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A Small-Scale Study on the Dissolution and Anion-Exchange Recovery of Plutonium from Rocky Flats Plant Incinerator Ash

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Los Alamos National Laboratory

A SMALL-SCALE STUDY ON THE DISSOLUTION AND ANION-EXCHANGE RECOVERY OF PLUTONIUM FROM ROCKY FLATS PLANT INCINERATOR ASH

by

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ABSTRACT

Incinerator ash has been processed successfully at plutohum production sites for many years, but not without some difficulty. Problems arise from complicated ash chemistries that adversely affect process operations and efficiencies. We performed small-scale experiments to examine four specific areas identified by Rocky Flats Plant personnel as sufficiently important to warrant further investigation to determine (1) the optimal feed fluoride concentration (the relationship between fluoride concentration and overall plutonium/ash dissolution in the nitric/hydrofluoric acid system), (2) the effect of free fluoride on anion exchange performance if aluminum nitrate is not added to ash filtrates as a complexing agent, (3) possible equipment corrosion problems resulting from potentially large quantities of uncomplexed fluoride, and (4) the effects of unburned carbonaceous material on anion exchange behavior. Our findings are reported.

INTRODUCTION

Combustible wastes are generated from plutonium processing activities across the defense programs complex. Because these residues typically do not meet discard limits, they are incinerated to reduce their volume and convert them to a form more suitable for reclamation of their plutonium content.

Ash from the incineration of these materials has proven to be one of the more difficult matrices from which to solubilize plutonium, often requiring three or more passes before a discardable residue is obtained. Poor dissolution characteristics coupled with low plutonium content have made ash one of the less desirable scraps to process. However, as plutonium streams become leaner, ash processing becomes increasingly important in the plutonium recovery cycle.

Although incinerator ash has been successfully processed at production sites for a number of years, there have been obstacles other than the difficulty of dissolution. Complications arise as a result of contaminants found in ash that adversely affect process operations and efficiencies. Two contaminants that have long plagued ash processing are silicon and carbon. Silica, which is abundant in ash, reacts with fluoride to produce $SiF_4(g)$ that reacts with water to form fluorosilicic acid and silicon solids that plug offgas systems.^{1,2} Additionally, aluminum nitrate, which is added to dissolver filtrates to complex free fluoride, complexes with silicon species present to form a troublesome gelatinous precipitate that readily plugs filters. If this gel is not completely removed, solids collect in storage tanks and diminished flow or plugging of ion-exchange systems occurs.

Carbon. reported as high as 36 wt % in Rocky Flats Plant ash,³ can most often be attributed to incomplete combustion of glove-box gloves and plastics during incineration. At Los Alamos, experience with processing off-site incinerator ash has shown that high-carbon ash presents a myriad of problems, including (a) excessive foaming during ash addition to the dissolver when using nitric/hydrofluoric acid mixtures. (b) peor filtration, (c) a viscous residue buildup on equipment, and (d) fouling of anion-exchange columns.⁴ Reburning incinerator ash at 600°C for a period of 4 hours has been shown to be a simple and effective method for carbon removal.⁵ Accordingly, Los Alamos has employed rotary calcination for this purpose and has successfully mitigated downstream problems associated with carbon. At Rocky Flats Plant (RFP), a major effort is under way to upgrade existing photonium processing facilities. Of particular interest is the ash processing area (PROVE) now under construction. Because PROVE has been selected as a test facility to verify the RFP approach to upgrading other process areas, representatives from sites across the production complex were called upon to perform a technology assessment on it. Their work culminated in the PROVE Technology Assessment Report (PTAR) that brought to light concerns warranting further investigation⁶ and resulted in the addition of an Incinerator Ash Processing Working Group (fAPWG) to existing technology exchange programs between Los Alamos and RFP.

This report is Part I of a three-part series documenting the results of a small-scale incinerator ash study conducted at Los Alamos by IAWPG members to address specific concerns outlined in the PTAR. Parts II and III will follow upon completion of production-scale verification testing and ash characterization studies in progress at Los Alamos and small-scale studies at RFP that parallel the work reported here.

EXPERIMENTAL

Four specific areas associated with ash processing were identified by RFP as sufficiently important to warrant further investigation: (1) the relationship between the calcium fluoride addition rate and overall plutonium/ash dissolution when dissolving ash in the nitric/ hydrofluoric acid system. (2) the effect of free fluoride on anion-exchange performance if aluminum nitrate is not added to ash filtrate as a complexing agent. (3) possible equipment corrosion problems resulting from potentially large quantities of uncomplexed fluoride, and (4) the effects of ash carbonaceous material on anion-exchange behavior.

