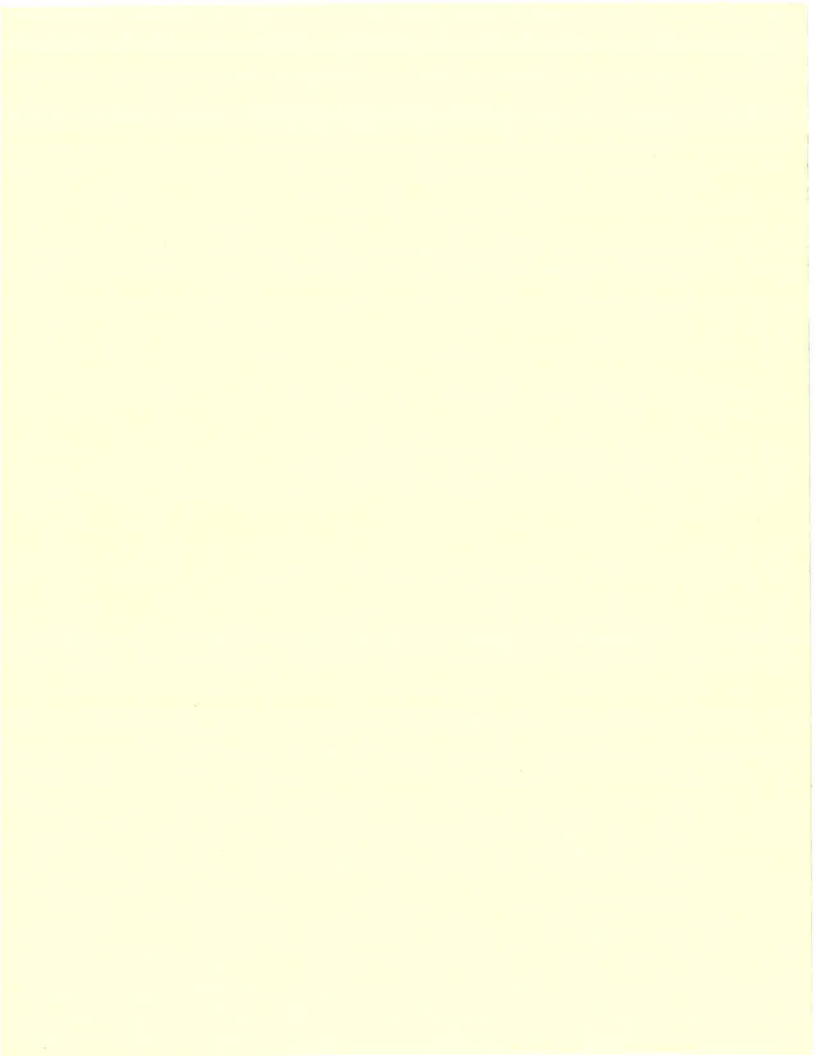


Bureau of Mines Report of Investigations/1986

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By J. C. Judd, R. G. Sandberg, and J. L. Huiatt





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UNITED STATES DEPARTMENT OF THE INTERIOR Donald Paul Hodel, Secretary

**BUREAU OF MINES**Robert C. Horton, Director

# Library of Congress Cataloging in Publication Data:

# Judd, J. C

Recovery of vanadium, uranium, and phosphate from Idaho phosphorite ores.

(Report of investigations; 9025)

Bibliography: p. 14-15.

Supt. of Docs. no.: I 28.23: 9025.

1. Vanadium-Metallurgy. 2. Uranium-Metallurgy. 3. Phosphates. 4. Phosphate rock-Idaho. I. Sandberg, R. G. (Richard G.); II. Huiatt, J. L. III. Title. IV. Series: Report of investigations (United States. Bureau of Mines); 9025.

TN23.U43 [TN799.V3] 622s [669'.732] 85-600353

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# UNIT OF MEASURE ABBREVIATIONS USED IN THIS REPORT

°C	degree Celsius	mV	millivolt
ft	foot	pct	percent
g	gram	ppm	part per million
g/L	gram per liter	psi	pound per square inch
h	hour	rpm	revolution per minute
L	liter	st	short ton
lb/st	pound per short ton	st/yr	short ton per year
min	minute	vol pct	volume percent
<u>M</u>	molar	wt pct	weight percent
mL	milliliter		

