	10	∽ 20
Time of exposure at temperature (salts molten-hr)	~ 90 (~ 30 with N <sub>S</sub> sparge) (~ 60 withouf N <sub>S</sub> spe	c 4145 .rge)	715	~ 1250
Salt composition (nominal mole %)	NaF-ZrF $_{\rm h}$ (50-50)	NaF-ZrF <sub>4</sub> (50-50)	NaF-ZrF <sub>4</sub> -UF <sub>4</sub> (48-48-4)	
Conditioning fluorine input (liters) <sup>a</sup>	None	35 in 14 hr	530 in 0.5 hr	565 in 14.5 hr
Operations fluorine,input (liters)	None	16 775 in 10 hr	40 830 in 51 hr	57 500 in 61 hr (7-33 liters/min)
UF <sub>6</sub> exposure (hr)	None	None	~ 20 hr	~ 20 hr
<sup>a</sup> These operat film of nickel flu	ions were done at 20-150 oride on the walls of th	"C for the purpose e fluorinator.	of inducing an ini	tial "protective"
PAn average c reduce the final v Cman Plance	of 3:1 mole ratio (F2:U) tranium concentration in	beyond theoretical the salt to a few	requirements was u parts per million.	sed in order to
Top Ilange I dSalts were u amounts of corrosi Studies with Nonra (May 12, 1958).	emoved, ~ > nr. sed previously in unirre on products as shown bel dive Salt, Volatility Pi	diated loop studie ow. Ref: C. L. W lot Plant Runs, C-	s and therefore con hitmarsh, <u>A Series</u> <u>9 Through C-15</u> , p.	tained significant of Seven Flowsheet 10, CF-58-5-113
Component: ( 0.002-3.4 wt % Si.	0.08-0.18 wt % Ni, 0.06-0	.10 wt % Cr, 0.01-	0.02 wt % Fe, 0.01-	0.60 wt % Ti.

Process Conditions for Mark I Volatility Pilot Plant Fluorina. (Unit Operations, Volatility Pilot Plant "M" and "C" Runs) Table I.

- 8 -



Fig. 2. Volatility Pilot Plant Flowsheet.



Fig. 3. Interior Piping, Gas Dispersion Assembly, and Placement of an Early Group of Corrosion Specimens in the Mark I VPP Fluorinator.



and  $1-2\% N_2$  and/or  $O_2$ . Prior to use of the fluorine in the VPP, the gas was passed through a fixed NaF pellet bed at approximately ambient temperatures. Under these conditions, the hydrogen fluoride content in the fluorine was lowered to approx 20 ppm. (ref 7)

After conditioning, the system was purged with commercial grade nitrogen dried to < 1 ppm  $H_2^{0}$ . The nitrogen contained approx 100 ppm  $O_2^{0}$  which was not removed. The fluorinator was heated to approx 600°C along with the salt freeze valve and salt inlet line. The latter two components were heated by autoresistance. Then a batch of fluoride salt was melted in the charge melt tank and drained by gravity flow into the fluorinator.

Fluorine was bubbled through the molten salt to convert any  $\rm UF_4$  in the salt to volatile  $\rm UF_6$ . During fluorination, the vessel operated with approx 25% of its volume filled with about 50 liters of fused salts. The remaining 75% of the volume contained variable quantities of fluorine, uranium hexafluoride, nitrogen, and various metal fluorides of high or intermediate volatility. During the process demonstration "C" runs, an average mole ratio of 3:1 (F<sub>2</sub>:U) beyond theoretical requirements was used in order to reduce the final uranium concentration in the salt to a few parts per million.

While the vessel wall in the salt-containing region of the fluorination vessel reached temperatures of 600-725°C, the upper vapor region remained at lower temperatures. The maximum temperature recorded on a thermocouple attached to the exterior wall of the fluorinator 12 in. down from the slip-on flange was 500°C. The average temperature in this same region was about 400°C.

After completion of fluorination, the waste salt left in the fluorinator was pressure transferred through a freeze valve into a waste container; and the gas from the fluorinator was passed through an Inconel trap, containing either nickel mesh or NaF pellets, which was maintained at approx 400°C. Prior to Run C-9, the trap contained nickel mesh for the purpose of collecting  $\text{ZrF}_4$ , "snow," and thereafter the unit contained NaF pellets to trap entrained salt, chromium, and zirconium fluorides. During and after Run C-9, the trap was termed a "CRP" or complexible radioactive products trap.

<sup>7</sup>F. W. Miles and W. H. Carr, <u>Engineering Evaluation of Volatility Pilot</u> <u>Plant Equipment</u>, CF-60-7-65, Section 15, p. 228.

- 11 -

Downstream from the Snow-CRP trap, the product stream was diverted through absorbers containing NaF at 65-150°C to absorb the UF<sub>6</sub>. The unabsorbed gas, mostly fluorine, was routed through a chemical trap (a NaF bed at ambient temperature) to retain any residual UF<sub>6</sub> and subsequently through a KOH gas disposal unit to neutralize the fluorine before being exhausted to the atmosphere. The product, UF<sub>6</sub>, was desorbed from the absorber bed by heating it to approx 400°C in a fluorine atmosphere and then passed through two cold traps maintained at -40°C and -55°C where the UF<sub>6</sub> condensed. The cold traps were isolated from the rest of the fluorination system and heated to approx 80°C to liquate the UF<sub>6</sub> which drained into a heated product cylinder.

#### C. Reaction to Environment

Ultrasonic-thickness measurements of the fluorinator were made with an "Audigage," an ultrasonic-thickness measurement device, after the Unit Operation's preliminary fluorination equipment studies. No detectable metal losses could be found in either the shell of the vessel or in the bottom head; this could be expected because no fluorine, uranium-bearing salts, or  $\rm UF_6$  was present during the short period of operation at elevated temperature and whatever attack occurred was so slight as to be undetected by the measuring equipment.

#### 1. Chemistry

During VPP Run C-6, a study was made of the interior deposits which formed on the wall of the fluorinator. Figure 4 shows the location and subsequent chemical analyses of these deposits. These data indicate a tendency for chromium, presumably from impure feed salts, and uranium to collect in the middle vapor region of the vessel. The values shown for nickel indicate that extensive corrosive attack had occurred in the system during operations.

After completion of the "M" and "C" runs described in Table I, the Mark I fluorinator was turned over to Metallurgy for corrosion evaluation. Figure 5 shows the interior of the fluorinator after retirement. Most of the interior walls of the vessel below the molten salt levels were free of surface UNCLASSIFIED ORNL-LR-DWG 49156

ANALYSES OF DEPOSITS FROM VPP MARK-I FLUORINATOR AFTER RUN C-6



Fig. 4. Analyses of Deposits from Mark I VPP Fluorinator After Run C-6.

- 13 -





- 14 -

deposits but the regions above the interfaces were covered with heavy scale and corrosion products. A solid ring of material, about 1 in. in cross section, was present on the interior of the vessel wall at about the same elevation as the exterior furnace seal. This was a few inches above the average elevation of the vapor-salt interfaces. Samples of some of these interior deposits were submitted for chemical analyses and identification by x-ray diffraction. The results are shown in Table II.

Table II. The Oak Ridge National Laboratory Analyses of Scale from the Volatility Pilot Plant Mark I Fluorinator after Run C-15<sup>a</sup>

		Con	ponent	t, wt	%		Approx Composition Indicated by X-ray
Origin of Sample	U	Na	Ni	Cr	Zr	F	Diffraction Intensities
Underside of Inconel slip-on flange	1.95	0.78	45.54	0.79	0.72	39.60	) 90% NiF <sub>2</sub> 10% NaF·NiF <sub>2</sub> ·2ZrF <sub>4</sub>
From A Nickel F <sub>2</sub> inlet tube, approx 21 in. below slip-on flange	0.98	5.10	33.76	0.09	0.64	41.15	5 60% NiF <sub>2</sub> 30% NaF·NiF <sub>2</sub> ·2ZrF <sub>4</sub> 10% β <sub>2</sub> ·2NaF·ZrF <sub>4</sub>
From A Nickel F <sub>2</sub> inlet tube, at vapor-salt interf	0.13 ace	6.64	8.36	0.02	1.18	43.30	) -

<sup>a</sup>C. L. Whitmarsh, <u>A Series of Seven Flowsheet Studies with Nonradive Salt</u>, Volatility Pilot Plant Runs, C-9 Through C-15, p.14, CF-58-5-113 (May 12, 1958).

Most of the salt deposits were removed by washing the interior of the vessel with a mixture of 0.7  $\underline{M}$  H<sub>2</sub>O<sub>2</sub>, 1.8  $\underline{M}$  KOH, and 0.4  $\underline{M}$  Na<sub>2</sub>C<sub>4</sub>H<sub>4</sub>O<sub>6</sub> at room temperature, aided by hand chipping. After cleaning, another visual inspection was made and the results are given as follows:

Region	Results
Vapor	Smooth, etched appearance near the top of the vessel with iso- lated, shallow pits. A yellow-to-green deposit encircled the vessel from a point approx 10 in. down to a point approx 20 in. from the top. Several small areas of flaking and scaling were noted at approx 16 in. from the top in the deposit zone. The area from 20 in. down to approx 24 in. from the top had a bluish cast and was rougher in texture than the top section.
Vapor-salt interface	Smooth metallic appearance with distinct indentations encircling the vessel at several levels.
Salt	Smooth metallic appearance. Flange-to-vessel weld not noticeably corroded.

In the middle vapor region, a tightly adherent, yellow-to-green deposit remained on the wall of the fluorinator. Samples of this deposit, surface, and subsurface millings, were removed and submitted for chemical analyses. Figure 6 details the results which indicate that chromium which had previously been found to collect in the upper vapor region of the fluorinator had penetrated into the vessel wall to some depth greater than 10 mils. This chromium concentration gradient was found both in the upper and middle vapor regions although higher concentrations were found in the former region. No excessive quantities of sulfur over that present in the base material were found.

#### 2. Dimensional Analysis

Micrometer measurements were taken in the three major regions of the fluorinator in all quadrants and show the greatest wall-thickness losses to be concentrated in the vapor region of the vessel shell. Figure 7 shows a schematic drawing of the vessel and denotes the sections that were removed from the vessel for these measurements and for metallographic study. The loss data are given in Table III. A full-length vessel section was removed from the northeast-by-east quadrant and micrometer measurements taken every vertical inch to establish a corrosion wall-thickness-loss profile. Figure 8 shows this plot and pinpoints the maximum metal loss of 47 mils as approx 12 in. below the bottom of the slip-on flange. UNCLASSIFIED ORNL-LR-DWG 49157



Analyses of Scale and Millings From VPP Mark I Nickel Fluorinator After Run C-15 and Vessel Decontamination. Fig. 6.



Fig. 7. Schematic of Mark I VPP Fluorinator Showing Areas Removed for Metallographic Examination and Micrometer Measurements.

- 18 -

Table III. Wall-Thickness Losses<sup>a</sup> on Coupons Removed from the Mark I Volatility Pilot Plant L Nickel Fluorinator<sup>b</sup>

	Location				
0 7 -	Elevation			Wal	l-Thickness
Number	(Inches below slip-on flange)	Quadrant	Region		Loss (mils)
N-l	12	North	Vapor		38
E-1	12	East	Vapor		29
S-l	12	South	Vapor		33
W1	12	West	Vapor		47
N-2	35	North	Vapor-salt	interface	6
E-2	35	East	Vapor-salt	interface	24
<b>S-</b> 2	35	South	Vapor-salt	interface	8
W-2	35	West	Vapor-salt	interface	10
N <b>-</b> 3	45	North	Salt		10
E-3	45	East	Salt		8
<b>S-</b> 3	45	South	Salt		11
W <b>-</b> 3	45	West	Salt		15

a By micrometer measurement.

<sup>b</sup>Original wall thickness 250 mils.



Fig. 8. Corrosion Profile and Typical Microstructures from the Mark I VPP Fluorinator. (Profile based on micrometer measurements from northeast quadrant full-length shell section.)

#### 3. Metallographic Study

Based on the corrosion wall-thickness-loss profile plot, areas were selected from the full-length section of the fluorinator and from previously trepanned samples for metallographic study. The location of these areas is shown in Fig. 8. In the as-polished condition, some slight roughening of the surfaces of the samples was noted. No grain-boundary modifications such as is common to intergranular corrosive attack could be found.

After etching with an 0.5% HCl, approx 60% HC<sub>2</sub>H<sub>3</sub>O<sub>2</sub> and approx 40% HNO<sub>3</sub> mixture, the grain boundaries of the interior surface samples appeared darkened below the exposed surfaces to depths varying from 4 to 25 mils. Figure 8 illustrates the typical structures found at varying elevations on the interior surfaces of the fluorinator wall. The deepest penetrations were found on samples from the middle vapor region of the vessel, the vapor-salt interface, and the salt region of the fluorinator. The exterior surface samples also showed intergranular attack varying from 3 to 8 mils in depth.

Widely variant grain sizes were found in the metallographic samples examined. A summary of these sizes, converted to average ASTM grain-size numbers, is shown in Table IV.

Distance down from slip-on flange (in.)	Region	ASTM Grain size number
1	Vapor	56
9	Vapor	2-3
12-1/2	Vapor	> 1
20	Vapor	1
24	Vapor	2
29-1/2	Vapor	5
35	Vapor-salt interface	5
43	Salt	5–6

Table IV. Summary of Grain Sizes in Samples Removed from the Mark I Volatility Pilot Plant Fluorinator<sup>a</sup>

<sup>a</sup>Grain sizes from interior wall and exterior wall samples were approximately equal. At a later date, in an attempt to determine the reasons for the variant grain size, diffractometer traces were obtained on selected samples removed from the wall of the Mark I fluorinator. Samples were taken from the 1-, 12-1/2-, and 43-in. levels, respectively, below the bottom of the vessel slip-on flange. The maximum amount of residual strain was noted for the 1-in. sample with little, if any, evidence of recrystallization having taken place during the Mark I operations. A very limited amount of recrystallization appeared to have occurred in the 12-1/2-in. sample since there seemed only slightly less strain present in this sample when compared to the one above. The 43-in.-sample traces showed considerably less residual strain present than either of the other samples. The indications were that partial, if not complete, recrystallization had taken place in the lower portion of the fluorinator wall.

Figure 9 shows a section through the girth weld which was located in the salt phase of the fluorinator. After etching, the specimen showed no corrosive attack at low magnification. At higher magnification, the weld deposit showed a grain-boundary attack similar to that found in the base metal, but to a lesser degree of severity.

Early work on L nickel corrosion rods placed in the Mark I fluorinator indicated sulfur contamination was probably a factor in the corrosive attack of the vessel.<sup>8</sup> In view of the lack of evidence from chemical analyses, other attempts were made to prove the presence of sulfur at the grain boundaries of interior wall specimens from the Mark I fluorinator.

The use of sulfur print papers did not provide positive evidence of the existence of sulfur compounds at the mating surfaces of the grains. One sample from the middle vapor region of the fluorinator did present the colors described as needed for the identification of nickel sulfide. Figure 10 shows the grain-boundary deposit at the 12-in. level below the bottom of the slip-on flange after etching with cyanide-persulfate and partial repolishing to show the deposit to its best advantage. However, duplicate results could not be obtained with the method.

<sup>8</sup>L. R. Trotter and E. E. Hoffman, <u>Progress Report on Volatility Pilot</u> <u>Plant Corrosion Problems to April 21, 1957</u>, ORNL-2495 (September 30, 1958).



Exterior (contact with air)

> Fig. 9. Macrostructure of Section Through Girth Weld 47 in. Below Top Flange of Mark I VPP Fluorinator. Etchant: Acetic, nitric, hydrochloric acid. 10 X.

Interior (contact with salt phase)



Fig. 10. Photomicrograph of Sample from Interior Surface of Mark I VPP L Nickel Fluorinator 12 in. Below Slip-on Flange (Vapor Phase) Showing Grain Boundary Deposit. Deposit was pale yellow under nonpolarized illumination and black under polarized light in accordance with Hall's method for identifying nickel sulfide deposits. Etchant: Potassium cyanide-ammonium persulfate, partially repolished. 1000X. Reference: A. M. Hall, "Sulfides in Nickel and Nickel Alloys," Trans. Met. Soc. AIME 152, 283, 1943.

Sulfur in almost any chemical form in contact with nickel at high temperatures will result in the formation of a low-melting nickel-nickel sulfide eutectic primarily along grain boundaries, leading to embrittlement of the material.<sup>9</sup> Consequently, samples of the fluorination vessel wall were bend tested. The samples did not show brittle behavior.

Two other techniques were considered as identification methods for the grain-boundary deposits described. One was the use of a microchisel, presently under development by the Metallography Group of the Metallurgy Division, by which some of the grain-boundary deposit could be removed and subsequently submitted for x-ray diffraction analysis. The very small size of the grain-boundary deposits prevented the microchisel's usage in this situation. The second technique considered was the use of an electron probe microanalyzer which could possibly identify a small portion of the deposit by x-ray diffraction analysis, <u>in situ</u>. Such an instrument is not yet available at ORNL and it was not possible to obtain service time on the few instruments currently in operation in this country. Consequently, the nature of the Mark I fluorinator grain-boundary deposits was left in doubt.

#### 4. Summary of Corrosive Attack

Table V summarizes the maximum corrosion losses of all types found in the three major regions of the VPP Mark I fluorinator. The maximum attack was calculated to be 46 mils/month based on exposure time to molten salts during the VPP "M" runs (1-48) and "C" runs (1-15) or 1.2 mils/hr based on fluorine sparge time during fluorination of molten salts. The maximum attack occurred in the middle vapor region.

#### D. Discussion of Results

## 1. Individual Actions of F<sub>2</sub>, UF<sub>6</sub>, and Fused Fluoride Salts

Major corrosive agents in contact with the VPP L nickel fluorination vessel, Mark I, were elemental fluorine, uranium hexafluoride, and

<sup>&</sup>lt;sup>9</sup>W. A. Mudge, "Nickel and Nickel-Copper, Nickel-Manganese, and Related High-Nickel Alloys," <u>The Corrosion Handbook</u>, (ed. by H. H. Uhlig), p. 679, John Wiley and Sons, Inc., New York, 1948.

Quadrant	
ole V. Summary of Maximum Corrosive Attack in Each Major Region and	of the Mark I Volatility Pilot Plant L Nickel Fluorinator
Tat	

	ur.			Thtergr	anular		Total Rate	Losses <sup>b</sup>
Flevation			Mall	Penetra	ation	Total	mils/	
(inches below slip-on			thickness Loss <sup>a</sup>	Interior wall	Exterior wall	Corrosive Attack	month (Molten	mils/hr (F <sub>2</sub> sparge
flange)	Quadrant	Region	(mils)	(mils)	(mils)	(mils)	salt time) <sup>C</sup>	<sup>c</sup> time)d
12	North	Vapor	38	54	m	65	<b>Т</b> +Ј	1.1
12	East	Vapor	29	20	4	53	33	0.9
12	South	Vapor	33	23	C)	59	37	1.0
12	West	Vapor	747	51	ſſ	73	94	1.2
35	North	Vapor-salt interface	9	18	2	31	20	0•5
35	East	Vapor-salt interface	4	17	9	27	17	0.4
35	South	Vapor-salt interfac <b>e</b>	ω	53	ſ	36	23	0.6
35	West	Vapor-salt interface	10	17	ω	35	55	0.6
45	North	Salt	10	25	ſ	38	54	0.6
45	East	Salt	ω	21	4	33	51	0.5
45	South	Salt	11	25	4	01	25	0.7
45	West	Salt	15	22	۔ ۲	42	26	0•7

<sup>a</sup>By micrometer measurement.

 $^{\rm C}{\rm Based}$  on molten salt residence time during VPP "M" (1-48) runs and "C" (1-15) runs. b<sub>l</sub>ncludes exterior intergranular penetration.

dBased on fluorine sparge time during fluorination of molten salts.

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- 26 -

molten fluoride salts, generally of the NaF-ZrF<sub>4</sub>-UF<sub>4</sub> type. The compatibility of each of these agents in contact with metals has received increased attention during the past decade, but a composite system has not been studied extensively.

Fluorine, a most active element, was known to react with virtually every metal under suitable conditions. Resistance to further attack was felt to be imparted by passive fluoride films which form on materials.<sup>10,11,12</sup> For nickel, the only known binary compound with fluorine was found to be nickelous fluoride, NiF<sub>2</sub>.<sup>(ref 13)</sup> The melting point of NiF<sub>2</sub> had been reported to be about 1000°C, well in excess of the operating temperatures (600-725°C) of the fluorinator.<sup>14</sup> The vapor pressure of NiF<sub>2</sub>, approx 1 x 10<sup>-5</sup> mm Hg at 650°C, appeared sufficiently low so that little volatilization of the protective film would occur.<sup>15</sup>

Recent experiments at the Argonne National Laboratory had indicated that the relative amounts of fluorine consumed by a nickel vessel and the change in rate-law behavior with temperature can be represented as shown in Fig. 11. At lower temperatures, 300 to 400°C, a logarithmic rate law appeared to hold, but at higher temperatures, 500 to 600°C, a parabolic behavior seemed prevalent.

<sup>10</sup> M. J. Steindler and R. C. Vogel, <u>Corrosion of Materials in the Presence</u> of Fluorine at Elevated Temperatures, Argonne National Laboratory Report, ANL-5662 (January, 1957).

<sup>11</sup>C. Slesser and S. R. Schram, <u>Preparation</u>, <u>Properties</u>, and <u>Technology of</u> <u>Fluorine and Organic Fluoro Compounds</u>, <u>National Nuclear Energy Series</u>, Div. VII, Vol. 1, pp.157, 173, McGraw-Hill, New York, 1951.