Although it was our desire to extend the scope of this study beyond the four areas identified, this was determined to be impossible if changes to the flowsheet were to be implemented in time for the scheduled PROVE startup. Therefore, we were limited to using as-designed PROVE operating parameters that may or may not be optimum for achieving maximum process efficiency or product quality.

The projected study was divided into two separate phases. In Phase I, both Los Alamos and RFP were to conduct a series of six small-scale scouting tests using varying levels of calcium fluoride to determine the optimum fluoride concentration for promoting plutonium/ash dissolution. The corrosion potential and effective capacity of the anion-exchange column were to be evaluated for each solution. To examine the effects of ash carbonaceous material on anion-exchange behavior, RI-P would use virgin ash tuntreated), whereas Los Alamos would use ash that had been calcined. In Phase II, conducted only at Los Alamos, production scale verification 'ests were to be performed using an optimum fluoride concentration determined from performance criteria evaluated in Phase I.

Equipment

Two-inch cascading air-lift dissolvers were designed and fabricated at Los Alamos for the dissolution of incinerator ash and a 3-in, anion-exchange column was assembled for the separation and recovery of plutonium from the ash filtrate. An identical set of equipment was transferred to RFP for use in parallel experiments.

The cascade dissolver (Fig. 1) was assembled using three 2-in. diameter by 12-in. long glass-pipe dissolvers with heads fabricated from chlorinated polyvinylchloride (CPVC) stock. The dissolvers, in series, were airsparged and fed by gravity from one to the other. A sampling valve was placed on the underside of the overflow pipe between dissolvers. Each dissolver was heated externally with a hot plate to an operating temperature of $100 \pm 5^{\circ}$ C. Dissolver leachate was circulated by drafttube (1-in. \times 17-3/4-in.) air action that facilitated particle suspension for maximum liquid/solid contact. Air. regulated through a supply manifold equipped with metering valves, was delivered independently to each dissolver at a rate of 600 cm³/min. Off-gas vapors were collected in a 3-in. diameter by 12-in. long flanged glasspipe scrubber, partially filled with water and fitted with a cooling coil supplied by a closed-loop negative-pressure chilled water system. Ash, calcium fluoride, and nitric acid were introduced through a feed inlet funnel mounted on the first dissolver. Ash and calcium fluoride were manually added, whereas nitric acid addition was controlled using a peristaltic pump between the acid feed tank and the first dissolver. Leachate exiting the third dissolver overflowed to a filterboar fitted with a 5-µm filtering cloth and was collected and stored in clean 6-in. glass receiver tanks.

The anion-exchange column (Fig. 2) was constructed using a 3-in. by 12-in. glass pipe and CPVC flanges fitted with a high-density polyethylene frit. Flow control was maintained using a peristaltic pump. Effluent, wash, and eluate streams were collected in clean 6-in. glass receiving tanks. All column runs were monitored using a computer-based on-line gamma monitor similar to that employed on our production columns.^{7,8}



Fig. 1. Cascade dissolver system used for small-scale openerator ash dissolution studies.

Ash Feed Preparation

The virgin ash used in this study was packaged in eight cans that had been shipped to and stored at Los Alamos in the early 1980s. When the cans were opened, three distinct color variations were observed. Because this might indicate a differing composition, and because we wanted to establish repeatability of results asing diftering ash feeds, we elected to combine similarly colored materials and divide the ash into three lots.

Two of the three feed lots resulting from this division were used in this study. These were individually calcined for a minimum of 4 hours at 600 C, pulverized, screened through a 40-mesh sieve, blended well with a V-blender, rescreened, and packaged for subsequent use. A reasonably high degree of confidence in the homogenicity of each lot following this procedure was supported by work performed at Los Alamos by S. D. Fink et al."

Anion-Exchange Feed Preparation

Before the anon-exchange puribeation step, ash filtrates were spectrophotometrically examined to determine if plutonium was in the preferred PurIVo oxidation state. Results indicated that no valence adjustment was required. As a final step in preparing feed solutions for amon exchange, act⁴ and phitomain concent; evens were adjusted to approximately 7.5 M and 1.5 g/L, respectively.

