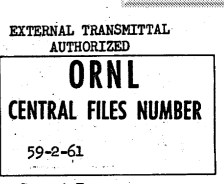
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SUBJECT:	Processing of Molten Salt Power Reactor Fuel
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FROM:	D. O. Campbell and G. T. Cathers

## ABSTRACT

-Fuel reprocessing methods are being investigated for molten salt nuclear reactors which use LiF-BeF<sub>2</sub> salt as a solvent for UF<sub>h</sub> and ThF<sub>h</sub>. A liquid HF dissolution procedure coupled with fluorination has been developed for recovery of the uranium and LiF-BeF<sub>2</sub> solvent salt which is highly enriched in Li-7. The recovered salt is decontaminated in the process from the major reactor poisons; namely, rare earths and neptunium. A brief investigation of alternate methods, including oxide precipitation, partial freezing, and metal reduction, indicated that such methods may give some separation of the solvent salt from reactor poisons, but they do not appear to be sufficiently quantitative for a simple processing operation.

Solubilities of LiF and BeF<sub>2</sub> in aqueous 70-100% HF are presented. The BeF<sub>2</sub> solubility is appreciably increased in the presence of water and large amounts of LiF. Salt solubilities of 150 g/liter are attainable. Tracer experiments indicate that rare earth solubilities, relative to LiF-BeF<sub>2</sub> solvent salt solubility, increase from about 10<sup>-4</sup> mole % in 98% HF to 0.003 mole % in 80% HF.

Fluorination of uranium from LiF-BeF<sub>2</sub> salt has been demonstrated. This appears feasible also for the recovery of the relatively small concentration of uranium produced in the LiF-BeF<sub>2</sub>-ThF<sub>h</sub> blanket.

A proposed chemical flowsheet is presented on the basis of this exploratory work as applied to the semicontinuous processing of a 600 Mw power reactor.

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#### INTRODUCTION

High temperature fluid fuel reactors using molten fluorides have been proposed for the production of nuclear power.<sup>1,2</sup> The core or blanket salt in this type reactor would be ideally a  $\text{Id}^7\text{F-BeF}_2$  mixture for optimum neutron moderation and economy. This material would act as a solvent or carrier for the fluorides of the fissile or fertile elements, uranium, plutonium and thorium. The feasibility and economic justification of such a reactor system depends on fuel processing, i.e., a processing method that will maintain the desired neutron economy and reactor operability at a reasonable cost. This paper presents a description of a new chemical process for this type of fluoride fuel based on two principles, namely, volatilization of UF<sub>6</sub> from the reactor salt by fluorination and recovery of the Id<sup>7</sup>F-BeF<sub>2</sub> solvent salt for reuse by a HF dissolution process.

## MOLITEN SALT REACTOR DESCRIPTION

The Aircraft Reactor Experiment, in which NaF-ZrF4-U<sup>235</sup>F4 fuel circulated through inconel tubing in BeO moderator, demonstrated the basic feasibility of a high temperature molten salt reactor<sup>1</sup>. Detailed calculations for a power reactor have been published on 600 Mw heat two-region homogeneous machines using 63 mole %Li<sup>7</sup>F--37 mole % BeF<sub>2</sub> with up to 1 mole % UF4 and ThF4 as a core salt and 71 mole %Li<sup>7</sup>F--16 mole % BeF<sub>2</sub>--13 mole % ThF4 as a blanket salt<sup>2</sup>. The reactor design used as reference in this paper has a uranium (U<sup>2</sup>33 or U<sup>2</sup>3<sup>5</sup>) inventory of 600-1000 kg (varying with time) with an 8-ft-dia core and a total fuel volume of 530 ft<sup>3</sup> (334 ft<sup>3</sup> external to the core). The core salt weight is approximately 75,000 lb, the blanket 150,000 lb.

In the reference reactor at least 90% of the power is produced in the core, yielding about 180 kg of fission products per year. After operation for one year without processing (except inert gas removal), the fission products absorb about 3.8% of all neutrons and  $U^{-36}$  and  $U^{-38}$  absorb about 3.9%. If the reactor fuel is processed for fission product removal at the rate of one fuel volume per year, after 10 years, the fission products would absorb 2.7% of the neutrons;  $U^{-34}$ ,  $U^{-36}$ , and  $U^{-38}$  would absorb 10.4\% (mostly  $U^{-36}$ ); and Np<sup>237</sup> about 0.9\%. Without fuel processing neutron absorption by fission products would continue to increase almost linearly with time, exceeding 10% after 10 years; Np<sup>237</sup> would build up at an accelerating rate, absorbing about 3% of the neutrons after 10 years; the amount of fissionable material required to keep the reactor critical would increase by about 200 kg/year; and the conversion ratio would decrease markedly.