<sup>12</sup>E. J. Barber and H. A. Bernhardt, K-1421 (April 9, 1959)(classified).

<sup>13</sup><sub>H</sub>. J. Emeleus, <u>Fluorine Chemistry</u> (ed. by J. H. Simons), <u>1</u>, 7, Academic Press, Inc., New York, 1950.

<sup>14</sup>Laurence L. Quill (ed.), <u>Chemistry and Metallurgy of Miscellaneous</u> <u>Materials: Thermodynamics</u>, National Nuclear Energy Series, Div. IV, Vol. 19b, p.202, McGraw-Hill, New York, 1950.

<sup>15</sup>M. Faber, R. T. Meyer, and J. L. Margrave, "The Vapor Pressure of Nickel Fluoride," J. Phys. Chem. 62, 883 (1948).



Fig. 11. Consumption of Fluorine by a Nickel Vessel. Reference: R. K. Steunenberg, L. Seiden, and H. E. Griffin, "The Reaction of Fluorine with Nickel Surfaces," <u>Argonne National Laboratory Chemical Engineering</u> <u>Division Summary Report</u>, July, August, September, 1958, ANL-5924, pp. 42-43.

UNCLASSIFIED ORNL-LR-DWG 55786

P.

More detailed work in this field has been reported by ORGDP.<sup>16</sup> Nickel was found to form a continuous, adherent fluoride film with an undiluted fluorine atmosphere at temperatures up to about 980°C. Electron microscopy indicated that the nickel fluoride films had few flaws in the individual crystals which would permit direct access of the fluoride to the nickel surface underneath. The studies also indicated that, as the nickel fluoride film thickened, more resistance to attack was obtained. However, considerable intergranular attack of the metallic nickel was noted at temperatures of approx 815 and 980°C (Fig. 12). The depth of intergranular penetration was estimated to be 5 to 8 times the average attack as calculated from scale formation. The report indicated that the primary mechanism of attack appeared to be diffusion along the crystal boundaries.

The second major corrosive agent placed in contact with the Mark I fluorination vessel during operations was UF<sub>6</sub>. The effect of this compound on metals had undergone investigation in connection with the design, construction, and operation of the ORGDP.<sup>17</sup> Some recent work at the same site indicated that on A nickel samples which were coated with nickel fluoride films of 37 000 and 7<sup>4</sup> 000 A, the average penetration of the nickel by UF<sub>6</sub> at about 815°C, calculated from the average nickel fluoride scale formation, appeared to be about one third of that experienced with fluorine at about 700°C. (ref 16) Later work by the same group indicated that at times one order of magnitude greater (hundreds of hours versus tens of hours) NiF<sub>2</sub> is lost by vaporization and/or a reaction process so that catastrophic attack can occur by additional UF<sub>6</sub> contact.<sup>18</sup>

The nickel exposed to the  ${\rm UF}_6$  at those elevated temperatures exhibited a grain-boundary attack beneath the fluoride scale quite similar to

<sup>17</sup>J. J. Katz and E. Rabinowitch, The Chemistry of Uranium, National Nuclear Energy Series, Div. VIII, Vol. 5, pp. 445-46, McGraw-Hill, New York, 1951.

<sup>&</sup>lt;sup>16</sup>C. F. Hale, E. J. Barber, H. A. Bernhardt, and Karl E. Rapp, <u>High</u> <u>Temperature Corrosion Study, Interim Report for the Period November, 1958</u>, <u>Through May, 1959</u>, KL-498 (July 28, 1959).

<sup>&</sup>lt;sup>18</sup> E. J. Barber, Technical Division, ORGDP, Jan. 7, 1960, Private communication.



Fig. 12. Microstructures of A Nickel Coupons (a) As Received; (b) After Exposure to  $F_2$  at 815°C for 4 hr and for 32 hr at Lower Temperatures; (c) After Exposure to  $F_2$  at 980°C for 35 hr and for 350 hr at Lower Temperatures. Etchant: Soldium cyanide, ammonium persulfate. 100X. Reference: C. F. Hale, E. J. Barber, H. A. Bernhardt, and Karl E. Rapp, <u>High Temperature Corrosion</u> Study, Interim Report for the Period, November, 1958, Through May, 1959, KL-498 (July 28, 1959).

- 30 -
that experienced in the exposures to fluorine. It was also observed that both nickel fluoride and  $\text{UF}_6$  corrosion products had appreciable vapor pressures at those test temperatures and were observed to migrate to the cooler portions of the reactor by vapor-phase transfer.

In addition to  $F_2$  and  $UF_6$ , fused-fluoride salts were also in contact with the Mark I VPP fluorinator. At ORNL, considerable progress has been made in determining material compatibility in various fused-fluoridesalt systems through the Aircraft Reactor Experiment and Molten-Salt Reactor Projects.<sup>19</sup> Nickel-base alloys were found to be, in general, superior to other commercial alloys for the containment of fluoride salt mixtures under dynamic flow conditions.

### 2. Collective Attack During Fluorination

The fluorination cycle of the fluoride volatility process produced greater corrosive attack by the collective system of  $F_2$ , UF<sub>6</sub>, and fused fluoride salts on nickel than had been reported for the individual constituents. During VPP operations, the Mark I L-nickel fluorinator displayed maximum corrosion rate losses of 1.2 mils/hr, based on  $F_2$  sparge time, or 46 mils/month, based on molten-salt residence time during VPP "M" runs (1-48) and "C" runs (1-15). These rates include wall thickness or metal losses as determined by dimensional analysis plus intergranular penetration as determined by metallographic examination. The rates are generally consistent with early bench-scale work on the volatility process.<sup>20</sup>

For convenience in reporting, the proposed reasons for the high corrosive attack on the fluorination vessel will be discussed under four major headings: (a) Interior Bulk Losses, (b) Interior Intergranular Attack, (c) Exterior Intergranular Attack, and (d) Grain-Size Variations.

<sup>19</sup>W. D. Manly et al., "Construction Materials for Molten-Salt Reactors," Fluid Fuel Reactors, (ed. by Lane, MacPherson, and Maslan) Chap. 13, Addison-Wesley, Reading, Mass., 1958.

<sup>20</sup>G. I. Cathers, "Fluoride Volatility Process for High Alloy Fuels," Symposium on the Reprocessing of Irradiated Fuels Held at Brussels, Belgium, May 20-25, 1957, TID-7534, pp. 571-572.

#### a. Interior Bulk Losses

The first portion of this section described "conditioning" treatments whereby nickel fluoride was induced on the walls of the Mark I fluorinator prior to actual high-temperature operations. This was in harmony with prevailing generalizations concerning passive fluoride films, which, when formed on exposed metal surfaces may inhibit further attack by elemental fluorine. However, based upon ORGDP and Argonne National Laboratory work, 16,21 it appears that passivation temperature should have been equal to, or greater than, the operating temperature, rather than 20-150°C, in order to induce greater film thicknesses. The work by Hale et al., indicated that while corrosive attack on nickel by fluorine did not cease after the nickel fluoride film thickened, additional resistance to attack was obtained at test temperatures up to 980°C. Considering the very high rate losses found on samples removed from the wall of the first VPP fluorinator, it would appear that conditions were present in the vessel which (1) did not allow sufficient film thickening to occur, (2) reduced the protectivity of thickened nickel fluoride films, or (3) permitted catastrophic losses of the nickel fluoride films.

Free fluorine was present, periodically, during all of the VPP operations in quantities above those amounts necessary to oxidize any  $UF_{l_1}$  in the fluoride salt mixtures to  $UF_6$ . Thus, even though catastrophic losses of old nickel fluoride films might occur, new films would have opportunity to form. Therefore, a continuous cycle of initial loss, reformation, and secondary loss of the nickel fluoride films forming on the walls of the fluorinator is proposed as the method whereby the large losses of bulk metal occurred.

This proposed loss cycle could be initiated and maintained in several ways depending on the region of the fluorination vessel under consideration. In the salt region, fused fluoride salt baths could dissolve the

- 32 -

<sup>&</sup>lt;sup>21</sup>R. K. Steunenberg, L. Seiden, and H. E. Griffin, "The Reaction of Fluorine with Nickel Surfaces," <u>Argonne National Laboratory Chemical Engineering</u> <u>Division, Summary Report, July, August, September, 1958, ANL-5924, pp. 42-43</u>.

nickel fluoride films until saturation of the baths with nickel fluoride occurred. It has been reported that NiF<sub>2</sub> is soluble in equimolar NaF-ZrF<sub>4</sub> to the extent of 1.8 wt % at 700°C. (ref 22)

At the vapor-salt interface, similar dissolution could have occurred to remove protective wall films. Also, a washing action caused by the fluorine sparge agitation and the rise and fall of the bath level could aid film removal.

In the vapor region, where maximum metal losses occurred, nickel fluoride films that have limited plasticity at the lower operating temperatures might be lost through cracking or rupturing. Also, the difference in linear coefficients of thermal expansion of nickel and nickel fluoride would be sufficient to exaggerate the spalling tendency. These actions may also occur in the other two major regions of the fluorination vessel, but the other loss mechanisms described for those areas would probably predominate.

Cathers has suggested an additional mode of film loss for the vapor region, i.e., dissolution of nickel fluoride in very highly corrosive liquids which condense in the cooler zones of the fluorinator.<sup>23</sup> Fluoride compounds of intermediate volatility such as those containing high-valent chromium, sulfur, titanium, and/or silicon, might be responsible. This condensate is pictured as being extremely variable in composition (possibly including HF and water vapor, if they were unintentionally admitted into the system) and in dissolution ability. After the condensate forms in a relatively cold region of the vessel, it would run down the wall toward a higher temperature zone, where it could then dissolve the nickel fluoride surface films. Progressing further down the walls of the vessel, it would eventually reach a temperature zone where it could reflux, and leave with the product stream or re-enter into the vessel corrosion environment.

<sup>22</sup>C. L. Whitmarsh, <u>Uranium Recovery from Sodium-Zirconium Fluoride-Salt</u> <u>Mixtures, Volatility Pilot Plant Runs L-1 Through L-9</u>, CF-59-9-2 (September 30, 1959).

 $^{23}$ G. I. Cathers, ORNL Chem. Tech. Div., Private communication.

Support is given to the Cathers' suggestion in view of the design used for the VPP fluorination vessel and the results of the chemical studies on the interior wall deposits from the vapor region of the vessel. The Mark I fluorinator heating system was designed to produce cooler temperatures in the upper regions of the vessel. The maximum temperature recorded on a thermocouple attached to the exterior vessel wall 12 in. below the slip-on flange was 500°C, while the average temperature in the same region was 400°C. The thought was that a cooler top vessel zone would permit deposition of low volatility compounds that might entrap uranium products. After some indefinite time of deposition, it was felt that these layers would gradually fall back into the salt baths and become available for further fluorination.

In support of the condensable corrosive liquid theory, chemical analyses of residual wall deposits and millings from the fluorinator vessel walls in the middle vapor region (Figs. 4 and 6) showed high concentrations of uranium and chromium. Thus, it would seem that entrainment of uranium and redeposition of materials containing uranium did occur and that chromium was a part of the entrapping agent or agents.

Rather than describing the agents which seem to have aided the corrosion progress in the middle vapor region as condensable liquids which dissolve nickel fluoride, favor is given to the idea that the nickel fluoride complexed with other more volatile fluorides, the combination of which have lower melting points than nickel fluoride. It has been reported that nickel fluoride complexes easily and that many combinations are known.<sup>24</sup> A complex of NiF<sub>2</sub> and high valent chromium fluoride appears to be a particularly good possibility.

Of interest in discussing the wall-thickness losses on the Mark I fluorinator is the region slightly above the salt-vapor interface where very low losses were found. In this same region, a ring of deposited material

- 34 -

<sup>&</sup>lt;sup>24</sup>H. J. Emeleus, Fluorine Chemistry (ed. by J. H. Simons), <u>1</u>, 7, Academic Press, Inc., New York, 1950.

was found after operations. Unfortunately, chemical analyses were not obtained on the deposit. However, the fact that this thick ring was present, probably continuously during operations, is believed to have prevented interactions with the nickel fluoride "protective" surface films underneath and thereby prevented high losses in that region. The reason for the formation and continued presence of the ring is associated with the low temperatures that predominated in this region. The location is between the furnace windings and the tubular-type heating elements attached to the upper portion of the vessel. Also, the presence of the furnace seal on the exterior of the fluorinator at this same elevation probably affected temperature by acting as a sink for conducted heat in this region.

## b. Interior Intergranular Attack

In the "as-polished" condition, interior surface samples from the Mark I fluorinator showed no sign of intergranular corrosive attack. After etching with a strong acid solution, a mixture of 0.5% HCl, approx 40% HNO<sub>3</sub>, and approx 60% HC<sub>2</sub>H<sub>3</sub>O<sub>2</sub>, the surface grain boundaries to depths of from 4-25 mils were attacked more severely and rapidly than grain boundaries closer to the interior of the specimens. Using a mild etchant, KCN-(NH<sub>4</sub>)<sub>2</sub>S<sub>2</sub>O<sub>8</sub>, and partial repolishing, a pale yellow deposit was observed at the mating surfaces of the grains. The appearance of the intergranular attack on the fluorinator walls was distinctly different than that produced in nickel by fluorine or UF<sub>6</sub> (with an initial nickel fluoride film) and reported by Hale et al.<sup>25</sup>

The formation of grain-boundary compounds or other changes in grain-boundary regions, such as those observed, and which are generally categorized as intergranular corrosion attack, often result in the loss or "sloughing" of entire grains from the exposed surface of the material. However, metallographic examinations of specimens from the fluorinator revealed relatively smooth surfaces.

<sup>25</sup>C. F. Hale, E. J. Barber, H. A. Bernhardt, and Karl E. Rapp, <u>High</u> Temperature Corrosion Study, Interim Report for the Period November, 1958, Through May, 1959, KL-498 (July 28, 1959). Some circumstantial evidence was available to indicate that sulfur contamination produced the grain-boundary modifications observed. In addition to previously reported work<sup>26</sup> where microstructures of nickel specimens purposely embrittled with sulfur were compared with early VPP L nickel corrosion specimens, a communication from ORGDP, which supplied the VPP with process fluorine, indicated that as much as 200 ppm of sulfur as sulfuryl fluoride had occasionally been detected in their manufactured fluorine.<sup>27</sup> Contamination from the commercial sodium fluoride used to strip hydrogen fluoride and water from the process fluorine could also be a source of sulfur. Sulfur could also be introduced into the system from impure feed salts or from trace quantities contained in corrosion test specimens.

Attempts to prove the presence of sulfur in the fluorinator wall samples in quantities greater than that present in the base metal by using sulfur print papers, chemical analyses of surface millings and wall deposits, bend tests, and various metallographic techniques either met with failure or produced inconclusive results. This was particularly frustrating in view of early reports that free Ni<sub>3</sub>S<sub>2</sub> had been identified in nickel microscopically when 0.05 and 0.005 wt % S, respectively, were present.<sup>28,29</sup> Examination of the nickel-sulfur constitution diagram indicated that Ni<sub>3</sub>S<sub>2</sub> would be the compound to seek in identifying low percentages of sulfur contamination in nickel.<sup>30</sup> It was considered that chromium and/or other system contaminants may have complexed the Ni<sub>3</sub>S<sub>2</sub> and thus prevented identification by some of the fluorinator samples, if sulfur were the major agent responsible for the intergranular attack, more positive indications should have been found.

<sup>26</sup>L. R. Trotter and E. E. Hoffman, Progress Report on Volatility Pilot Plant Corrosion Problems to April 21, 1957, ORNL-2495 (September 30, 1958).

<sup>27</sup>H. J. Culbert, Process Engineering Division, ORGDP, Private communication. <sup>28</sup>G. Masing and L. Koch, Z. <u>Metallkunde</u> 19, 278-279 (1927).

<sup>29</sup>P. D. Merica and R. G. Waltenberg, <u>Trans. Met. Soc. AIME</u> 71 709-716 (1925); National Bureau of Standards Technical Paper 281, 155-182 (1925).

<sup>30</sup>M. Hansen and K. Anderko, <u>Constitution of Binary Alloys</u>, p. 1035, McGraw-Hill, New York, 1958. The nature of the grain-boundary deposits may not be determined until they are studied by an electron probe microanalyzer or some other precise tool. Nevertheless, because of the potential effect on corrosion and the cumulative and irreversible embrittling tendencies of the nickel-nickel sulfide eutectic, reduction of sulfur in VPP nickel fluorinator's environments to the lowest possible levels is essential.

# c. Exterior Intergranular Attack

As described, the exterior of the Mark I fluorinator shell also suffered intergranular attack, but to a lesser degree than the interior. Penetration on the exterior surfaces varied from 3-8 mils. The appearance of this attack at high magnification, the exterior environment, and the presence of NiO<sub>2</sub> found on the second VPP fluorinator operated under similar conditions, indicate typical high-temperature intergranular oxidation of nickel.

## d. Grain-Size Variations

Metallographic examination of samples removed from the shell of the Mark I fluorinator disclosed widely variant grain sizes. Small, uniform grains of 5-6 average ASTM grain-size number prevailed in the salt and saltvapor interface regions, while much larger grains were observed in most of the vapor region. The largest grains found in the vapor region were at an elevation of approx 12 in. below the bottom of the slip-on flange. The average ASTM grain-size number at the 12 in. elevation was greater than 1. Diffractometer traces were obtained on vessel samples removed from the 1-, 12-1/2-, and 43-in. levels in order to compare the residual strain remaining in the samples. The results indicated approximately no recrystallization had occurred at the 1-in. level, only a very limited amount of recrystallization took place at the 12-1/2-in. level, and partial, if not complete, recrystallization occurred in the salt-phase sample.

The classical factors which determine crystallite growth upon heating are amount of prior cold deformation, annealing temperature, and annealing time. Generally, larger amounts of cold deformation provide for

- 37 -

smaller resultant grain sizes after isothermal and constant time annealing. Higher temperatures and, to a lesser degree, longer times result in larger grain sizes on material containing some fixed amount of cold work.

In many materials a small amount of prior cold deformation results in the production of exaggerated grains after annealing at ordinary times and temperatures. In these cases, the term "critical strain" has been given to the quantity of cold work originally introduced. The mechanisms involved here have been studied and reported in detail.<sup>31</sup>

The VPP Mark I fluorinator was fabricated from L nickel by converting a flat, annealed, 1/4-in. plate into a 14-in.-diam right cylinder by roll forming. During forming, cold deformation was induced in the outermost fibers of the shell to a calculated maximum of approx 2%. This amount of cold work is at the right level to be termed "critical strain" for most metals and presumably was present along the entire length of the fluorinator shell. However, exaggerated grain sizes were observed only in a portion of the vapor phase of the vessel after pilot plant operations. Therefore, an additional variable, rate of heating after prior deformation, has been suggested as the most influential factor in producing the grain sizes found in the Mark I fluorinator.<sup>32</sup>

The first time the fluorinator was heated was during preliminary fluorination equipment studies. During those cycles, the lower 23 in. of the vessel was surrounded by a high heat-flux 30-kw electricresistance furnace which raised the temperature of the enclosed portion of the vessel to approx 450°C. No external heat or insulation was in close proximity with the upper portion of the fluorinator. Later cycles during the same preliminary studies were done with a glass-lined heating mantle covering the top portions of the fluorinator. Temperatures in the lower regions of the vessel reached 600-700°C, while temperatures in the upper regions of the vessel were much lower.

<sup>&</sup>lt;sup>31</sup>J. E. Burke, "The Fundamentals of Recrystallization and Grain Growth," <u>Grain Control in Industrial Metallurgy</u>, American Society for Metals, Cleveland, Ohio, 1949.

<sup>&</sup>lt;sup>32</sup>L. K. Jetter and C. J. McHargue, ORNL Metallurgy Division, Private communication.

In view of the initial Mark I heating cycles, and confirmed by the diffractometer traces, the salt region of the fluorinator appears to have experienced thermal levels where recovery and at least partial recrystallization occurred. The recrystallization temperature for nickel varies from approx 600°C for A nickel, 99.4 wt % (Ni + Co) to 370-470°C for various purities of electrolytic nickel, 99.9<sup>+</sup> wt % (Ni + Co) after 50% reduction by cold rolling and annealing for 1 hr at the temperature indicated.  $^{33}$  Jetter and McHargue have suggested that during the initial heating the rapid rate of heating did not allow complete relief of internal stresses which at higher temperatures served as nucleation sites for recrystallization. The rate of appearance of these nuclei is known to increase with time, exponentially with temperature, and with increasing amounts of prior cold deformation.<sup>31</sup> After nucleation, stable nucleus growth occurred and the presence of so many grains growing in a fixed volume limited the salt region grain size. Later heat cycling of the fluorinator was done at approximately the same temperature levels as the highest original heating (600-725°C) so that probably little, if any, grain-boundary migration occurred after recrystallization to increase the grain size in the lower portions of the vessel.

The top half of the Mark I fluorinator sustained lower temperatures during the first heat cycles as the result of heat losses by radiation and convection from the upper portions of the vessel wall. A steady state temperature of < 200°C at the 12-in. level has been estimated.<sup>34</sup> The upper portions of the vessel also experienced sluggish heating rates when compared to that portion of the vessel enclosed by the heating furnace. It has been suggested that the latter influenced the grain size in the vapor region by permitting more complete recovery and resulting in few nucleation sites.

During later operations in the pilot plant, rod-type electric heating elements with a total rating of 9 kw were in contact with the vapor

<sup>33</sup>E. M. Wise and R. H. Shafer, "The Properties of Pure Nickel - I, II, III," <u>Metals and Alloys</u>, <u>16</u> Sept., p. 424, Nov., p. 891, Dec., p. 1067, 1942.