Feed and Flow Rates

With the exception of calcium fluoride, which was controlled to achieve a specific feed fluoride concentration, feed and solution flow rates were proportionately adjusted to as-designed PROVE process operating parameters. In PROVE, nitric acid and incinerator ash will be delivered to the dissolvers at 18 L and 1.2 kg per hour. Amon-exchange feed, wash, and clutriani solutions will be delivered at 150, 100, and 85 L per hour, respectively. (It is important to note that Los Alamos maintains a position different from that of RFP on how to "proportionately" adjust anion-exchange operating parameters. This will be discussed further under Anion Exchange in the Results and Discussion section.)

Anion-Exchange Resin

The anion exchanger used was Lewaiit MP-500-FK. 40-70 mesh, macroporous resin. The resin was converted from the chloride to the nitrate form using Los Alamos/MST-12 Standard Operating Procedure 461-REC-R02.¹⁰



Fig. 2. Anion-exchange column used for small-scale incincrator ash studies.

Assays

During or following each processing step, samples were analyzed for plutonium or elemental impurities All solid materials, except for the ash feed, which was blended well, were mixed just before sampling. Accordingly, the sample taken may not have been representative and is subject to error greater than that inherent in the method of analysis. Where an analytical technique required that a solid sample first be solubilized, a sealedreflux dissolution was performed.¹¹ With the exception of the elements noted below, all elemental values reported were determined by using DCarc emission spectroscopy. This technique has a factor of 2 accuracy and a precision of 50% relative standard deviation (RSD). Plutonium values were determined by radiochemical methods having a stated accuracy of 2-5% RSD. Fluoride and chloride concentrations were determined using ion-selective electrodes having an estimated precision of 5% RSD for either element. For comparison, fluoride concentrations were also determined using on-site analytical capabilities. Samples were analyzed using ion-selective electrodes and a standard addition method of analysis. The uncertainty of measurement for this technique is 10% at the 95% confidence level.

Carbon and hydrogen were determined by mass spectrometry. The accuracy for this method is typically 0.25%. Nitrogen was determined by the Kjeldahl-Nessler method.

Plutonium concentration adjustments on anion-exchange feeds were based on gamma assays for plutonium using an in-line solution assay instrument. Operating experience has demonstrated an accuracy of approximately 5% for this instrument. The adjusted feed was then resampled for plutonium concentration determination by codiochemistry.

Experimental Procedure

Dissolution. A series of six separate small-scale dissolutions were performed using feed fluoride concentrations of 0.4, 0.1, 0.25, 0.35, 0.5, and 0.4 M respectively. Ash from blend lot 1 was exhausted after run 4. and ash from blend lot 2 was used for the remaining two dissolutions. Before each experiment, ash and calcium fluoride were weighed and packaged in aliquots sized to achieve the desired hourly addition rate. Because the scaled-down equipment did not permit using an auger leeder, we manually added one quarter of an aliquot every 15 min. Bulk concentrated nitric acid was diluted to 9 M with deionized water and fed continuously to the dissolvers during the run. At termination, a 2-L aliquot was collected for corrosion testing. The remaining filtrate was transferred to and processed through anion-exchange before proceeding with the next dissolution. The dissolvers were disassembled for cleanout, gloveboxes were cleaned, and a material balance was established.

Corrosion Studies. Because the aluminum nitrate addition step was eliminated, there was concern over the resistance of 304L stainless steel to general and localized fluoride-induced corrosion. A cyclic potentiodynamic polarization scan was performed on each filtrate

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solution generated at the varying feed thiorid- addition levels. From these scans and measured polari, stion resistances, localized corrosion and inst-ataneous general corrosion rates were determined.

Anion-Exchange. A comparative determination of column capacity for each solution was the method of choice for evaluating the effects of uncomplexed thioride on anion-exchange behavior. As noted by Marsh and Gallegos,¹²

> The effective capacity of an anion exchange column may be determined in various ways. Some investigators have chosen to report the quantity of plutonium retained on the ion exchange column when the plutonium exiting the column (breakthrough) reaches predetermined levels...

> We chose instead to determine the saturation capacity of the anion exchange column by loading each column with a volume of feed solution that contained an excess of plutomum.

Although we agree with their approach, the limited quantity of ash feed available restricted us to using the "breakthrough" technique. Accordingly, we chose to base all column capacity determinations on a 5% level of plutonium breakthrough.

