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SUMMARY OF 234-5 DEVELOPMENT PROGRESS  
JANUARY, 1951 - JUNE, 1959

By

H. W. Crocker

234-5 Development Operation  
Research and Engineering Operation  
Chemical Processing Department

July 29, 1959

HANFORD ATOMIC PRODUCTS OPERATION  
RICHLAND, WASHINGTON

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SUMMARY OF 234-5 DEVELOPMENT PROGRESS  
JANUARY, 1951 - JUNE, 1959

INTRODUCTION

This summary of laboratory investigations has been assembled to provide a directory for source data of past laboratory activities. The dates listed in the outline are the operational periods covered by Separations Technology Section and Research and Engineering monthly activity reports. The monthly activity reports or specific program reports are to be consulted for valid detailed information on the indicated topics. A list of the program documents and activity reports is included. This document supersedes HW-49123 RD, "Summary Of Development Laboratory Progress, 1951 - 1954", by W. L. Lyon.

A. WET CHEMISTRY

1. Plutonium(III) Oxalate

1951

July Study of process variables.

August Study of process variables; coupling to Redox and Recuplex.

1957

March Precipitation of continuous ion exchange product at 55 C. 1 M oxalic acid, filtrate 0.2 g/l Pu with solid formation.

2. Plutonium Peroxide

1951

August Coupling studies.

September Coupling studies - acid neutralization; plutonium peroxide from AT-solution - 0.5 M sulfate.

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October Variations in bulk densities; purities from single peroxide strike on 3BP- for dry chemistry and reduction; effect of fluoride upon lanthanum separation.

November Bulk density variations; disposal of methanol; Redox coupling.

December Bulk density variations; reaction of supernatants with alcohols; lanthanum separation.

## 1952

January Neutralization of 3BP by various agents; precipitation from concentrated plutonium solutions.

February Effect of total nitrate concentrations; temperature variations.

March Purity data for peroxides.

April Decontamination factors for peroxide precipitations.

September Precipitation from cleanout solutions containing manganese ion.

## 1953

February Decontamination factors for peroxide precipitation. Work on high recycle.

March High solubility in MRC's; effect of manganese ion.

## 1954

February Effect of fluoride and phosphate on formation of peroxy-complex.

August Sulfate elimination: addition of phosphate, oxalate; temperature and acid effects.

## 1956

March Continuous precipitation from 1.3 M HNO<sub>3</sub>, 0.05 M sulfate, 50 g/l Pu solution at 20 C feasible with 50 percent H<sub>2</sub>O<sub>2</sub>. Effects of sulfate concentration.

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### 3. AT-Solution Studies

1952

June                      Sulfate in AT still.

1953

April                      Plutonium sulfate in AT-solution (more work: brown vs. pink sulfates).

June                        Gas evolution.

November                Gas evolution from AT solution.

December                Gas evolution from AT solution.

1954

January                    Plutonium(IV)-(VI) change in evaporation as a function of nitric acid concentration; sulfate studies - dissolution of red sulfate. Unreported - gas evolution studies.

1956

May                        Prevention of solid precipitation in solution shipment.

4. Plutonium(IV) Oxalate1951

- September Preliminary entries.
- October More preliminary work.
- December Preliminary work (supernatant destruction by evaporation).

1952

- January Precipitation from 3BP solution; also from concentrated plutonium solution.
- February Precipitation from synthetic 2BP; filter boat tests, cake-washing, filtration rates, etc.
- April Precipitation in 231 Bldg.; decontamination factors.
- May Coupling studies solubility vs acidity; plutonium-uranium separation; supernatant treatment - boiling, platinum, hydrogen peroxide.
- June Purity data, effect of filtration; UX1 separation; supernatant kill - platinum.
- August Filter tests; effect of impurities on waste losses.
- September Effect of oxalic acid concentration.
- October Rate of oxalic acid addition; separation from lanthanum, iron, nickel.
- November Orange oxalate(VI?).
- December Precipitation from concentrated solutions, greater than 100 grams per liter.

1953

- January  
February Mixed reagent addition.
- March Decontamination factors.
- April Valence adjustment.
- June Task I prototype studies.
- August Temperature effects; aluminum addition; 0.06 M oxalate prior to pre-reduction; (pre-reduction-precipitation

effects, explanation?); Task I prototype, supernatant destruction - oxalate kill by permanganate as function as temperature, 15 to 65 C; use of ozone.

October

Effect of anions; effects of stirrer speed; precipitation of solution after partial neutralization; filter cloth tests.

1954

February

Glycol in the oxalate precipitation; dissolution of plutonium oxalate in aluminum and iron systems

June

Agitation studies including use of air lift; interference by manganese ion; filter cloth evaluation; and mercury separation (factors for dry chemistry, reduction).

November

Continuous filter - Task I; gas evolution from supernatants; supernatant treatment - kills with permanganate and dichromate.

December

Supernatant treatment: valence adjustment after kill. Unreported effect of ANN on plutonium oxalate solubility; mercury separation factors.

1955

February

Batch precipitation of concentrated Redox and AT solutions in Task I prototype.

March

Hg limits on Task I nitrate feed of 1,000 ppm, corrosion effect of Hg on gold components of filter boat.

May

DF's of 6.6 and 10 for Zr-Nb and Ru-Rh from Redox PR solution in oxalate precipitation. Stability of Dynel SD-9 cloth to oxalate precipitation chemicals.

June

Batch precipitation studies. Continuous filter and calciner fabrication.

August

Initial continuous oxalate precipitation studies.

September

No effect on batch precipitation characteristics by increasing Al to 100,000 ppm and Fe to 70,000 ppm in feed.

November

Continuous oxalate precipitation studies at ambient temperature process operation.

December

Continuous precipitation, filtration, calcination studies at 0.5 kg/hr rate, 4 percent loss to filtrate.

1956

- January Continuous Task I runs, effect of process rates, slurry holdup on waste losses.
- February Continuous precipitation studies, separation factors equivalent to batch process, effects of washing on filter cake quality. Life test (500 hours) for Dynel SD-9 filter cloth in continuous oxalate processes. Dissolution of Pu(IV) oxalate by aqua regia (without metallic contamination) for preparation of spectrographic analysis samples.
- March Effect of Zr, Versene, fluoride on Zr-NB DF's across batch oxalate precipitation. Continuous Task I studies, 1 kg/hr rate with 3 percent filtrate loss. Effects of agitation speed, slurry residence, filter cloth deterioration. Capacity of 3" diameter by 36" long calciner. Continuous prereduction of plutonium nitrate feed. Continuous precipitation from dilute (10 g/l) solutions with low filtrate loss.
- April Calciner operating temperature limits. Effect of temperature on calciner capacity.
- May Degradation of Dynel SD-9 filter cloth in Task I process solutions.
- November Continuous Task I operation with 120 g/l Pu in 5 M HNO<sub>3</sub> containing 0.65 M (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> (negligible effect).
- December Effect of 0.65 M (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> on sulfate precipitation in Task I feed solutions on standing at room temperature.

1957

- January Purex product 146 g/l Pu in 6 M HNO<sub>3</sub>, 0.8 M SO<sub>4</sub><sup>2-</sup> did not show precipitation after one-week storage. First cold runs in new continuous Task I unit used cerous nitrate (50 g/l Ce) feed, 1 M oxalic acid strike solution. Calcined at 300 C, with 15 minutes' residence. Throughput 250 g/hr.
- February Continuous Task I cold run was made under identical conditions of the first run. Rocker arm agitator modifications.
- May Operability and capacity of continuous Task I unit demonstrated in hot runs. Purex 2BP feed 75 - 100 g/l Pu in 4 M HNO<sub>3</sub>, 500 g/hr, 1 M oxalic acid, loss to filtrate 1 - 3 percent. Precipitation directly in filter pan.

- June Continuous Task I runs using Redox 3BP as feed  
95 g/l Pu in 4.5 M HNO<sub>3</sub>, 500 g/hr, 1 M oxalic acid,  
loss to filtrate 5 percent.
- Calciner residence time studies at 350 C. Fe im-  
purities 300 ppm.
- July Continuous Task I runs. Continuous oxalate chemical  
kill of filtrates.
- August Operation of continuous Task I on extended period to  
supplement Building production. 25 kg Pu, 10 percent  
waste loss, Dynel SD-9 cloth superior to 904 in runs.  
Material acceptable for fluorination and reduction.
- September Spectrographic results of the oxide produced in the  
sustained runs of August demonstrate the higher quality  
of oxide produced in continuous processing equipment.
- Runs demonstrate use of continuous Task I precipitation  
reactor with overflow of slurry into filter pan. Normal  
flow sheet. Effects of flow control on cake quality.
- October Continuous Task I runs demonstrate suitability of  
simultaneous reduction-precipitation process.
- Calciner surge capacity tests.
- November Calciner surge tests were continued. Surge capacity  
70 percent above normal steady state operating capacity.
- Test of cyclone separator and filter on calciner off-  
gas system.
- 1959
- March Effect of titanium and vanadium on Z Plant processing.
- Aluminum oxide filter media tested with plutonium(IV)  
oxalate slurry.
- Study of plutonium oxalate and filtrate densities.
- April Particle size distribution in plutonium(IV) oxalate  
slurry. 10 - 100 micron size range.
- Water content of filtered plutonium oxalate cake.
- May Purification study of the continuous oxalate process  
flow sheet using 200 g/l Pu in feed.

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## 5. F-10-P And 3BP Studies

### 1952

April Red precipitate in P-1 solution - analyses of solution and of solid; abnormal beta, gamma activities in sample cans and 3BP: solids in 3BP.

May Solids in 3BP and P-1.

### 1953

March Solids and Redox PR solution.

October Sludge in Redox PR can - high radiation.

### 1954

February Plutonium phosphate in F-10-P.

December F-10-P filtration and stabilization.

Unreported ANN in 3BP - concentration effects.

### 1955

May Filtration studies on Redox PR solution - no decontamination from FP's across solution filter.

### 1956

February Effects of filtration and precipitation on removal of Zr-Nb from Purex PR solution. DF of 1.2 obtained.

March Studies on use of silica gel or Dowex-1 column for Zr-Nb removal from Purex PR solution (DF's of 3).

Effect of versenes and fluoride on Zr-Nb DF across across oxalate precipitation.

May Solids in concentrated Purex PR solution - Pu polymer, silica, iron, phosphorus. Dissolution procedure.

Concentration of Purex 2BP.

July Solids in Redox L-3 to L-4 valve. Ru, Zr, Nb, Al, Cr, Cu, Fe, Ni identified.

August Study on solids in Purex tube bundle - primarily silica.

Study on precipitation in Redox E-1, E-3 vessels.

Studies on solids in Redox L-3 to L-4 valve.