# RECOVERY OF VANADIUM, URANIUM, AND PHOSPHATE FROM IDAHO PHOSPHORITE ORES

By J. C. Judd, <sup>1</sup> R. G. Sandberg, <sup>2</sup> and J. L. Huiatt <sup>3</sup>

#### **ABSTRACT**

The Bureau of Mines investigated the recovery of vanadium, uranium, and phosphorus from low-grade phosphorite ores and concentrate by leaching with  $\rm H_2SO_4$ . Samples containing 0.12 to 0.23 pct  $\rm V_2O_5$ , 0.009 to 0.012 pct  $\rm U_3O_8$ , and 18 to 30 pct  $\rm P_2O_5$  were treated by batch-agitation leaching, roasting followed by acid leaching, salt-roasting followed by acid leaching, and leaching under pressure.

Maximum recoveries using leaching alone ranged from 78 to 98 pct of the  $V_2O_5$  and 74 to 88 pct of the  $U_3O_8$ . Leaching under pressure resulted in extractions of 88 to 95 pct of the  $V_2O_5$  and 84 to 89 pct of the  $U_3O_8$ . Leaching following roasting—with or without NaCl—produced extractions of 97 to 99 pct of the  $V_2O_5$  and 86 to 91 pct of the  $U_3O_8$ . Depending on the technique(s) employed, phosphate extractions ranged from 89 to 99 pct.

Solvent extraction laboratory tests were conducted to selectively recover vanadium and uranium values from acid leach solutions. An average of 95 pct of the  $\rm U_3O_8$  was recovered using di(2-ethylhexyl)phosphoric acid, plus trioctylphosphinic oxide. Essentially all of the  $\rm V_2O_5$  was recovered with octylphenyl acid phosphate plus isodecanol. The phosphate remained in the raffinate.

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#### INTRODUCTION

Vanadium is classified as a strategic and critical material because of its significant import dependence and essential use in equipment for defense, energy, and transportation (1).4 It is principally consumed by the iron and steel industry ferrovanadium or related vanadiumcarbon ferroalloys. Vanadium cantly increases strength and improves toughness and ductility of carbon steels, which are used for constructing high-rise buildings, bridges, and pipelines. important applications for vanadium are as additives to titanium-based alloys, which are consumed in jet engines other aircraft parts, and as catalysts, which are used in the manufacturing of sulfuric acid and other chemicals.

Vanadium is widely distributed throughout the earth but in rather low concentration. In the United States, particularly in the West, vanadium is found in uranium-bearing minerals and in certain phosphatic shales and phosphate rocks. One of these phosphatic deposits, located in Idaho, represents a large domestic vanadium reserve. The phosphorite beds, generally 3 to 20 ft thick, average 20 to 32 pct  $P_2O_5$  and 0.14 pct  $V_2O_5$ . At least 125,000 short tons (st) of coproduct vanadium is contained in these deposits. An additional 1.6 million st of coproduct vanadium occurs in marginal reserves of phosphatic shale and low-grade phosphorite dispersed in the northern Rocky Mountains (1).

The western phosphate industry processes 2.6 million st/yr ore (2). Following beneficiation, the plus 325-mesh concentrate (30 to 32 pct  $P_2O_5$ ) is leached with  $H_2SO_4$ , and the minus 325-mesh material is rejected to the tailings. A significant amount of vanadium and uranium is lost with the rejected tailings. Phosphoric acid  $(H_3PO_4)$  is produced from concentrates by either the dihydrate process (at temperatures of  $<80^{\circ}$  C) or the hemihydrate process (at

>80° C). The  $P_2O_5$  content of the acid is 28 to 30 pct with the dihydrate process and 45 to 54 pct with the hemihydrate process. If necessary, evaporation is used to increase the  $P_2O_5$  content to 50 to 54 pct, regardless of which process is used (3-4). The chemical equation for the dihydrate (A) and hemihydrate (B) reactions are given below (4-6):