Even-numbered uranium isotopes, particularly  $U^{236}$ , are the worst poisons, but their removal is beyond the scope of chemical reprocessing. Of the 180 kg of fission products per year 22 atom % with half-lives of more than 78 min are

<sup>1</sup>R. C. Briant, A. M. Weinberg, et al, "The Aircraft Reactor Experiment," Nucl. Sci. and Eng., 2, 797-853 (1957).

<sup>2</sup>J. A. Lane, H. G. MacPherson and F. Maslan, "Fluid Fuel Reactors," Addison-Wesley Publishing Co., Inc., 1958. subject to removal from the reactor as rare gases; these would contribute 26% of the fission product poisoning for 100 ev neutrons. About 26 atom % of the long-lived fission products are rare earths, which contribute 40% of the total fission product poisoning. The rest of the fission products consist of a wide variety of elements, no one of which is outstanding from the nuclear poisoning point of view. After reasonably long operation Np<sup>237</sup> is the worst individual poison other than the rare earths.

## FLUORIDE VOLATILITY PROCESS FOR MSR FUEL

Fluoride volatilization processing for uranium recovery appears feasible for molten salt reactor (MSR) fuel on the basis of laboratory studies. It is based on direct fluorination of the fuel salt to convert UF<sub>4</sub> to UF<sub>6</sub> with attendant volatilization and recovery. Similar volatility processes have been proposed and developed for zirconium alloy reactor fuel elements after dissolution in fused salt.<sup>3,4</sup> One of these, the ORNL Volatility Process, was successfully used for recovery and decontamination of uranium from the NaF-ZrF<sub>4</sub>-UF<sub>4</sub> salt fuel of the Aircraft Reactor Experiment.<sup>5</sup> The MSR volatilization process would differ, however, from other volatility processes in that complete decontamination of the product UF<sub>6</sub> would not be essential, since it could be remotely reduced to UF<sub>4</sub> and reconstituted into reactor salt.

A series of small-scale fluorinations was carried out with a 48 mole %LiF--52 mole % BeF<sub>2</sub> eutectic mixture containing about 0.8 mole % UF<sub>4</sub>. (MSR fuel would contain 0.25 to 1.0 mole % UF<sub>4</sub>, depending on the operating time.) The eutectic salt was used instead of the fuel salt in order to investigate lower temperature operation. In fluorinations at 450, 500, and 550°C, the rate of uranium removal increased with the temperature (Table 1).

The thorium-containing blanket salt cannot be processed for uranium recovery at as low a temperature as that used to process the fuel salt.

<sup>3</sup>G. I. Cathers, "Uranium Recovery for Spent Fuel, "Nucl. Sci. and Eng. 2, 768-777 (1957).

<sup>4</sup>H. H. Hyman, R. C. Vogel and J. J. Katz, "Proceedings of International Conference on Peaceful Uses of Atomic Energy," Vol. 9, pp. 613-626, United Nations, New York (1956).

<sup>9</sup>G. I. Cathers et al., "Recovery of Uranium from Highly Irradiated Fuel," United Nations Paper 535, 2nd International Conference on Peaceful Uses of Atomic Energy (1958).

## Table 1. Effect of Fluorination Temperature on the Fluorination of Uranium from LiF-BeF<sub>o</sub> (48-52 mole %)\*

Fluorination	Uranium in	Salt after Tre	atment, wt %
<u>Time, hr</u>	At 450°C	At 500°C	At 550°C
- 1910 - 1910 <b>O</b>	3.39**	5.10	4.91
0.5	<b>1.9</b> 6	0.20	0.55
1.0	0.39	0.17	0.20
1.5	0.21	0.12	9.06
2.5	0.32	0.11	0.05

\* No induction period before uranium evolution.

 $^{**}$ 5 wt% added; some of the uranium probably precipitated as oxide.