To minimize variability introduced when processing anion-exchange feeds with differing plutonium values, we adjusted feed plutonium concentrations to approximately 1.5 g/L. Grab samples were taken from the effluent stream at 2.5- to 5.0-L intervals and analyzed for plutonium. From the feed plutonium concentration, plutonium in the effluent, sample number, and corresponding effluent volume collected, we were able to determine the column capacity at 5% breakthrough.

In all tests, the column was loaded and washed (7.5 M nitric acid) in the upflow direction, and eluted using a downward flow of 0.35 M nitric acid.

RESULTS AND DISCUSSION

Ash Feed

Table I shows the composition of ash before and after rotary calcination and of the ash heel following each of the six dissolutions. A loss in net weight was seen and is attributed to impurities that react with oxygen during burning (rotary calcination) and are removed in the off gas. Conversely, one would expect the nonvolatile impurities to become more concentrated, but many of these also show a decrease in relative concentration following ¹ stary calcinate on. This indicates the humanions of emission spectroscopy and accordingly, the results presented should be interpreted as qualitative only. It is intereating to note, however, that with the exception of phitonium, eb⁴ rine, and possibly sodium, no preferential dissolution is evident.

Dissolution Studies

To sharming the level of thioride best suited for plutonium dissolution, criteria other than the amount of plutonium solubilized were considered; for example, the ratio of plutonium to bulk ash dissolved, which if in proper proportions, could result in a discardable heel. Also, one would hope to minimize the formation of SiF_3 and plugging of the off-gas system by limiting the amount of fluoride added to the minimum amount needed for optimum plutonium dissolution.

Inevitably some spillage occurs when processing ruclear materials in a glove box. Because of the difficulty in determining the amount of plutonium or bulk ash lost to a spill, we elected not to consider these losses when calculating the plutonium and bulk ash dissolution efficiencies. Accordingly, the computed efficiencies may be slightly higher than the values reported.

We computed plutonium (P_d) and ash (A_d) dissolution efficiency as follows:

$$P_{d} = \frac{P_{1} + P_{1}}{A_{p}} \times 100$$

and

$$A_d = 1 - \frac{(F - F_p) + (H - H_p)}{A - A_p} \times 100$$

where

A = net weight of ash feed, granis;

 A_n = weight of plutonium in ash feed, granis:

= weight of filter residue. grams:

 F_p = weight of plutonium in the filter residue, grams:

H = weight of ash heel. grams:

 H_p = weight of plutonium in the ash heel. grams;

 P_{f} = weight of plutonium in the ash filtrate, grams; and

 P_t = weight of plutonium in the leachate. grans.

As shown in Fig. 3, the amount of plutonium and bulk ash dissolved increased with increasing fluoride concentrations up to 0.35 M and leveled off thereafter. The one exception is the marked difference in bulk dissolution from fot 1 to lot 2 at 0.4 M fluoride. As a result of this difference, the heel left after processing ash from lot 2 was discardable.

Plugging of the off-gas system was encountered when the feed fluoride concentration exceeded 0.4 M. Because

Table I. Composition of Rocky Haw Plant incinerator ash before and after rotary calcination, and ash heef following dissolution at fluoride concentrations of 0.1 to 0.5 M