$$3Ca_{3}(PO_{4})2 \cdot CaF_{2} + 10H_{2}SO_{4} + 20H_{2}O$$

$$\iff 6H_{3}PO_{4} + 2HF + 10(CaSO_{4} \cdot 2H_{2}O) \qquad (A)$$

$$3Ca_{3}(PO_{4})2 \cdot CaF_{2} + 10H_{2}SO_{4} + 5H_{2}O$$

$$\iff 6H_{3}PO_{4} + 2HF + 10(CaSO_{4} \cdot 1/2H_{2}O) \qquad (B)$$

The approximate amount of H<sub>2</sub>SO<sub>4</sub> used for maximum conversion to H<sub>3</sub>PO<sub>4</sub> is the stoichiometric amount required to convert the CaO content of the concentrate to  $CaSO_4$ , plus an additional 2 to 3 pct. Acid concentration is very critical because, if the free sulfate content is too low, the gypsum (CaSO<sub>4</sub> •2H<sub>2</sub>0) crystals do not grow properly; this affects the rate of filtration (6). Also, the HPO4 ions are substituted for SO<sub>4</sub> ions in the CaSO<sub>4</sub> lattice; this represents a significant P205 loss. The extent of substitution decreases with H<sub>2</sub>SO<sub>4</sub> excess and increases with increased P<sub>2</sub>O<sub>5</sub> concentration. stitution of P2O5 in gypsum is also related to solids content in suspension. If the  $H_2SO_4$  concentration is too high, an insoluble CaSO4 layer forms around the particles to be digested (6).

Acid processes for recovery of uranium and vanadium from the usual western resources are similar to those processes used in the production of  $\rm H_3PO_4$ , except the feed generally has not been beneficiated prior to  $\rm U_3O_8$  and  $\rm V_2O_5$  recovery. One benefit of processing unbeneficiated phosphorite ore is the elimination of one solid waste stream and the significant problems in its handling (7-8). A second benefit is higher recoveries, as value losses during beneficiation are generally significant. A third benefit is conservation of the resource, as more

<sup>&</sup>lt;sup>4</sup>Underlined numbers in parentheses refer to items in the list of references at the end of this report.

phosphate should be available per short ton of phosphorite ore mined. Therefore, processing of phosphorite to recover vanadium and uranium requires that phosphorus also be recovered, in order to offset the costs of additional  $\rm H_2SO_4$  consumption.

Because of the strategic importance of vanadium the Bureau of Mines conducted an investigation to recover vanadium, uranium, and phosphorus from unbeneficiated Conda and Fort Hall (Gay Mine) ores, a Conda high-magnesium material, and a Conda concentrate (Simplot Mill). Extraction techniques investigated were batchagitation leaching alone; roasting or

salt-roasting followed by leaching; and under pressure. leaching Tests were performed to determine acid addition, temperature, solids content, and time requirements for optimum extractions. Complementary solvent extraction tests were conducted to selectively extract the vanadium and uranium from the leach solutions. Solvent extraction was selected because it has been used successfully in the past to recover uranium (4) and vanadium (9) with trioctylphosphinic oxide (TOPO) and to recover both vanadium and uranium (10) with trialkylphosphine oxide plus an alkyl-substituted diaryl phosphoric acid.

#### SAMPLE DESCRIPTION AND PROCEDURES

#### SAMPLE DESCRIPTION

The four samples used in this study were designated as Conda, Conda (high-Mg), and Fort Hall ores, and Conda concentrate. Conda and Fort Hall ore samples were from ore deposits that are presently being processed to produce phosphate fertilizer. The Conda concentrate, as received, was a deslimed, calcined product. Ninety-seven percent of this material was minus 35 mesh. Conda (high-Mg) ore sample represents material that is presently being rejected because its high magnesium content causes processing problems. Chemical and inductively coupled plasma (ICP) analyses of the four samples are listed in tables 1 and 2, respectively.

Microscopic and X-ray diffraction examination identified the samples as calcium phosphate shale or phosphorite. X-ray diffraction also revealed that (1) Conda and Fort Hall samples contained illite clay, quartz, clinozoisite-epidote (a mineral with composition intermediate to these two minerals), limonite, calcite, plagioclase, pyrite, and apatite, (2) the Conda (high-Mg) sample contained dolomite, apatite, quartz, calcite, plagioclase. Neither fluorescence a Geiger counter could detect any uranium in the four samples, nor could specific vanadium or uranium minerals be identified.