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The uranium concentration in the blanket, however, is very low; it has been estimated that with continuous processing at the rate of one blanket volume per year, the blanket salt (LiF-BeF<sub>2</sub>-ThF<sub>4</sub>, 71-16-13 mole %) will contain approximately 0.004 mole % UF<sub>4</sub> (140 ppm) after one year and 0.014 mole % UF<sub>4</sub> after 20 years. Fluorinations of two such mixtures at 600°C for 90 min gave uranium concentrations in the salt of 1-2 ppm, the lowest uranium concentrations ever obtained in fused salt laboratory fluorinstions. Over 90% of the uranium was removed in 15 min. It is concluded, therefore, that fluorination of uranium from blanket salt can be accomplished.

The behavior of protactinium in the blanket salt during fluorination is of interest, although the protactinium is not lost, in any case, since the salt is returned to the reactor. A LiF-BeF<sub>2</sub>-ThF<sub>4</sub> (71-16-13 mole %) mixture containing sufficient irradiated thorium to give a Pa<sup>2</sup>33 concentration of 5.5 x 10<sup>-9</sup> g per gram of salt was fluorinated for 150 min at 600<sup>o</sup>C; there was no measurable decrease in protactinium activity in the salt. Protactinium volatilization in the process seems to be unlikely. However, the protactinium concentration in the blanket of the reference design reactor is higher (~10<sup>-4</sup> g per gram of salt).

117F-BeF, SALT RECOVERY WITH HF

Experimental work has demonstrated that the LiF-BeF<sub>2</sub> salt can be processed by dissolution in anhydrous or nearly anhydrous liquid hydrogen fluoride. Decontamination of the LiF-BeF<sub>2</sub> salt from the major neutron poisons, rare earths and neptunium, is achieved due to the relative insolubility of the fluorides of these elements in such solutions. The LiF-BeF<sub>2</sub> salt is recoverable from the HF solution by evaporation.

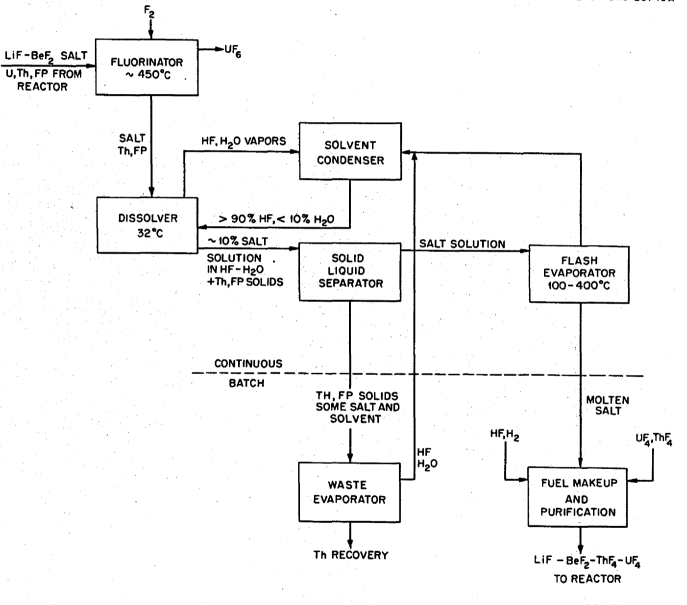
It is well known that BeF<sub>2</sub> is very soluble and LiF is rather insoluble in water in contrast to liquid HF where the reverse is true. In anhydrous HF the polyvalent element fluorides generally exhibit low solubilities.<sup>6</sup> Initial consideration of the problem suggested that use of aqueous HF (greater than 80% HF) would give sufficient solubility of both LiF and BeF<sub>2</sub> to meet process objectives. The solubility studies were therefore carried out over the range of 70-100% HF. However, in addition to the mixed solvent effect there existed also the possibility that anhydrous HF would be suitable as a solvent for the LiF-BeF<sub>2</sub> salt complex in the analogous sense that cryolite, Na<sub>3</sub>AlF<sub>6</sub>, is quite soluble whereas AlF<sub>3</sub> is insoluble in HF.<sup>7</sup> Some emphasis was therefore placed on using "anhydrous" HF with the material being obtained by vapor transfer from a commercial tank (nominally containing less than 0.1½ water).