<sup>34</sup>S. H. Stainker, ORNL Chemical Technology Division, Private communication.

region exterior surfaces of the fluorinator. A thermocouple at approximately the 12-in. level indicated average temperatures of 400°C and a maximum of 500°C. Such thermal levels permitted only a small amount of recrystallization to occur, as indicated by the diffractometer studies. An extremely coarse grain size resulted from the described conditions.

Examination of samples removed from the Mark I fluorinator revealed that the metal-loss profile and grain-size profile closely paralleled one another except for the region just below the top flange of the vessel. Because of the evidence of corrosion by intergranular attack during exposure to the volatility fluorination environment, a causal relationship is suggested. Thus, loss of a large grain by effectively losing grain-boundary material until the grain sloughs off would mean greater losses than for the same procedure happening in fine-grain regions. However, sloughing of grains was not observed and only a slight widening of the boundaries was noted at the junction of the boundaries and the exposed metal surfaces. Also, the depth of intergranular penetration in coarse-grain regions was about the same as in most of the fine-grain regions. The conclusion is that metal loss and grain size do not seem to be interrelated in the corrosion of the Mark I fluorinator.

However, for two reasons, the authors recommend using finegrained material for fluorination vessels. One, a higher strength level will be achieved in nickel, a material not noted for superior strength; and two, one variable will be removed in a system replete with variables. Fine-grained material can be provided by modification of the specifications of purchased stock and by revising the annealing cycle after fabrication.<sup>35</sup>

#### II. Mark II Volatility Pilot Plant L Nickel Fluorinator

#### A. Material Selection and Fabrication-Design Changes

While L nickel seemed deficient in its corrosion resistance for longtime service, no other commercially available material had, at that time, proved

<sup>35</sup>"Annealing of Nickel, Monel, and Inconel," <u>Tech. Bull. T-20</u>, The International Nickel Company, Inc., New York, April, 1953. itself on the basis of scouting or other tests to be worthy of immediate substitution as a fluorinator material of construction. Moreover, the heavy vapor-phase attack on the Mark I fluorinator was still under investigation, so that the use of the same construction material seemed especially appropriate to ascertain more about this corrosion problem. Consequently, the Mark II fluorinator was fabricated in the ORNL shops from the same heat of 1/4-in.thick L nickel as was used in the previous pilot plant vessel. Analogous fabrication techniques were used. Figure 13 shows the second VPP fluorinator, the vessel furnace, and the complexible radioactive products (CRP) trap.

Certain design changes were incorporated in the second pilot plant fluorinator. In order to provide for remote handling and to allow desirable upward flow through the CRP trap, its location was changed from the original side position shown in Fig. 1 to an elevation completely above the Mark II fluorinator. Other process design changes included some modification of the draft tube assembly, repositioning of the off-gas line, and installation of a small port in the top blind flange of the fluorinator for observation and entry.<sup>36</sup>

B. Operational History

The Mark II fluorinator was placed in service at the beginning of the "E" runs during which time fuel used in the Aircraft Reactor Experiment was reprocessed.<sup>37</sup> The use of the vessel was continued during the "L" (spiked) runs and during the gas-entrainment studies, M-62 through M-64. The process history, summarized in Table VI, has been divided into three phases for convenience in reporting the Vidigage inspections on the vessel at the completion of Runs L-4 and L-9.

The "L" or spiked runs were made in the VPP to extend process development data and used a nonirradiated salt with fully enriched uranium.  $^{38}$  Some

<sup>36</sup>F. W. Miles and W. H. Carr, Engineering Evaluation of Volatility Pilot Plant Equipment, CF-60-7-65, pp. 43-45.

<sup>37</sup>W. H. Carr, <u>Volatility Processing of the ARE Fuel</u>, CF-58-11-60 (November 14, 1958).

<sup>38</sup>C. L. Whitmarsh, Uranium Recovery from Sodium Zirconium Fluoride Salt Mixtures, Volatility Pilot Plant Runs L-1 Through L-9, CF-59-9-2 (September 30, 1959).



Fig. 13. Mark II VPP Fluorinator and CRP Trap.

	игг т, цо апо	M (0			
	Phase I Runs E-1 Through E-6; L-1 Through L-4	Phase II Runs L-5 Through L-9	Phase III Runs M-62 Through M-64	Total	
Temperature, °C Thermal cvcles	540-730 10	550-700 4	600-690 17	540-730 31	
Time of exposure at temper-	~ 900	~ 310	~ 725	~ 1935	
avure, sarus muruen-mr Fused salt, nominal mole %	El-E6 NaF-ZrF <sub>4</sub> -UF4 (c, d) (48-49.5-2.5) Ll-L4 NaF-ZrF,-UF, (e,f)	NaF-ZrF <sub>4</sub> -UF <sub>4</sub> (g) (52-45.5-2.5)	NaF-ZrF <sub>4</sub> (50-50)		
Conditioning fluorine <sub>a</sub> input, standard liters <sup>a</sup>	(54-41-5) 4535 in 18 hr (2.8-6.9 standard liters/min)	750 in 3 hr (2.5-7 standard liters/min)	none	5285 in 21 hr (2.5-7 standard 11ters/min)	
Operating fluorine input, standard liters <sup>b</sup>	44 470 in 74 hr (1.6-20.2 standard liters/min)	16 000 in 18 hr	none	60 470 in 92 hr (1.6-20.2 standard liters/min)	- 43 -
UF <sub>6</sub> exposure, hr	~ 25	~ 10	none	~ 35	
aThese operations were nickel fluoride on the walls	done at 20-150°C for the of the fluorinator.	purpose of inducing	an initial "r	protective" film of	
<sup>D</sup> An average of approx 3 used in order to reduce the	:l excess F2 over that qu final uranium concentrati	antity necessary fo on in the salt to a	r theoretical few parts per	requirements was million.	
CUranium was fully enri 0.031-0.056 wt % Cr, 0.017-C	ched. Feed salts contain .138 wt % Ni, 0.033-0.078	ed the following range $\psi t \ \% \ Fe$ , 0.0015-0.	nges of impuri 080 wt % Si, <	ties: Component: c 0.001- < 0.005 wt % M	Mo.
dC. L. Whitmarsh, <u>Repro</u> (May, 1959) and <u>Reprocessing</u>	cessing of ARE Fuel, Vola of ARE Fuel, Volatility	tility Filot Plant, Filot Plant, Runs E	Runs E-l and E-6, CF	2, CF-59-5-108, 59-8-73 (August, 1959	.(63
<sup>e</sup> Ten milligrams of PuF <sub>4</sub> Component: 0.0183-0.0345 wt < 0.001- < 0.0052 wt % Mo, $\varepsilon$	added in Run L-3-4. Fee % Cr, 0.035-0.236 wt % N nd ~ 0.010 wt % Ti for L-	d salts contained t i, 0.0133-0.030 wt 1,-3, and -5 only.	he following 1 % Fe, < 0.001-	anges of impurities: 0.005 wt % Si,	
<sup>f</sup> C. L. Whitmarsh, <u>Urani</u> Plant, Runs L-1 Through <u>L-9</u> ,	um Recovery from Sodium-Z CF-59-9-2 (September 30,	irconium Fluoride S 1959).	alt Mixtures,	Volatility Pilot	
<sup>g</sup> Uranium was fully enri Same feed salt impurities ar	ched. One gram of $PuF_{l\mu}$ a d reference as detailed i	dded in Run L-6; 10 n (d) above.	grams of PuFl	. added in Run L-8.	

43

of the runs were spiked with high burnup salt, thus the name of this series. During three of the "L" runs, selective additions of  $PuF_4$  were added to the feed salt so as to gather information on the behavior of plutonium in the volatility process.<sup>39</sup> Previous experiments had indicated that some volatilization of  $PuF_6$  would occur. However, greater than 99% of the plutonium remained in the salt in the fluorinator during the "L" runs.

Several months after the "L" runs were completed, the Mark II fluorinator was used to study gas-phase entrainment problems in the VPP.<sup>40</sup> These runs, M-62 through M-64, investigated  $\text{ZrF}_4$  solid condensation in the vapor phase. Barren equimolar NaF-ZrF<sub>4</sub> salt was used in these studies and fluorine was not present.

#### C. Reaction to Environment

## 1. Visual and Vidigage Inspections

During the first portion of the "L" runs, 1 through 4, persistent line plugging occurred. Presumably this was due to nickel contamination in the feed salts plus corrosion losses which allowed the concentration of nickel fluoride in the salt to exceed the solubility limit. After run L-4, the plant was shut down and approx 200 lb of a high-melting salt complex was found in the bottom of the Mark II fluorinator. See Fig. 14. Petrographic and x-ray diffraction analyses showed the mass to contain 20-30 wt % NiF<sub>2</sub> and 50-60 wt % NaF-NiF<sub>2</sub>-2ZrF<sub>4</sub>. In order to continue using the vessel, the frozen salt was chipped out with a pneumatic hammer.

Results of a visual inspection of the Mark II VPP fluorinator after exposure to "E" and L-1 through L-4 runs and cleanup operations are cited:

Vessel walls Free of salt on the interior. Exterior surfaces were covered with a dull gray film. Those portions of the wall near the girth weld had scars on the interior surfaces presumably caused by glancing blows of the pneumatichammer blade.

<sup>40</sup>J. B. Ruch, Volatility: Fluorinator Design FV-100, Zr-U Fuel Element Processing Phase, CF-59-5-89.

<sup>&</sup>lt;sup>39</sup>R. A. Cross and C. L. Whitmarsh, <u>Plutonium Behavior in the Fluoride</u> Volatility Process, CF-59-9-5.



Fig. 14. Interior of Mark II VPP Fluorinator After Run L-4. (Fluorine inlet cut off prior to taking of photograph.)

- Longitudinal Interior surfaces were sharply defined by corrosive attack. seam weld Corrosion of the weld metal (INCO 61) appeared to have been as severe or more severe than that of the wall (L nickel).
- Girth weld Interior surfaces were sharply defined by corrosive attack. Corrosion of the weld metal (INCO 61) appeared to have been as severe or more severe than that on the wall or the dished head.
- Dished head Interior surfaces exhibited many scars on sides and bottom of the head, presumably caused by the pneumatic-hammer blade. Exterior surfaces were bulged approx 1/16 in. in about six places corresponding to the deeper internal scars.

Vidigage (ultrasonic thickness) measurements were made to assess the damages during salt removal plus the wall-thickness losses that had occurred during previous process cycling. A maximum wall-thickness loss of 68 mils was found in the salt-containing region of the fluorinator. Complete results of the Vidigage examination are shown in Table VII along with other data.

Despite the high-corrosion losses indicated, a decision was made to use the vessel for an additional five runs so as to complete the scheduled process development studies. No plugging difficulties were encountered during the subsequent "L" runs 5 through 9. This was accomplished by diluting the feed salt with an equal amount of nickel-free barren salt.

Figure 15 shows the interior of the Mark II fluorinator after run L-9. The results of a visual inspection after hand chipping and decontamination were as follows:

Vessel walls	Free of salt on the interior. Vapor region appeared rough and pitted and had an adherent green deposit in the region 10-20 in. below the slip-on flange. Exterior surfaces were covered with a dull gray film. The scars caused by the pneumatic-hammer blows after Run L-4 were still present.
Longitudinal seam weld	Interior surfaces were more sharply defined by corrosive at- tack than previously noted. Corrosion of the weld metal appeared to have been more severe than that of the base metal.
Girth weld	Interior surfaces were more sharply defined by corrosive at- tack than previously noted. Corrosion of the weld metal appeared more severe than that of the base metal.
Dished head	Interior and exterior scars from previous chipping operations were still present.

		S.W. Qui	adrant	N.W. Qua	drant	N.E. Qua	drant		S.E. Que	adrant	
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le/	Jə	0.226	ı	0.224	0.224	0.221	0.218	0.228	0.230	0.230	0.229
33	դս Հ—	0.210	ı	0.207	0.206	0.210	0.204	1	0.226	0.222	0.220
35	I ~	ł	0.236	0.196	0.236	0.198	0.196	0.202	0.210	0.210	0.210
37		0.198	0.230	0.196	0.230	0.192	0.212	0.198	0.206	0.195	0.194
39	·	0.194	0.227	0.198	0.234	0.189	0.215	0.193	0.194	0.192	0.187
+ 		0.194	0.224	0.204	0.242	0.187	0.210	0.189	0.102	701.0	0.187
5 5	Γ¤ι	0.206	0.227	0.206	0.248	0.184	0.202	0.186	0.184	0.190	0.186
10 10	<u> </u>	0.210	0.242	0.208	1	0.188	0.204	0.186	0.184	007.0	0.132
1	~	0.238	0.246	0.240	0.252	0.192	0,204	0.198	0.194	0.196	0.196
aAccura	acy o	f reading	s estimat	ed to be	± 0.003 ±	n.					a a substantia da companya

Table VII. Comparison of Vidigage Thickness Readings, in Inches, on the Volatility Pilot Plant Mark II L Nickel Fluorinator<sup>a</sup>

- 47 -



Decontamination was done by alternate washings using a mixture of 0.7  $\underline{M}$  H<sub>2</sub>O<sub>2</sub>, 1.8  $\underline{M}$  KOH, and 0.4  $\underline{M}$  Na<sub>2</sub>C<sub>4</sub>H<sub>4</sub>O<sub>6</sub> at room temperature and 4  $\underline{M}$  (NF<sub>4</sub>)<sub>2</sub>C<sub>2</sub>O<sub>4</sub> at 100°C. At that time Vidigage readings were obtained and are reported in Table VII, along with the previous readings obtained after Run L-4. No significant additional wall-thickness losses were found.

The corrosion conditions established during the gas-entrainment studies, Runs M-62 through M-64, which were done at a later date using the Mark II fluorinator, were not felt to be sufficiently serious to warrant another complete Vidigage inspection.

## 2. Chemistry

Samples of the dull gray film present on the exterior wall of the Mark II fluorinator after Runs L-4 and L-9 were analyzed by x-ray diffraction techniques to be  $\text{NiO}_2$ . In addition, samples of the adherent green deposit which remained on the interior wall of the vessel in the vapor region were submitted for x-ray diffraction analyses. The presence of NiF<sub>2</sub>, NiO, and ZrO<sub>2</sub> was noted.

Surface and subsurface millings were taken from the interior wall of the vessel in the region where the green deposit seemed thickest, 15-17 in. below the bottom of the slip-on flange, and submitted for additional chemical analyses. For comparison purposes, subsurface millings were also taken from the exterior vessel wall at the same elevation. The results of the metal chemistry are shown in Fig. 16, and indicate an increase in chromium concentration in the millings near the surface of the interior wall of the vessel. No significant increase in sulfur in the vapor region of the fluorinator over that quantity present in the base material was found. However, on specimens removed from the salt region of the Mark II fluorinator, which were later analyzed by Battelle Memorial Institute (BMI), an increase in the sulfur content of the interior wall specimens was noted. Figure 16 also shows these results. UNCLASSIFIED ORNL-LR-DWG 49159



Fig. 16. Analyses of Millings from Mark II VPP L Nickel Fluorinator After Run L-9 and Vessel Decontamination. •

## 3. Dimensional Analysis

After decontamination and other cleanup operations, samples were trepanned from the Mark II fluorinator and sent to BMI for dimensional analysis and metallographic examination. The location of these coupons is displayed in Fig. 17. In addition, a full-length section from D-E area, southeast quadrant, was removed and micrometer measurements taken at ORNL to plot a complete corrosion profile of wall-thickness loss. This plot is shown in Fig. 18 and incorporates BMI micrometer measurements on the trepanned samples from the other quadrants of the fluorinator. For comparison, Vidigage readings of the D-E area after runs L-4 and L-9 are also plotted in Fig. 18. The maximum wall-thickness losses found were in the salt region of Mark II fluorinator. In a region 40-43 in. below the bottom of the slipon flange of the vessel, a wall-thickness loss of 82 mils was recorded. Fair correlation between the Vidigage readings and the micrometer measurements was noted.

# 4. Metallographic Study

A metallographic study was made on the trepanned samples from the Mark II fluorinator by personnel of the BMI Corrosion Research Division.<sup>41</sup> At low magnification, 20X, the surfaces from the top vapor (T) sections showed a light etch and some outline of grain boundaries. The middle vapor region specimens (la and lb) were etched to a greater extent and were covered with a thin green crystalline deposit. Previous chemical analyses of this deposit at ORNL indicated that the scale contained NiF<sub>2</sub>, NiO, and ZrO<sub>2</sub>. The vapor-salt interface (2a and 2b) and salt region specimens (3a and 3b) seemed to have been severely etched, and, in the latter, areas of intergranular attack could be seen.

Typical photomicrographs of sections from the Mark II fluorinator can be seen in Figs. 19 through 23. Intergranular attack was prevalent in all

- 51 -

<sup>&</sup>lt;sup>41</sup>Letter Report from F. W. Fink, Battelle Memorial Institute, to R. P. Milford, ORNL Subcontract No. 988 (October 7, 1959).



Fig. 17. Location of Specimens Removed from the Mark II VPP L Nickel Fluorinator for Dimensional Analysis and Metallographic Study.

- 52 -



Fig. 18. Corrosion Profile of Mark II VPP L Nickel Fluorinator.



Fig. 19. Microstructure of Sample from Interior Surface of Mark II VPP Fluorinator 3 in. Below Slip-on Flange (Vapor Phase, Area A) Etchant: Nitric-acetic acid. 100X.



Fig. 20. Microstructure of Sample from Interior Surface of Mark II VPP Fluorinator 19 in. Below Slip-on Flange (Vapor Phase, Area A). Etchant: Nitric-acetic acid. 100X.



Fig. 21. Microstructure of Sample from Interior Surface of Mark II VPP Fluorinator 34 in. Below Slip-on Flange (Vapor-Salt Interface, Area A). Etchant: Nitric-acetic acid. 100X.



Fig. 22. Microstructure of Sample from Interior Surface of Mark II VPP Fluorinator 42 in. Below Slip-on Flange (Salt Phase, Area C). Etchant: Nitric-acetic acid. 100X.



Fig. 23. Microstructure of Sample from Interior Surface of Mark II VPP Fluorinator Approx 47 in. Below Slip-on Flange (Salt Phase, Area A) Vicinity of Girth Shell to Bottom Head Weld. Etchant: Nitric-acetic acid. 100 X.

sections in all regions but penetration was deepest in the samples from the interface and salt regions. Maximum grain-boundary attack was found to be 33 mils in the salt-phase sample A-3b which was in the vicinity of the shell to head girth weld.

Some slight variation in grain sizes was found in the metallographic samples. The largest grains were in the salt region of the fluorinator. Table VIII summarizes the grain-size data in terms of average ASTM grain-size number.

Location (inches down from bottom of slip-on ASIM grainflange) Region size number Vapor 3 3-5 15 Vapor 3-4 19 3-4 Vapor 29 Vapor-salt interface 3-5 34 Vapor-salt interface 3-4 14S Salt 2-4 47 Salt 2-4

Table VIII. Summary of Grain Sizes in Samples Removed from the Mark II Volatility Pilot Plant L Nickel Fluorinator<sup>a</sup>

<sup>a</sup>Grain sizes from interior wall and exterior wall samples were approximately equal.

A pitting attack was found on the interior wall near Area C in the vapor region, about 3 in. down from the bottom of the slip-on flange. Figure 24 shows the appearance of the surface at 3X, and Fig.25 shows a metallographic section through this pitted area. The latter indicates a level of intergranular penetration similar to that found in other upper-vapor regions. The depth of intergranular attack did not appear to vary from top to bottom of the pits.

Unclassified BMI N59673



Fig. 2<sup>1</sup>. Photograph of Sample from Interior Surface of Mark II VPP Fluorinator 3 in. Below Slip-on Flange (Vapor Phase, Area C) Showing Pitting-Type Attack. 3X.



Fig. 25. Microstructure of Sample from Interior Surface of Mark II VPP Fluorinator 3 in. Below Slip-on Flange (Vapor Phase, Area C) in Region of Pitting-Type Attack. Etchant: Nitric-acetic acid. 100X. In the salt region of Area C, a few small intergranular cracks were found on the interior of the vessel wall. A typical crack is shown in the unetched condition in Fig. 26 and in the etched condition in Fig. 22. Very pronounced darkening of the grain boundaries to approximately the same depth as the crack was evident after etching.

Cross sections from the fluorinator shell longitudinal weld are shown in Fig. 27 (Samples B-T and A-3b) which depicts sections in the vapor and salt phases, respectively. The weld joint from the salt region shows what appears to be an increased attack at the weld root. This confirms the visual examinations of the Mark II fluorinator. Examination of the weld from the salt section at high magnification revealed pronounced grain-boundary darkening after etching to a maximum depth of 10 mils. Similar, deeper grainboundary darkening occurred in the base metal adjacent to the weld metal, extending to a maximum depth of 33 mils.

The exterior of the vessel displayed a corrosive attack which appeared to be intergranular in nature. Penetration varied from 1-6 mils, the maximum occurring at or below the vapor-salt interface (Fig. 28, Sample A-3b). However, there was one exception to this pattern. A grain-boundary penetration of approx 12 mils in depth was found on the exterior surface opposite the pitted region in the vapor near Area C.

## 5. Summary of Corrosive Attack

Table IX summarizes the corrosion losses of all types found in the three major regions of the Mark II VPP L nickel fluorinator. The maximum attack was calculated to be 60 mils/month based on exposure to molten salts during the VPP "E" runs (1--6) and "L" runs (1--9) or 1.1 mils/hr based on fluorine sparge time during fluorination of molten salts. The maximum attack occurred in the salt region.