October

Study of solids found in Purex concentrator. Contained high silica and iron. Si/Pu ratio of 1/5 in solids.

1957

March

Purex concentrator solids. Black flint-like material from Purex product concentrator contained mostly Si and Fe. Cr. Ni also detected, Pu was 1%.

1958

April

Solids in Purex ion exchange concentrator. Pu(IV) polymer and resin fines identified.

6. 224 Building Studies

1951

September

Barium sulfate scavenging.

October

Scavenging of barium.

1953

May

Calcium separation; tolerance to aluminum.

July

Tolerance to aluminum.

1954

April

Decontamination factors for various steps + omission of lanthanum fluoride by-product.

7. Double Salt Studies

1953

November Precipitation studies; large-scale strike.

December Precipitation studies.

1954

January Double salt precipitation; 300 gram reduction.

February Calcium-thorium fluoride precipitation.

March Separation factors for the calcium double salt.

April  
May Large-scale operations.

June Plant scale operations.

Unreported Decontamination factors for zirconium, niobium, and separation from sulfate.

## 8. Miscellaneous

### 1951

October Disposal of methanol.

### 1952

August Evaluation of filter aid for N-1 filter.

### 1953

August Stability of hydrogen peroxide; spectrophotometric determination of plutonium valence.

October Plutonium(III) oxalate precipitation - hydrogen peroxide, sulfamic acid reduction of Redox product.

### 1954

February Plutonium fluoro oxalate.

October Specific gravity measurements for plutonium nitrate - nitric acid solutions.

Unreported Specific gravities for plutonium polymer. Effect of fluoride on plutonium for light absorption.

### 1956

September Titanium corrosion in 8 M HNO<sub>3</sub>, 130 g/l Pu(IV), at B.P. 0.0003 ipm.

October Corrosion of tantalum, titanium, 304L stainless steel in HNO<sub>3</sub>-Pu solutions. Effect of sulfate ion. Ta and Ti superior.

December Recovery of Pu from sandpaper and sweeping by ashing at low temperature followed by 6 M HNO<sub>3</sub> - 0.25 M HF leaching at B.P. Pu recovery of 90%.

304-L corrosion rate in boiling solution of 0.003 M oxalic acid - 5 M HNO<sub>3</sub> containing 1.5 g/l Cr<sup>+3</sup>, 0.00018 ipm.

1958

January

Spectrophotometric method for nitrate to determine small amounts of nitric acid in presence of acids or salts yielding hydrogen ions.

August

Neptunium oxide ( < 200 ppm metallic impurities) produced by oxalate precipitation and calcination. Bulk density was 2.7 g/cc. Effect of radiation buildup noted.

1959

June

Initial studies of direct calcination of plutonium nitrate (275 g/l) to plutonium oxide and subsequent chlorination to  $\text{PuCl}_3$ .

## 9. Plutonium Trifluoride Precipitation

1956

April

Continuous precipitation and filtration of  $\text{PuF}_3$  from  $\text{HNO}_3$  solution using HF as precipitant. Effects of alcohol wash and filtrate loss. Sulfamic reductant.

Attempted precipitation of  $\text{PuF}_3$  from  $\text{HNO}_3$  solution using fluosilicic acid. No precipitation noted. Effect of complexing strengths of Pu(III) and Pu(IV) on fluoride.

June

Studies on  $\text{PuF}_3$  precipitation from high purity plutonium nitrate using HF precipitant.

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- September      Precipitation of  $\text{PuF}_3$  by HF from an  $\text{HNO}_3$  solution. Effects of reducing agents, and precipitation data included.
- October         Continuous precipitation from nitrate solution using 3 M HF. Feed reduced with  $\text{SO}_2$  + ascorbic acid. Filtrate losses 8 - 17%. Separation factors 10, 20, 10, 4 for Fe, Ni, Cr, Mg.
- December        $\text{PuF}_3$  dried at 600 C showed no weight change on standing 17 days.

1957

- March            Continuous ion exchange product subjected to  $\text{PuF}_3$  precipitation using 1 M excess fluoride. First strike slow filtering, 0.2 g/l Pu in filtrate. Second strike, spiked 1000 ppm Dow Separan 2610, fast filtration, 0.05 g/l Pu in filtrate.
- April            Small-scale continuous precipitation runs, feed 50 g/l Pu, 5 M  $\text{HNO}_3$ , precipitant 3 M HF. Valence adjustment with semi-carbozide. Separation factors 125, 147 for Al and Fe. Slurry degrades upon standing overnight.

10. Plutonium Tetrafluoride Precipitation

1955

- October         Aqueous precipitation of  $\text{PuF}_4$  from Pu(IV) nitrate solution using HF and ammonium fluosilicate.
- November       Precipitation of  $\text{PuF}_4$  from Pu(IV) nitrate solution using hydrofluosilic acid and HF.

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1956

January

Precipitation of  $\text{PuF}_4$  from 40 g/l Pu and 5 M  $\text{HNO}_3$  with 4 M HF. Fe and Al separation factors of 10 - 20 at initial impurities of 10,000 and 100,000 ppm respectively. Effects of washing on cake quality.

February

Studies on drying of the  $\text{PuF}_4$  precipitate.

## 11. Oxalate Kill

1955

September

Chemical kill of plutonium oxalate demonstrated with  $\text{KMnO}_4$  and  $\text{H}_2\text{O}_2$ .

1956

May

Investigation of conditions for batch and continuous killing of oxalate filtrate.

Studies on stability of Dynel SD-9 filter cloth in oxalate precipitation chemicals.

June

Studies on continuous evaporation of killed oxalate filtrates.

Studies on evaporative heat kill of oxalate filtrates.

July

Prototype continuous oxalate filtrate kill equipment installed. Studies on continuous oxalate kill.

Studies on evaporative heat kill of oxalate filtrate.



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November Adaptability of Recuplex dissolvers for evaporative oxalate kill process.

1957

January Pu nitrate solutions containing sulfamic acid or semi-carbazide, reoxidized and concentrated by a factor of five in a continuous kill. Feed solution was 6 M HNO<sub>3</sub>, 0.05 M semi-carbazide, 11-50 g/l Pu. Feed solution slowly added to boiling solution to safely accomplish the kill.

July Filtrate from continuous oxalate precipitations treated by continuous chemical kill using KMnO<sub>4</sub> at 45 C. Permanganate leg of the "U" column operated satisfactory. Peroxide leg (1" diameter) reaction too vigorous at 10% or 2-1/2% peroxide concentration. Increased diameter of peroxide leg is required.

October Evaporative kill experiments indicate manganous ion addition may be needed as catalyst, due to low amount of iron in filtrate.

November Evaporative kill studies indicate a 21-minute decomposition half-time. Small amount of solids can be tolerated in the evaporative kill process.

## 12. Plutonium Polymer

1956

August Limiting conditions for formation of plutonium polymer in nitric acid solution.

November Formation of plutonium polymer in nitric acid solutions. Absorption peaks of polymeric solutions. Effect of nitric acid concentration and HNO<sub>3</sub>/Pu ratio on polymer formation. Temperature effects.

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December Study of polymer formation in Pu(IV) nitrate solutions of low acidity. Effects of nitric acid concentration, and temperature. Absorption spectra of Pu(IV) from 0.02 M HNO<sub>3</sub> to 14 M HNO<sub>3</sub>.

1957  
February An E-7 sample from Redox studied spectrophotometrically, found to have a spectrum characteristic of polymeric plutonium. Depolymerization in 4 M HNO<sub>3</sub> produced plutonium(IV) spectrum.

1958  
February Study of polymer formation as a purification process. Separation factors of 3300, 40, 1000, 5, 100, 100 for Al, Cr, Cu, Fe, Mg, Ni based on initial impurity concentrations of 10,000 ppm. Effect of precipitation on fine particle formation.

### 13. Ion Exchange

1956

November Initial testing of semi-continuous Higgins-type columns.

December Equipment testing results. Flow control problems on continuous equipment.

1957

January Mechanical testing of hydraulic ram, conductivity probe, and pumps.

February First cation hot run completed. Steady state not achieved because of excessive gassing. Run conditions:

XAF 35 ml/min	Dowex 50W-X8 resin
XCS 2.1	
XAS 2.0	10-minute cycle
XCIS 5.0	Feed concentration 8.3 g/l Pu
XAX 2.0	Product concentration 25 - 70 g/l Pu
	Waste losses ~ 1%

Effects of feed reductants discussed.  
 Elutant was 6.4 M HNO<sub>3</sub>, 0.3 M sulfamic or semicarbazide.

March

Workable flow sheet was demonstrated in a two-week-long "hot" run of cation columns. Run conditions:

XAF 39.5 ml/min	10-minute cycle
XCIS 5.0	Dowex 50W-X8 resin
XAS 2.0	Feed 2.5 - 7 g/l Pu
XCS 2.1	Product 35 - 75 g/l Pu
XAX 2.0	Waste loss 0.1%

Most promising feed reductant was 0.1 M NH<sub>4</sub>HSO<sub>3</sub>.

Effects of plutonium polymer formation during feed preparation.

Most promising elutant 5 M HNO<sub>3</sub>-0.33 M sulfamic acid.

Higher product concentration may require cooling of XC column.

Stability of cation exchange product: oxidation of XCP dependent on solution acidity. Oxidation rate 0.2 - 5. %/day. Effects of other reagents discussed.

Adjustment of Purex 2BP to 7 M HNO<sub>3</sub> resulted in oxidation to Pu(IV) within two minutes, for anion exchange feed.

April

Tentative Purex flow sheet (HW-49524) was demonstrated in a hot anion exchange run.

Run conditions:

Resin Dowex 1-X4 (nitrate form) 5-minute cycle  
 Feed concentration 6 g/l  
 Product concentration 45 g/l Pu  
 Column temperature 50 C

Effect of column temperature, feed flow rates, etc., discussed.

May

Continuous runs with Purex 2BP as feed solution showed anion exchange to be superior to cation process for

purification and decontamination.

1. Anion Zr-Nb DF of 28, Ru-Rh DF of 2.0  
Cation Zr-Nb DF of 3, Ru-Rh DF of 3.0
2. Total gamma DF of 3 for both anion and cation.
3. Anion reduces silicon 10 - 20 fold, silicon not removed in cation process.
4. Anion: separation factors for (Cr, Cu, Fe, Mn, Ni, Pb) of 5 - 20 and (Al, Ca, K, Mg, Na) of 100.

Cation removed only slight amounts of light metals, none of the heavy metals.

June

Changes made in flow, and temperature control to permit sustained anion exchange runs at high throughput with Dowex 1-X4 anion resin.