Initial measurements of the solubilities of LiF and BeF<sub>2</sub>, separately, in aqueous HF solutions indicated that both materials are soluble to an appreciable extent in solutions containing 70 to 90 wt % HF (Tables 2 and 3). In general,

<sup>6</sup>A. W. Jache and G. W. Cady, J. Phys. Chem., <u>56</u>, 1106-1109 (1952).

<sup>1</sup>L. F. Audrieth and J. Kleinberg, "Non-Aqueous Solvents," Chapter 10, John Wiley and Sons, Inc., New York (1953).

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Fig. 2. Tentative Flowsheet for Fluoride Valatility and HF Dissolution Processing of Molten-Salt Reactor Fuel.

		LiF in Solu	ution, mg/g of	solution	
Temperature,	73.6 wt % HF	82.3 wt % HF	90.8 wt % HF	96.2 wt % HF	100 wt % HF
12	10.7	31.7	58.4	74.8	88.2
62*	20.6			·	
47*		41.0			
37*		· · · · · · · ·	62.8	80.4	
32.5*					91.4

## Table 2. Solubility of LiF in Aqueous HF Solutions

\* Reflux temperature.

## Table 3. Solubility of BeF2 in Aqueous HF Solution

Temperatu °C	re,	72.8 wt % HF	BeF2 in Solu 78.3 wt % HF	ution, mg/g ( 90.8 wt % HF	of solution 95.2 wt % HF	100 wt % HF
12	•	45.8	26.3	9.2	2.8	0.0012
-60	•				3.8	

LiF was more soluble at higher temperatures if water was present in the solvent, but the effect of temperature on  $BeF_2$  solubility was not definitely established. The LiF solubility decreased rapidly as water was added to anhydrous HF, and the BeF<sub>2</sub> solubility increased from near zero; the solubilities were roughly the same in 80 wt % HF, 25 to 30 g/kg. The BeF<sub>2</sub> was glassy in pature and dissolved slowly. The solubility values reported for LiF in Table 2, except at 12°C, were obtained after refluxing HF over the salt for 3 hr. No further measurements were made with LiF or BeF<sub>2</sub> alone, since the solubilities of the two components together were of primary importance.

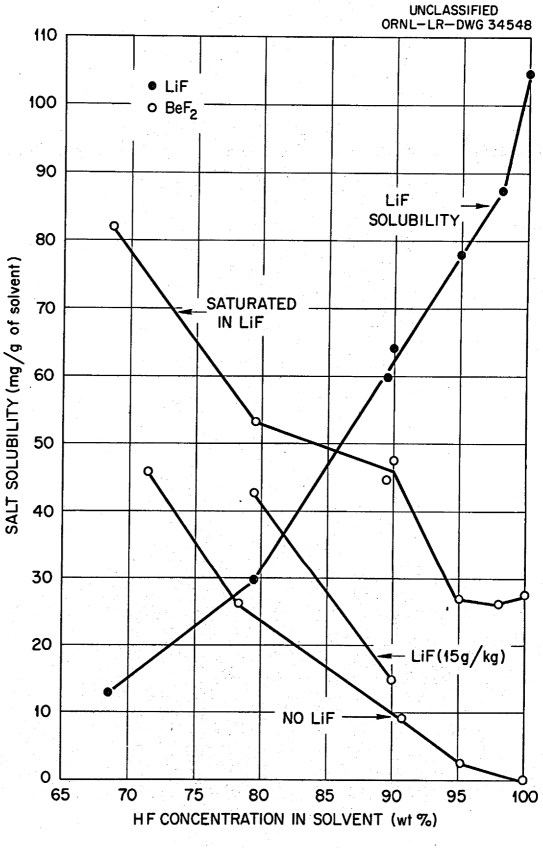
The solubility of BeF<sub>2</sub> was also measured at various HF concentrations and temperatures in the presence of LiF. The results generally confirmed that the presence of LiF as well as water increases the BeF<sub>2</sub> solubility. A plot of solubility values at one temperature  $(12^{\circ}C)$  illustrates both effects (Fig. 1). These and data at other temperatures are presented in Tables 4, 5, and 6. The favorable effect of water on BeF<sub>2</sub> solubility was enhanced further when 70 and 80% HF solutions (Table 4). BeF<sub>2</sub> solubility was enhanced further when 70, 80, and 90 wt % HF solutions were saturated with LiF (Table 5). Because of the increase in BeF<sub>2</sub> solubility at higher HF concentrations when LiF is present, the 90-100% HF range was studied further (Table 6). The solubilities in approximately 100% HF appear sufficiently high for process purposes.