#### D. Discussion of Results

The Mark II VPP L nickel fluorinator displayed a maximum corrosion attack during the described VPP runs of 1.1 mils/hr, based on  $F_2$  sparge time during fluorination, or 60 mils/month, based on molten-salt residence time





Fig. 26. Photomicrograph of Sample from Interior Surface of Mark II VPP Fluorinator <sup>1</sup>42 in. Below Slip-on Flange (Salt Phase, Area C) Showing Crack Easily Visible Before Sectioning. As-polished. 100X.

Unclassified BMI N59674



Fig. 27. Photomacrographs of Longitudinal Weld Sections Through Wall of Mark II VPP Fluorinator (a) From Vapor Phase 3 in. Below Slip-on Flange, (b) From Salt Phase 47 in. Below Slip-on Flange. Etchant: Nitric-acetic acid. 5X.

## Unclassified BMI C618



Fig. 28. Microstructure of Sample from Exterior Surface of Mark II VPP Fluorinator Approx 47 in. Below Slip-on Flange (Salt Phase, Area A) Showing Result of Air Oxidation. Etchant: Nitric-acetic acid. 100X.

co <b>nve</b> rted t time <sup>b</sup>	mils/hr <sup>d</sup> (F <sub>2</sub> sparge time)	0.1	0.4	0.4	0°3	0.4	0.8	0.8	0.75	1.7	6.0	0.75	0.1
Total losses to mils/uni	mils/month <sup>c</sup> (molter salt time)	5	20	22	18	23	7t5	46	Ω;∔Ω	60	50	L,4	54
Total Total	Corrosive attack (mils)	œ	33	36	30	39	75	2.2	69	100	84	69	66
nular ation	Exterior wall (mils)	-1	-t-	m	CU	Ŋ	9	4		CJ	10	1	с
Intergra Penetr	Interior Wall (mils)	m	2	ω	Ø	Ō	9T	77	16	16	33	16	23
TTBW	Thickness loss (mils)	4	22	25	50	28	53	59	52	82	54	52	64
	Region	Vepor	Vapor	Vapor	Vapor-salt interface	Vapor-salt interface	Vapor-salt interface	Salt	Salt	Salt	Salt	Salt	Salt
Location Elevation	low Quadrant	S.W.	S.W.	S.W.	S.W.	N M	ນ. ພ.	S.W.	N. W.	N E	S.W.	N.W.	M.E.
	(inches be. slip-on flange)	m	15	19	29	29	34	t,	D;D	1;2	7:47	747	h+7

Table IX. Summary of Corrosive Attack in Each Major Region of the Mark II Volatility Filot Plant L Mickel Fluorinator

<sup>a</sup>By micrometer measurement.

bIncludes exterior intergranular penetration.

 $^{\rm C}Based$  on molten salt residence time during VPP E(1-6) runs and L(1-9) runs.

d<sub>B</sub>ased on fluorine sparge time during fluorination of molten salts.

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during VPP "E" runs (1-6) and "L" runs (1-9). Maximum attack occurred in the salt-phase region of this vessel whereas the first VPP fluorinator experienced maximum losses in the vapor-phase region. The rates of attack for both vessels were of the same orders of magnitude. The corrosive attack in the Mark II fluorinator, as was the case for the Mark I vessel, can be categorized into bulk metal losses from the interior wall of the vessel and intergranular attack on both the interior and exterior walls of the unit.

#### 1. Interior Bulk Losses

Bulk metal losses are believed to be the result of continuous loss and reformation of "protective" NiF<sub>2</sub> films on the interior wall of the Mark II fluorinator. As discussed in Section I, the nickel fluoride films formed on the walls of the volatility fluorinators during conditioning and fluorination operations could be removed by three and possibly four methods. These are: removal (1) by rupturing or spalling, (2) by a fluxing action of the fluoride salt baths, (3) by a washing action of the melts, and (4) by dissolution in certain highly corrosive liquids condensable in the cooler regions of the vessel.

Maximum losses occurred in the salt region of the Mark II vessel which seems to indicate that the fluxing action of the fluoride baths was the predominant method of removing the protective films from the walls of the fluorinator. The high vapor regions losses described for the first VPP fluorinator were partially attributed to the presence of corrosive liquids which condensed in the cooler regions of that vessel and some discussion of whether similar liquids were present in the second vessel seems in order.

Evidence of the presence of these condensable liquids in the middle vapor regions of both vessels were (1) the tenacious wall deposits, (2) the segregation of chromium in surface and subsurface layers, and (3) the bulk metal loss maxima. However, the bulk metal loss maximum in the second vessel occurred a few inches below that of the initial fluorinator.

One major dissimilarity found was the lack of uranium segregation in the vapor phase of the Mark II vessel which was in contrast to the behavior of the first fluorinator. This latter fact may suggest the reason for the lesser middle vapor region attack in the second fluorinator when compared with the first vessel. Perhaps the additional operating experience of the VPP personnel plus the modifications in the vessel's interior plumbing and exterior appendages permitted  $\text{UF}_4$  to remain in the salt baths until more or less complete oxidation to volatile  $\text{UF}_6$  had occurred.

The wall-thickness-loss profile of the Mark II fluorinator shows an additional, smaller, maximum at a point 25 in. below the bottom of the slip-on flange. This additional peak in the upper interface region is believed to have been induced by low operational temperatures a few inches below the 25-in. level. There was no direct method of heat at 28 in. below the bottom of the slip-on flange. This region was between the furnace windings used to heat the salt region and the rod-type heating elements used to heat the vapor region. In addition, the furnace seal was present at this point, resulting in a built-in heat sink. Thus, the corrosion profile curve probably would have continued in a smooth curve from the 25-in. elevation on down toward the salt region except for this low-temperature region cited. A similar low-corrosion area was found in the wall of the Mark I fluorinator at approximately the same elevation.

The corrosion losses found in the salt region of the Mark II fluorinator were the maximum found in the system and on the order of twice that noted for the first fluorinator in the same region. Two reasons are proposed for the higher salt-phase losses in the second fluorinator. First, the fluoride salt baths in contact with the lower regions of the second fluorinator contained uranium for a period approx 67% longer than for the first fluorinator. As will be described in a later section, the presence of uranium in fused fluoride salt systems during bench-scale volatility corrosion studies has enhanced corrosive attack. Second, although the total quantity of fluorine sparged during Mark II was only slightly higher than that used during operations with the Mark I vessel, the total time of fluorination for the Mark II vessel was approx 50% longer than for the first fluorinator. Since in both cases an average of 3:1 excess fluorine over that quantity necessary for theoretical reaction was used, fluorine probably had opportunity to remain in the salt longer during Mark II operations. This would allow the corrodent a longer period of time to attack the fluorinator wall.

## 2. Interior Intergranular Penetration

Upon etching samples removed from the wall of the Mark II fluorinator, the grain boundaries appear heavily darkened to various depths. In the vapor region, a maximum depth of 8 mils penetration was noted. The vaporsalt interface region showed a maximum of 16 mils while the salt region displayed penetration proceeding to a maximum depth of 33 mils.

Suspecting that the intergranular attack might be the result of sulfur contamination, personnel at BMI attempted two tests to make positive identification of the grain-boundary deposits. The sulfur-print technique, whereby acidified photographic paper is pressed against the metal surface, was used. Sulfides could not be detected using this method. Next, the re-action of a solution of lead nitrate and nitric acid upon constituents at the grain boundaries was observed under the microscope. The precipitate formed appeared to be similar to but weaker than that observed on a nickel tube known to be grossly contaminated with sulfur. Battelle Memorial Institute also indicated that the structures observed at the grain boundaries resemble those found in L nickel rods tested and studied at ORNL.<sup>1</sup>/42

A section from the Mark II fluorinator wall which had been exposed to the salt phase was analyzed for sulfur content at several depths by EMI personnel, (Fig. 24). Twice the weight percent of sulfur was found in the interior wall at an O to 5-mil depth as that found at the same depth on the exterior wall, 240 vs 120 ppm. Sulfur analyses conducted at ORNL on wall material removed from various depths in the middle vapor region of the vessel (Fig. 16) did not disclose any increase in sulfur content on subsurface interior samples when compared to subsurface exterior samples, 20 vs 40 ppm.

Unfortunately, the poor correlation obtained on sulfur analyses from BMI and ORNL on exterior wall samples plus the weak case for sulfur

<sup>&</sup>lt;sup>142</sup> L. R. Trotter and E. E. Hoffman, Progress Report on Volatility Pilot Plant Corrosion Problems to April 21, 1957, ORNL-2495, pp. 22, 26, 29 (September 30, 1958).

contamination presented by other test methods dilutes any blanket statement which could be made regarding the role that sulfur played in the interior intergranular attack of the Mark II fluorinator. However, the serious embrittling and potential corrosive effects of sulfur in contact with nickel at high temperatures are definite and known facts. Therefore, sulfur contamination should be stringently avoided in any of the VPP's nickel process equipment.

#### 3. Exterior Intergranular Attack

The exterior of the Mark II fluorinator shell also underwent intergranular attack. This was noted by BMI personnel in their examination of samples removed from the vessel. The general depth of the attack, 1-6 mils, was of the same order as that reported for the first VPP fluorination vessel. In the analyses of scale from the exterior wall of the Mark II, NiO<sub>2</sub> was found indicating that the exterior intergranular attack on the second fluorinator was due to air oxidation.

# 4. Grain-Size Variations

Grain sizes found in samples removed from the wall of the Mark T fluorinator varied from an average ASTM number of 5-6 to > 1. The large sizes occurred exclusively in the vapor region of the vessel. The second VPP fluorinator showed a different grain-size pattern. Average ASTM grain-size numbers in this vessel varied from 3-5 to 2-4, the largest occurring in the salt region.

Although the second fluorinator was fabricated from the same heat of L nickel and in a similar manner as the first vessel, initial thermal conditions were quite different in the respective vapor regions. Initial heatup for the Mark I was done without the benefit of an external heat source in the vapor region while the Mark II had rod-type heating elements with a total rating of 9 kw attached to the upper half of the vessel. Heavy refractory insulation covered the rod-type elements.

From a classical metallurgical viewpoint, it appears that many more nucleation sites were present in the vapor region of the second fluorinato

- 70 -

as the result of the initial rapid heatup and more or less incomplete recovery when compared with the first vessel. The many sites affected the production of many grains in constant volume which could only result in comparatively smaller grain sizes.

The somewhat larger grain sizes found in the salt region of the Mark II fluorinator when compared to the same region of the first vessel and to the resulting grain sizes noted in the vapor region of the Mark II seem to be the result of some coalescence during operations. It will be recalled that the first fluorinator was at 600-725°C for approx 1250 hr while the second vessel was at about the same temperature range for over 1900 hr.

The specific effect of grain size on corrosion in the fluorination system has been reported in Section I as being conjectural. However, maximum bulk losses on the walls of both fluorinators occurred at locations where the largest grain sizes predominated. As such, the production and stabilization of small grain sizes in nickel fluorination vessels seem a reasonable precaution. Open annealing cycles for L nickel to ensure small grain sizes have been reported.<sup>43</sup>

#### E. Corrosion of Internal Components from the Mark II VPP Fluorinator

Several internal components of the Mark II fluorinator have been examined and an evaluation of the corrosive attack on these parts made by personnel of the Corrosion Research Division, EMI.<sup>(ref 44)</sup> Figure 3 shows the location of most of these components with respect to the top flange of the fluorinator. Of particular interest are the draft tubes and fluorine inlet tubes which, by virtue of their position in the system, were subjected to initial contact by fluorine, and could be expected to sustain the greatest corrosive attack. Two draft tubes, Mark IIA and Mark IIB, were used in the Mark II fluorinator during Phase I and Phase II of the fluorinator's lifetime, (Table VI). The draft tubes were fabricated at ORNL from A nickel. Results

<sup>43</sup>"Annealing of Nickel, Monel, and Inconel," <u>Tech. Bull. T-20</u>, The International Nickel Company, Inc., New York, April, 1953.

<sup>44</sup>Letter Report from F. W. Fink, Battelle Memorial Institute, to R. P. Milford, ORNL Subcontract No. 988 (October 7, 1959). of the examinations of the draft tubes and fluorine inlet lines are schematically displayed in Figs. 29 and 30 and show the configuration details of the assemblies. Supporting photomicrographs are presented in Figs. 31 and 32.

Considerable metal loss was found on the Mark IIA draft tube, but little visible intergranular attack was noted. Conversely, the Mark IIB draft tube showed little dimensional change but a rather deep intergranular attack. The etching characteristics and general microscopic appearance were also quite different. While the Mark IIB tube etched normally for nickel using a nitric-acetic acid mixture, the other draft tube did not, and it was necessary for EMI personnel to use an HCL-HNO<sub>3</sub> (3:1) etch, which they commonly use for Inconel to develop grain-boundary detail. However, spectrographic analysis at EMI showed the Mark IIB draft tube construction material to be A nickel.<sup>44</sup>

The maximum bulk metal loss on the Mark IIA draft tube by micrometer average measurement was 77 mils and occurred near the upper support plate of the assembly. Comparison of the measured outside diameter of the tube with the original diameter indicated that most of the metal loss occurred on the outer surfaces of the unit. Micrometer measurements on the Mark IIB draft tube showed maximum bulk losses of only 6 mils but a maximum total intergranular penetration on both the interior and exterior wall of 29 mils. The fluorine inlet lines associated with the Mark IIB draft tube assembly showed similar corrosive behavior when compared to the draft tube bodies. However, the Mark IIA fluorine inlet line showed a much more severe intergranular attack on the inside of the pipe when compared with the outside.

A high-probe line and a thermocouple well, both made from 3/8-in. sched-40 A nickel, were inspected also by BMI personnel for corrosive attack. Both components operated at temperatures essentially the same as those of the fluorinator wall. The high-probe line was a high-pressure liquid level and density-probe combination which extended from the top flange of the fluorinator beneath the molten salt level. The line was open at the bottom to allow contact with the fused fluoride salt baths, but pressurized nitrogen, inside the tube, for the most part prevented the salts from entering the tube.



Scale: Approximately 1/2 size

Fig. 29. Corrosion Losses on the Mark IIA Draft Tube and Fluorine Inlet from the Mark II VPP Fluorinator. Considerable metal loss has occurred at all areas but most of the intergranularly attacked grains appear to have sloughed off.



Fig. 30. Corrosion Losses on the Mark IIB Draft Tube and Fluorine Inlet from the Mark II VPP Fluorinator. Metal loss was not as severe as in the case of the Mark IIA assembly and more intergranular penetration was noted.

- 74 -

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Fig. 31. Microstructures of Samples from A Nickel Draft Tubes Used in Mark II VPP Fluorinator. (a) and (b) exterior and interior surfaces of Mark IIA draft tube at Section 21B; (c) and (d) exterior and interior surfaces of Mark IIB draft tube at Section 11F. Etchants: (a) and (b) Hydrochloric-nitric acid, (c) and (d) Nitric-acetic acid. 100X.



Fig. 32. Microstructures of Cross Sections Through A Nickel Fluorine Inlet Tubes from (a) Mark IIA Draft Tube (b) Mark IIB Draft Tube. Etchants: Nitric-acetic acid. 70X.

The thermocouple well was positioned in a similar manner to that described for the probe line but the end was sealed with an A nickel plug. Table X cites the corrosive losses found on the high-probe line and thermowell as well as giving more detail on the losses found in the fluorine inlet lines.

The internal components from the Mark II VPP fluorinator sustained total corrosion losses comparable to the walls of the fluorination vessel. No increased attack was noted on the fluorine inlet lines or on the draft tube bodies as the result of the proximity to elemental fluorine during operations.

Of interest is the indication of much more corrosive conditions present during the L-1 through L-4 runs when compared to E-3 through E-6 or L-5 through L-9 runs. Extended times of service at high temperatures for the early "L" runs may explain these differences. Corrosion control specimens of L nickel in place during the run groups mentioned, and reported in Section IV, corroborate the variable corrosive behavior present during these different operation groups.

#### III. Bench-Scale Fluorination Corrosion Studies

The Volatility Studies Group, Chemical Development Section A of the Chemical Technology Division, has continued to study process chemistry in connection with the volatility process since their early work indicated the latter's feasibility. These studies have included the gathering of corrosion data from small-scale experiments.

As stated in Section I, nickel-base alloys have shown superior corrosion resistance to fused fluoride salts under dynamic flow conditions, and nickel and nickel-base alloys exhibit generally good resistance to elemental fluorine and  $\rm UF_6$ . In this connection, commercial purity nickel and Inconel have been the primary materials of construction for facilities using the individual corrodents mentioned above. Extensive studies at ORNL on Inconel in contact with fused fluoride salts have shown that appreciable quantities of chromium were removed through reaction with UF<sub>h</sub> and other

- 77 -

Specimen	Exposure	stance from Bottom of slip-on Flange (in.)	Nominal Outside I (mil Maximum M	)iamd s) finimum	Intergrar Penetra (mile Inside (	nular clon <sup>e</sup> s) Dutside	Wall Thickness losses <sup>f</sup> (mils)	Total Corrosion (mils)
High-Probe Line								
正形 <b>-</b> 2D 1子 2下	Vapor Interface	ಥ ಸ ೧೧	429 77	673 667	- <del>1</del> M.	4 0 1	M [	107
3F 3F(weld) 3F(tip)	Liquid Liquid Liquid	48.5 49 50.5	696	<b>1</b> 1 663	ан ф	4 1 0 0 1	м <b>т</b>	
Thermocouple Well <sup>t</sup> FF 6F 6F (weld)	Vapor Interface Liquid Liquid	18 18 19 19 19 19 19 19 19 19 19 19 19 19 19	679 679 •	676 677 666	エヤヤキ	4 0 0 0 4 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	HMO I	607 -
Fluorine-Inlet Tut Mark IIB <sup>c</sup> 7F 8F 9F 10F 12F	e Vapor Liquid Liquid Liquid	<b>*</b> 45 *45 %	829 829 829 829 821 829 821 821 821 821 821 821 821 821 821 821	887 856 857 857 857 857 857 857 857 857 857 857	5 - 1 0 0 t	200000000 1111	よの375	1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
Fluorine-Inlet Tut Mark IIA <sup>c</sup> 5A 5B 5C 	e Liquià Liquià Liquià Liquià	~38 ~422 ~455 ~145	756 751 776 789	756 776 789	16 14 14 7	m m m m	1;4 1;4 30 30 30	50700 50700
aletter Report fro bConstructed from cConstructed from dBased on micromet eBased on optical	m F. W. Fin 3/8 insch 1/2 insch cer measuren microscopi(	nk, BMI, to R. hed-40 nickel p md-40 nickel pi ments. c measurements.	P. Milford ipe; nomina pe; nomina	, ORML S' Blo.d., Lo.d., {	ibcontract 675 mils, 340 mils,	No. 983 nominal nominal w	(October 7, wall thickr all thickne	1959). Less 91 mils. Ss, 109 mils.

Table X. Measurements of the High-Probe Line, Thermocouple Well, and Fluorine-Inlet Tubes From the Mark II Volatility Pilot Plant Fluorinator<sup>a</sup>

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fBased on optical microscopic measurements and subtracted from nominal original wall thicknesses.

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- 78 -

oxidizing impurities.<sup>45</sup> The chromium removal was accompanied by the formation of subsurface voids in the metal. Substitutions of molybdenum in an approximate ratio of 2:1 (Mo:Cr) for about half of the chromium content in a typical chromium-containing alloy, plus other modifications, provide an alloy, INOR 8, which has exhibited no measurable attack when in contact with molten fluoride salts at temperatures up to approx 700°C. Also, INOR 8 has been used as the material of construction for the VPP Dissolver-Hydrofluorinator, the major vessel for the head-end cycle of the Volatility Program.

For these reasons, a 2-in.-diam Inconel fluorinator, ten l-in.-diam A nickel fluorinators, and four l-in.-diam INOR 8 fluorinators were tested and subsequently examined for comparative corrosion behavior.

#### A. Inconel Fluorinator

#### 1. Test Method

The Inconel fluorinator was used in hot-cell studies and was fabricated from 0.065-in. stock at ORNL. Prior to the exposure detailed in Table XI, fluorine flowed into the fluorinator as it was raised from ambient

Table XI. Exposure Conditions for the Inconel Fluorinator Vessel

	and the second secon
Temperature, °C	600–800
Time of exposure at temperature, hr with salt molten	187
Thermal cycles	70
Fused salt, nominal mole $\%$	NaF-ZrF <sub>4</sub> -UF <sub>4</sub> (48-48-4) <sup>a</sup>
Fluorine input, standard liters <sup>a</sup>	294 in 49 hr

<sup>a</sup>Uranium irradiated, not enriched.

<sup>45</sup>W. D. Manly et al., "Metallurgical Problems in Molten Fluoride Systems," Progress in Nuclear Energy, Series IV, Vol 2 - Technology, Engineering, and Safety, pp.164-179, Pergamon Press, London, 1960. room temperature to operating temperature. This was done for leak-testing purposes although, at the same time, the interior vessel walls were probably "conditioned" by producing films of fluorides. The total time involved for this testing-conditioning treatment was about 2 hr.

After exposure operations, areas from the vessel were selected for micrometer measurements and metallographic examination. Figure 33 shows a cross section of the vessel and the location of the sample areas removed for study. A summary of corrosive attack found is shown in Table XII where total losses are reported in mils per hour, based on fluorine sparge time, and mils per month, based on residence time in molten salts. Representative photomicrographs are grouped in Fig.  $3^{l_4}$ .

#### 2. Discussion of Results

Maximum corrosive attack in the Inconel fluorinator was encountered at the vapor-salt interface and occurred at a rate of 0.24 mils/hr, based on fluorine sparge time, or 48 mils/month, based on molten-salt residence time. Intergranular penetration seemed the predominant mode of attack in the salt and vapor regions but no evidence of intergranular corrosion was found at the vapor-salt interface.