July

Operation of the anion exchange system on a Purex flow sheet in the temperature range of 50 - 70 C and feed acidity range of 5 - 7.2 M HNO<sub>3</sub> was satisfactory. Flow rates up to 30 percent greater than the 54 mg Pu/mir/cm<sup>2</sup> required by flow sheet were used. Product concentration was near the 58 g/l equilibrium value. The ratio of XSW to XAX appears to depend on the volume of resin pushed. Satisfactory resin push was attained with 10 psi pressure. Zr-Nb DF = 250, Ru-Rh DF = 30 when adequate scrub flows are used. Appreciable FP's accumulate on the resin and are difficult to remove. Metallic separation factors varied from 5 - 100. Effects of temperature control, acidity, and column dimensions are recorded.

## 14. Metal Dissolution

1957

September

Plutonium dissolved in 15 weight percent sulfamic acid solution. Dissolution not as rapid as  $\text{HNO}_3$ -HF, no residue left as in  $\text{HNO}_3$  - sulfamic mixtures. Concentrations 100 g/l Pu, in 0.3 - 1 M acidity, stable for three weeks.

1958

March

Initial tests of continuous dissolution of plutonium metal (10 g scale) in  $\text{HNO}_3$ . Effect of  $\text{PuO}_2$  formation.

1959

April

Study of massive plutonium metal dissolution using sulfamic acid. Effects of final oxidation adjustment, plutonium sulfate and plutonium polymer formation.

Study of reactive alpha Pu skull passivation by caustic solution.

May

Study of reactivity of Pu-Al skulls during caustic passivation.

June

Problem of unusual reactivity of temperature cycled alpha Pu skulls.

## 15. Solution Concentration

### 1955

September Concentration limits of oxalate filtrates containing 200,000 ppm Fe, 10,000 each Al, Cr, and Ni was a factor of three. Oxalate destroyed by concentration.

October Concentration studies for Redox 3BP containing 100,000 ppm Al. ANN precipitation noted at 130 g/l Pu concentration.

### 1957

March Concentration of cation exchange product, XCP (0.33 M sulfamic) forms sulfate precipitation after concentration by a factor of two.

April Cation exchange product:

Solution of 65 - 80 g/l Pu, 5.4 - 6.8 M HNO<sub>3</sub>, 0.5 M SO<sub>4</sub><sup>=</sup> stable at room temperature.

Solutions up to 150 g/l Pu, 7 M HNO<sub>3</sub>, 0.8 M SO<sub>4</sub>=  
stable at boiling temperature.

16. Valence Adjustment

1956

- April Pu(IV) to Pu(III) reduction studies using H<sub>2</sub>O<sub>2</sub>, hydroxylamine hydrochloride, with sulfamic acid as holding reductant. Solutions were 17 - 70 g/l Pu and 0.8 - 2.5 M HNO<sub>3</sub>.
- June Studies of Pu(IV) to Pu(III) reduction in HNO<sub>3</sub> solution using sulfamic acid and ascorbic acid. Effects of solution concentration.
- September Ascorbic acid stability tests. Effect of sulfamic acid, nitric acid, and plutonium concentration.
- October Ascorbic acid stability  
  
Use of ascorbic acid as Pu(IV) to Pu(III) reductant for HNO<sub>3</sub> solution feed, ion exchange process. Effect of ascorbic and sulfamic acid. Adaptability of ascorbic-sulfamic bearing filtrate solutions to batch and continuous kills temperature effects.
- November  
  
Effect of semi-carbazide and aminoguanidine as reducing agents for ion exchange process. Effects of sulfamic acid on reduction stability.

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Preliminary study on reduction of Purex 2B stream with sulfurous acid, ascorbic acid, and hydroxylamine sulfate. Initial studies with sulfurous acid fail to reduce the Pu in the 2B stream.

December

Effectiveness of hydroxylamine, ascorbic acid, amino-guanidine, and semi-carbazide as reductants in high acid solutions. Effects of time and oxidation. Semi-carbazide is slower, but more stable than others.

TBP strip experiments on Purex 2BP show ascorbic acid and hydroxylamine as effective reductants. Sulfurous acid ineffective.

1957

March

Study of reduction of cation exchange feed to Pu(III). 0.05 M hydroxylamine sulfate, 0.05 M sodium bisulfite, and 0.1 M semicarbazide required 24 hours, 2.5 minutes and 0.5 minute, respectively, for reduction. Hydroxylamine sulfate reduced solution stable for two weeks, others for four days.

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1959

January

Test of reducing agents for Pu reduction to (IV) valence in hydrolysis-precipitation recovery method for nickel-coated MgO crucibles.

Large-scale (one crucible and slag) dissolver has been installed for chloride slag and crucible recovery development.

February

Study of valence adjustment and precipitation parameters for hydrolysis-precipitation method of reprocessing nickel-coated MgO crucibles.

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B. DRY CHEMISTRY

1. Processing Of Metal Turnings

1951

August                      Hydrofluorination.  
September

2. Hydrogen Fluoride Absorption

1951

September                      Miscellaneous.  
October

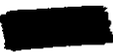
1952

February                      Dorex cannister capacity.

June                              Caustic scrubber for HF absorption.  
July

September                      Calcium carbonate.

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3. Hydrofluorination Of Plutonium Peroxide

1951

October      Effect of air leakage.

4. Fluoride Analysis

1952

January  
February      Analysis by conversion to oxide.

November      Analysis by conversion to oxide.

5. Filter Boats

1952

May              Losses in drying - filter boat.

June             Filter boat leak tests.

July              Filter paper liners.

October         Filter boat cleanout - use of oxalic acid or hydrogen peroxide.

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6. Freonation

1952

July First work.  
December Additional results.

1953

February Two deep cake runs.  
April Work abandoned: corrosion; polymerization.

7. Calcination

1952

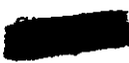
September Drying of plutonium(IV) oxalate: 150 C, lumps; 300 C for calcination; thermal decomposition of plutonium(III) and (IV) oxalate.  
October Plutonium(IV) oxalate drying procedure - glowing boat; temperature measurements.  
December Drying of plutonium(IV) oxalate (130 C - hexahydrate).

1953

February Plutonium(IV) oxalate decomposition - exothermic reaction.  
May  
June Calcination of plutonium(IV) oxalate; reactivities of oxides (HCl-KI).  
July PuO<sub>2</sub> reactivities (wet).  
September Calcination of plutonium oxalate.  
October Thermal decomposition of plutonium oxalate.

1954

February Infra-red drying and calcining of oxalate.

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March                    Decomposition of plutonium(IV) oxalate; reactivity of PuO<sub>2</sub>.

1956

April                    Operating temperature range of 300 - 400 C for continuous Task I calciner determined from laboratory tests.

8. Hydrofluorination

1952

July                    Summary of dry chemistry and reduction data.  
September            Cake depth vs. hydrofluorination rates.  
October                Tolerance to laboratory air - 37 percent.  
November             Hydrofluorination of erupted boat load; tolerance to laboratory air.

1953

January  
February                Preparation of reduction of low temperature fluorides.  
September             Direct hydrofluorination.  
October                Direct hydrofluorination of oxalate - PuF<sub>3</sub> production.

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1954

November CeF<sub>3</sub> production; conversion of blue powder to PuF<sub>4</sub>.

December Conversion of blue powder to PuF<sub>4</sub>.

1955

February Continuous PuO<sub>2</sub> hydrofluorination studies with 97 percent conversion to fluoride; metal reduction yields were 98 percent.

May Continuous PuO<sub>2</sub> hydrofluorination studies.

June Continuous PuO<sub>2</sub> hydrofluorination studies.

August Study on SiC filters as replacement of platinum frits for Task II.

December Continuous PuO<sub>2</sub> hydrofluorination studies, processing rate 0.4 kg/hr, PuF<sub>3.72</sub>, reduction yield of 98.6 percent on PuF<sub>4</sub> product.

1956

March Continuous PuO<sub>2</sub> hydrofluorination studies, 100 percent conversion to PuF<sub>4</sub>.

May Operating temperature range for continuous hydrofluorinator determined to be 450 - 550 C.

September Simulated operation of plant reactor tested by conveying powder in a vibrating oval tray.

1957

January A plastic (Teflon and fluorothene) rotary feed valve replaced inconel feed screw to eliminate corrosion in continuous hydrofluorinator.

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## 9. Chlorination

1958

February

Initial  $\text{PuCl}_3$  preparation by  $\text{CCl}_4$  (in argon carrier) at 450 C in Pyrex.

$\text{PuCl}_3$  hydration path and rate study. Effect of closed container storage on hydration.

March

Plutonium trichloride hydration study resume'. Effects of atmosphere and powder storage.

$\text{PuCl}_3$  preparation (600 g) by  $\text{CCl}_4$  treatment at 450 C resulted in Cl/Pu molar ratio of 2.94.

Study on drying of  $\text{PuCl}_3 \cdot 6\text{H}_2\text{O}$  by heating.

$\text{PuCl}_3$  process off-gas line plugging problems.

Neutron count study of  $\text{PuCl}_3$  shows greater reduction in neutrons from  $\text{PuCl}_3$  than  $\text{PuF}_4$ .

April

Chlorination rate study for conversion of  $\text{PuO}_2$  to  $\text{PuCl}_3$  with  $\text{CCl}_4$ . Effect of temperature on rate and powder density.

May

$\text{PuCl}_3$  prepared in large-scale batches by  $\text{CCl}_4$  in argon carrier at 500 C.

Batch tests of  $\text{PuO}_2$  chlorination by  $\text{COCl}_2$  gave 97 percent conversion, decreased reagent consumption, more desirable off-gas by-products.

Small Pyrex vibratory tube chlorinator assembled to permit continuous process study.

June

Chlorination rate study for conversion of  $\text{PuO}_2$  to  $\text{PuCl}_3$  with  $\text{CCl}_4$ . Reaction products identified as  $\text{C}_6\text{Cl}_6$ ,  $\text{C}_2\text{Cl}_4$ ,  $\text{C}_2\text{Cl}_6$ ,  $\text{CO}_2$ ,  $\text{CO}$ ,  $\text{Cl}_2$ ,  $\text{COCl}_2$ .

Unsuccessful attempts to produce  $\text{PuOCl}$  at 225 - 400 C.  
By-products from the reaction are  $\text{Cl}_2$ ,  $\text{HCl}$ ,  $\text{CO}_2$ .

Chlorination of  $\text{PuO}_2$  by  $\text{COCl}_2$  at 350 - 550 C;  $\text{Cl}_2$ ,  
 $\text{HCl}$ ,  $\text{CO}_2$  off-gas products.

Direct chlorination of plutonium(IV) oxalate by  $\text{COCl}_2$   
studied; 175 - 450 C on 100 g scale. Reaction rate and  
powder density comparison.

Preliminary testing of continuous 1" diameter chlorina-  
tion reaction used cerium oxide.