The results presented here must be considered preliminary. Beryllium solubilities, in particular, may be generally higher than indicated because of analytical problems. Difficulty was also encountered in determining accurately the water content in highly concentrated HF. This was partly a problem of sampling the volatile solutions and partly the result of determining the small amount of water by the difference of the sample weight and the total weight of LiF, BeF<sub>2</sub>, and HF.

Solubility measurements were made also on 63-37 mole % LiF-BeF<sub>2</sub> salt (48-52 wt %) in the 80 to 98 wt % HF range (Table 7). This salt contained about 0.1 mole % ZrF<sub>h</sub>; 0.2 mole % mixed rare earth fluorides (Lindsay Code 370), and trace fission products (see following section). The salt was crushed but not ground to a powder; average particles were flakes about 10 mils thick and 50 to 100 mils across. The solutions were sampled 15 and 60 min after salt addition to permit an estimate of the rate of dissolution. The results indicated that an appreciable concentration is reached rapidly, but that the solutions are not saturated, especially with respect to BeF<sub>2</sub>, in less than several days. They also demonstrate that the fused salt behaves similarly to the two components added individually.

## FISSION PRODUCT DECONTAMINATION IN THE HF SOLUTION PROCESS

The aqueous HF solutions of the salt from the preceding experiment were filtered through sintered nickel, and radiochemical analyses were made to determine the fission product solubilities or decontamination effect. The results (Table 8) show that rare earths, as represented by cerium, were relatively insoluble in the HF solution and were therefore effectively separated from the salt.





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		BeFo in Solution, mg/g of solution			
Temperature,	LiF Added, mg/g_of_solvent	68.6 wt % HF	79.5 <u>wt % HF</u>	90.0 <u>wt % HF</u>	
12	7.0	68.8	,		
12	15.0		42.8	15.6	
-60	7.0	65.8			
-60	15.0		38.2	16.5	

## Table 4. Solubility of BeF2 in Aqueous HF Containing LiF

## Table 5. Solubility of LiF and BeF<sub>2</sub> in Aqueous HF Saturated with Both Salts

Temperature,	68.6 wt	Salt in % HF	n Solution, 79.5 v	ng/g of a rt % HF	90.0 wt	% HF
<mark></mark>	LIF	BeFe	LIF	BeF <sub>2</sub>	LIF	BeF2
12	12.9	82.4	29.9	53.8	64.2	48.2
-60	8.5	76.8	22.6	54.2	46.0	58.2

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	Salt in Solution, mg/g of solution							
Temperature,	89.5	<u>wt % HF</u>	95	wt % HF	<u>98. v</u>	rt % HF	100 w	
<u> </u>	LIF	BeFe	LIF	BeFe	Idf	BeFe	LIF	BeFe
12	60	45	78	27	92	26	107	28
-60	50	24	60	33	62	32	90	30
43*	69	68						
40 <del>×</del>			87	66				
<b>3</b> 6*	-				98	46		
33*					· ·		105	49
¥. 5. •		:						

Table 6.	Solubility	of	IJF	and	BeFo	in	Solvents	Containing
		89	.5 to	<b>)</b> 100	) wt (	βH	7	

Reflux temperature.

Table 7. Solubility in Aqueous HF Solution of LiF and BeF<sub>2</sub> from

Salt Mixture Contain:	ing 63 mole 9	LIF	and 37	mole 9	BeF.

•		Salt in Solution, mg/g of solution								
	Temperature,		wt % IIF			95 w			-98 wt % HF	
Time	<u>o</u> C	<u>Fit</u>	BeFe	LIF	BeFe	IIF	BeFe	LIF	BeFe	
15 min	12	.28	33	50	44	40	17	28	4	
l hr	and a second second Second second second Second second	29	35	51	50	43	17	22	3	
20 hr		31	58	68	74			• •	· · ·	
4 days		32	68*	79	82*	63	28	70*	22	
5 hr	-60	21	65	74	102	70	36	71*	22	
20 hr	12	34	100	76	92	88*	53	60*	22	

Essentially all the component indicated had dissolved and therefore the solubility may be higher than the value given. Insufficient salt was added to some of the solutions to ensure an excess of both components.