In this connection, the intergranular penetration seemed dissimilar to that found on the interior walls of the VPP fluorinators. That is, on the Inconel vessel there was evidence of the sloughing of whole grains of material, a condition not observed on the full-size L nickel vessels. The depth of the intergranular penetration on the Inconel vessel was only a single grain deep. The VPP fluorinators demonstrated intergranular modifications many grains in depth on samples removed from corresponding service regions containing similar grain-size material.

In the Inconel bench fluorinator, it appears that after intergranular penetration had proceeded to about one grain in depth, many of the affected grains could not be retained in position by the remainder of the grain-boundary material and sloughed off, leaving new material ready for corrosive attack. This method of metal loss would seem devastating to the vessel

- 80 -



Fig. 33. Cross Section of the Chemical Development Section A Inconel Fluorinator. Metallographic specimens were removed as shown.

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Chemical Development	
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ximum Corrosion Results	Inconel Fluorinator
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Summary	
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Table	

onverted to ait Time	th mils/hr n- (F <sub>2</sub> sparge e) <sup>2</sup> time)	0.06	0.12	0.20	0.24	0.22
Losses Co Mils/U	mils/mon (molten salt time	קר י	S <sup>†</sup> D	40	4,8	7 <u>;</u> 24
	Total Corrosion (mils)	m	רי† -יז	IO	75	TT T
granular tration	Exterior wall (mils)	I	Ъ	1	I	8
Interg Penet	Interior wall (mils)		m	N	ı	CU
Wall	Thickness Loss (mils) <sup>a</sup>	ć	ŝ	ω	12	0
stimateč	Wall Temp (°C)	00T	250	500	600-800	600-300
E	i- Specimen Location	top vapor	upper vapor	lower vapor	vapor-salt interface	salt
	Spec men No.	22 15	TO	Ţ,	CU	-1

<sup>a</sup>Four micrometer measurements taken in each area. <sup>b</sup>Does not include the exterior attack found on the vessel wall.

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INNER SURFACE, SALT AREA

Fig. 34. Typical Microstructures of Samples Removed from Chemical Development Inconel Fluorinator. Etchant: Glyceria regia. 250X. Reduced 32%.

in service, but the anomaly of the situation is that maximum wall-thickness losses occurred in the Inconel vessel at the vapor-salt interface and at that point no evidence of intergranular attack could be found.

No explanation can be given for the interior wall interface anomaly described nor can one be given for the heavy exterior intergranular attack which has localized in a cooler region of this Inconel fluorinator (see Fig. 34).

### B. A Nickel Fluorinators

#### 1. Test Method

In addition to the Inconel fluorinator reported upon in Section IIIA, ten A nickel miniature fluorination vessels, each 1-in.-o.d. and 0.035-in.-wall thickness, have been operated by the Volatility Studies Group, Chemical Development Section A, in order to compare fluorinator corrosion using different process flowsheets. Figure 35 shows the units in test position, while Table XIII is a summary of the imposed test conditions.

The vessels, in all cases, were charged with 75 g of fluoride salt that had been ground and classified to -8 +20 mesh. The vessels were then placed in split tube-type furnaces and brought to the specified temperature under a nitrogen purge of 0.05 liters/min. At temperature, the nitrogen was bypassed and fluorine was allowed to bubble through the salt at the same flow rate. In those cases where uranium was added to the melt, 0.50 g increments of UF<sub>4</sub> were added at intervals of 1 hr or more. As shown, some reactors were placed in series both for convenience and to minimize the fluorine consumption. The total elemental fluorine exposure in each case was 50 hr at a rate of 0.05 liters/min. During these 50-hr exposures on test reactors Nos. 5, 6, and 10, twenty-five UF<sub>4</sub> additions and UF<sub>6</sub> volatilizations were made.

In all tests, the salts were kept molten for over 200 hr while under a nitrogen purge. This was done to facilitate the test procedure, that is, avoid remelting salts for the next fluorine test period. Figures 36 and 37 illustrate the corrosion results obtained by micrometer measurements and

- 84 -



Fig. 35. Apparatus for Comparison of Fluorination Corrosion on A Nickel Miniature Reactors.

Vessel No.	Fluoride Salt	Temperature (°C)	Hr of Molten Salt Exposure With N <sub>2</sub> Sparging	Hr of Molten Salt Exposure With F <sub>2</sub> Sparging	Position in Test Series
1	*	450	222	50	l
4	*	525	238	50	l
7	×	525	240	50	1
8	*	525	240	50	2
2	*	600	222	50	2
5	2 + 0.5% U	600	238	50	2
9	<del>* *</del>	525	240	50	3
10	3 + 0.5% U	525	240	50	1
3	<del>***</del>	600	222	50	3
6	1 + 0.5% U	600	238	50	l

Table XIII. Process Conditions for A Nickel Bench-Scale Test Fluorinators

\*26-37-37 mole % LiF-NaF-ZrF<sub>4</sub>: Composition 31, plus LiF addition. \*\*31-24-45 mole % LiF-NaF-ZrF<sub>4</sub>: Composition 31, plus LiF and ZrF<sub>4</sub>

additions.

\*\*\*50-50 mole % NaF-ZrF<sub>4</sub>: Composition 31, as received.



€.0 0.275 0.250 - INTERGRANULAR ATTACK / METALLOGRAPHIC EXAM. - METAL LOSS/MICROMETER MEASUREMENT 0.225 MAXIMUM LOSS SHOWN IN EACH PHASE V - VAPOR i - VAPOR SALT INTERFACE S - SALT 0.2 LEGEND 0.125 0.150 0.175 mis/hr (BASED ON FLUORINE E XPOSURE TIME) ;; 0 0.075 (SLIGHT PITTING NOTED) 0.050 0.025 0 > -S > S S S S > S S > v. > TEST SALT TEMP NO. NO. (°C) 525 450 525 525 600 600 525 600 600 525 ⊂ % 0:5 % 0.5 % 0.5 % 0.5**%** ∪ N N N ю ---N N ი 9 4 ထ 2 ഹ φ -~ б

Fig. 37. Summary of Corrosion on A Nickel Miniature Fluorinators Using Different Process Flowsheets.

- 88 -

# UNCLASSIFIED ORNL-LR-DWG 49164

metallographic examinations of specimens removed from the vessels. The losses are reported both in mils per hour of fluorine sparge and mils per month based on total residence time in molten salts for comparison with other studies. Figures 38 through 42 show representative photomicrographs from this test series.

2. Discussion of Results

The A nickel fluorinators showed widely varying corrosion rates as anticipated in planning this test series.

Comparison of the results on test vessel No. 6 with those of vessel No. 3 indicated that uranium additions to equimolar NaF-ZrF<sub>4</sub> resulted in increased corrosion occurring as intergranular attack at the vapor-salt interface. A rate of 0.1 mils/hr based on fluorine sparge time was noted at the interface of No. 6.

Comparison of vessel No. 2 with vessel No. 3 indicated that the addition of 26 mole % of LiF caused increased attack at the vapor-salt interface, as evidenced by bulk metal losses. Upon metallographic examination, no evidence of intergranular attack was found in either vessel.

The addition of uranium to the LiF-NaF-ZrF<sub>4</sub> (26-37-37 mole %) salt in vessel No. 5 resulted in significantly increased metal losses plus intergranular attack at the vapor-salt interface. Additional intergranular attack was noted in the vapor phase of this vessel.

Lowering the temperature of the reactor containing the same LiF-bearing salt, LiF-NaF-ZrF<sub>4</sub> (26-37-37 mole %), to 525°C produced erratic corrosion results. Two of the vessels, Nos. 4 and 7, showed uniform, comparatively small metal losses in all regions while vessel No. 8 showed an increased attack, especially at the vapor-salt interface. Intergranular attack was also exhibited by vessel No. 8. As shown in Table XIII, vessel No. 8 was in a downstream position from vessels Nos. 4 and 7, and the deviation in behavior may, therefore, have been the result of carry-over and collection of certain constituents conducive to greater corrosive attack.

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INNER SURFACE, SALT AREA

Fig. 38. Typical Microstructures of Samples from A Nickel Miniature Test Fluorinator No. 1. Etchant: Acetic-nitric-hydrochloric acid. 200X.



INNER SURFACE, SALT AREA

Fig. 39. Typical Microstructures of Samples from A Nickel Miniature Test Fluorinator No. 5. Etchant: Acetic-nitric-hydrochloric acid. 200X.



Fig. 40. Typical Microstructures of Samples from A Nickel Miniature Test Fluorinator No. 6. Etchant: Acetic-nitric-hydrochloric acid. 200X.



INNER SURFACE, SALT AREA

Fig. 41. Typical Microstructures of Samples from A Nickel Miniature Test Fluorinator No. 8. Etchant: Acetic-nitric-hydrochloric acid. 200X.



INNER SURFACE, SALT AREA

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Fig. 42. Typical Microstructures of Samples from A Nickel Miniature Test Fluorinator No. 10. Etchant: Acetic-nitric-hydrochloric acid. 200X. Lowering the process temperature to  $450^{\circ}$ C and utilizing the same lithium-bearing salt (No. 2) reduced corrosive attack to the lowest levels found in this test series. These results were provided by vessel No. 1 where the approximate losses were 0.02 mil/hr, based on fluorine sparge time. This is especially significant since lithium-sodium-zirconium fluoride salt mixtures can be used in the volatility process at lower operating temperatures than the NaF-ZrF<sub>4</sub> composite system because of the lower liquidus line of the lithium-bearing system.

Tests Nos. 10 and 9 at 525°C used higher LiF- and  $\text{ZrF}_4$ -content salts, 31-24-45 mole % LiF-NaF-ZrF<sub>4</sub>, with and without uranium, respectively. Vessel No. 9 showed slightly increased attack over vessels Nos. 4 and 7 which were operated at the same temperature. Metallographic examination of vessel No. 10 revealed that intergranular penetration was present in all phases of the interior wall.

The A nickel miniature fluorinators demonstrated generally lower rates of corrosive attack in simulated fluorination environments when compared to the full-sized L nickel vessels or the latter's A nickel internal components which were exposed to pilot plant fluorination conditions. The following three reasons can probably account for most of this deviation: (1) the temperatures of fluorination generally were somewhat lower in the bench-scale work than during pilot plant operations; (2) somewhat better control over thermal cycling and other process conditions was obtained during bench-scale work; (3) the feed salts used in the pilot plant work were contaminated by having been used in previous loop studies, the ARE, or having been contained for long periods of time in the pilot plant type 347 stainless steel charge melt tank.

The A nickel miniatures also demonstrated greater resistance to corrosive attack during fluorination than the Inconel vessel used in Chemical Development hot-cell studies.

C. INOR-8 Fluorinators

1. Test Method

Four INOR-8 test fluorination vessels, each 1-in.-o.d. and 0.065-in.-wall thickness, and of similar design to the A nickel reactors

reported in Section IIIB were also fabricated at ORNL for bench-scale volatility corrosion studies. The composition of INOR-8 used in test fluorinators was 74.5 wt % Ni-15.3 wt % Mo-6.5 wt % Cr-3.7 wt % Fe-0.02 wt % C.

A summary of the test conditions for the INOR-8 miniature fluorinators, as shown in Table XIV, indicates that only vessel No. 3 contained

Vessel No.	Fluoride Salt	Temperature (°C)	Hr of Molten Salt Exposure With N <sub>2</sub> Sparging	Hr of Molten Salt Exposure With $F_2$ Sparging	Position in Test Series
1	×	450	236	50	l
2	<del>* *</del>	600	236	50	2
3	2 + 0.5% U	450	290	50	1
4	×	600	290	50	l

## Table XIV. Process Conditions for INOR-8 Bench-Scale Test Fluorinators

\*26-37-37 mole % LiF-NaF-ZrF<sub>4</sub>: From addition of LiF to Composition 31 salt.

\*\*50-50 mole % NaF-ZrF $_{
m h}$ : Composition 31, as received.

uranium. Vessel No. 3 received 25  $\mathrm{UF}_4$  additions which were subsequently fluorinated to  $\mathrm{UF}_6$  in like manner as the A nickel miniatures previously reported. Figures 43 and 44 illustrate the corrosion losses obtained by micrometer measurements and metallographic examinations on specimens removed from the walls of the vessel. The losses have been converted to mils lost per unit time for comparison purposes. Figures 45 through 48 show typical etched microstructures found in the INOR-8 fluorinator specimens.

## 2. Discussion of Results

The INOR-8 test fluorinators showed a variety of corrosion manifestations depending on the test conditions. Considering the two vessels which operated at 600°C, vessel No. 2, which contained equimolar NaF-ZrF<sub>4</sub>, showed less attack than vessel No. 4, which contained LiF-NaF-ZrF<sub>4</sub> (26-37-37 mole %).





TEMP (°C)

TEST SALT NO. NO.





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- 98 -

UNCLASSIFIED ORNL-LR-DWG 49166



Fig. 45. Typical Microstructures of Samples from INOR-8 Miniature Test Fluorinator No. 1. Etchant: Modified aqua regia. 200X.

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Fig. 46. Typical Microstructures of Samples from INOR-8 Miniature Test Fluorinator No. 2. Etchant: Modified aqua regia. 200X.



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Fig. 47. Typical Microstructures of Samples from INOR-8 Miniature Test Fluorinator No. 3. Etchant: Modified aqua regia, 200X.



Fig. 48. Typical Microstructures of Samples from INOR-8 Miniature Test Fluorinator No. 4. Etchant: Modified aqua regia. 200X.
Maximum attack in the latter vessel occurred at the vapor-salt interface at a rate of 30 mils/month, based on residence time in the molten salts. This was the maximum attack rate for the entire INOR-8 series.

Reducing the operating temperature to 450°C and using the same lithium-bearing salt significantly reduced attack as indicated for vessel No. 1 (Figs. 43 and 44). The maximum attack found in this vessel was also at the vapor-salt interface at a rate of 5 mils/month. The addition of uranium to the lithium-bearing salt described plus operation at 450°C more than doubled the corrosive attack when compared to the nonuranium-containing salt. This occurred in vessel No. 3.

Examination of the metallographic specimens in the as-polished state, Fig. 49, disclosed spongy layers on the surfaces of the vapor region specimens from vessels Nos. 2 and 4. These two vessels operated at the highest temperatures of the test series. Upon etching with a modified aqua regia solution, 5:1, HC1:HNO<sub>3</sub>, the spongy regions were for the most part destroyed. Figures 46 and 49 display the microstructures for specimens from the two vessels and show that in only one region, the upper vapor area of vessel No. 2 (Fig. 46) did significant amounts of the spongy region remain.

Careful repolishing on the upper vapor area sample from vessel No. 2 and examination of the surface disclosed a corrosion product which resolved itself into two distinct layers, as shown in Fig. 50. Just above the INOR-8 base metal was a spongy region which was composed of voids and solid metal intermixed. Many of the voids appeared to be partially filled with nonmetallic-appearing compounds. Above the spongy region, on the outermost surface of the specimen, was an irregular layer that had the angular appearance of metal crystals.

Upon etching the specimen with a mixture of 10% KCN and 10% (NH<sub>4</sub>)<sub>2</sub>S<sub>2</sub>O<sub>8</sub>, 1:1 ratio in water, the outermost layer of the surface showed different characteristics than the INOR-8 base metal. The comparatively mild etchant delineated grain boundaries in the outer layer but left the base metal unaffected, as displayed in Fig. 51.



Fig. 49. Typical Spongy Surface Layers from Vapor Region Samples of (a) INOR-8 Test Fluorinator No. 2 (b) INOR-8 Test Fluorinator No. 4. As-polished. 500X.

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Fig. 50. Surface Layers on Sample from Vapor Region of INOR-8 Miniature Test Fluorinator No. 2. As-polished. 500X.



Fig. 51. Surface Layers on Sample from Vapor Region of INOR-8 Miniature Test Fluorinator No. 2. Etchant: Potassium cyanide-ammonium persulfate. 500X.

Samples of the outer corrosion product layer and of the spongy subsurface region were obtained by mechanical milling and scrapping. Most of the material comprising the outer layer was found to be strongly ferromagnetic in contrast to the base metal. These samples plus millings removed from the exterior diameter of the No. 2 vessel wall, 5-10 mils below the surface, were submitted for spectrochemical analyses. All of the samples were quantitatively analyzed for Cr, Mo, Fe, and Ni by emission spectroscopy using a porous cup electrode method of sample excitation. The results are shown in Table XV.

Table XV. Analyses of Corrosion Products and Base Metal Removed from INOR-8 Miniature Fluorinator No. 2

Sample Description and Location	Chromium (wt %)	Iron (wt %)	Molybdenum (wt %)	Nickel (wt %)
Outer corrosion product layer	3.9	3.3	8.4	84.3
Spongy subsurface layer	7.6	3.8	16.3	69.4
Base metal	6.6	3•75	15.45	75.45

Examination of Table XV shows that the outer corrosion product layer was 13 wt % richer in nickel when compared to the base metal, INOR-8, while the chromium, molybdenum, and iron contents in the corrosion product decreased 40, 45, and 11 wt %, respectively. The spongy subsurface layer showed generally small weight percent increases for the chromium, molybdenum, and iron, respectively, when compared to the base metal analysis. Nickel in the subsurface layer was approx 8 wt % less than that found in the base metal.

The mode of corrosive attack on INOR-8 in contact with the simulated volatility process fluorination environment at 600°C appears to involve selective losses of chromium, molybdenum, and iron from the nickel solid solution. This may occur by formation of metal fluorides on the bulk metal surface which subsequently volatilize, or by the various methods of fluoride film losses described in Section I.

The reason(s) for the slightly higher minor element concentrations in the spongy subsurface corrosion layer when compared with the base metal appear(s) anomalous to normal diffusion processes. Concentration gradients produced by the initial selective losses should become the driving force for diffusion of the minor alloy elements toward the exposed surface of the reactor and lower than base metal concentrations of chromium, molybdenum, and iron would be expected in the subsurface region. It may be that complex fluoride compounds are formed in the subsurface region which do not volatilize under the test conditions and thus simply tie up higher concentrations of chromium, molybdenum, and iron. However, since x-ray diffraction patterns on samples from the sublayer disclosed only the presence of a face-centered cubic material very similar to the pattern obtained on the base metal, supporting evidence for this theory has not been obtained.

Detailed treatment of the corrosive attack on INOR-8 deserves separate study which may or may not be warranted considering the ultimate goal of the entire test series covered in this section; that is, comparative behavior of A nickel, Inconel, and INOR-8 in contact with a volatility process fluorination environment. In regard to the latter, A nickel had the best resistance to attack at pilot plant fluorination temperatures of 600°C; but at 450°C INOR-8 presented favorable competition to the nickel. There are several advantages attendant to using INOR-8 as a fluorinator construction material including its high strength and oxidation resistance. Also, since the construction material for the VPP hydrofluorinator is INOR-8, it might be possible to consolidate both hydrofluorination and fluorination operations in a single vessel.

### IV. Volatility Pilot Plant Scouting Corrosion Tests

# A. Material Selection

In order to take advantage of the service conditions provided by the VPP process runs and to achieve insight into the resistance of other materials to the complex fluorination environment, a series of corrosion specimens was located in the Mark I and Mark II fluorinators and examined at convenient intervals. As mentioned in Section I, resistance to further attack by fluorine on metals was felt to be imparted by passive fluoride films which form on the metal. Protection has been shown to be dependent on the properties of the fluoride films, especially volatility, adherence to the substrate metal, mechanical and thermal stability, and thickness.

For the scouting corrosion tests described in this section, selection of materials which contained constituents known to form low volatile fluorides seemed especially appropriate. As a guide to volatility, the melting and boiling points of the common ingredients in many commercial materials of construction were reviewed. A partial listing taken from Brewer<sup>46</sup> is shown in Table XVI. It can be seen that chromium and molybdenum, commonly added for improved resistance to air oxidation and/or improvement of hightemperature properties, form highly volatile fluorides, especially at their higher oxidation states. Nevertheless, because of the previous use of nickelbase alloys containing chromium (Inconel) and/or molybdenum (INOR-8) in the Aircraft Reactor Experiment (ARE) and the Molten-Salt Reactor (MSR) studies, alloys containing these two constituents were included in the materials selected for the scouting tests.

Table XVII lists the specimen materials, their trade names, where applicable, and the general alloy classifications to which they belong. Most of the materials were nickel-rich alloys. The 90 wt % Ni-10 wt % Co and 80 wt % Ni-20 wt % Co alloys were nonproprietary and were fabricated from melts made in Metallurgy Division facilities. The remaining materials were obtained from commercial suppliers. The platinum specimen served as an electrode probe in an attempt by VPP operating personnel to investigate the electrochemical effects during the fluorination process. However, current did not flow through the probe and corrosive losses were recorded in a regular manner for the platinum specimen. Table XVIII shows the nominal compositions for all scouting test materials.

<sup>46</sup>L. Brewer, "The Fusion and Vaporization Data of the Halides," <u>The Chemistry</u> and <u>Metallurgy of Miscellaneous Materials</u>: <u>Thermodynamics</u> (ed. by Laurence L.Quill) National Nuclear Energy Series, Div. IV 19B, McGraw-Hill, New York, 1950.