Ore samples were prepared for leaching by pulverizing to minus 35 mesh and

TABLE 1. - Chemical analyses of phosphorite samples, percent

	Phosphorite samples						
Material	Conda   Fort Hall		Conda	Conda			
			(high-Mg)	concentrate			
V <sub>2</sub> O <sub>5</sub>	0.164	0.227	0.105	0.12			
U <sub>3</sub> O <sub>8</sub>	.009	.010	•007	.012			
P <sub>2</sub> O <sub>5</sub>	24.9	24.7	18.8	30			
Fe <sub>2</sub> 0 <sub>3</sub>	1.28	1.24	1.27	•66			
MgÖ	2.51	1.08	5.58	.61			
CaO	35.8	35.4	33.6	44.7			
SiO <sub>2</sub>	17.4	19.8	14.7	12.2			
Organic carbon	2.2	3.4	.7	•05			

TABLE 2 Analy	ses of o	re and	concent	trate samples
by inductively	coupled	plasma	(ICP)	spectrometry
percent				

Element	Conda	Fort Hall	Conda	Conda
			(high-Mg)	concentrate
A1	1.7	1.3	0.93	0.39
Ba	.01	.014	.01	.007
Cd	.01	.014	<.01	.002
Co	.001	.002	<.001	.003
Cr	.08	.09	.06	.033
Cu	.008	.009	.007	.009
Mn	.009	.01	•01	•01
Мо	.008	.013	<.01	•006
Ni	.025	.018	.01	.012
Pb	.001	•005	<.01	.001
Th	.03	.03	<.05	.06
Ti	.11	•11	•054	.169
Zn	.15	.19	.10	.066

TABLE 3. - Wet-screen analyses of Conda ore

Mesh size	Wt pct	Ana	lysis, p	ct	Distribution, pct		
		V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> 0 <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>	V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> O <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>
Minus 35 plus 48	7.9	0.093	0.011	32.5	4.9	9.5	10.8
Minus 48 plus 65	10.8	.095	.011	29.4	6.9	13	13.5
Minus 65 plus 100	12.1	.090	.012	32	7.4	15.4	16.4
Minus 100 plus 150	7.7	.080	.012	32	4.2	9.5	10.4
Minus 150 plus 200	7.7	.075	.013	31.6	3.9	10.3	10.2
Minus 200	53.8	•20	.007	17	72.7	42.3	38.7
Calculated head	100	.148	.0094	23.7	100	100	100
Assay head	NAp	.164	.0091	24.9	NAp	NAp	NAp
NAp Not applicable.							

splitting into 100-g samples. Concentrate samples were screened through minus 35 mesh; the oversize was pulverized to minus 35 mesh. The screen fractions were recombined and split into 100-g samples.

A wet-screen analysis was performed to determine the distribution of vanadium, uranium, and phosphate in the screen fractions from each of the samples. Results of the wet-screen analysis are listed in tables 3-6. The results show 59 to 73 pct of the  $V_2O_5$ , 27 to 42 pct of the  $U_3O_8$ , and 20 to 39 pct of the  $P_2O_5$  in the ores were distributed to the fines (minus 200-mesh fraction). The Conda concentrate was not considered, as it had already been beneficiated and calcined. The analyses indicate that significant amounts of vanadium, uranium,

and phosphate could be lost if the fines are removed during beneficiation.

The Conda concentrate contains less vanadium but more phosphate than the Conda and Fort Hall ore samples. Vanadium and phosphate content are lowest in the Conda (high-Mg) ore. The magnesium content of the Conda (high-Mg) ore is twice the amount in the Conda ore, and five times the amount in the Fort Hall ore.

#### EXPERIMENTAL PROCEDURES

Leaching procedures used in this study were as follows: batch-agitation leaching; roasting, with and without salt addition, followed by leaching; and leaching under pressure. The lixiviant in all

TABLE 4. - Wet-screen analyses of Fort Hall ore

Mesh size	Wt pct	Ana	lysis, p	ct	Distribution, pct			
		V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> 0 <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>	V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> 0 <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>	
Minus 35 plus 48	13.6	0.17	0.012	28.3	10.1	15.8	16.5	
Minus 48 plus 65	15.5	.16	.011	29.2	10.8	16.5	19.4	
Minus 65 plus 100	16.5	.14	.011	30.5	10.1	18.9	21.4	
Minus 100 plus 150	9.1	.13	.012	31.8	5.2	10.9	12.4	
Minus 150 plus 200	8	.14	.012	29.7	4.9	10	10.2	
Minus 200	37.3	.36	.007	12.6	58.9	27.9	20.1	
Calculated head	100	.228	.0099	23.4	100	100	100	
Assay head	NAp	.227	.010	24.7	NAp	NAp	NAp	

NAp Not applicable.