Table 8. <u>Fission Product Solubilities in Aqueous HF Solutions</u> LiF-BeF<sub>2</sub> (63-37 mole \$) + ~0.2 mole \$ rare earth fluorides + trace fission products between 1 and 2 years old

Fission Product*	Activity in Original Salt and in HF Solution, counts/min/g of salt							
	Original Salt	79.5 10t % HF	89.5 wt <u>% EF</u>	95 wt % HF	98 wt % HF			
Gross B	745 x 10 <sup>4</sup>	$230 \times 10^4$	$200 \times 10^4$	225 x 10 <sup>4</sup>	250 x 10 <sup>4</sup>			
Gross 7	228	293	213	244	282			
Cs 7	174	251	<b>19</b> 6	223	258			
Srβ	104	73	67	72	75			
TRE B	650	97	92	94	101			
Cēβ	510	7.9	3.6	1.3	0.28			
<sup>у90</sup> в	105	73	66	74	81.5			

<sup>\*</sup>Zr and Nb precipitated from molten salt before this experiment and were not present in significant concentration. The solubility decreased as the HF concentration was increased. The total rare earth (TRE) and trivalent rare earth (except cerium) analyses do not show this separation because of the presence of the yttrium daughter of strontium. No rare earth activity other than cerium (and yttrium) was detected in the HF solutions. The slight apparent decontamination from strontium is not understood; strontium is expected to be fairly soluble in these solutions. Cesium is known to be soluble, and all cesium in the salt added apparently dissolved.

Thus the rare earths, as represented by cerium, are removed from LiF-BeF salts by dissolution of the salt in an aqueous HF solution, and strontium and cesium are not. The rare earth solubility in these HF solutions saturated with LiF and BeF, increased from about  $10^{-4}$  mole % in 98 wt % HF to 0.003 mole % in 80 wt % HF, based on the amount of LiF + BeF, dissolved. Reactor fuel with a l-year fuel cycle will contain rare earths at a concentration of about 0.05 mole%.

#### SOLUBILITIES OF HEAVY ELEMENTS IN HF

Neptunium is the most serious nuclear poison other than the rare earths in a molten salt reactor, burning  $U^{235}$ .

The neptunium solubility in 80 to 100% HF saturated with LiF and BeF<sub>2</sub> was found sufficiently low to permit its removal along with the rare earths (Table 9). For these determinations small quantities of neptunium (aqueous Np<sup>+4</sup> nitrate solution) were added incrementally to HF solutions saturated with LiF and BeF<sub>2</sub>. The concentration of nitrate so added was ~0.02% of the fluoride concentration. Solubilities are reported in mg Np per g solution and in mole % of neptunium relative to dissolved LiF-BeF<sub>2</sub> salt.

The neptunium was determined by alpha counting with pulse analysis to distinguish neptunium from plutonium activity present as an impurity. The plutonium appeared to be carried with the neptunium to a considerable extent; the ratio of the amount of plutonium in solution to the amount undissolved was within a factor of 2 of the ratio for neptunium although the plutonium concentration was smaller by a factor of 500.

Addition of iron and nickel metal to the HF solutions resulted in a significant reduction in the gross  $\alpha$  activity, probably as the result of reduction to Np(III) and Pu(III). This has not yet been verified by pulse analysis. The trivalent state might be expected to behave in somewhat the same way as the rare earths; the observed solubilities are in the same range as those reported previously for rare earths. It is expected that the rare earths, neptunium, plutonium, and possibly uranium will behave as a single group and therefore exhibit lower solubilities when present together than the values reported here for the separate components.