Corrosion tests by D&CCO show Hastelloy B and C to be  
suitable for  $\text{COCl}_2$  system at 400 - 600 C. Effect on  
other materials recorded.

July Large-scale batch chlorinations of  $\text{PuO}_2$  by  $\text{COCl}_2$  con-  
tinued. Effect of calcination temperature of  $\text{PuO}_2$  on  
chlorination rate. High temperature chlorination (600 C)  
introduced unknown residues in off-gas.

Plastic materials' resistance tested in phosgene atmosphere.

August  $\text{PuCl}_3$  production by  $\text{COCl}_2$  chlorinating agent.  
Conversion of  $\text{PuO}_2$  to  $\text{PuCl}_3$  by  $\text{H}_2$  and  $\text{HCl}$ . Effect of  
temperature on chlorination rate. Formation of  $\text{PuOCl}_2$ .

September  $\text{PuO}_2$ , previously heated to 900 C, was successfully chlorina-  
ted with  $\text{COCl}_2$  at 500 C.

Report, HW-57498, was issued to cover the status of the  
 $\text{PuCl}_3$  program. Flow sheets for preparation of the com-  
pound, reduction to metal, and recovery schemes are  
included.

October  $\text{PuCl}_3$ , prepared at 600 C, was a dark-green crystalline  
material in contrast to the blue-green powder produced  
at 500 C.

Continuous 1" diameter chlorinator operated with  $\text{PuO}_2$   
and  $\text{COCl}_2$  for  $\text{PuCl}_3$  conversion at 500 C.

November  $\text{PuO}_2$ , previously heated to 1800 C, was chlorinated to  
 $\text{PuCl}_3$  in five hours using  $\text{COCl}_2$  at 450 C.

Study of hydration rates of  $\text{PuCl}_3$  completed over range  
2 - 75 percent relative humidity. A discussion and  
graph are included.

December Description of 1" diameter continuous chlorinator system.

Continuous 1" diameter chlorinator operated at 34 g/hr on  
extended run conditions. Chlorination of  $\text{PuO}_2$  by  $\text{COCl}_2$ .

was complete.

Samples of  $\text{PuCl}_3$  from batch and continuous processing were found to have four and six micron particle size respectively.  $\text{PuO}_2$  powders for the conversion were 0.2 micron size.

Large-scale batch conversions of  $\text{PuO}_2$  to  $\text{PuCl}_3$  by  $\text{COCl}_2$  were made for reduction studies.

Off-gas studies for  $\text{COCl}_2$  chlorination completed over 275 - 600 C range. Products are  $\text{COCl}_2$ , CO, HCl, and  $\text{Cl}_2$ .

## 1959

### January

Continuous chlorination of  $\text{PuO}_2$  by  $\text{COCl}_2$  in the 1" diameter tube operated at 70 g/hr on an extended run.

Run conditions:

Powder bed depth: 3/8"  
Residence time in furnace: 45 minutes  
 $\text{COCl}_2$  flow: 200 percent excess  
Temperature: 500 C

Effects of temperature and flow rates on the conversion are included.

Pu/Cl ratio = 2.7  
 $\text{COCl}_2$  retention in cake:  $0.02 \frac{\text{mols } \text{COCl}_2}{\text{mol } \text{PuCl}_3}$

Corrosion studies in  $\text{COCl}_2$  system over 100 - 500 C range. Hastelloy B and C most promising materials.

Off-gas analysis study indicates gas composition to be dependent on temperature, phosgene flow, and powder bed composition. Elimination of off-gas solid deposits of metal chlorides discussed.

### February

Lumping problems of  $\text{PuCl}_3$  in the 1" diameter continuous chlorinator.

Study of brown solid deposits on chlorination tube.

### March

Initial runs in the large Hastelloy continuous chlorinator at 90 g/hr Pu rate. Mechanical difficulties encountered.

Off-gas filter problems.

Document HW-59759, "The Analysis Of Exhaust Products In

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The Chlorination Of Plutonium Oxide With Phosgene",  
was issued to summarize off-gas studies.

April

Large continuous chlorinator operated at 295 g/hr Pu  
rate. Phosgene flow, cake lumping problems.

May

Off-gas filtration development.

June

Chlorination of PuO<sub>2</sub> from direct calcination of plutonium  
nitrate solution.

Elimination of lump formation in the large continuous  
chlorinator.

Chlorination of PuO<sub>2</sub> by CO plus Cl<sub>2</sub>.

Test of Davis phosgene detector.

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## C. METAL REDUCTION

### 1. Reduction Of Off-Standard Fluorides (Blue And Low Temperature)

#### 1951

July Reduction of  $\text{PuF}_3$ .

#### 1952

July Sintered  $\text{PuF}_4$  reduction.

October Reduction of  $\text{PuF}_3$  from freon halogenation.

#### 1953

April Reduction of low temperature fluorides.

May Blue fluoride reductions.

July  
August Blue fluoride reductions - plant scale.

#### 1957

July Nine hundred grams of "blue fluoride" was received from the processing operation. The material was divided into two batches and reduced.  $\text{I}_2/\text{Pu}$  was 0.8, excess Ca was 0.6. Yields were 98 percent, metallic impurities < 550 ppm. This indicates poor reduction from wet precipitated fluorides are probably due to oxide formation on the powder surface.

### 2. Sulfur Booster

#### 1952

April Melting points of slags; addition of calcium iodide; uranium reductions; use of mixed magnesium and calcium; addition of  $\text{I}_2$ .

May Reduction of PuF<sub>4</sub>; addition of sodium fluoride; calcium-magnesium, calcium-strontium alloys.

### 3. Button Reactivities

1954

August  
September Miscellaneous entries.  
November

### 4. Miscellaneous

1951

September Bomb fusion of turnings.

1952

April Use of gallium oxide.

July Air reductions.

August Air reductions; preparation and use of gallium oxide.

September Use of gallium oxide.

1953

June Use of single crucible for different batch sizes.

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November Large-scale reduction of calcium plutonium fluoride; small reductions; reduction of calcium fluoride, plutonium fluoride mixtures.

1954

January Test of new Task III pressure vessel.

Unreported Spectrochemical analysis of metal from double salt reduction.

1955

July Studies on hot water pickling of reduction buttons.

August Evaluation of Nelco Calcium without redistillation or grinding for Task III.

December Oxidation of calcium metal in unreacted  $\text{PuF}_4$ -Ca mixtures to eliminate  $\text{H}_2$  evolution during powder dissolution.

1956

January Reduction of  $\text{CeO}$  to metal using Ca plus  $\text{H}_2$  booster showed cerium formation. Effects of slagging agents noted.

May Reduction of  $\text{PuO}_2$  by Ca resulted in no segregation or coalescence of metal.

Cerium oxide reduction by Ca resulted in metal beads (cerium) up to 1/4" diameter.

June Attempt to reduce 40-gram  $\text{PuO}_2$  resulted in no visible metal formation.

1958

January Study of calcium reactivity and pressure buildup associated with reduction in Task III. Effect of moisture and  $\text{I}_2$  on  $\text{PuF}_4$  and Ca mixtures at room temperature.

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5. Metal Recovery

1951

September Bomb fusion of turnings.

1955

September

October Controlled oxidation of metal turnings in argon and oxygen, subsequent fluorination in Freon-12.

Direct conversion of metal turnings to PuF<sub>3</sub> with argon and Freon-12.

December

1956

January

March

1957

November Recovery of plutonium from skulls by vacuum melting - impractical on small or badly-oxidized fragments (10 percent yields). Vapor phase reduction with calcium was unsuccessful. Experiments started on melting skulls in CaCl<sub>2</sub> - CaF<sub>2</sub> eutectic.

December Glascast crucibles fired at 2300 F will contain the CaCl<sub>2</sub> - CaF<sub>2</sub> melt at 1000 C. Magnesia, CaF<sub>2</sub>, quartz, sodium silicate impregnated magnesia were ineffective. Study of Ca reduction of PuO<sub>2</sub> in a CaCl<sub>2</sub> - CaF<sub>2</sub> melt.



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1959

January

Test on recovering Pu from skulls was successfully made by melting skulls with Ca-CaCl<sub>2</sub> at 1050 C. Recovery was >90 percent.

February

Recovery of skulls by calcium chloride salt bath.

6. High Purity Plutonium Metal

1955

September

0.5 kg high-purity Pu prepared.

1957

January

900-gram pure metal has been made. Total impurities of most recent batch is < 500 ppm.

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## 7. Plutonium Trichloride

1958

March            Reduction of  $\text{PuCl}_3$  by Ca. Effects of moisture concentration.

Study of chloride slag soaking into MgO crucibles. Use of Glascast reduction liners, and effect of calcium and plutonium on Glascast.

Test of Al slag and crucible cans exposed to  $\text{CaCl}_2$ ,  $\text{CaI}_2$ , and  $\text{CaF}_2$  at room temperature. Physical strength of cans adequate after 30-day exposure.

April            Small-scale (30-g)  $\text{PuCl}_3$  batch reduction. Effects of water, iodine, and calcium concentrations were investigated. No serious crucible (MgO) sticking problem on 30-g scale.

Effect of Si and B pickup by Pu from Glascast crucibles.

May              Large reductions (0.6 - 1.0 kg) of  $\text{PuCl}_3$  investigated. Button quality acceptable. Slag soaking problem in large reduction. Effect of dual crucibles as slag container.

June             Nickel-coated MgO crucibles prevent slag soaking and crucible sticking in reduction of  $\text{PuCl}_3$  to Pu.

Acceptable reductions made with  $\text{PuCl}_3$  containing up to one percent water.

$\text{PuCl}_3$  reduction yields are >96 percent.

Effect of  $\text{PuCl}_3$  storage on reduction chemistry.

- July Effect of batch reduction compounds on various crucible materials studied.
- Study initiated to determine optimum nickel-coat thickness for MgO reduction crucibles.
- A phase diagram for  $\text{CaCl}_2 - \text{CaI}_2$  system was determined.
- Effect of plutonium trichloride produced from plutonium oxalate on the batch reduction characteristics.
- August Large-scale batch reductions of  $\text{PuCl}_3$  to Pu in nickel-coated MgO crucibles. Yields and purity satisfactory when crucible does not break.
- Coating thickness study continued.
- $\text{PuCl}_3$  stored 260 hours at 8.5 percent relative humidity was successfully reduced to metal.
- September  $\text{PuCl}_3$  stored 1300 hours at 8.5 percent relative humidity was reduced to metal with a 72 percent yield.
- October  $\text{PuCl}_3$  formed at 600 C required additional Ca-I<sub>2</sub> booster for good reduction to metal.
- A mixture of 55 percent  $\text{PuCl}_3$  - 45 percent  $\text{PuOCl}$  was reduced to metal by calcium, with a 95 percent yield.
- Study of optimum amount of water tolerated in  $\text{PuCl}_3$  to give 96 percent reduction to metal. 0.6 percent water can be tolerated..
- November Study to determine feasibility of eliminating the chemical mixing step in  $\text{PuCl}_3$  batch reductions. Effects of chemical segregation are discussed.
- Use of salt-impregnated MgO crucibles was tested in 30-g scale  $\text{PuCl}_3$  batch reductions.
- Effect of substitution of  $\text{CaF}_2$  for MgO sand in  $\text{PuCl}_3$  reductions.
- December Study of  $\text{PuCl}_3$  reduction with elimination of the mixing step
- 1959
- January Reduction of  $\text{PuCl}_3$  from continuous chlorinator was success-

ful. Powders used had bulk densities of 2.0 - 2.9 g/cc, no differences in reduction characteristics noted.