TABLE 5. - Wet-screen analyses of Conda (high-Mg) material

Mesh size	Wt pct	Analysis, pct			Distribution, pct		
		V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> O <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>	V <sub>2</sub> O <sub>5</sub>	U308	P <sub>2</sub> O <sub>5</sub>
Minus 35 plus 48	17.2	0.07	0.008	22.4	11.7	18.7	20.8
Minus 48 plus 65	15.4	.07	.009	23	11.1	21	19.1
Minus 65 plus 100	12.6	.07	.010	22.3	9.3	17.6	15.1
Minus 100 plus 150	6.4	•06	.008	23.4	4.2	7.9	8.1
Minus 150 plus 200	7.1	•06	.007	23.7	4	7.4	9
Minus 200	41.3	.14	•005	12.5	59.7	27.4	27.9
Calculated head	100	.097	.007	18.5	100	100	100
Assay head	NAp	.105	.007	18.8	NAp	NAp	NAp

NAp Not applicable.

TABLE 6. - Wet-screen analyses of Conda concentrate

Mesh size	Wt pct	Ana	lysis, p	oct	Distr	ibution	, pct
		V <sub>2</sub> 0 <sub>5</sub>	U <sub>3</sub> O <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>	V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> 0 <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>
Minus 35 plus 48	4	0.07	0.012	32.3	2.6	4.0	4.1
Minus 48 plus 65	6.1	.08	.012	35	4.5	6	6.6
Minus 65 plus 100	10.4	.08	.012	35.5	8.1	10.6	11.5
Minus 100 plus 150	9.7	.08	.013	34.1	7.7	10.2	10.4
Minus 150 plus 200	15.6	.09	.013	34	12.7	16.2	16.5
Minus 200 plus 325	17.5	.09	.012	32.6	15.4	17.5	17.8
Minus 325	36.7	.14	.012	28.8	49	35.5	33.1
Calculated head	100	.11	.012	32	100	100	100
Assay head	NAp	.12	.012	30	NAp	NAp	NAp

NAp Not applicable.

tests was reagent-grade sulfuric acid  $(H_2SO_4)$ .

Batch-agitated leach tests were performed using 100 g of minus 35-mesh samples. The samples were leached for 3 to 24 h in a 1,000-mL tall beaker equipped with a reflux tube and stopper. Agitation was accomplished using a custom-fabricated impeller rotated at sufficient speed to maintain particle suspension.

A hotplate equipped with a temperature probe was used to keep leach temperatures within  $\pm 1^{\circ}$  C. After leaching, the slurry was filtered through a Buchner funnel and the residue washed three times using about 50 mL of wash solution per wash. The first wash solution was a dilute solution of  $\rm H_2SO_4$  (pH adjusted to 1.5). Deionized water was used for the second and third washes.

Roast-leach tests were performed by preroasting the ore sample, mixed with or without NaCl, in a muffle furnace. The samples were heated at a specific temperature for 4 h, removed from the furnace, cooled to ambient temperature, and leached for 18 h at 75° C using H<sub>2</sub>SO<sub>4</sub> solution.

Pressure-leach tests were performed on 50-g charges of minus 35-mesh material using a glass-lined pressure reaction apparatus equipped with stirrer and temperature probe. The reactor was purged for 5 min with oxygen before heating to the operating temperature. At the end of

testing, the reactor was quenched in cold water. The slurry was filtered through a Buchner funnel and washed using a procedure similar to that described for the leach-only tests, except 25 mL of wash solution was used per wash.

The pH and emf values were measured on the terminal leach solution (leach filtrate before wash water was added). The emf values were determined with platinum and saturated-calomel electrodes; the highest values indicate the most highly oxidized solution. Leaching efficiency was assessed by determining and comparing  $V_2O_5$ ,  $U_3O_8$ , and  $P_2O_5$  extractions.

#### LEACH TESTS

#### BATCH-AGITATION LEACHING

Batch-agitation leach tests were performed using various acid concentrations, solids contents, temperatures, and leaching times. Good agitation with a well-designed impeller was important to obtain reproducible results. Improper mixing during leaching caused temperature gradients, which decreased mass transfer rates between ore particles and leach solutions.