Measurements of the solubilities of thorium and uranium fluorides in 90 to 100% HF indicate that uranium and thorium are relatively insoluble. All  $ThF_{l}$ , determinations showed less than 0.03 mg of thorium per gram of solution,

## Table 9. Np(IV) Solubility in LiF-BeF, Saturated Aqueous HF

4	Solubility in Solvent, mg/g							
Salt	~80 wt % HF	~90 WE % HF	~94 <b>vt % HF</b>	~100 wt <b>%</b> HF				
LIF	29	64	96	112				
BeF <sub>2</sub>	111	70	60	40				
Np	0.026	0,011	0.0086	0.0029				
Np solubility relative to salt, mole %	0.0031	0.0012	0.00072	0.00024				

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the limit of detection. Better analytical methods are available for  $UF_{h}$ , and solubilities were generally in the range 0.005-0.010 mg of uranium per gram of solution. In the presence of fresh iron or nickel metal solubilities were somewhat lower, 0.002-0.005 mg/g, indicating perhaps a higher solubility for higher oxidation states. Relative to dissolved salt these solubilities are the order of 0.001 mole %.

## SOLUBILITY OF CORROSION PRODUCT FLUORIDES IN HF SOLVENT

Measurements were made of the solubilities of corrosion products fluorides (iron, chromium, and nickel) in HF solutions saturated with LiF. Chromium fluoride was relatively soluble, with values of about 8, 12, and 18 mg of chromium per gram of solution in 100, 95, and 90 wt % HF, respectively. Tron and nickel fluorides were less soluble. Measurements of the iron fluoride solubility varied from 0.08 to 2 mg iron per gram of solution; the average and average deviation neglecting extreme values was  $0.14 \pm 0.10$ . Measurements of the nickel fluoride solubility varied from 0.01 to 1.3 mg of nickel per gram of solution, the average being  $0.15 \pm$ 0.06. The results were so inconsistent that no trend with HF concentration could be established. When iron, nickel and chromium fluorides were present together, the solubilities of all appeared to be somewhat lower.

Similar measurements in 90 and 100% HF (no dissolved salt) fell in the same range, but in 80 wt % HF the iron and nickel solubilities appeared to be significantly higher.

## RECOVERY OF SALT FROM SOLUTION

The LiF and BeF, salt can be recovered from the HF solutions by evaporation. There is some question of hydrolysis of the salt in recovery, the result particularly of the existence of a high-boiling azeotrope at 38 wt % HF, which would be obtained if some water was present initially and if flash evaporation was not used. A 90% HF solution saturated with LiF and BeF, was slowly evaporated to dryness and heated to 450°C. X-ray diffraction indicated that the resulting salt was about 90% 2LiF-BeF2 and 10% BeF2, and petrographic examination indicated less than 3% of material other than these fluorides. A second sample was evaporated after addition of a little NH<sub>4</sub>F (which would decompose and perhaps hydrofluorinate the salt at elevated temperatures) and fused at 700°C. This salt appeared to be entirely the binary compound, and the x-ray pattern was somewhat cleaner than in the first case. However, hydrolysis did not appear to be a significant factor in any event.

### PROCESS FLOWSHEET FOR REACTOR PROCESSING

A tentative flowsheet for application of the fluoride volatility and HF dissolution processes to molten salt reactor fluids has been prepared (Fig. 2). In this system the uranium is separated from the salt as  $UF_6$  before HF dissolution of the salt, although the reverse might be feasible in some circumstances. The LiF-BeF<sub>2</sub> salt is then dissolved in concentrated HF (>90% HF) for separation primarily from the rare earth and neptunium neutron poisons. The salt is re-formed in an evaporation step, from which it would proceed to a final makeup and purification step. The latter would perhaps involve the H<sub>2</sub> and HF treatment now believed necessary for all salt used in a reactor. The UF<sub>6</sub> produced in the volatility process would be converted to UF<sub>h</sub> for reuse in salt by hydrogen reduction, or alternately it could possibly be reduced in situ in the fused salt by essentially the same method. Although the volatility process achieves high decontamination, and the dissolution process leads to the elimination of the rare earths and neptunium activities, the salt recycle would require shielding and remote operation.

The scale of the process with a 1-year cycle, relative to the reactor, is indicated in Fig. 3. The daily core salt processing rate would be about 125 kg containing about 4 kg of uranium. Assuming a 10% solubility for LiF-BeFo in the HF solution the solution processing rate would be less than one liter per minute for the reference reactor (260 Mw electric). The daily blanket processing rate would be about 250 kg containing only about 0.1 kg of U<sup>233</sup> which would be recovered by the fluoride volatility process. The blanket salt would then be returned to the reactor, after treatment to remove corrosion products, if necessary. Infrequent processing of the blanket salt to remove fission products would be required since at most a few percent of the fissions occur in the blanket. Although more development of some of the process chemistry is certainly needed, the principal features of the process appear to be suited to the objectives for processing molten salt reactor fuel. Much more work will be needed for developing chemical flowsheets for specific reactor systems.