	i Ashfutt			Fafter Residues				Entrar Providence		
	Lucalement	Calenced	Run 1	Run 2	Run i	Run 4	Uncalcined	Calcined	Ruo 5	Ruo 6
Net Weight	5031 Sg	4824 ⁻ ¥					2898.8g	2762.6g		•
Plutonium	0.138g/g	0145g/g	0.04 1 4g/g	019)g/g	0.0621g/g	0.4892.2	0.086225	0.09072/2	1111412/2	0.01666/2
Carbon	* **;	0.26%	0.407	0.354	•	terner e		0.32%	0.719	(174%
Hydrogen	061	0.16%	0.27%	1 80°7		0.28%	0.50%	ecour;	1.40%	1.603
Nurogen	370						280			
Chlorine	j 2.517	294	0.139	(1(17);	031971	0.064	3.34		0.0762	0,0877
Fluorac	1		<u>2.2067</u>	1.60%	Lorri	t stra			1.60%	1.70%
Calcium	15%	617	29	.)* e		217	812	ב ,	217	1,50%
tron	159	511	517	512	5%	511	51;	512	N' i	49
Magnessum	1 1463	14	114	14	19	1.1	343	15	-408AD	3000
Phosphorus	્રાત્રપ્રદ	300N1	્ર સારમા	< 3000	311081	< 3000	-C SORRI	50XXI	(XXF, >	< 1000
Tdancum	5.	3	44	¥.,	2%	4%	5%	44	5.1	44
Zirconium	E SOUXE	2000	3CH M 3	200	200	5000	~100	500	<u>20081</u>	1500
Aluminuoi	1 59	517	317	<u>24</u> :	1.5%	34	54;	54	31 ;	44
Cadmium	1000	500	þи	ton	tori	100	500	400	200	400
Gathum	2000	2000	2000	30.83	,WH1	500	500	2000	281	< 100
Manganese	3.4	70081	5000	66833	5000	5000	te	5000	30001	2(XX)
Lead	2%	14	3000	3EN NI	3000	3000	44	217	4090	5000
Boron	i 1000	1000	1000	500	800	1000	2000	(D)(D)	700	500
Silcon	2077	74	t0%	1597	בi	tari	159	ter;	tar;	10%
Barrum	i 29	8000	3000	4000	4000	1.59	1.5%	177	40001	5000
Sodium	314	1.59	2000	2000	1000	1000	29	1.59	2500	5000
Berythum	1500	2000-	-400	300	200	200	3000	ACKAD	200	500
Chromum	34	29	29	347	29	24	297	2%	1.59	17
Potassium	39	5%	8000	7000	4000	5000	20%	107	7000	19
Cupper	1.59	7000	3000	5000	4000	4000	27	ts	3000	3000
Nickel	377	1.5%	29	24	2%	247	29	217	177	5000
Zinc	49	172	19	5000	5000	5000	34	24	5000	5000
Sdver	500	< 100	<100	500	500	50	<30	< 100	200	300
Vanadium	<100	< t00	<100	<100	<100	< 100	< t00	<100	< 100	< 100
Cerium	<500	<500	<500	<500	< 500	< 500	<500	< 500	<500	<500
Germanium	<100	< 100	<100	<100	< 100	<100	<100	< 100	< 100	< 100
Molybdenum	5000	4000	2000	1000	1000	3000	<300	2000	1000	800
Tungsten	<3000	< 3000	< 3000	<3000	<3000	<,3000	<3000	<3000	<,3000	< 3000
Cobah	<100	<100	< t00	< 300	<300	<300	<100	<100	< 100	< 300
Hafnium	<300	<500	<500	<300	< 300	<300	<.300	< 500	<300	<.300
Tin	< 300	<300	< 300	<300	<300	<300	4000	<300	1500	<.300
Niobium	< 300	<.300	<300	<300	< 300	<300	<300	<300	<300	<300
Strontium	2000	1000	tono	200	200	400	3000	1000	300	500
Bismuth	<100	<100	< 100	<100	< tub	<100	< t00	<100	< t00	< 100
Lahium	500	300	100	100	<100	< 100	< 200	200	<100	<100
Yitnum	500	1000	100	.30	30	100	<t00< td=""><td>100</td><td>< 100</td><td>300</td></t00<>	100	< 100	300



Fig. 3. Single-pass plutonium and bulk ash dissolution efficiencies at fluoride concentrations of 0.1-0.5M.

we were first inclined to attribute this to a change in feed (blend lot 2 was used for the 0.5 M test), we elected to repeat dissolution at 0.4 M using feed from lot 2.

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When plugging was not observed in the repeated run, we assumed that fluoride added in excess of 0.4 M increased $SiF_4(g)$ formation.

Corrosion Studies

A portion of the 2-L aliquots collected from five of the six dissolution filtrates were tested with solutionannealed or sensitized 304 L stainless steel coupons. Table II shows the general corrosion rates found for the five ash filtrates tested. An increase in the rate of corrosion was observed at elevated temperature; however, results were within the acceptable limits established at RFP. Solutions retested following aluminum nitrate addition showed no significant difference in the rate of corrosion. No localized attack was found at either room or elevated (90–95°C) temperatures.

Anion Exchange

Earlier we stated that "flow rates were proportionately adjusted." However, scale reduction based on column diameter proportionately adjusts the feed flowrate img Pu min⁻¹ cm⁻² of resmit, whereas scale reduction by bed depth proportionately adjusts the volume flow rate or bed residence time [ml min⁻¹ cm⁻² of resin or column volumes tevo/hour]. It is believed by Los Alamos investigators that volume flow rate is a concern of importance when processing and proportionally adjusting plutoniumi-lean solutions.