Tests of reduction with all of Ca and I<sub>2</sub> on top of PuCl<sub>3</sub> have resulted in 75 percent yields. Method was successful when all of I<sub>2</sub> and 2/3 Ca were on top and remainder of Ca below the PuCl<sub>3</sub>.

A copper-coated MgO crucible was used successfully in a 700-gram PuCl<sub>3</sub> reduction.

A 30-g button was recovered when 50 g of Pu as PuCl<sub>3</sub> were dropped with calcium into a crucible at 800 C. The semi-continuous reduction atmosphere was argon.

February

Batch studies of PuCl<sub>3</sub> reductions without mixing of chemicals.

Reaction bomb loading safe temperature limits.

Reduction of lumps chlorides.

Tests of dual reduction liner technique in PuCl<sub>3</sub> reduction.

March

Study of steel-coated MgO crucible in PuCl<sub>3</sub> batch reductions.

Batch studies of PuCl<sub>3</sub> reduction without mixing chemicals, 98 percent reduction yield.

April

Reduction of PuCl<sub>3</sub> from large continuous chlorinator runs were successful.

Study of tantalum liners for batch reductions.

Phase diagram study for CaI<sub>2</sub> - CaCl<sub>2</sub> slag summarized in HW-60151, "Calcium Iodide - Calcium Chloride System".

May

Test of reusable NaCl-KCl impregnated MgO crucible for PuCl<sub>3</sub> batch reductions 20 - 200 gram scale.

Test of reusable nickel-coated MgO crucible on 700-gram PuCl<sub>3</sub> batch reductions.

June

Reusable crucible - studies.

PuCl<sub>3</sub> reduction studies in tantalum crucibles.



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D. RECUPLEX1. Slag And Crucible Dissolution1951

August Odd experiments.

October Comparison of Los Alamos and Hanford procedures.

November Flow Sheet No. 1.

December Additional dissolvings.

1952

January Dissolution of sulfur slags.

May Corrosion rates for dissolution.

June Analysis of filter cakes; removal of iodate-sodium nitrite addition.

July Silica analysis.

August Plutonium holdup from dissolution.

September Slag and crucible dissolution.

1953

May  
June Filtration of slag crucible solution - alundum ( $Al_2O_3$ ) plate.

November Dissolution of calcium fluoride - calcium iodide - calcium sulfide slag.

1954

January Silica in off-gas.

March Aluminum-to-fluoride studies; plutonium(IV) in SCF.

April Iodate formation.

June Silica in off-gas.

October Slag and crucible dissolution - SCX-1 corrosion rates, Haynes 25 and stainless steel in dissolution; iodate method.

1955

- May                    Studies on dissolution of reduction slags, effects of  $CaI_2$  content.
- July                    Studies on hydrogen formation in dissolver reactions.
- August                Studies on dissolver off-gas. Analysis of silica cake slurry.

1958

- February            Study of steel slag and crucible can dissolution in 8 M  $HNO_3$  - 0.3 M HF.

2. Miscellaneous Feed Preparation

1951

- November            Clarification of F-10-P filtration; use of carbon  
December            tetrachloride.

1952

- September        Use of nitrite for feed adjustment (3BF, F-10-P).

1954

- August              Precipitation in F-10-P solution.
- September        Valence adjustment.

October F-10-P and solvent extraction - greases; valence adjustment and F-10-P, 3BP.

1955

March Valence adjustment studies on oxalate filtrates.

May Valence adjustment studies on oxalate filtrates.

June Effects of ferric ion on valence adjustment studies for oxalate filtrates.

August Studies on reactivity of H<sub>2</sub>O<sub>2</sub> as associated with Recrux feed makeup.

3. Extraction Studies

1952

February Disengaging times for CAF-CAX as a function of specific gravities.

June Extraction coefficient values.

July Extraction coefficient values.

August Effect of iodate, sulfate, hydrogen peroxide - adjustment on extraction coefficients.

September Extraction coefficients.

1953

August Life test of coated plates; coefficients for HW-10.

October Plate testing solvent stability.

November Plate testing.

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1954

January Flow Sheet EX-1: effect of oxalate; solvent decompositions.

April EX-1 batch, counter-current runs; effect of oxalate - function of plutonium concentration.

July Coefficients for Recuplex system; total reflux.

August Coefficients for plutonium(VI).

September Coefficients for 20 percent TBP; organic phase saturation; small column studies.

December Coefficients for CC column - function of time.

1955

January Study of decrease of  $E_g^0$  in the Recuplex stripping column when CCX is heated to 50 C.  
  
Proposed uranium-plutonium partition flow sheet for Recuplex. H-1 and H-2 columns or batch processing in the solvent treatment tank.

February Solvent decomposition studies, effect of plutonium concentration and solvent age.

March Solvent damage studies, effects of  $CCl_4$ , formation of  $Cl^-$  corrodant.  
  
Plutonium distribution studies and an equation for scrub section of H-1 column.

May Recuplex flow sheets C-9 and C-11 tested in laboratory columns.

June Recuplex flow sheet C-11 studies. Solvent damage studies.

August Valence studies on Recuplex concentrated COP stream.

1957

August Dibutyl butylphosphonate (DBBP) results in more favorable extraction of the plutonium into the organic phase than TBP. However, stripping from DBBP has been ineffective. DBBP is more stable than TBP in aqueous  $HNO_3$  solutions. Tetrabromoethane (TBE) will be tested as a density-increasing agent. Charts relating density to composition for the TBP-TBE- $CCl_4$  system have been prepared.

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Solvent density increase will permit plant capacity increase.

September Further work confirms more favorable distribution coefficients for DBBP over TBP. Satisfactory stripping could not be attained by acid adjustment. Hydroxylamine sulfate (0.1 M) in CCX resulted in satisfactory stripping.

October Satisfactory extraction and stripping was observed in batch tests using DBBP-TBE- $\text{CCl}_4$  solvent. CCX stream was 0.05 M  $\text{HNO}_3$  - 0.05 M  $(\text{NH}_2\text{OH})_2\text{SO}_4$ .

December Further extraction tests of the DBBP-TBE- $\text{CCl}_4$  solvent shows a decrease in distribution coefficient due to presence of TBE.

Pentachloroethane, a high-density solvent (possible replacement for  $\text{CCl}_4$ ) exhibited much lower extraction coefficients for Pu(IV) with TBP and with DBBP than found with  $\text{CCl}_4$ .

Trimethoxyboroxine (TBM) was subjected to Recuplex-type extraction and stripping. 90 percent of TBM went to raffinate. TBM is a possible extinguishing agent for Pu fires.

1958

January Extraction study shows TBE in the DBBP- $\text{CCl}_4$  system decreases distribution coefficient of unsalted feeds and in stripping. No material effect on highly-salted feeds. Pentachloroethane behaves similarly when in place of  $\text{CCl}_4$ .

Effect of TBE and pentachloroethane on solvent degradation.

Corrosion study of stainless steel extraction plate attacked by TBE-DBBP- $\text{CCl}_4$ , TBE,  $\text{CCl}_4$ , and pentachloroethane.

February Investigation of Recuplex extraction column emulsification, stripping column precipitation. Effect of dispersed silica and TBP degradation products on extraction. Effect of fluoride on stripping.

Extraction coefficient studies on solutions from dissolution of steel slag and crucible cans.

June Stripping from Recuplex DBBP solvent resulted in much poorer results than previous work showed. Degradation

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products from DBBP act similar to DBP in TBP system.

August

Addition of "Mistron 28" to Recuplex sump cleanout solutions eliminated column emulsification problems during cleanout runs. Solvent was TBP-CCl<sub>4</sub>.

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## 4. Product Concentration

1953

August            Plutonium - red oil reactions.  
October            Evaporation of CCP.

1954

January            Red oil and precipitation of peroxide or oxalate.

## 5. CAW Treatment

1952

April            Neutralization of CAW; viscosities; setting characteristics.  
May              Viscosity of neutralized CAW.  
November        Neutralization of CAW.

1954

October          Protective coatings for CAW crib.

## 6. Miscellaneous

1951

December        First cycle waste evaporation; corrosion and material testing.

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## 1952

<u>January</u> <u>February</u>	Caustic scrubber solutions; cannisters.
<u>April</u>	First cycle waste evaporation; corrosion and material testing.
<u>July</u>	Physical properties of process streams.
<u>August</u>	Solvent treatment; first cycle waste evaporation; corrosion and material testing.
<u>September</u>	Solvent treatment.
<u>October</u>	Sintered stainless steel - porosities; resistance of rubber to nitric acid, hydrogen peroxide.  Corrosion of various materials in hydrofluorination off-gases.
<u>November</u>	Dissolution of fluorides in ANN-nitric acid.  Porosities of platinum discs.
<u>December</u>	Corrosion of alloys in HF-HNO <sub>3</sub> mixes; G-9 asbestos gaskets in iodine.

## 1953

<u>February</u>	Skull dissolution; platinum liner destroyed.
<u>May</u> <u>June</u>	Dissolution of outstanding reduction charges (caustic dissolution tried).
<u>August</u>	Chemical resistance of Homalite.
<u>November</u>	Corrosion tests - metals in nitric - ANN solutions; corrosion in freon.
<u>December</u>	Corrosion in freon.

## 1954

<u>January</u> <u>March</u> <u>November</u>	Filtration of aluminum nitrate.
<u>February</u>	Fluoroethene stability in process streams.
<u>September</u>	Use of Haynes 25 for HNO <sub>3</sub> - HF systems; Elgiloy in

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HNO<sub>3</sub> - HF vapors.

October Corrosion of Haynes 25 in HNO<sub>3</sub> - HF solutions.

December Filter cloth or aluminum nitrate filtration.

Stainless steel corrosion in slag and crucible solutions.

1958

March Scavenging of Recuplex solvent by precipitation of PuF<sub>4</sub> from CAX (15% TBP in CCl<sub>4</sub>) by HF or HF and NaF. Residual Pu in organic ~ 0.01 g/l.