Element	Fluoride	Melting Point (°C)	Boiling Point (°C)
Ni	NiF	1027 <sup>b</sup>	1627 <sup>b</sup>
Fe	FeF	1.102	1 <b>8</b> 27 <sup>b</sup>
	FeF	1027	1327 <sup>b</sup>
Co	CoF	1202	1727 <sup>b</sup>
	CoF	1027 <sup>b</sup>	1327 <sup>b</sup>
Cu	CuF	Disproportionates	Ъ
	CuF <sub>2</sub>	947	1377
Al	Alf	827 <sup>b</sup>	1357 <sup>6</sup>
	Alf <sub>3</sub>	> 1272	1272
Mg	MgF	1263	2227
Mn	MnF	856	2027 <sup>b</sup>
	MnF <sub>3</sub>	1077 <sup>b</sup>	1327 <sup>b</sup>
$\mathrm{Ti}$	TiF <sub>3</sub>	1227 <sup>b</sup>	1427 <sup>b</sup>
	$\operatorname{TiF}_{h}$	427 <sup>b</sup>	284
Cr	CrF2	1102	2127 <sup>b</sup>
	$\operatorname{CrF}_{3}$	1100	1427 <sup>b</sup>
	$\operatorname{CrF}_{h}$	277 <sup>b</sup>	297 <sup>b</sup>
	CrF	102 <sup>b</sup>	117 <sup>b</sup>
Мо	MoF <sub>5</sub>	77 <sup>b</sup>	227 <sup>b</sup>
	MoF	17	36
Pt	$PtF_{3}$	> 727	> 727 <sup>b</sup>
	$\operatorname{PtF}_{\underline{\mu}}$	> 627	727 <sup>b</sup>
	$\operatorname{PtF}_{C}$	Decomposes at $280^{\circ}C^{(c)}$	
Au	AuF <sub>3</sub>	727 <sup>b</sup>	> 727 <sup>b</sup>

## Table XVI. Melting and Boiling Point Data of Various Metal Fluorides Constituents of Materials Used in the Volatility Pilot Plant Scouting Corrosion Tests<sup>a</sup> (Converted from °K)

 <sup>a</sup>L. Brewer, "The Fusion and Vaporization Data of the Halides," <u>The Chemistry</u> and Metallurgy of Miscellaneous Materials: <u>Thermodynamics</u> (ed. by Laurence L. Quill) National Nuclear Energy Series, Div. IV 19B, McGraw-Hill, New York, 1950.
<sup>b</sup>Estimated or Obtained by Extrapolation of Experimental Data by L. Brewer.
<sup>c</sup>Bernard Weinstock and J. G. Malm, "Some Recent Studies with Hexafluorides," <u>Basic Chemistry in Nuclear Energy</u> 28, pp.125-129, 2nd United Nations International Conference on the Peaceful Uses of Atomic Energy, Geneva, 1958.

Material	Classification	
L Nickel	Ni	
INCO 61 Weld Wire	l <u>N</u> .i.	
Gold-plated L Nickel	Ni	
90 wt % Nickel—10 wt % Cobalt	Ni-Co	
80 wt % Nickel-20 wt % Cobalt	Ni-Co	
Cobanic	Ni-Co	
Monel	NiCu	
D Nickel	Ni-Mn	
Nimonic 80	Ni-Cr	
Inconel	Ni-Cr-Fe	
Waspalloy	Ni-Co-Cr (+ Ti, Al, and Fe)	
INCO 700	Ni-Co-Cr	
Hymu 80	Ni-Fe-Mo	
Hastelloy B	Ni-Mo-Fe	
Hastelloy W	Ni-Mo-Fe	
INOR-2	Ni-Mo-Cr	
INOR-8	Ni-Mo-Cr-Fe	
Hastelloy X	Ni-Mo-Cr-Fe	
OFHC Copper	Си	
Platinum	Pt	

Table XVII. Corrosion Scouting Test Specimen Materials Used in the Volatility Pilot Plant Mark I and Mark II Fluorinators

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								Nominal Com	osition	1 wt %						
Material	Ni	ы е	Co	£	Al	MM	Pt	Τi	Mo	сr	C	N M	Zr	n	S. <sup>1</sup>	E
L Nia	99.47	0.11				0.17				0	023			0.0075	< 0,01	
INCO 61 weld wire	93.00 (min)	ਾ. ਜ		0.25	1.5	0.4		2.0-3.5		0	.15			10°0	0.75	
Au-plated L Ni <sup>b</sup>	Golå plat	ie = ~ 0	.0015 ir	1.												
D NI DFHC Cu	95.õe	0.05		0.02 04 0.02		4.75				0	T.			0.005	0.05	
Hymu 80	62	16 (bal)		•		0.50			h.o	0	•02				0.15	
Cobanic	55		45							0						
INCO 700	45	1.5	29		S.S			2.3	3.0	16 O	.08					
80 Ni-20 Co	a 79.59		20.13	~						0	.026					
90 Mi-10 Co	a 90.02		9.6	ŝ						0	.020					
Monel <sup>a</sup>	67.66	1.40	0.47	7 30.30		0.98				0	.19			0.01	0.11	
Inconel	77°	2.		0.1		0.20				15 O	.06			0,007	0.2	
Hastelloy X <sup>c</sup>	48 (bal)	18.76	10.1			0.64			8.82	21.81 0	, 1	0.20		0,008	0.78 0.(	11
Waspalloy	57 (bal)	2 (max)	15-15	0.1 (max)	5.11	0.5 (max)		2.75-3.25	3.5-5	18-21 0 (m	г. (хг		0.1	0.03 (max)	0.75 (max)	0.008 (max)
Hastelloy B	65	ĥ							28	0.4 0	1					
Hastelloy W	60	5.5	5.0			1			25	0	दा	•• •	0			
INOR-2 INOR-8 <sup>ä</sup>	79 69.8	5.1				0.83		0.08	16 16.5	5 6.9 0	بر. 80			0.005	0.14 0.0	80
Nimonic 80 <sup>c</sup> Platinum	: 74 <b>.</b> 28	0.6		0.01	1.42	0.5	+6.66	2.42		20.38 0	.05			0,007	0.28	
<sup>a</sup> oRNL Chemi	cal analy	sis. b	Base met	al same	e as L N	i above.	cven.	dor chemical	analysi	s. <sup>d</sup> Wei	stinghc	nuse che	emical	analysis	(Heat 8M3)	
eIncludes c	obalt.															

Table XVIII. Nominal Composition of Corrosion Specimens from the Volatility Pilot Plant Fluorinator

- 112 -

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#### B. Test Method

The scouting specimens consisted of lengths of rod, sheet, or split pipe stock, or combinations thereof, each approx 48-56 in. long. They were held in place in the fluorination vessels by two methods. During the Mark I fluorinator's operations, all specimens were welded to the top blind flange of the vessel as shown in Fig. 3. When the Mark II vessel was in use several specimens were welded to the inspection port flange in a similar manner while the remaining test rods were held in place by metal connectors gripping a fitting prevelded to the end of the test rod. The end fittings were A nickel tubing 1/2-in.-o.d. x approx 3 in. long and seal welded to prevent the escape of process gases. Figure 52 shows a cut-away view of the Mark II VPP fluorinator illustrating the placement of corrosion specimens and a typical test specimen prepared for insertion.

A total of 31 test specimens was exposed to the fluorination environments. In each test grouping at least one L nickel specimen was included as a control. Figure 53 shows a top view of the fluorinators indicating the placement of the specimens in terms of polar geometry, the testing order and identification number, and corrosion specimen numbers. The corrosion specimen number identifies particular fluorinator runs when specimens were in place and also indicates the total number of specimens tested simultaneously. The major test groupings were fluorination runs M-21 through M-48, C-9 through C-15, E-3 through E-6, L-1 through L-4, and L-5 through L-9. Figure 54 details the process cycling for each run group. Tables I and VI summarize the other process details during these scouting runs.

## C. Reactions to Environments

After exposure, the test specimens were removed from the fluorination vessels and subjected to dimensional analyses and metallographic study. Figures 55 and 56 summarize the maximum corrosive losses found in each major exposure region as determined by micrometer measurements and optical microscopy. Figure 55 details the loss data as a plot of mils per month, based on individual molten-salt residence times, while Fig. 56 shows the loss data in terms of



Fig. 52. View of Mark II VPP Fluorinator Vessel Showing Methods of Corrosion Rod Placement.



\*Example: M21-48 refers to runs when specimen was exposed; (:1) refers to local group number, that is, four specimens tested simultaneously during the "M" runs.

Fig. 53. Top View of VPP Fluorinators Showing Placement of Corrosion Test Specimens, Testing Order, and Specimen Number.





- 116 -

Fig. 55. Summary of Corrosion on Specimens Exposed in VPP Fluorinators.



UNCLASSIFIED ORNL-LR-DWG 57784

- 117 -



Summary of Corrosion on Specimens Exposed in VPP Fluorinators. Fig. 56.

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mils per hour, based on operational fluorine sparge time. In both figures dotted and cross hatched bars represent bulk metal losses and visible intergranular attack, respectively, while solid bars indicate a total loss of the specimen occurred at the point indicated.

Several specimens registered lower rates of maximum corrosive attack than the L nickel control rods. These specimens included Hymu 80, INOR-2, INOR-8, Hastelloy W, Hastelloy X, Waspalloy, and the 90 wt % Ni-10 wt % Co alloys. In particular, Hymu 80 was superior to all test specimens in all runs, although the specimen registered a maximum bulk loss rate of 11 mils/month, based on molten-salt residence time, at the vapor-salt interface. Total loss of corrosion specimens, indicating corrosion rates of > 105 mils/month based on molten-salt time, was found for one L nickel specimen, one INCO-61 weld wire specimen, and for the Hastelloy B, Nimonic 80, Inconel, Monel, copper, and platinum specimens.

Selected photomacrographs and photomicrographs of samples from many of these corrosion specimens are presented in Appendix A. In most cases, the areas of maximum attack are shown.

#### D. Discussion of Results

A wide variation in resistance to the fluorination environment was found on corrosion specimens used in the compatibility testing described. The variations in corrosion noted were anticipated because of the widely divergent process conditions. Because the VPP has essentially been operated for feasibility studies, demonstration runs, and runs designed to recover uranium from available fluoride salt mixtures, the determination of optimum process conditions rather than the gathering of corrosion data has been the prevailing philosophy.

In many groups of runs a wide variety of scouting materials was used which were in proximity. This was recognized as being far from ideal for corrosion testing, but little choice was available under the circumstances. However, Run Group M-21-48 did contain only the reference material, L nickel, and the results were almost as divergent as the data obtained in run groups where as many as seven different materials were present in the same system.

The L nickel reference specimen's mean rate of corrosive attack was similar to that found on the walls of the corresponding pilot plant fluorinators, although large deviations from the mean existed. While maximum corrosive attack occurred in the vapor region on the Mark I fluorinator and in the salt region of the Mark II vessel, in almost all cases the L nickel specimens showed maximum attack at the point of vapor-salt interface contact. A notable exception was specimen M-21-48:4, tested in the Mark I fluorinator, which exhibited failure in the upper vapor region. The addition of chromium fluoride during the M-21-M-48 Run Group may have produced the vapor region failure through mechanisms proposed in Section I.

The increased attack on the L nickel specimens at the vapor-salt interface possibly resulted from salt agitation at the center of the vessel induced by the fluorine and nitrogen sparging through the draft tube. This agitation could result in increased erosion and vibration of the specimens with resultant loss of protective films. In addition, the location may have allowed primary contact of the specimens with the fluorine sparge before the latter reached the vessel walls. However, the increased corrosion to be expected from this effect was not noted on the A nickel fluorine inlet tubes, the draft tube walls, or other fluorinator internal components discussed in Section II.

Tests on INCO-61 weld wire were intended to provide data on a nickelrich alloy with additions not readily available in commercial alloys. Besides nickel, this material contains small amounts of Al, Ti, Fe, Mn, Si, and Cu (Table XVIII). Since this material was used in fabricating the fluorinators, these tests provided an indication of weld metal behavior, although the weld metal after deposition would have a cast structure compared to the wrought structure of the weld wire. The INCO-61 specimens showed generally analogous bulk metal loss behavior when compared to the L nickel specimens. However, no intergranular attack was found in the INCO-61 specimens. The behavior of the gold-plated L nickel corrosion specimens indicated that no protection was provided by the plating. Comparison with L nickel specimen number L-1-L-4:6 in Fig. 55 shows comparable losses.

On the basis of single exposures of the 90% Ni-10% Co and 80% Ni-20% Co specimens, cobalt additions seemed to improve the resistance of commercial nickel to the fluorination environment with the one exception of the high vaporsalt interface attack found in the 80% Ni-20% Co specimen. The cobanic alloy, containing 45% Co, was tested only in the fluorination vapor phase and showed greater losses when compared with L nickel. Thus, further investigation of nickel-base alloys with less than 20% Co seems warranted. Because the most corrosion-resistant Ni-Co alloys are experimental, the investigation should ultimately include the determination of mechanical and physical properties and fabricability.

The nickel-rich copper specimen, Monel, provided poor resistance to the fluorination environment during a single test run. The specimen failed completely at the vapor-salt interface.

The D nickel, containing about 5% Mn, and developed for improved resistance to sulfur attack in oxidizing atmospheres at elevated temperatures, presented somewhat higher loss rates than the L nickel specimen simultaneously exposed. The D nickel specimen presented a different geometry to the corrodents, sheet vs rod, but that difference is not believed to have been significant in producing the higher rates of attack.

As expected, the Nimonic 80 alloy, containing 20% Cr and approx 2.5% Ti, exhibited substantially greater bulk losses than the corresponding L nickel specimen. Reference to Table XVI shows that chromium and titanium can form highly volatile fluorides at their higher valence states.

A high-temperature, oxidization-resistant, nickel-chromium-iron alloy, Inconel, has been mentioned as being useful in contact with fused fluoride salts. During the single test in the fluorination environment, the specimen completely failed at the vapor-salt interface, and also exhibited a rather high vapor-phase attack. The Ni-Co-Cr alloys, Waspalloy and INCO 700, differed significantly in attack. Samples of INCO 700 were available only for a salt-phase test (L-1-L-4:4) and in this region had a bulk loss rate > 100 mils/month which considerably exceeded the L nickel specimen exposed simultaneously. The Waspalloy specimen showed a rate of attack slightly less than that of a corresponding L nickel specimen. Further investigation of Waspalloy seems warranted, although it should be remembered that the alloy ages at 750°C, a temperature only slightly higher than the fluorinator operating range. Thus, temperature excursions, which may be encountered in extended operations, might drastically reduce the ductility of the material.

The Hymu-80 specimen, Ni-Fe-Mo, presented the best resistance to corrosion of any of the specimens tested including the L nickel reference material. This alloy is commercially available and six miniature fluorinators have been fabricated for compatibility testing with  $NaF-ZrF_{l_4}$  and  $LiF-NaF-ZrF_{l_4}$  salt mixtures by the Volatility Studies Group of Chemical Development Section A.

The essentially Ni-Mo-Fe alloys, Hastelloy B and W, again showed the variant behavior characteristic of this test series. The Hastelloy-B specimen had extremely poor resistance and failed completely 22 in. above the static salt level. Since the specimen was exposed during the Mark I operation, the high vapor-phase attack evidenced at that time may well have influenced the behavior of Hastelloy B. The Hastelloy-W specimen, in place during a relatively low corrosion run series, L-5-L-9, exhibited slightly lower losses when compared with the L nickel control specimen. This was because of the intergranular attack experienced by the L nickel.

One of the early manifestations of the Ni-Mo alloy series developed at ORNL for fused-fluoride salt use was INOR 2. The single INOR-2 specimen tested had corrosion resistance superior to either of the L nickel specimens exposed concurrently. Lack of additional material has hindered further corrosion testing.

The Ni-Mo-Cr-Fe alloys tested, INOR 8 and Hastelloy X, had corrosion resistance comparable to that of the L nickel rods exposed simultaneously. As described in Section III, miniature fluorinators were fabricated from INOR 8 and tested. These tests did not indicate any superiority over commercial nickel.

A single copper specimen, L-3-L-6:7, was exposed to the Mark II environment. The specimen failed completely in the middle vapor region. Cupric fluoride was considered most likely to form under the excess fluorine conditions present during testing. Although  $CuF_2$  has a melting point of approx 950°C and should have afforded protection to the parent metal, copper's lack of resistance to oxidizing environments at fluorination temperatures may explain its poor performance.

The platinum specimen was completely severed by corrosive attack in the upper vapor region. This wire was intended to serve as an electrode probe, but never carried current. The reason for the extreme rate of attack on platinum seems to be the instability of  $PtF_6$  which has been reported to decompose at 280°C (Table XVI).

Whereas the L nickel specimens exhibited intergranular attack, very few instances of this mode of attack were noted on the other specimens tested. Sections I and II describe the mechanisms presumably operating on L nickel.

### E. Future Studies

The corrosion rates for all scouting materials tested to date appear to be excessive for long-time use. Therefore, additional fluorination scouting corrosion tests of the type described have been planned. Particular emphasis has been placed on binary nickel-rich alloys containing various amounts of Fe, Co, Mn, Mg, and Al. The latter two elements have not been considered in previous experiments because of the fabrication, age hardening, or other difficulties characteristic when alloyed in appreciable quantities with nickel. Only a few nickel-base commercial alloys containing magnesium and aluminum in combination with other ingredients have been available and still fewer have been commercial as Ni-Mg or Ni-Al binaries. Yet, aluminum and magnesium are known to form fluorides at their highest valence states, which have higher melting points than  $NiF_0$ .

To date, Fe, Co, Mn, Mg, and Al in the quantities shown in Table XIX have been added to induction melts of nickel and cast into 1-in. rounds. After suitable homogenization treatments of the castings, the rounds were cold swaged

- 123 -

			Nomina	l Compos (wt %)	sition	
Alloy Group	Ni	Fe	Со	Mn	Mg	A.J.
NiFe	95	5				
	90	10				
	80	20				
NiCo	95		5			
	90		10			
Ni <b>M</b> n	98			2		
NiMg	99•95				0.05	
	99•9				0.1	
	99				1.0	
NiAl	99					1.
	97					3

Table XIX.	Proposed Noncommercial Binary Alloys for Corrosion Testing
	in the Volatility Process Fluorination Environment

to 1/4-in.-diam rods and hydrogen annealed. The rods have been cut to proper lengths, end fittings attached by seal welding, and are presently awaiting placement in a VPP fluorinator.

In addition, high-purity vacuum-melted nickel has been obtained and fabricated into the proper test specimen shape for examining the resistance of that material to the fluorination environment.

#### V. Argonne National Laboratory Fluorination Corrosion Studies

The Chemical Engineering Division of Argonne National Laboratory has done extensive work on nonaqueous processing of irradiated fuels.<sup>47</sup> A portion of their effort has been on studies of fluoride volatility processes. A review of one of the Argonne National Laboratory studies on materials compatibility in a simulated fluorination environment is included here for comparison with ORNL data.

#### A. Test Method

Fluorination corrosion tests were conducted on coupons of L nickel, D nickel, Hastelloy B, and INOR 1. (ref 48) The coupons were contained in a vessel which had an inner liner and internal piping fabricated from A nickel. The coupons, each 8 in. long x 0.5 in. wide x 0.032 in. thick, were wired together to form a simulated draft tube of square cross section. The composition of the wire was not indicated. The nominal composition of the first three materials listed above has been given in Table XIX. Another of the early experimental alloys, INOR 1, studied in the development of INOR 8, had a nominal composition of 78 wt % Ni-20 wt % Mo-0.5 wt % Mn-0.5 wt % Si-0.25 wt % Fe-0.01 wt % C.

The simulated draft tube was partially immersed in a bath of equimolar NaF-ZrF<sub>h</sub> so that the vapor-salt interface was approx 3 in. up from the

<sup>47</sup>R. C. Vogel and R. K. Steunenberg, "Fluoride Volatility Processes for Low Alloy Fuels," <u>Symposium on the Reprocessing of Irradiated Fuels Held at</u> Brussels, Belgium, May 20-25, 1957, Book 2, Session IV, pp.498-559, TID-7534.

48 L. Hays, R. Breyne, and W. Seefeldt, "Comparative Tests of L Nickel, D Nickel, Hastelloy B, and INOR 1," Chemical Engineering Division Summary Report, July, August, September, 1958, ANL-5924, pp.49-52. bottom of the coupons. The bath was held at 600°C while fluorine or helium was introduced centrally beneath the melt surfaces at rates of approx 0.1 standard liters/min. During this test series, the fluoride salts were kept molten for a total of 216 hr and for 63 hr of that time fluorine was sparged into the bath. The process gases were introduced on a cyclic basis, i.e., for 7 hr/day fluorine was sparged while helium was sparged for the remaining 17 hr of a day.

Figures 57 and 58 illustrate the corrosion losses obtained during this test series. Figure 57 is a bar graph where bulk metal losses as determined by micrometer measurement and additional losses as determined by metallographic examination have been converted to mils/month of molten-salt residence time. Figure 58 is a similar graph plotted as mils/hour of fluorine sparge time. Rates have been plotted instead of original data for comparison with ORNL data.

## B. Discussion of Results

The corrosion losses in this Argonne National Laboratory test series suggest that INOR 1 was the most resistant to the test environment, while the L nickel reference material fared poorly. This is attributed to the extensive intergranular attack on the latter material, particularly at the vapor-salt interface.

Comparison of the Argonne National Laboratory corrosion results with those obtained at ORNL is difficult because of differences in test procedures which include fluorine flow rates, specimen location and geometry, and salt compositions. However, Argonne's results on L nickel fit the corrosion limits determined for the ORNL scouting corrosion reference specimens in the salt and at the salt-vapor interfaces. Vapor-phase specimens are not readily comparable, since Argonne specimens were only 3 to 4 in. above the salt surface.