Table II. General corresion rates for 3041, stanless steel exposed to inbrate solutions from the dissolution of inciderator ash in 9 M mirric acid with 0.1 to 0.5 M duoride.

	GENERAL CORROSION RATES onlyyear				
Coupon Type	Room Temperature	Flevated Temperature			
Solution Annealed	0.07 to 0.54	18.2 to 46.5			
Sensitized	0.63 10 33.1	151 to 169			

In our study, scale reduction was performed based on column diameter only. Therefore, conservatively estimating that the resin occupies 50% of the total column volume, the feed residence time was approximately 1 min. Because plutonium absorption is kinetically limited, a substantial reduction in bed residence time could impact resin performance—even Lewatit, which is known for its outstanding sorption kinetics. As shown in Table III, our volume flow rate was about 10 times greater than that planned in PROVE. (The PROVE anion-exchange system is composed of two 6-in, diameter by 60-in, long columns.)

 Table III. Anion-exchange operating parameters used in small-scale studies and PROVE.

	Small-Scale Studies	PROVE	
Flow (L/hour)	37.5	150	
Feed (Pu in g/L)	~1.5	~1.5	
Feed Flow rate 1mg Pu min ⁻¹ cm ⁻²) 22.6	22.6	
Column Volume (L)	1.39	150	
Volume Flow rate (cv/h)	27.0	2.75	

Table IV shows the plutonium, aluminum, and fluoride concentrations of anion-exchange feed before adjustment and the column capacity at the varying feed fluoride levels used at dissolution. As can be seen, the column capacity at 5% plutonium breakthrough was not determined for runs 1 and 2. Although the column appeared to be loaded and breakthrough was presumed to have occurred, the lag time between effluent sample submittal and the reporting did not lead to early recognition of this error.

Table IV. Anomerschange column supreny at 5% photonium breakthough using ash filtrate feeds containing varying ratios of 1° MPu in 7.5 M HNO3.

		Amon Exchange Leed (non adjusted)					Capacity		
Ron	Pugli	Alight	hgh	†∙AI	Al-Pu	F Pu	(-sluam	Operating	
t	53	to	22	3.1	1.7	52	261	>44	
2	51	1.5	1.9	§ 8	2.6	4.*	2.62	>44	
1	7.5	1.2	2.5	3.0	14	42	74,1	515	
4	110	27	2.3	1.2	22	27	41.8	66.1	
5	4,8	•	2.9	•	•	76	846	610	
6	3.0	•	4.7	•	•	15.4	-1.5	\$3.6	

Anion-exchange feed from dissolution using 0.35 M fluoride had the lowest F:Al:Pu ratio and the highest colunin plutonium loading at 5% breakthrough. In contrast, we found the next highest column plutonium loadings were attained using feed having the greatest Pu:F ratios. However, the data are incomplete because of an oversight in requesting aluminum for feeds 5 and 6. This was not recognized until after the sample was discarded.

It is important to note that the fluoride values shown in Table IV do not necessarily indicate the level or presence of free fluoride. In fact, we suspect that the abundance of silicon. aluminum, and tantalum present in ash is sufficient to complex any fluoride present over the entire range added at dissolution.

Ryan et al.¹³ determined the equilibrium and operational capacity of Lewatit MP-500-FK resin from 7M HNO₃ as a function of solution-phase plutonium concentration. Using a 1.08 g Pu/L feed at a flowrate of 16 mg Pu min⁻¹ cm⁻² (17.8 cv/hour), they estimated the equilibrium capacity to be 86.5 g Pu/L resin. With the same feed, they recommend a maximum operating capacity of about 67 g Pu/L resin for a primary column of a two-column series.

In our small-scale system where the feed flow rate was approximately 23 mg Pu min⁻¹ cm⁻² (27 cv/hr), a wetted resin capacity of 54 to 66 g Pu/L was realized at 5% plutonium breakthrough.

Marsh¹⁴ found that the distribution coefficient (K_d) of plutonium was measurably suppressed at all levels of fluoride tested, nc matter how much aluminum was added, for dynamic contact periods from 10 to 60 min. From the data presented in Fig. 4 having F:Al:Pu ratios that nearly approximate ours, it would appear that the K_d of plutonium may not be significantly affected at contact periods of less than 1–2 min.