## E. MISCELLANEOUS

1952

November  
December

Nickel carbonyl analysis.

1953

December

Reaction of plutonium metal in nitrogen or in CO<sub>2</sub>; ferrous sulfamate preparation by ion exchange.

1955

February

Nickel carbonyl coating studies. Eleven uranium pieces coated to one mil thickness.

March

Nickel carbonyl coating studies.

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- April Nickel carbonyl coating studies, experimental data submitted for statistical analysis.
- June Use of ultrasonics in nickel dissolution.
- December Dissolution of nickel coatings from rejected shapes using Enstrip "S". Cost is 1/6 that of HNO<sub>3</sub> treatment and Pu waste loss is 0.6 g per piece treated.

## 1956

- January Removal of nickel coating from reject shapes by using Enstrip "S", including waste losses, time cycle, solvent disposal.
- Absorption spectra of chromic, nickelous, ferrous, ferric, permanganate, calcium, lanthanum, aluminum, and their interaction in nitrate solution investigated.
- Extinction coefficients for Pu in various valence states have been determined.

## 1957

- May Optimum nickel coating removal was accomplished with 6.5 M HNO<sub>3</sub> at 70 C in 15 minutes. Additional effects of acid concentration, time, and temperature are included.
- April A two-gram sample of PuO<sub>2</sub>-UO<sub>2</sub> mixed crystal prepared by H<sub>2</sub> reduction at 900 C of ammonium diuranate - plutonium hydroxide formed by an ammonium hydroxide precipitation from a mixed nitrate solution. Pu/U = 1/5.

## 1958

- March Test of Eastman Adhesive 910 for attaching Plexiglas port rings to Homalite CR-39 plastic.
- New dry chemistry hoods installed in Rooms 185, 186, and 187.
- May Eastman 910 adhesive in use for attaching Plexiglas to Homalite CR-39 (Hood 9).
- August Shipment of PuO<sub>2</sub>, <1400 ppm impurities, to Monsanto Co.
- December Test of radiation dose increase with plutonium concentration for solutions in polyethylene bottles. Effect of lead shields recorded.

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1959

March

Study of equipment and technology for critical mass preparations, PuO<sub>2</sub> suspensions in hydrogeneous materials.

May

Preparation of CeO - paraffin test cylinders, H/Pu - 15, .by atmosphere and pressure casting.

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F. CERAMIC DEVELOPMENT

1955

- April Test of MgO - 5% CaF<sub>2</sub> crucible.  
Slip casting studies.
- June Graphite mold preparations.  
MgO - 5% CaF<sub>2</sub> preparations.
- July Coated graphite molds prepared by spraying of MgO slurry.
- August Preparation of Al<sub>2</sub>O<sub>3</sub>, MgO, zirconia slip cast crucibles.

1956

- November Fabrication study on straight thermocouple wells produced from MgO. Essential dimensions and technique.
- December Flow sheet for improved method of preparing magnesium oxide slip.

1957

- February Nine-inch, coarse-mix, MgO chimneys made without use of hardened dies or other costly equipment. CaF<sub>2</sub> slip casting.
- May Development of routine method of alkaline slip casting of CaF<sub>2</sub> crucibles was demonstrated. Crucibles were smooth, and free of blisters. Effect of calcining temperature on slurry pH was investigated. Low CaO impurities did not affect stability of crucible.
- June Doughnut-shaped setter plates show promise in reducing CaF<sub>2</sub>. Crucible distortion experienced with circular setter plates.
- July CaF<sub>2</sub> (-48 mesh, calcined at 1200 C) required 12-hour ball milling to attain best slip casting properties.  
  
Patterns for ceramic reduction cells are being prepared in support of continuous Task III studies.  
  
A guillotine button cutter has been tested using aluminum buttons 1-1/8" thick. Twenty tons' force is required to cut the button.
- August Work is in progress on a four-compartment vibration-

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cast crucible. Cracks formed during firing at 1760 C. Slip cast shells (from the I-4 pattern) appear to be more promising and will be tested.

The guillotine button cutter was put into use. Cutter was operated by a one-ton press, and worked well until damaged by maloperation. A new model was built, but does not have suitable springs or button holder.

September Acceptable four-compartment I-4 vibration-cast crucibles were produced by eliminating Carbowax from the formulation and substituting magnesium sulfate solution.

Blister-free slip cast  $\text{CaF}_2$  crucibles were achieved by de-airing the slip by means of an evacuated desiccator and lowering the sintering temperature from 1100 C to 1000 C.

Magnesia crucibles were impregnated with molten  $\text{CaF}_2$  at 1400 C. The crucibles thus treated were impervious to water, while ordinary crucibles are not.

October Difficulties in fabricating four-place magnesia casting mold (CS-201) were overcome.

A magnesia pouring crucible was successfully coated with 1/32" nickel.

Glascast was used successfully for slip casting tensile specimen molds.

November Plutonium tensile specimens cast in Glascast molds are fine-grained, superior strength. Surface impurities of B, 4 ppm and Si, 200 indicate promise of Glascast in plutonium casting.

December Studies initiated to form Glascast shell mold for Pit 65.

Glascast crucibles fired at 2300 F developed for skull recovery employing  $\text{CaCl}_2$ - $\text{CaF}_2$  melt at 1000 C.

## 1958

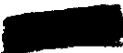
January Problems of removing Glascast Pit 65 casting molds from plaster form overcome.

$\text{CaF}_2$  crucibles fired using green MgO setter plates to prevent cracking.

March MgO crucibles nickel-coated externally for batch and

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continuous Task III.

April

Method for slip cast MgO continuous reduction cell with integral drain tube and riser.

June

July

Tests of impregnating MgO crucibles with salts to reduce porosity.

Initial fabrication of MgO liners for Rokide Z-coated pressure vessel.

Nickel-coated MgO crucible study.

September

A paper, "Slip Casting Of High Purity Calcium Fluoride Crucibles", was prepared.

October

Test of impregnating MgO crucibles with salts to reduce porosity.

1959

May

Test of H-7 pressed sintered high-silica and I-10 MgO-CaF<sub>2</sub> investment-cast pouring crucibles.

June

Results of the H-7 and I-10 crucible tests in pouring applications. H-7 most promising.

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G. PYROCHEMICAL PROCESSING

1. Continuous Task III

1958

March

Preparation of document on continuous electrodeposition of Pu metal--development equipment.

April

Container studies started to determine suitable material for the NaCl-KCl-BaCl<sub>2</sub> melt at 800 C. Zircon, stabilized zirconia, high-density alumina, nickel-coated MgO show promise. Studies used argon and H<sub>2</sub>-HCl atmospheres.

May

June

Initial solubility study for PuCl<sub>3</sub> in NaCl-KCl-BaCl<sub>2</sub> melt.

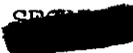
Container study continued using system of CeCl<sub>3</sub>, NaCl-KCl-BaCl<sub>2</sub> melt, and Ce in argon at 800 C. Effect of melt penetration and sticking noted.

Installation of continuous reduction cell started in Hood 31.

July

Thoria, pyroceram, and molybdenum crucible tests in the NaCl-KCl-BaCl<sub>2</sub> melt, argon, H<sub>2</sub>, HCl system at 800 C.

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August

Description of newly-installed continuous Task III equipment.

Tests of crucibles in a mixture of plutonium metal,  $\text{BaCl}_2$ - $\text{KCl}$ - $\text{NaCl}$  melt, and  $\text{PuCl}_3$  in argon at 800 C. Alumina crucible was most promising. Effect on other materials are recorded.

2. Miscellaneous

1958

August

Reduction of  $\text{CeO}$  to  $\text{Ce}$  with calcium in  $\text{CaF}_2$ - $\text{CaCl}_2$  eutectic.

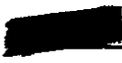
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H. CASTING

1956

January

September

November

1957

January

February

May

June

November

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J. HIGH-EXPOSURE PLUTONIUM

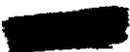
1958

- February Radiation dosage study on high-exposure plutonium. Effect of increase in hard gamma radiation on dosage rate.
- March Documents on capability and program of high-exposure plutonium button line.
- June Design of loading areas, powder trap, calciner-chlorinator combination, and product hood are completed. Chemical flow sheet and flow diagram have been issued.

1959

- February High exposure button line program cancelled. Details in notes of H. W. Crocker.
- March Initial design of laboratory equipment for processing PRTR spike fuel was started.
- May High-exposure plutonium processing summarized in EW-60297, "234-5 Development Processing Facility For PRTR Plutonium".

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## Z. DOCUMENTS ISSUED

### PERIODICAL SEPARATIONS TECHNOLOGY AND RESEARCH AND ENGINEERING REPORTS

#### 1951

January	HW-20161 H
February	HW-20438 H
March	HW-20671 H
April	HW-20991 H
May	HW-21260 H
June	HW-21506 H
July	HW-21802 H
August	HW-22075 H
September	HW-22304 H
October	HW-22610 H
November	HW-22875 H
December	HW-23140 H

#### 1952

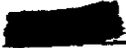
January	HW-23437 H
February	HW-23698 H
March	HW-23982 H
April	HW-24337 H
May	HW-24605 H
June	HW-24928 H
July	HW-25227 H
August	HW-25535 H
September	HW-25781 H
October	HW-26047 H
November	HW-26376 H
December	HW-26720 H

#### 1953

January	HW-27522
February	HW-27739
March	HW-27624 H
April	HW-28355
May - June	HW-28695
July	HW-29035
August	HW-29504
September	HW-29692
October	HW-30158
November	HW-30486
December	HW-30677

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## 1954

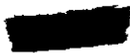
January	HW-31078
February	HW-31336
March	HW-31732
April	HW-31956
May	HW-32070
June	HW-32189
July	HW-32818
August	HW-33150
September	HW-33750
October	HW-34072
November	HW-34161
December	HW-34147 H

## 1955

January	HW-34631 H
February	HW-35530 H
March	HW-35891 H
April	HW-36440 H
May	HW-36928 H
June	HW-37658 H
July	HW-38375 H
August	HW-38828 H
September	HW-39260 H
October	HW-39751 H
November	HW-40182 H
December	HW-40692 H

## 1956

January	HW-41205 H
February	HW-41702 H
March	HW-42219 H
April	HW-42626 H
May	HW-43137 H
June	HW-43938 H
July	HW-44580 H
August	HW-45115 H
September	HW-45707 J
October	HW-46432 J
November	HW-47056 J
December	HW-47675 J

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1957

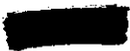
January	HW-48132 J
February	HW-48835 J
March	HW-49503 J
April	HW-50089 J
May	HW-50584 J
June	HW-51211 J
July	HW-51802 J
August	HW-52353 J
September	HW-52864 J
October	HW-53449 J
November	HW-53967 J
December	HW-54319 J