#### HIGH-TEMPERATURE PROCESSING

The direct removal of rare earths from molten fluorides was investigated briefly using two methods--oxide precipitation and partial salt crystallization. These results were generally not favorable and were not pursued further. An alternative approach which appears to have considerable promise consists of direct equilibration of the salt with a bed of CeF<sub>3</sub> to remove the worse poisons, the rare earths.<sup>0</sup>

Oxide precipitation, achieved by small additions to the fused salt of water or CaO, was tried with 50-50 mole % NaF-ZrF<sub>4</sub> and 11.5-42-46.5 mole % NaF-KF-LiF since it had not been determined at the time that LiF-BeF<sub>2</sub> salt would definitely be preferred. Trace fission products were added. Oxide addition results in precipitation of primarily zirconium in the case of zirconium-bearing salt without too much effect

<sup>8</sup>J. H. Schaeffer, N. V.Smith, R. A. Strehlow, W. P. Ward and G. M. Watson, "High Temperature Processing of Molten Fluoride Reactor Fuels." In press - Chemical Engineering.

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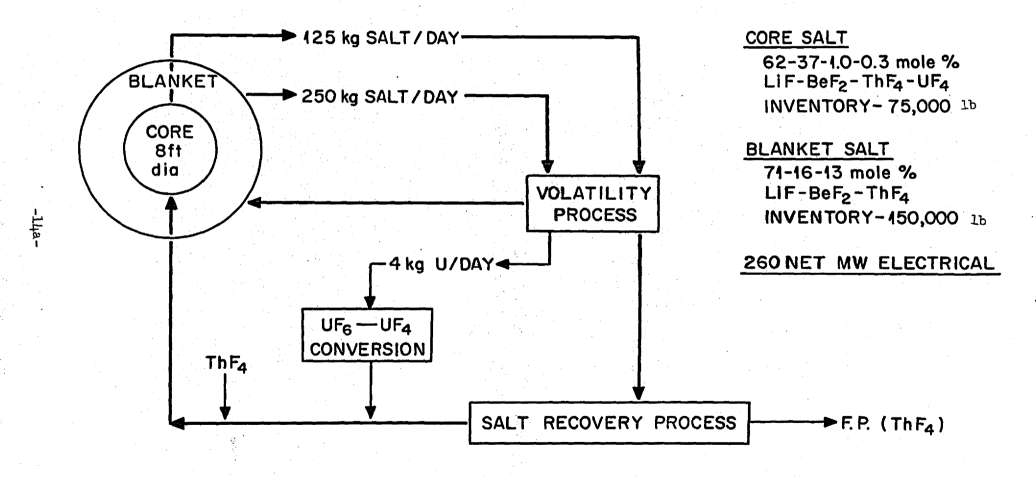


Fig. 3. Molten-Salt Reactor Processing.

on fission product activities (uranium if present would behave similarly to zirconium). However, in NaF-KF-LiF salt, the rare earth, zirconium, and niobium activities were effectively removed since a large amount of zirconium (or other oxide forming material) was not present. In the case of LiF-BeF<sub>2</sub> (63-37 mole %and 50-50 mole %) salt, CaO or water vapor addition caused precipitation of zirconium and niobium activities, with little effect on the rare earths or other activities.

Partial freezing by slow cooling of both NaF-KF-LdF and NaF-ZrF<sub>4</sub> salt containing 0.2 mole % rare earth fluoride resulted in a concentration in the unfrozen liquid of most fission products, including cesium, strontium, and rare earths. The concentration factors were generally less than 2 and always less than 3. The solubilities of these components are considerably higher than their concentrations in these salts. However, partial freezing of LdF-BeF<sub>2</sub> (50-50 mole % and 63-37 mole %) and NaF-BeF<sub>2</sub> (63-37 mole %), each containing 0.2 mole % rare earth fluorides, 0.1 mole % ZrF<sub>4</sub> and trace fission products, resulted in a decrease in rare earth activity in the unfrozen liquid. This is the result of the solubility being less at the melting point than the 0.2 mole % rare earth concentration.

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