Fig. 4. Sorption of Pu(IV) on Lewatit MP-500-FK anion exchange resin from 7 M nitric acid that contains varying ratios of aluminum to plutonium for F/AI = 1.5 and 3.0, as a function of dynamic contact time. Data from Marsh (1987).

CONCLUSIONS

It would be purely speculative at this point to draw any conclusions until our counterparts at RFP have completed small-scale parallel studies and we have conducted production-scale verification testing. However, a few preliminary observations may be in order.

- 1. The quantity of plutonium solubilized from incinerator ash tested is comparable at added fluoride levels of 0.35, 0.40, and 0.5 M.
- It would appear possible to obtain a discardable ash heel following single-pass dissolution by adding an amount of fluoride just sufficient to promote maximum plutonium dissolution while leaving the major fraction of the bulk ash behind. For some types of incinerator ash. 0.4 M fluoride addition may be well suited.
- For ash tested, as feed fluoride is increased above 0.4 M. off-gas plugging becomes a concern.

- 4. All dissolver filtrates tested were within RFP acceptable limits of corrosion. It would appear that the aluminum nitrate addition may be elinanated without severely affecting the lifetime of equipment.
- 5. Throughout our studies we observed no discoloration of anion exchange resin. However, RFP has reported a darkened resin bed and a diminished column capacity after one run. Experience at Los Alamos coupled with the preliminary findings at RFP indicate that the presence of carbon in ash adversely effects anionexchange performance.

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REFERENCES

- R. L. Ames, "Demonstrate the Operability of the Ash Dissolution System," Final Report, Revision 1, Rockwell International, Rocky Flats Plant internal report PPC 88-003 (March 1988).
- T. C. Johnson, "Recovery of Plutonium from Incinerator Ash at Rocky Flats Plant," Rockwell International, Rocky Flats Plant report RFP-2520 (Dec, 1976).
- G. P. Hajela and A. Delgado, "PROVE Mass Balance Code," Rockwell International, Rocky Flats Plant support document 169ER0000111 (Sept. 1986).
- 4. J. J. Balkey, personal communication (April 1988).
- C. H. Delegard, "Laboratory Tests on Plutonium Recovery from Rocky Flats Ash Using Nitrie Acid-Calcium Fluoride Leaching," Rockwell Hanford Operations, support document SD-CP-DTR-005 (March 1984).
- W. E. Harris, T. W. Blum, R. C. Hoyt, W. R. Dworzak, J. L. McAtec, and C. A. Ostenak, "Technology Assessment of the Plutonium Recovery Option Verification Exercise (PROVE) Project at the Rocky Flats Plant," committee report (Sept. 1988).
- S. F. Marsh and M. C. Miller. "Plutonium Process Control Using an Advanced On-Line Gamma Monitor for Uranium. Plutonium. and Americium." Los Alamos National Laboratory report LA-10921 (May 1987).

- 8 N. G. Pope and S. F. Marsh, "An Improved, Computer-Based, On-Line Gamina Monitor for Plutonium Anion Exchange Process Control," Los Alanios National laboratory report EA-10975 (June 1987).
- S. D. Fink, J. H. Gray, S. J. Kent, S. A. Apgar III, R. G. Behrens, and N. D. Stalnaker, "Fluorination of Incinerator Ash by Hydrofluorination or Ammonium Bifluoride Eusion for Plutonium Recovery," Savannali River Laboratory report DPST-89-313 (Feb. 1989)
- A. E. Nixon, "Nitrate Anion Exchange," Los Alanios National Laboratory MST-12 Standard Operating Procedure 461-REC-R02 (Nov. 1988).
- J. W. Dahlby, R. R. Geoffrion, and G. R. Waterbury, "The Scaled-Reflux Dissolution System" Los Alamos Scientific Laboratory Report LA-5776 (Jan. 1975).
- S. F. Marsh and T. D. Gallegos, "The Influence of Plutonium Concentration and Solution Flow Rate on the Effective Capacity of Macroporous Anion Exchange Resin," Los Alamos National Laboratory report LA-10990 (July 1987).
- J. L. Ryan, L. A. Bray, and E. J. Wheelwright, "Development and Demonstration Support of the Aqueous Process Flowsheet for Recovery of Plutonium from Special Isotope Separation Process Residues," Pacific Northwest Laboratory report PNL-6597 (July 1988).
- S. F. Marsh. "The Effects of Fluoride and Aluminum on the Anion Exchange of Plutonium from Nitric Acid." Los Alamos National Laboratory report LA-10999 (July 1987).

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