The Argonne National Laboratory Hastelloy B and the D nickel specimens generally had less attack than scouting specimens at ORNL. The differences may be attributed to more severe operating conditions in the ORNL pilot plant fluorinator. Since no INOR-1 specimen had been tested at ORNL, no comparisons could be derived. UNCLASSIFIED ORNL-LR-DWG 49169

20 /// - INTERGRANULAR ATTACK/METALLOGRAPHIC EXAM - SUBSURFACE VOIDS/METALLOGRAPHIC EXAM 65 - METAL LOSS/MICROMETER MEASUREMENT V – VAPOR I – VAPOR SALT INTERFACE S – SALT MAXIMUM LOSS SHOWN IN EACH PHASE 30 LEGEND (BASED ON SALT RESIDENCE TIME) 25 mits/mo 20 5 <u>0</u> ŝ 0 -- v > > --ഗ > -S > S ----HASTELLOY B L NICKEL D NICKEL INOR 1

Fig. 57. Summary of Corrosion on Coupons Exposed to a Fluorination Environment by the Argonne National Laboratory.





Fig. 53. Summary of Corrosion on Coupons Exposed to a Fluorination Environment by the Argonne National Laboratory.

Close cooperation has been maintained with the Argonne National Laboratory on the selection and testing of candidate materials of construction for use in the fluoride volatility process. In addition to materials research as a means of containing the fluorination environment, Argonne National Laboratory has suggested three other approaches. These have been the use of (1) cold wall fluorination vessel, (2) a spray tower fluorination, and (3) lower melting salts.<sup>49</sup> Further work on these approaches has been covered in the Argonne National Laboratory Chemical Engineering Division Summary Reports.

### VI. Supplementary Volatility Pilot Plant Equipment

The operation of the VPP necessarily included the use of a number of auxiliary components such as traps for radioactive products, absorbers, valves, fittings, fluorine disposal systems, and piping. Each of these are subject to various corrosive conditions including fluorine, uranium hexafluoride, fused fluoride salts, etc. Thus, successful operation of the volatility process requires that materials be selected appropriate for the particular conditions of service. Construction of the present system was predicated on the knowledge available at the time and as operating experience has been gained these parts have been examined to verify the original assumptions. In most cases the metal selected appears to give satisfactory service and failure analyses have suggested alternate metals in the case of those which proved unsuitable. The details of the findings of this investigation are presented in Appendix B.

#### ACKNOWLEDGMENT

The authors wish to acknowledge the assistance given by the Metallography Section of the Metallurgy Division, by personnel of the Spectrochemical Group and the Special Analyses Laboratory of the Analytical Chemistry Division, and by the Graphic Arts and Photography Departments.

A portion of the corrosion evaluations reported in this document and the accompanying illustrations are the work of the Corrosion Research Division, Battelle Memorial Institute, and their contribution deserves special mention here.

<sup>49</sup>A. Goldman, <u>Trip Report, ANL</u>, to W. D. Manly, June 12, 1958.

Thanks are due to the many personnel of the Chemical Technology Division working on the ORNL Fluoride Volatility Process who aided in gathering process data or in reviewing this report.

Special thanks are due J. H. DeVan, E. E. Hoffman, and D. A. Douglas, Jr., for their constructive appraisal of portions of this report and to the Metallurgy Division Reports Office for their patient typing and careful preparation of the material for reproduction.

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G. I. Cathers, "Fluoride Volatility Process for High-Alloy Fuels," Symposium on the Reprocessing of Irradiated Fuels, Held at Brussels, Belgium, TID-7534, Book 2, pp. 560-573, May 20-25, 1957.

G. I. Cathers, M. R. Bennett, and R. L. Jolley, The Fused Salt Fluoride Volatility Process for Recovering Uranium, ORNL-2661 (April, 1959). APPENDIX A

PHOTOMICROGRAPHS OF VPP SCOUTING CORROSION TEST SPECIMENS

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Fig. 59A. Cross Section of Interface Region Sample from L Nickel Corrosion Specimen (Runs L-5-L-9:3). Interior circumference of white circle indicates original size. As-polished. 10X.



Fig. 59B. Portion of Fig. 59A. Typical Surface and Microstructure. Note intergranular attack. Etchant:  $\text{KCN-(NH}_4)_2 \text{S}_2 \text{O}_8$ . 200X.



Fig. 60A. Cross Section of Salt Region Sample from Gold Plated L Nickel Corrosion Specimen (Runs L-1-L-4:2). Interior circumference of white circle indicates original size. As-polished. 15X.



Fig. 60B. Portion of Fig. 60A. Typical Surface and Microstructure. Intergranular attack. Etchant:  $HNO_3$ -H<sub>2</sub>SO<sub>4</sub>. 200X.



Fig. 61A. Cross Section of Interface Region Sample from INCO-61 Weld Wire Corrosion Specimen (Runs L-3-L-4). Interior circumference of white circle indicates original size. As-polished. 15X.



Fig. 61B. Portion of 61A. Typical Surface and Microstructure. Etchant: Concentrated HNO<sub>3</sub>. 200X.



Fig. 62A. Cross Section of Interface Region Sample from INOR-2 Corrosion Specimen (Runs C-9-C-15:1). Interior circumference of white circle indicates original size. 85X.



Fig. 62B. Portion of Fig. 62A. Typical Surface and Microstructure. Etchant: Chromium regia, 200X.



Fig. 63. Section of Vapor Region Sample from Hastelloy B Corrosion Specimen (Runs C-9-C-15:4) Showing Typical Surface and Microstructure. Etchant: Chromium regia. 200X.



Fig. 64. Section of Vapor Region Sample from Inconel Corrosion Specimen (Runs E-3-E-6:2) Showing Typical Surface and Microstructure. Etchant: Modified aqua regia. 200X.


Fig. 65. Section of Vapor Region Sample from Monel Corrosion Specimen (Runs E-3-E-6:3) Showing Typical Surface and Microstructure. Etchant:  $HC_2H_3O_2$ : HNO<sub>3</sub>: HCl. 200X.



Fig. 66. Section of Vapor Region Sample from OFHC Copper Corrosion Specimen (Runs E-3-E-6:7) Showing Typical Surface and Microstructure. Etchant:  $NH_4OH: H_2O_2$ . 200X.



Fig. 67A. Cross Section of Interface Region Sample from 80 Ni-20 Co Corrosion Specimen (Runs E-3-E-6:4). Interior circumference of white circle indicates original size. Combination of grain size and etchant presents a relief appearance because of light. Etchant:  $HNO_3: H_2SO_4$ . 15X.



Fig. 67B. Portion of Fig. 67A. Typical Surface and Microstructure. Etchant: Concentrated HNO<sub>3</sub>. 200X.



Fig. 68A. Cross Section from Interface Region Sample from 90 Ni-10 Co Corrosion Specimen (Runs E-3-E-6:6). Interior circumference of white circle indicates original size. Etchant:  $HNO_3: H_2SO_4$ .



Fig. 68B. Portion of Fig. 68A. Typical Surface and Microstructure, Etchant: Dilute HNO<sub>3</sub>. 200X.



Fig. 69. Longitudinal Cross Section of Vapor Region Sample from Platinum Corrosion Specimen (Run E-6) Showing Typical Surface and Microstructure. Cathodic vacuum etch. 200X.



Fig. 70A. Cross Section of Interface Region Sample from D Nickel Corrosion Specimen (Runs L-5-L-9:1). Interior of white lines indicates original size. As-polished, 10X.



Fig. 70B. Portion of Fig. 70A. Typical Surface and Microstructure. Etchant:  $KCN-(NH_4)_2S_2O_8$ . 200X.

- 144 -



Fig. 71A. Cross Section of Salt Region Sample from INCO-700 Corrosion Specimen (Runs L-1-L-4:4). Interior circumference of white circle indicates original specimen size. As-polished. 5X.



Fig. 71B. Portion of Fig. 71A. Typical Surface and Microstructure. Cathodic vacuum etch. 200X.

- 145 -



Fig. 72. Cross Section of Vapor Region Sample from Cobanic Corrosion Specimen (Runs L-1-L-4:5). Interior circumference of white circle indicates original size. Etchant: Concentrated HNO<sub>3</sub>. 100X.



Fig. 73A. Cross Section of Salt Region Samples from INOR-8 Corrosion Specimen (Runs L-1-L-4:1). Interior circumference of white circle indicates original size. As-polished. 15X.



Fig. 73B. Portion of Fig. 73A. Typical Microstructure and Surface Showing intergranular attack. Etchant: Chromium regia. 200X.



Fig. 74A. Cross Section of Interface Region Sample from Waspalloy Corrosion Specimen (Runs L-5-L-9:4). Interior circumference of white circle indicates original size. As-polished. 5X.



Fig. 74B. Portion of Fig. 74A. Typical Surface and Microstructure. Etchant: Aqua regia. 200X.



Fig. 75A. Cross Section of Interface Region Sample from Hastelloy X Corrosion Specimen (Runs L-5-L-9:5). Interior of white lines indicates original thickness. As-polished. 10X.



Fig. 75B. Portion of Fig. 75A. Typical Surface and Microstructure. Etchant: Chromium regia. 200X.



Fig. 76A. Cross Section of Interface Region Sample from Hastelloy W Corrosion Specimen (Runs L-5-L-9:6). Interior of white lines indicates original thickness. As-polished. 10X.



Fig. 76B. Portion of Fig. 76A. Typical Surface and Microstructure. Etchant: Chromium regia. 200X.



Fig. 77A. Cross Section of Salt Region Sample from Nimonic 80 Corrosion Specimen (Runs L-5-L-9:7). Interior of white lines indicates original thickness. As-polished. 10X.



Fig. 77B. Portion of Fig. 77A. Typical Surface and Microstructure. Etchant: Aqua regia. 200X.



Fig. 78A. Cross Section from Interface Region Sample from Hymu 80 Corrosion Specimen (Runs L-5-L-9:8). Interior circumference of white circle indicates original size. As-polished. 5X.



Fig. 78B. Portion of Fig. 78A. Typical Surface and Microstructure. Etchant: Aqua regia, 200X.

APPENDIX B

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# SUPPLEMENTARY VPP EQUIPMENT

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## Supplementary Volatility Pilot Plant Equipment

Sections I and II have described reactions to the process environment of VPP L nickel fluorination vessels. Other important components used in the fluorination cycle of the fluoride volatility process have been examined for corrosion resistance to their local environments. The components include a CRP trap, a waste-salt line, the absorber vessels, valves and fittings, components in the fluorine disposal system, and selected sections from the process gas lines. Figure 79 illustrates the relative positions of these components in the pilot plant.

# Complexible Radioactive Products Trap

To date, two CRP traps have been utilized in the VPP. The trap examined was the second used and was in service during the Mark II fluorinator's lifetime, as has been described previously in Section IIA. This vessel was fabricated from a 5-in. sched-40 Inconel pipe, 31 in. long and it contained NaF pellets to scrub the fluorinator effluent gas. The trap was designed to remove  $\text{ZrF}_4$  and  $\text{CrF}_5$  and provides a certain amount of radioactive decontamination. Table XX summarizes the exposure conditions for the trap. A complete description of the performance of the CRP trap has been published.<sup>50,51,52</sup>

After the Mark II fluorinator was taken off stream, trepanned sections were removed from the CRP trap and sent to BMI for corrosion analyses. Figure 80 illustrates the sections removed.

Macroscopic examination of the samples at 20X revealed outlining of the Inconel grains and an etched surface with increased attack noted at Section B-T.

<sup>50</sup>C. L. Whitmarsh, <u>Reprocessing of ARE Fuel</u>, Volatility Pilot Plant Runs, <u>E-1 and E-2</u>, CF-59-5-108 (May 11, 1959).

<sup>51</sup>C. L. Whitmarsh, <u>Reprocessing of ARE Fuel</u>, <u>Volatility Pilot Plant Runs</u>, <u>E-3 through E-6</u>, CF-59-8-73 (August 26, 1959).

<sup>52</sup>C. L. Whitmarsh, <u>Uranium Recovery from Sodium Zirconium Fluoride Salt</u> <u>Mixtures, Volatility Pilot Plant Runs, L-1 through L-9</u>, CF-59-9-2 (September 30, 1959).





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- 156 -

Table XX. Summary of Process Conditions for Volatility Filot Plant Complexible Radioactive Products Trap During VPP Runs E-1 through E-6 and L-1 through L-9

	and a first state of the second		Time of Expos	sure During Ope	rations
Region	Exposure	Operating Temperature (°C)	Room Temperature to Operating Temperature (hr)	At Operating Temperature With N <sub>2</sub> Flow (hr)	At Operating Temperature With F2 Flow <sup>a</sup> (hr)
Top (outlet)	) <sup>F</sup> 2, UF6, <sup>W</sup> 2	305-395	00†~~	~ 1400	~ 100
Middle	F <sub>2</sub> , UF <sub>6</sub> , N <sub>2</sub> , NaF (solids)	~ 500 <sup>b</sup>	~ 400	~ 400	~ 100
Bottom (inlet)	F <sub>2</sub> , UF6, N2 (soličs)	385-500	400	001 ~	v 100

<sup>a</sup>For approx 35 hr UF<sub>6</sub> was added to F<sub>2</sub> stream. <sup>b</sup>Temperatures recorded only during 2 runs, E-1 and E-2.



Fig. 80. Location of Specimens Trepanned from the Inconel CRP Trap.

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Dimensional analyses were performed on the sections as well as a microscopic examination. A summary of the corrosion losses found by BMI personnel is presented in Table XXI. Included in the table is a column converting the maximum total losses to a mils per hour rate based on fluorine exposure time at operating temperature for the vessel. A maximum total corrosion attack rate of 0.46 mils/hr occurred in the lower section of the trap which sustained the higher operating temperatures.

Optical microscopic examination of sections removed from the trap revealed intergranular attack on both the interior and exterior surfaces of the vessel in the lower regions. The upper region showed intergranular attack only on the vessel's interior wall. A typical cross section of the interior wall of the CRP trap is shown in Fig. 81.

Comparison of the corrosive attack on the Inconel CRP trap and the Inconel bench-scale fluorinator described in Section III indicated maximum rate losses for the trap to be approximately twice that for the bench-scale fluorination vessel. The most similar areas with respect to operating environments were the lower vapor region of the fluorinator and the lower region of the CRP trap. At those locations, similar temperatures (approx 500°C) and process gases  $(F_2, UF_6)$  were present. In addition, however, sodium fluoride pellets were in contact with the wall of the CRP trap. The maximum bulk metal losses in the two regions were in a ratio of 3:1 (Trap:Fluorinator) while the maximum interior intergranular penetration ratio was > 5:1. These differences are difficult to reconcile in view of the 2:1 ratio for fluorine contact time at elevated temperatures for the two vessels.

No appropriate reason can be advanced to explain the variations cited in corrosive behavior for the CRP trap and the Inconel bench-scale fluorinator. However, the results present additional evidence that Inconel should be used with caution as the construction material for certain fluoride volatility process components.

### Waste-Salt Line

The removal of waste fluoride salt from the VPP fluorination vessels was accomplished by pressure transfer through a freeze valve and waste-salt line

- 159 -

Specien	- Location	Distance from Top Flange Weld (in.)	Wall Thickn Loss Maximum	ess esa <u>Minimum</u>	Intergr Penetr (mi Interior Surface	anular ation <sup>b</sup> <u>1s)</u> Surface	Total Maximum Corrosion Losses (mils)	Maximum Corrosion Rate <sup>C</sup> (mils/hr)
				111 <b>0</b> 111111	) ) 1 1 2 2			/ דדר /מיד דווו <i>\</i>
년 년 년	Top	ŝ	2	0	m	None	TO	0.1
1-1 N	Middle	8.5	ı	ı	년 년	Γ	I	I
В <b>-</b> Т	Bottom	26	24	б	17	11	46	0,46
	a Based on	12 mirrometer mea	surements.	taken on e	ach cracim	בין האר היה היה	א מטאלי הס+הפמ מואלי הס+הפמ	omi vol

Table XXI. Summary of Corrosion Losses on Specimens Trepanned from the Inconel CRP Trap

TRUTTURT E nan ortar ona alic THOM 222 original wall thickness of 258 mils.

bAs determined by optical microscopy.

<sup>c</sup>Based on fluorine exposure time at operating temperature.

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Fig. 81. Typical Microstructure of Sample B-T from the Interior Wall of the Inconel CRP Trap Showing Intergranular Attack. Etchant: Hydrochloric-nitric acid. 100X. into a low-carbon steel waste container. Pressures of approx 5 psig in the fluorinator were normally used to start the waste-salt flow. The salt then flowed through the waste line by siphon effect (See Fig. 79). The salt line reported here was used during the VPP L Runs 1-9 and for a single transfer at the end of Run M-64.

The subject waste-salt line was 3/8-in. sched-40 Inconel pipe. The line was held at temperatures of 550-835°C for approx 22.5 hr and exposed to flowing molten salts for about 2.5 hr during the pressure transfers.<sup>52</sup> It was probable that residual salts remained in static contact with the interior wall of the piping for all operations.

After Run M-64, the piping was removed from the pilot plant, sectioned in nine places and specimens sent to BMI for corrosion analyses. Figure 82 shows a schematic drawing of the waste line and the location of the specimens. Table XXII lists the corrosion data established at BMI by micrometer measurements and metallographic examinations. Maximum wall-thickness losses of 21 mils were found in specimen IX which was at the exit end of the waste-salt line.

Metallographic examination of the specimens showed slight intergranular attack of 1/2-mil penetration depth had occurred on the interior wall of all the samples with the exception of specimen IX. In the latter, a 3.5-mil thick corrosion product layer was found on the interior surface of the specimen.

Figure 83 shows the corrosion product layer and the subsurface structure. The product layer was spongy in character and similar in appearance to surface layers found on sections removed from early test Inconel freeze valves which had been in contact with NaF,  $\text{ZrF}_4$ ,  $\text{UF}_4$  (50-46-4 mole %) at 650°C. (ref 53)

Personnel at BMI suggested that the product layer might be the result of selective leaching of chromium by the fused salt. They indicated that the appearance was quite similar to layers found on Inconel in contact with hydrogen fluoride and fused fluoride salts where chromium leaching had been proven. Analyses of the metallic spongy deposit found on sections removed from the Inconel freeze valve referenced above indicated considerable losses of chromium

<sup>&</sup>lt;sup>53</sup>L. R. Trotter and E. E. Hoffman, Progress Report on Volatility Pilot Plant Corrosion Problems to April 21, 1957, ORNL-2495, pp.14-16 (Sept. 30, 1958).



- 163 -

# Table XXII. Corrosion Loss Data for Waste-Salt Line Material: 3/8-in. sched-40 Inconel pipe

Specimen	Distance from Top of Waste-Salt Container (in.)	Wall Thickness Losses <sup>a</sup> (mils)	Intergranular Penetration (Interior Surface) <sup>b</sup> (mils)	Total Corrosion <sup>c</sup> (mils)
I	191.5	4	2	6
ΤI	170	6	2	8
TIT	159.5	4	2	6
IV	146	24	1	5
V	110.5	4	1	5
VI	7.1	3	1.	4
VII	40.5	3	l	24
VIII	39.5	6	1	7
ΤX	0	21	0	21

<sup>a</sup>Based on metallographic measurements and subtracted from nominal original wall thickness of 91 mils.

 $^{\rm b}{\rm No}$  intergranular attack was observed on the outer surface of the pipe.

<sup>c</sup>Total corrosion equals the assumed original thickness of 91 mils minus the measured sound metal remaining.

Unclassified BMI C623



Fig. 83. Microstructure of Sample IX from the Interior Wall of Inconel Waste Salt Showing Corrosion Product Layer.

and iron in the corrosion product layer when compared with the nominal analysis of Inconel, i.e., 0.41% Cr and 0.46% Fe in the product layer.<sup>54</sup>

Since air, probably containing water vapor, could enter the exit line at the point of preferential losses, the mechanism involved appears to be an effect of a strong oxidant in the fluoride salt contacting the Inconel. In such cases, it has been reported that the nickel ion rapidly goes into solution and then is reduced by chromium resulting in leaching of chromium from the alloy.<sup>55</sup>

The attack described emphasizes the importance of preventing air and/or moisture from entering the fluorination system.

### Absorbers

The absorbers in the VPP are used to decontaminate the VF $_6$  product by sorption and desorption on NaF beds.<sup>56</sup> Incomel was used for the construction of these vessels.

Absorber Gas-Distribution Ring Failure. The Inconel gas-distribution ring in the first absorber failed during VPP operations. The failure was discovered after Run M-52 but is believed to have occurred during Run M-47.

A schematic view of the absorber and the gas-distribution ring is shown in Fig. 84. As shown on the photographs, Fig. 85-a and -b, approx 3 in.<sup>3</sup> of Inconel was melted in an area roughly 180 deg from the junction of the distribution ring and the inlet pipe. The melted area was about 5 in. long and the distribution ring was nearly severed by this action.

The fused salt visible in Fig. 85-b around the metal nodules which were formed as a result of melting was found by analyses to contain 4 wt % Cr as  $CrF_3$ , 10 wt % Ni as NiF<sub>2</sub>, and 3 wt % U as  $UF_{l_1}$ . Process chemicals normally

<sup>55</sup>F. F. Blankenship, The Effect of Strong Oxidants on Corrosion of Nickel Alloys by Fluoride Melts, CF-60-3-125 (March 30, 1960).

<sup>56</sup>R. P. Milford, Engineering Design Features of the ORNL Fluoride Volatility Pilot Plant, CF-57-4-18, pp. 3-4.

<sup>&</sup>lt;sup>54</sup>Trotter and Hoffman, <u>Op.</u> <u>Cit.</u>, p. 16.



Fig. 84. Schematic View of VPP Inconel Absorber and Gas-Distribution Ring Illustrating Area of Failure.