1958

January	HW-54821 J
February	HW-55215 J
March	HW-55571 J
April	HW-55914 J
May	HW-56218 J
June	HW-56602 J
July	HW-56972 J
August	HW-57328 J
September	HW-57640 J
October	HW-58051 J
November	HW-58305 J
December	HW-58711 J

1959

January	HW-59079 J
February	HW-59434 J
March	HW-59849 J
April	HW-60236 J
May	HW-60559 J
June	HW-60915 J
July	HW-61366 J
August	
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November	
December	

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PROGRAM DOCUMENTS

<u>DOCUMENT NUMBER</u>	<u>TITLE</u>	<u>AUTHOR</u>	<u>ISSUE DATE</u>	<u>CLASSIFICATION</u>
HW-18738	Use Of AT Solutions Without Evaporation	JF Facer WL Lyon	9/1/50	Secret
HW-18796	Direct Hydrofluorination Of Plutonium(III) Oxalate	JF Facer WL Lyon	9/11/50	Secret
HW-19122	Recovery Of Plutonium From Skulls	JF Facer WL Lyon	10/12/50	Secret
HW-19771	Gas Composition For The Direct Hydrofluorination Of Pu(III) Oxalate	JF Facer WL Lyon	12/26/50	Secret
HW-19882	Filtration Of Plutonium Peroxide Slurry	WL Lyon	1/5/51	Secret
HW-20164	The Preparation Of Plutonium Tetrafluoride By Direct Hydrofluorination Of Pu(III) Oxalate	JF Facer WL Lyon	2/1/51	Secret
HW-20229	Report Of Invention - Direct Hydrofluorination Of Pu(III) Oxalate or Pu(IV) Oxalate	JF Facer WL Lyon	2/9/51	Secret
HW-20621	Report Of Invention - Oxidation Of Plutonium Metal Using Water At 140 C To Form A Soluble Powder	JF Facer WL Lyon	3/26/51	Secret
HW-20622	Report Of Invention - Dissolution Of Plutonium Metal Using 16 M HNO <sub>3</sub> - 0.04 M HF	JF Facer WL Lyon	3/26/51	Secret
HW-21082	Treatment Of 234-5 Building Wet Chemistry Waste Solutions For R Recycle	RE Falkowski RA Pugh	5/23/51	Secret
HW-21616	Treatment Of 234-5 Building Concentrated Oxalate Waste Solutions For Recycle	RA Pugh	7/6/51	Secret
HW-21691	Laboratory Evaluation Of One And Two Cycle Peroxide Processing Of F-10-P Solutions On The Basis Of Metal Purity Obtained	WL Lyon	7/19/51	Secret

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<u>DOCUMENT NUMBER</u>	<u>TITLE</u>	<u>AUTHOR</u>	<u>ISSUE DATE</u>	<u>CLASSIFICATION</u>
HW-21751	The Use Of Plutonium Peroxide For The Precipitation Of Plutonium Tetrafluoride	WL Lyon B Weidenbaum	7/26/51	Secret
HW-21897	Production Of Plutonium Tetrafluoride To Metal In Charges Containing Calcium And Sulfur	WL Lyon	7/26/51	Secret
HW-22136	The Recovery Of Plutonium From Metal Wastes	JF Facer B Weidenbaum	9/12/51	Secret
HW-22232	Preliminary Memorandum On Processing Of Plutonium Metal Turnings	WL Lyon	9/24/51	Secret
HW-22262	Report Of Invention - The Conversion Of Plutonium Metal Turnings To Massive Metal By A Bomb Fusion Process	WL Lyon	9/26/51	Secret
HW-22394	Report Of Invention - The Dissolution Of Plutonium From Metal Processing Wastes	JF Facer	9/18/51	Secret
HW-23019	Removal Of Hydrogen Fluoride At Low Concentrations In Air Streams By Activated Charcoal And Sodium Fluoride	GE Benedict KM Harmon	12/5/57	Restricted
HW-23132	234-5 Building Program Committee Meeting Minutes	KM Harmon	12/29/51	Secret
HW-24495	Removal Of Hydrogen Fluoride From An Air Stream By Adsorption In A Dorex Charcoal Filter	KM Harmon	5/16/52	Secret
HW-25401	Filter Aids For Isolation Building Plant Use	JF Facer KM Harmon	8/15/52	Secret
HW-25402	Addition Of 70-58 Oxide In Task III	KM Harmon WL Lyon MN Myers	8/15/52	Secret
HW-25468	Further Evaporation Of Residual Solution From The First Cycle Waste Evaporator	GE Benedict	8/28/52	Secret

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<u>DOCUMENT NUMBER</u>	<u>TITLE</u>	<u>AUTHOR</u>	<u>ISSUE DATE</u>	<u>CLASSIFICATION</u>
HW-25562	Processing Of Isolation Building Cleanout Solution	JF Facer KM Harmon	9/16/52	Secret
HW-25771	Capacity Of Marble Chips To Absorb HF Gas	KM Harmon	9/29/52	Unclassified
HW-25779	Basis And Program For Development Work On Plutonium Reduction-Casting	WL Lyon	9/26/52	Secret
HW-25844	Filter Media For Plutonium(IV) Oxalate	JF Facer MN Myers	9/11/52	Secret
HW-25955	Valve Stem Cones For Motor-Operated Throttling Valves To Give Close Remote Control Of Task II Furnace Gases	GE Benedict	10/17/52	Secret
HW-25986	Recovery Of Plutonium From Bismuth Phosphate Decontamination Runs	KM Harmon RA Pugh	10/21/52	Secret
HW-26092	Use of Counter-Current Caustic Scrubber Column For Removal Of HF From Hydrofluorination Off-Gases	GE Benedict	11/3/52	Secret
HW-26758	Observations On Mixing Of Redox And Uranium Recovery Plant Neutralized Aqueous Wastes	KM Harmon RA Pugh	1/6/53	Secret
HW-27488	Laboratory Evaluation Of Double Batch Size Reductions	KM Harmon WL Lyon	3/27/53	Secret
HW-27562	Laboratory Batch Evaporation Of Redox PR Solution	RL Beede JF Facer RA Pugh	4/3/53	Secret
HW-27563	The Precipitation Of Plutonium(IV) Oxalate In Task I	JF Facer KM Harmon	4/3/53	Secret
HW-27837	Procedures For Hydrogen Peroxide Addition In Valence Adjustment For Plutonium(IV) Oxalate Precipitation	JF Facer	7/29/53	Secret
HW-27881	Laboratory Evaluation Of The Use of Sulfur In Reduction	WL Lyon	5/1/53	Secret

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<u>DOCUMENT NUMBER</u>	<u>TITLE</u>	<u>AUTHOR</u>	<u>ISSUE DATE</u>	<u>CLASSIFICATION</u>
HW-28168	The Precipitation Of Plu- tonium(IV) Oxalate In Task I, Nitric Acid Tolerances, Reactor Dimensions	KM Harmon	5/26/53	Secret
HW-28177	Preliminary Memorandum On Combined Reduction And Casting Of Plutonium	WL Lyon	6/1/53	Secret
HW-28357	The Preparation Of Plutonium Alloy Castings In The Reduction Crucible	WL Lyon	10/1/53	Secret
HW-28789	Recovery Of Plutonium From Off-Standard Fluorides	RL Beede KM Harmon	6/29/53	Secret
HW-28838	Report Of Invention - Re- duction Casting Development	WL Lyon	7/27/53	Secret
HW-29200	Plutonium Purification And Fabrication Manual (Chemical Technology Sections)	WL Lyon	4/1/54	Secret
HW-30015	Dissolution Of Slag And Cruci- ble Residues For Plutonium Recovery	RA Pugh WL Lyon	11/1/53	Secret
HW-30036	Shipment Of Laboratory Slag And Crucible Solution	KM Harmon	11/18/53	Secret
HW-30188	Trip Report: Iowa State College	WS Figg	11/20/53	Secret
HW-30558	Laboratory Study Of Causes For High Solubilities Of Plutonium Peroxide	RA Pugh	1/6/54	Secret
HW-31186	Precipitation Pu(IV) Oxalate	JF Facer KM Harmon	3/30/54	Secret
HW-31628	Progress Report: Continuous Task II Program	WS Figg	4/13/54	Secret
HW-32100	Notes Pertaining To Recuplex Product Evaporation	RA Pugh	4/28/54	Secret
HW-33640 RD	Plutonium Requirements For 234-5 Development Program	KM Harmon	11/3/54	Secret - Rough Draft

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<u>DOCUMENT NUMBER</u>	<u>TITLE</u>	<u>AUTHOR</u>	<u>ISSUE DATE</u>	<u>CLASSIFICATION</u>
HW-33666 RD	Supernatant Kill In Task I	KM Harmon	11/4/54	Secret - RD
HW-34385	Recycle Solution Composition - Z Plant To Redox Plant	RA Pugh	1/10/55	Secret
HW-34540	Chemical Reactivity Of Plu- tonium Metal	KM Harmon RC Smith	1/24/55	Secret
HW-35727	Plutonium(IV) Oxalate Pre- cipitation In The Task I Prototype	JF Facer	3/7/55	Secret
HW-35767	F-10-P Solution Studies	GE Benedict MN Myers RA Pugh	3/14/55	Secret
HW-36173	Continuous Task II Program - Plutonium(IV) Oxalate Cal- cination And Hydrofluorination Studies	WS Figg	4/12/55	Secret
HW-37114	Trip Report: Los Alamos Scientific Laboratory	RC Smith	6/7/55	Secret
HW-37115	Trip Report: Los Alamos Scientific Laboratory	RC Smith	6/7/55	Secret
HW-38555	Direct Concentration Of Redox Product Solution	RA Pugh	10/19/55	Secret
HW-38637	Media For Filtration Of Aluminum Nitrate Solution	RL Beede	8/12/55	Secret
HW-38753	Hydrogen Production In The Calcium And Magnesium - Nitric Acid Reactions	MN Myers	8/19/55	Official Use Only
HW-39012	Laboratory Evaluation Of Nelco Calcium For Plutonium Tetrafluoride Reduction	RL Beede	9/15/55	Confidential
HW-39080	Effect Of Ultrasonics On Dissolution Of Nickel In Nitric Acid	RC Smith R Wirta	9/20/55	Unclassified
HW-39328	Preliminary Flowsheet - Continuous Tasks I and II	HH Hopkins, Jr.	10/5/55	Secret