- 167 -



Fig. 85. Photographs of Failed Region from the Inconel Gas-Distribution Ring of Absorber, (a) Top View (b) Bottom View, Showing Molten and Recast Nodules and Adhering Salt.

present in this area of the absorber were NaF pellets,  $F_2$  and  $UF_6$ . Table XXIII shows the prior history of the ring up to the time of failure. Three unusual plant operating procedures were a part of this prior history:<sup>57</sup> (1) substitution of hydrogen fluoride for low temperature fluorine conditioning, (2) allowing fluorine to enter the absorber at approx 600°C rather than 375°C as specified in the standard operating procedures (the higher temperature was used to minimize hydrogen fluoride from the NaF and to reduce the sodium fluosilicate content of the NaF). (3) use of an accelerated cooling rate after desorption.

Only (2) is felt to be pertinent toward causing the failure described.

Metallographic examination and chemical analysis of the samples cut from the failed pipe, revealed only that the Inconel was normal in chemistry and microstructure. The microstructure was equivalent to as-received stock. The molten and recast nodules exhibited a typical cast structure.

The Inconel gas-distribution ring appears to have failed as the result of ignition with fluorine. Ignition is believed to have been initiated by an unknown foreign substance, such as dirt or grease, introduced while the absorber was open. The failure points up the necessity for extreme cleanliness when handling fluorine.

Inspection of Absorbers. Following fluorination Run L-9, the absorbers were visually inspected and thickness measurements taken with a "Vidigage." The ultrasonic device was operated by members of the Nondestructive Testing Group of the Metallurgy Division. Results of those examinations are cited in Figs. 86 and 87. The maximum detectible losses found were approx 30 mils in the upper regions of the vessel, a rate of approx 0.06 mils/hr, based upon fluorine residence time.

Based on these results, it is believed that the vessels can continue to be used in pilot plant operation.

<sup>57</sup>F. W. Miles and W. H. Carr, <u>Engineering Evaluation of Volatility Pilot</u> Plant <u>Equipment</u>, CF-60-7-65 (September 30, 1960) pp.95-96. Table XXIII. Exposure History for Absorber Containing Failed Inconel. Gas-Distribution Ring from Absorber FV-120

Twelve (12) absorption-desorption runs using NaF which typically 1. included: a. Absorption at 65-150°C; 2 hr  $F_2$  at ~ 15 standard liters/min UF<sub>6</sub> + 3 NaF -----> UF<sub>6</sub>· 3 NaF b. Purge-sparge with  $\rm N_{\rm p}$  at 20 standard liters/min for 1.5 hr c. Desorption at 100-425°C; 12-14 hr  $F_{\odot}$  at ~ 18 standard liters/min  $UF_6$ · 3 NaF ----->  $UF_6$  + 3 NaF 2. Vessel was removed from service and left exposed while the remainder of the system was water washed, dried, and treated with  $F_{2}$ . Vessel was then placed back in service. 3. One (1) special run whose purpose was preparation of NaF from  $NaHF_2$ . a. Purge-sparge with  $\rm N_{2}$  at  $\sim$  3 standard liters/min for 12 hr 25-125°C in 7 hr at 125°C for 5 hr b. Preparation of NaF under  $N_{2}$  at 3 standard liters/min by  $\operatorname{NaHF}_2$  +  $\Delta \xrightarrow{}$  NaF + HF 125-600°C in 9 hr at 600°C for 4 hr Conditioning of absorber at 635°C with  $F_2^*$ с. at ~ 15 standard liters/min for 0.5 hr

\*Failure is thought to have occurred at this time.

# - 171 -

#### FV-120 FIRST ABSORBER

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#### Exterior

Shell --- Covered with a thin black adherent film, presumably oxide.

Bottom - Covered with a thin black adherent film, presumably oxide.

Welds - All beads oppeared to be in good condition.

Vidigage Inspection: (Nondestructive Test Section operators expressed "no confidence" in front and back readings due to the wall deposit and operational difficulties.)

#### Wall Thickness in Inches

Distance <sup>1</sup>	Location Remarks	Front Series	Back Series <sup>2</sup>	Bottom Series <sup>3</sup>
2	Neck	0.226	0.235	0.370
5	Reducer	0.222	0.220	0.374
8	Shell	0.236	0.224	0.370
10	Shell	0.223	NA	
12	Shell	0.230	NA	
15	Shell	0.236	NA	
18	Shell	0.240	NA	
20	Shell	0.238	0.235	
23	Shell	0.241	0.238	
25	Shell	0.242	0.240	
28	Shell	0.244	0.234	

1. In inches, from bottom of flange.

2. These readings taken 180° from front series.

3. Random readings, ±0.002 in.

NA: Not Available

Fig. 86. Results of Visual and Vidigage Inspection on VPP First Absorber.

#### FV-121 SECOND ABSORBER



#### Exterior

- Shell -- Covered with a thin black adherent film, presumably oxide.
- Bottom Covered with a thin black adherent film, presumably oxide.
- Welds All beads appeared to be in good condition.

#### Vidigage Inspection: (Nondestructive Test Section operators expressed "no confidence" in front and back readings due to wall deposit and operational difficulties.)

Wall Thickness in Inches

D	istance <sup>1</sup>	Location	Front Series	Back Series <sup>2</sup>	Bottom Series <sup>3</sup>
	2	Neck	0.228	0.232	0.370
	5	Reducer	0.222	0.230	0.374
	8	Shell	0.220	0.230	0.372
	10	Shell	0.222	0.230	
	12	Shell	NA	0.228	
	15	Shell	NA	0.228	
Ciah	18	Shell	NA	0.226	
Wala	20	Shell	NA	0.230	
wera	23	Shell	0.228	0.237	
	25	Shell	0.232	0.240	
	28	Shell	0.238	0.238	

1. In inches, from bottom of flange.

2. These readings taken 180° from front series.

3. Random readings,  $\pm 0.002$ .

NA: Not Available

Fig. 87. Results of Visual and Vidigage Inspection on VPP Second Absorber.

# Valves and Fittings

Aside from sporadic leakage and some plugging by unidentified deposits of valves carrying UF<sub>6</sub>, valves and fittings in the VPP performed satisfactorily.<sup>57</sup> Many screwed fittings carrying nitrogen were back-welded after various amounts of plant operational time to eliminate such leakage.

Fluorine carrying values of various body materials, i.e., Monel, nickel, Inconel, steel, types 316 and 347 stainless steel, brass, and copper, were welded or brazed into the associated piping. At temperatures greater than 150°C, only nickel and Inconel values were in contact with fluorine under normal operating conditions.

One failure, originally thought to be in a valve but later discovered to be in the adjacent fittings, has been selected for treatment here because of the materials involved and their reaction to service conditions.

Following Run L-4 in the VPP, leaded, yellow brass valve fittings of the composition 60-63% Cu-2.5-3.7% Pb-0.35% Max Fe-0.5% other-bal Zn (vendor's analysis) were removed from a nitrogen purge line, PE-34, because of defective valve operation.

The inlet side of the valve, a packless diaphragm-seal valve with an Inconel diaphragm, Duranickel stem point, and Monel body and trim, was placed in conjunction with the NaF absorber outlet piping which carried  $F_2$  and  $UF_6$ . Under normal operating conditions, the valve was not in contact with process gases, but there was continuous nitrogen contact. The valve was shut only when nitrogen pressure was low and there was the possibility of process gases diffusing past the valve to the instrumentation units.

During the early L-runs, persistent system plugs and low pressure in the nitrogen line probably permitted  $F_2$  and UF<sub>6</sub> to contact the value and its fittings for an unknown period of time. The operating service time on the value and its fittings was approx 6 weeks.

Visual examination of the value disclosed no obvious defects. Therefore, it was subjected to several times operating pressure (15 psig) by VPP personnel and found to function satisfactorily without leakage. The value was subsequently placed in stock to be used as an emergency replacement. Visual observation of the fittings revealed that several components had ruptures running the complete length of the walls. The fissures were normal to stresses induced during installation. Figure 88 and Table XXIV show and describe the defects noted. Metallographic examination of the fittings revealed the cracks to be intergranular in nature. Dezincification apparently occurred along the inside edges of the fissures, as evidenced by copper-rich deposits which lined the fissures. (See Fig. 89.) Copper deposits were also evident in several grain boundaries radiating outward from the flaw. General corrosive attack was noted on the outside of the inlet union, Fig. 90.

Although all evidence indicates that failure was by selective intergranular dezincification, the stresses induced during installation were believed to have provided a strong directional driving force. The use of stock Monel fittings, because of its increased corrosion resistance, was recommended for use in the environment described.

The associated Monel valve was found to be in good operating condition, appeared satisfactory, and can continue to be used for the mentioned service conditions.

This failure serves to point out the necessity for constant vigilance in selecting materials and components for a highly corrosive process scheme. This includes the selection of piping and joining couples which should, if at all possible, match the components in corrosion-resistant properties.

# Fluorine Disposal System

Excess fluorine and nitrogen from the VPP have been disposed of by cocurrent contact with an aqueous KOH solution in a gas-disposal unit. (See Figs. 79 and 91.) The KOH solution was prepared in a caustic surge tank, shown in Figs. 79 and 92, to a concentration of 2-10% KOH and pumped to the disposal unit. The caustic solution was then sprayed into the disposal unit as the excess process gases entered the vessel. Fluorine in the waste gases formed potassium fluoride in water solution according to

> $2F_2 + 2KOH \longrightarrow F_2O + 2KF + H_2O$  $F_2O + 2KOH \longrightarrow 2KF + O_2 + H_2O$


(Note Fig. 88. Photograph of Gas Purge Line (PE-34) Valve and Attached Fittings. fractures in inlet union, inlet mut, and inlet front ferrule.)

- 175 -

# Table XXIV. Defects Found in Leaded Yellow Brass Fittings Subjected to ${\rm F_2}$ and ${\rm UF_6}$

Part Name	Location	Defect
Union	Inlet side of valve	* Wall fracture * through threaded end covered by hex nut. Fracture extended through hex body and part of opposite threaded end.
Hex Nut	Inlet side of valve	Wall fracture * running entire length of ferrule.
Front Ferrule	Inlet side of valve	Wall fracture * running entire length of ferrule.
Back Ferrule	Inlet side of valve	$Crack^{\star}$ partially through wall.
Union		Wall fracture partially through threaded end covered by hex nut. (Not shown in Fig. 88 because of photo orientation.)

\* All fractures and cracks were essentially in line with one another (see Fig. 88).

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Fig. 89. VPP Leaded, Yellow Brass Inlet Back Ferrule Showing Intergranular Fracture Lined with Copper-Rich Deposits Indicative of Dezincification. Etchant: NH<sub>4</sub>OH-H<sub>2</sub>-H<sub>2</sub>O. 250X.



Fig. 90. VPP Leaded, Yellow Brass Inlet Union Showing General Corrosive Attack. Etchant:  $NH_4OH-H_2O_2-H_2O$ . 100X.

## - 179 -

UNCLASSIFIED ORNL-LR-DWG 49175



Distance <sup>1</sup>	SHE Circumf Seri	erential es <sup>2</sup>	SHELL Inter-KOH Nozzle Series	SHELL F <sub>2</sub> Inlet-Impingement Series <sup>3</sup>	HEAD Series
1	0.119	0.118	No. 12	0.126	Dished Top
	0.118	0.118	0.125	0.126	0.182
	0.118	0.119	0.126	0.127	0.184
			0.127	0.126	0.184
9	0.124	0.125	0.127	0.127	0.186
	0.126	0.126		0.127	0.183
	0.125	0.125	No. 2-3	0.128	0.182
			0.128	0.128	
24	0.130	0.125	0.128	0.128	Flanged Side
	0.129	0.127	0.126	0.128	0,185
	0.128	0.130		0.128	0.184
	477125			0.129	0.189
39	0.125	0.126	No. 3-4		0.190
	0.127		0.124		0.189
			0.128		0.186
54	0.130	0.126			
-	0.129	0.128	No. 4-5		
	0.127	0.129	0.129		
			0.128		
69	0.127	0.129			
•	0.129		No. 5-6		
			0.124		
			0.125		
84	0.128	0.128			
-	0.129	0.128			
	0.128	0.129			
99	0.130	0.128			
	0.129				

1. Distance, vertically down, in inches starting at top head to shell weld -- reference only to shell

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District, veniculty down, in these starting a top field to shart need a terrative entry to circumferential series.
 Readings taken at 60° or 120°; rotated ~ 20° for each successive group.
 Readings at 1 in. intervals starting 13 in. below the head to shell weld 180° from F<sub>2</sub> inlet.
 All readings, ±0.002 in.

Fig. 91. Results of Visual and Vidigage Inspection on VPP Gas Disposal Unit.





FV-152 SURGE TANK .

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- 180 -

which subsequently was discharged as chemical waste while the nitrogen remaining in the excess gases was exhausted to the atmosphere. The entire fluorine disposal system was fabricated from Monel.

After VPP Run L-9, visual and ultrasonic thickness measurements were taken on the gas-disposal unit and the surge tank to ascertain their suitability for additional VPP service. The respective service conditions for the two vessels and the results of those inspections are presented in Figs. 91 and 92. A maximum wall-thickness loss of 5 mils was found near the top head of the gas-disposal unit while no detectable metal losses were found in the surge tank.

Previous studies have indicated that heavy corrosive attack can occur on the process gas inlet pipe inside the gas-disposal unit.<sup>58</sup> Therefore, an inlet pipe fabricated from 1-in. sched-40 Monel pipe in service for approx 4900 hr was removed from the disposal unit and subjected to visual and metallographic examination. Figure 91 shows the location of the gas inlet pipe.

Visual examination of the exterior of the pipe inlet disclosed the presence of two holes, approx 1/32 in. and 3/32 in. in diameter, respectively, on the upper half of the pipe. The holes completely penetrated the pipe wall and were 1-1/2 in. and 2-1/2 in. from the exit end of the pipe. Inspection of the interior of the nozzle showed an extremely rough surface and a severe pitting attack extending from the end of the nozzle inwards for approx 3 in. A white crystalline deposit encircled the interior at a point 4 in. from the end of the pipe. Subsequent x-ray diffraction analysis indicated this deposit to be wholly  $KHF_0$ .

Metallographic examination of sections removed from the pipe showed a roughened surface and metal losses concentrating in circular cavities but no evidence of intergranular attack. The holes described above were simple extensions of the localized attack. Figure 93 shows a magnified view of the pipe sectioned through the center of the largest hole found in the inlet pipe. A photomicrograph of the surface at the base of one of the pits is shown also in this figure.

<sup>58</sup>L. R. Trotter and E. E. Hoffman, <u>Progress Report on Volatility Pilot</u> Plant Corrosion Problems to April 21, 1957, ORML-2495 (September 30, 1958).





Fig. 93. Cross Sections of Gas Disposal Unit Inlet Pipe (a) Photomacrograph Showing Severe Pitting Attack. 15X. (b) Portion of Fig. 93A showing typical surface at base of pit. 500X. Etchant: Acetic-nitric-hydrochloric acid.

- 182 -

Corrosion seemed to have been directional from the interior toward the exterior of the pipe. Examination of the fluorine-nitrogen entrance end of the pipe disclosed no noticeable or measurable attack.

Intermittently during operations, the gas-inlet nozzle plugged (presumably by KF and/or  $\text{KHF}_2$  deposits) and this condition occurred with such frequency during the "E" and early "L" runs that a secondary nozzle was inserted in place of a lower KOH inlet. Examination of this secondary nozzle, which had approx 100 service hours, disclosed a very slight attack similar to that found on the primary pipe.

The presence of  $\text{KHF}_2$  inside the main gas-inlet pipe is believed to occasion the high localized corrosion found in that component. It would appear that an easy way for  $\text{KHF}_2$  to form in the pipe would be by the alkali deficient reaction of KOH with HF as

 $2HF + KOH \longrightarrow KHF_2 + H_2O.$  (ref 59)

The formation of hydrogen fluoride seems possible from the reactions of fluorine monoxide or fluorine with the available water of solution present. Investigators have reported the hydrolysis reaction of  $F_00$ 

F<sub>2</sub>O + H<sub>2</sub>O -----> O<sub>2</sub> + 2HF

to be very slow at ordinary temperatures but a steam- $F_2^0$  reaction occurs so rapidly that it can be considered explosive.<sup>60</sup> For reasons unknown, the reaction of fluorine with water has been reported as not always occurring.<sup>61</sup> One source has indicated that fluorine reacts with water easily, giving oxygen mixed with ozone, hydrogen peroxide, and other oxidizing substances.<sup>62</sup>

If hydrogen fluoride is present in the gas-inlet pipe, then the high degree of localized attack on the nozzle may be explained as the result of

<sup>59</sup>G. I. Cathers, Chem. Tech. Div., ORNL, Private communication.

<sup>60</sup>A. B. Burg, "Volatile Inorganic Fluorides," <u>Fluorine Chemistry</u> (ed. by J. H. Simons) p.83, Academic Press, New York, 1950.

61. R. Landau and R. Rosen, "Industrial Handling of Fluorine," Preparation Properties and Technology of Fluorine and Organic Fluoro Compounds (ed. by C. S. Lesser and S. R. Schram) NNES VII-I, p.154, McGraw-Hill, New York, 1951.

<sup>62</sup>N. W. Sidgwick, <u>The Chemical Elements and Their Compounds</u>, Vol. II, p. 1101, Oxford University Press, London, 1950. hydrogen fluoride corrosion highly accelerated by the presence of oxygen. Also, at particular concentrations of hydrogen fluoride in water, close to the azeotropic composition (38 wt %), one investigator has reported corrosion rates on Monel of 145 mils/yr at 120°C. (ref 63) It may be that either or both of these reactions may be responsible for the localized attack described.

Considering the easy replacement of the gas-inlet nozzle and the major corrosion problems remaining to be solved in the VPP, no development work has been planned toward finding an improved material of construction.

#### Process Gas Lines

Sections of Monel pipe and copper tubing near the VPP absorbers, chemical traps, and cold traps were removed from the process gas piping system after Run M-64 and sent to BMI for corrosion analyses. A schematic drawing showing the location of these sections is presented in Fig. 94, and a description of the piping and the individual process environments can be found in Table XXV.

The wall thicknesses of the sections were measured by BMI and no indication of significant metal losses were found. The metallographic examinations conducted on the specimens disclosed no serious corrosive attack. Almost all of the specimens showed isolated pitting but the pits were less than one grain deep. Slight indication of selective attack at the grain boundaries was observed on specimens 1-1, 1-2, 1-3, 2, and 3 which were located close to the absorbers and the first cold trap. However, the intergranular attack was never deeper than one grain, and for the most part, was less than 1 mil deep.

<sup>63</sup>W. Z. Friend, "Nickel-Copper Alloys," Corrosion Handbook (ed. by H. H. Uhlig) p. 273, John Wiley and Sons, New York, 1948.



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Fig. 94. Diagrammatic Flowsheet Showing Location of Process Gas Line Specimens Removed from the VPP System.

				Expo	sure	
	Specimen E	Description	Fused-Salt	C	Cooling After	-
Number	Location	Material	Fluorination-UF <sub>6</sub> Sorption	Vesorption	Desportion	roduct Kemova
1-1 1-2 1-3	Ahead of first absorber	$1\frac{1}{4}$ and $1\frac{1}{2}$ in. sched-40 Monel pipe	200 hr, 110-140°C, F2 and UF6 (30 hr with 90% UF6)	350 hr, 110-140°C, F_2	350 hr, 110-140°C, N₂	None
2	Outlet from second absorber	$1 ight)_4$ in. sched-40 Mone! pipe	200 hr, 75-150°C, F <sub>2</sub>	350 hr, 75-350°C, F 2 and UF 6	350 hr, 75-350°C, N2	None
ო	Iniet to first cold trap	$1  rac{1}{3}$ and 1 in. sched-40 Monel pipe; 90 deg 'ell' and reducer	200 hr, 140-165°C, F 2	350 hr, 140-365°C, F <sub>2</sub> and UF <sub>6</sub>	350 hr, 140-165°C, N <sub>2</sub>	190 hr, 140-1 <b>3</b> 5°C, 0-55 psia, UF <sub>6</sub>
4	Outlet from first cold trap	$ ho_2$ in. sched-40 Monel pipe	200 hr, 110-135°C, F <sub>2</sub>	350 hr, 110-135°C, F <sub>2</sub>	350 hr, 110-135°C, N₂	190 hr, 110-135°C, 0-55 psia, UF <sub>6</sub>
Ś	Bypass vaive of second cold trap	$^{ m N}_2$ in. sched-40 Monel pipe	200 ir, 110-135°C, F <sub>2</sub>	350 hr, 110-135°C, F <sub>2</sub>	350 hr, 110-135°C, N <sub>2</sub>	190 hr, 110-135°C, 0-55 psia, UF <sub>6</sub>
Ŷ	Ahead of chemical trap	1/2 and 1 in. sched-40 Monel pipe; straight 'T' reducer and 90-deg 'ell'	200 hr, 100-150°C, F_2	350 hr, 100-150°C, F_2	350 hr, 100-150°C, N <sub>2</sub>	190 hr, 100-150°C, 0-55 psia, UF <sub>6</sub>
7	Outlet from chemical trap	1 in. sched-40 Monel pipe	200 hr, 35-60°C, F <sub>2</sub>	350 hr, 35-60°C, F_2	350 hr, 35-60°C, N <sub>2</sub>	None
8-2 8-3 3-2	Inlet side of product receiver	3, in. copper tubing	None	None	None	190 hr, 66-100°C, ⊍F <sub>6</sub>
9-1 9-2 9-3	Outlet side of product receiver	$rac{3}{8}$ in. copper tubing	None	впе	None	190 հւ, ઠઠ-100°C, ՍԲ <sub>ઠ</sub>

TABLE XXV. LOCATION, DESCRIPTION, AND EXPOSURE CONDITIONS FOR SPECIMENS FROM PROCESS GAS LINES

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