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<u>DOCUMENT NUMBER</u>	<u>TITLE</u>	<u>AUTHOR</u>	<u>ISSUE DATE</u>	<u>CLASSIFICATION</u>
HW-39513	Concentration Of Redox Product With Increased Aluminum Concentration	MN Myers	10/12/55	Secret
HW-39581	Preliminary Coupling Flow-sheet, Redox And Purex, For Task I Replacement	RA Pugh	12/28/55	Secret
HW-39880	Controlled Oxidation Of Plutonium Metal Turnings	RC Smith DR Doman	11/10/55	Secret
HW-40178	Plutonium Content Of Recuplex Slag And Crucible Filter Slurries	MN Myers	11/29/55	Confidential
HW-40274	Separan 2610 As Task I Flocculant	RL Beede	12/5/55	Confidential
HW-40275	Redox Product Coupling To Task I and 231 Building Processes	RL Beede	12/5/55	Confidential
HW-40781	Permanent Mold Casting Program	RC Smith	12/21/55	Secret
HW-40996	Report Of Invention: Use of Fluosilicate For The Precipitation Of Plutonium Fluoride	RL Beede	1/19/56	Secret
HW-41100	Recommendation For Inclusion Of Continuous Task I In Scoping Of Continuous Task II	HH Hopkins, Jr.	2/22/56	Confidential
HW-41286	Recommendations For Study Of Reflux And Modified Plutonium Concentrator Operation	HH Hopkins, Jr.	2/2/56	Secret
HW-41464	Evaluation Of Enstrip "S" For Stripping Nickel	RL Beede CC Wheeler, Jr.	2/17/56	Secret
HW-41635	Technical Specification Letter No. 2 - Continuous Task II - Calcination Reactor Substitution	WS Figg HH Hopkins, Jr.	2/23/56	Secret
HW-42660	Technocal Specification Letter #1 Permanent Mold Casting	HH Hopkins, Jr.	4/25/56	Secret

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<u>DOCUMENT NUMBER</u>	<u>TITLE</u>	<u>AUTHOR</u>	<u>ISSUE DATE</u>	<u>CLASSIFICATION</u>
HW-43953	The Use Of Freon-12 In the Conversion Of Plutonium(IV) Oxalate To Plutonium Tri-fluoride	MN Myers	6/27/56	Confidential
HW-44258	Technical Specification Letter No. 2 Continuous Task I	HH Hopkins, Jr. RL Beede	7/11/56	Confidential
HW-44744	Absorption Spectra Of Plutonium And Impurity Ions In Nitric Acid Solution	MN Myers	7/31/56	Declassified
HW-44987	Reduction Of Plutonium(VI) With Hydrogen Peroxide	MN Myers	8/17/56	Confidential
HW-45700	Continuous Plutonium(IV) Oxalate Precipitation, Filtration, And Calcination Process	RL Beede	9/27/56	Secret
HW-45917	Conversion Of Plutonium Tri-fluoride To Plutonium Tetra-fluoride With Oxygen-Hydrogen Fluoride Mixtures	RL Beede	10/4/56	Secret
HW-46711	Evaporative Kill Of Oxalate Supernatants	CJ Berglund	11/30/56	Confidential
HW-46779	Corrosion Of Stainless Steel, Titanium, And Tantalum In Plutonium-Nitric Acid Solutions	A Brunstad	11/27/56	Unclassified
HW-48914	Continuous Plutonium Reduction--Processes Recommended For Development	MH Curtis	1/16/57	Secret
HW-49121 RD	Plutonium Distribution In Recuplex Systems	WL Lyon	Undated	Secret - Rough Draft
HW-49122 RD	Annual Report - CY 1956, 234-5 Development	HH Hopkins, Jr.	1/1/57	Secret - Rough Draft
HW-49124-RD	Thermometallic Reduction	MH Curtis	10/29/56	Confidential - RD
HW-49125 RD	Possible Continuous Plutonium Reduction Methods	MH Curtis	10/29/56	Confidential - Rough Draft
HW-49126 RD	Thermo-Decomposition Of Plutonium Iodide	MH Curtis	10/19/56	Confidential - Rough Draft

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<u>DOCUMENT NUMBER</u>	<u>Title</u>	<u>AUTHOR</u>	<u>ISSUE DATE</u>	<u>CLASSIFICATION</u>
HW-49138 RD	Research And Development Study TS-R, 234-5 Development	HH Hopkins, Jr.	6/8/56	Secret - Rough Draft
HW-49147	Continuous Plutonium Tri-fluoride Precipitation Progress Report	CJ Berglund	3/18/57	Confidential
HW-48900	Casting Of Plutonium In Copper Molds - Interim Report	RC Smith	1/17/57	Secret
HW-49464 RD	Research And Development Study TS-5, 234-5 Development FY-58	HH Hopkins, Jr.	4/1/57	Confidential - Fough Draft
HW-49699	Laboratory Plutonium Purity	HH Hopkins, Jr.	4/17/57	Secret
HW-49740	Trip Report - Knolls Atomic Power Laboratory, Rocky Flats Plant, Savannah River Plant	HW Crocker MH Curtis	4/19/57	Secret
HW-50280	Continuous Concentration Of Cation Exchange Product	RE Latta HH Hopkins, Jr.	5/20/57	Secret
HW-50597	Coating Removal With Dilute Nitric Acid	A Brunstad	7/31/57	Secret
HW-51276	Trip Report - Rocky Flats Plant, Savannah River Laboratory, Oak Ridge National Laboratory, And Argonne National Laboratory	HH Hopkins, Jr.	7/8/57	Secret
HW-51655	Oxidation Of Plutonium(III) By Sodium Nitrite	A Brunstad	7/19/57	Confidential
HW-52796	Reduction And Stabilization Of Plutonium Nitrate In Cation Exchange Feed And Product Solutions	A Brunstad RC Smith	9/24/57	Confidential
HW-53339	Trip Report, Argonne National Laboratory and Rocky Flats Plant, 9/2/57 - 9/5/57	RC Smith	10/24/57	Secret
HW-54203	Polymerization And Precipitation Of Plutonium(IV) In Nitric Acid	A Brunstad	12/17/57	Unclassified

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<u>DOCUMENT NUMBER</u>	<u>TITLE</u>	<u>AUTHOR</u>	<u>ISSUE DATE</u>	<u>CLASSIFICATION</u>
HW-54405	The Stability Of Anhydrous Plutonium Trifluoride To Saturated Water Vapor At Room Temperature	RE Latta	1/7/58	Confidential
HW-54578	Laboratory Production Of Pure Plutonium	RE Latta	1/15/58	Secret
HW-54610	Annual Report 234-5 Development Operation - CY 1957	HH Hopkins, Jr.	1/20/58	Confidential
HW-54613	R&E Study 234-5 Development - CY-58 (July 1957 - December 1958)	HH Hopkins, Jr.	1/20/58	Confidential
HW-54874	A Plan For Plutonium Trichloride Process Development	HH Hopkins, Jr. by HW Crocker	2/6/58	Secret
HW-55072	Continuous Electrodeposition Of Plutonium Metal Developmental Equipment Considerations	MH Curtis	2/21/58	Confidential
HW-55349	Turbidimetric Microdetermination Of Sulfate In Plutonium Solution	HD Warren A Brunstad	3/14/58	Unclassified
HW-55357	Solubility Of Plutonium(IV) Sulfate In Nitric Acid Solutions	HD Warren A Brunstad	3/14/58	Unclassified
HW-55369	Capability Of Proposed Development Button Line For High-Exposure Plutonium	HH Hopkins, Jr.	3/18/58	Confidential
HW-55434	Program - Development Laboratory High-Exposure Button Line	HH Hopkins, Jr. by Members of 234-5 Development Laboratory	3/21/58	Confidential
HW-55674	Emulsification And Precipitation in the Recuplex Columns	A Brunstad	4/9/58	Confidential
HW-55713	Trip Report - Los Alamos Scientific Laboratory and Rocky Flats Plant, April 7 - 9, 1958	HW Crocker HH Hopkins, Jr.	4/14/58	Secret - Atomic Weapon Data
HW-56606	RD Laboratory Evaluation Of Dibutyl Butylphosphonate	A Brunstad	7/1/58	Confidential - Rough Draft

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<u>DOCUMENT NUMBER</u>	<u>TITLE</u>	<u>AUTHOR</u>	<u>ISSUE DATE</u>	<u>CLASSIFICATION</u>
	And Tetrabromoethane For The Recuplex Solvent System			
HW-57362	Neptunium Oxide Shipment	HH Hopkins, Jr.	9/5/58	Secret
HW-57498	Status Of Plutonium Trichloride Process Development	HH Hopkins, Jr.	9/17/58	Secret
HW-58021	Precipitation Method For The Recovery Of Plutonium From Chloride Slag And Crucible - Progress Report	RW Henkens	10/30/58	Confidential
HW-58095	Recovery Of Plutonium From Chloride Slag And Crucible Solutions	RW Henkens	11/6/58	Confidential
HW-58763	Annual Summary And Program Review, 234-5 Development Operation	HH Hopkins, Jr.	1/6/59	Confidential
HW-59065	234-5 Development Support Of Prototype Button Line - Summary For January	HH Hopkins, Jr.	1/30/59	Confidential
HW-59749	The Analysis Of Exhaust Products In The Chlorination Of Plutonium Oxide With Phosgene	LF Lust	3/24/59	Confidential
Undocumented	Trip Report - Technical Meeting Sessions Of The Metallurgical Society, San Francisco, 2/15 - 19, 1959	RC Smith	3/25/59	Unclassified
HW-59796	Effect Of Titanium In Z-Plant Process	HH Hopkins, Jr. by HW Crocker	3/27/59	Confidential Undocumented
HW-60151	Calcium Iodide - Calcium Chloride System	TS Soine	4/24/59	Unclassified
HW-60153	Trip Report - Ames Laboratory, Rocky Flats Plant, And Los Alamos Scientific Laboratory	HH Hopkins, Jr.	4/24/59	Confidential
HW-60297	234-5 Development Processing Facility For PRTR Plutonium	HH Hopkins, Jr. by HW Crocker	5/4/59	Confidential
HW-60353	Trip Report - Rocky Flats	RS Rosenfels	5/13/59	Secret

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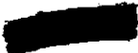
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	Plant And Los Alamos Scientific Laboratory			
HW-60492	234-5 Development Processing Facility For Critical Mass Fuel Preparation	HH Hopkins, Jr. by HW Crocker	5/27/69	Unclassified
HW-61564 RD	Batch Reduction Of Plutonium Trichloride	TS Soine	8/14/59	Secret - Rough Draft
HW-61565 RD	The Continuous Chlorination Of Plutonium Dioxide	MJ Rasmussen	8/14/59	Confidential - Rough Draft
HW-61570	Status Of Plutonium Trichloride Process Development (II)	HH Hopkins, Jr.	8/14/59	Secret
HW-61577	Recovery Of Plutonium From Chloride Slag And Crucible	RC Smith	8/14/59	Confidential

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