# Conda Ore

The best extractions using Conda ore were 97, 87, and 99 pct of the  $V_2O_5$ ,  $U_3O_8$ , and  $P_2O_5$ , respectively. These results were achieved by leaching the ore at 40 pct solids for 18 h at 90° C using 2,000 lb/st  $H_2SO_4$  (table 7). Most wetprocess phosphoric acid plants operate at a temperature between 75° and 85° C; a range which is most practical for leaching the phosphorite ores (3). Leaching

TABLE 7. - Effect of temperature on acid leaching of Conda and Fort Hall ore samples 1

TI CO	C-1445	Tomo	Tr-		Ent.	ation			
$H_2SO_4$ ,	Solids,	Temp,		rminal		ction,			
lb/st	pct	°C	pН	emf, mV	V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> 0 <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>		
CONDA ORE									
1,500	40	65	1.1	330	84	86	98		
•		75	1.3	370	86	88	97		
		85	1.1	320	<b>9</b> 0	87	92		
		90	1.1	300	92	87	93		
	50	65	1.2	340	86	88	99		
		75	1.1	360	90	87	98		
		85	1.2	330	92	87	99		
	,	90	1.3	323	89	87	95		
2,000	40	65	.4	360	85	86	94		
		75	.5	345	94	88	98		
		85	-4	370	95	87	97		
	ļ	90	.5	360	97	87	99		
		FORT	HALL	ORE		-			
1,500	40	65	0.7	320	74	77	92		
-		75	.9	340	78	74	89		
		85	.7	290	84	78	96		
		90	.9	350	89	89	98		
1-							• •		

<sup>1</sup>Conditions: 100 g of minus 35-mesh samples leached for 18 h.

Conda ore at 75° C for 18 h at 40 pct solids using 2,000 lb/st  $\rm H_2SO_4$  extracted 94, 88, and 98 pct of the  $\rm V_2O_5$ ,  $\rm U_3O_8$ , and  $\rm P_2O_5$ , respectively (table 8). Increasing the temperature to 85° C had little effect on extraction.

Experiments performed at 75° C and 18-h leaching time (table 8) showed that increasing pulp density from 33-1/3 pct solids to 40 pct solids significantly improved  $\rm V_2O_5$  and  $\rm P_2O_5$  extraction from Conda ore for a given acid concentration, but had little effect on  $\rm U_3O_8$  extraction. Increasing pulp density to 50 pct solids had no significant effect on extraction.

# Fort Hall Ore

The best extractions using Fort Hall ore were 89, 89, and 98 pct of the  $V_2O_5$ ,  $U_3O_8$ , and  $P_2O_5$ , respectively. These results were achieved by leaching the ore at 40 pct solids for 18 h at 90° C using 1,500 lb/st  $H_2SO_4$  (table 7). Higher temperatures and longer leaching times generally improved extraction from this ore (tables 7 and 9). However, leaching at 85° C lowered  $V_2O_5$ ,  $U_3O_8$ , and  $P_2O_5$ extractions to 84, 78, and 96 pct, respectively, and leaching at 75° C produced still lower results of 78, 74, and 89 pct  $V_2O_5$ ,  $U_3O_8$ , and  $P_2O_5$ , respectively. Increasing the acid addition (table 10) from 1,500 to 2,500 lb/st  $H_2SO_4$  (at 75° C) increased phosphate extraction from 89 to 93 pct but in turn reduced  $U_3O_8$  extraction from 74 to 71 pct.

In general, temperature and time had greater effects on extraction than did acid addition. (See tables 7 and 9.) The ore was more difficult to agitate at  $2,000~1b/st~H_2SO_4$  than at either  $1,500~or~2,500~1b/st~H_2SO_4$ , which may account for the lower extractions at  $2,000~1b/st~H_2SO_4$  (table 10). The Fort Hall ore could not be agitated at 50 pct solids.

TABLE 8. - Effect of acid concentration and solids content on acid leaching of Conda ore samples 1

H <sub>2</sub> SO <sub>4</sub> ,	Te	rminal	Extra	ction,	pct					
lb/st	pН	emf, mV	V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> O <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>					
33-1/3 pct SOLIDS										
1,000	1.7	350	63	62	71					
1,500	1.4	330	79	85	84					
2,000	•6	350	89	86	92					
2,500	•5	350	81	72	82					
3,000	.2	370	91	86	96					
	4	0 pct SOL	IDS							
1,000	1.6	330	65	63	70					
1,500	1.3	370	86	88	97					
2,000	•5	350	94	88	98					
2,500	.2	370	90	75	98					
3,000	.0	380	90	52	90					
	5	0 pct SOI	IDS							
1,000	2.0	310	62	64	76					
1,500	1.1	360	90	87	98					
2,000	.7	360	91	88	97					
2,500	.3	360	92	75	99					

<sup>1</sup>Conditions: 100 g of minus 35-mesh sample leached at 75° C for 18 h.

TABLE 9. - Effect of time on acid leaching of Conda and Fort Hall ore samples 1

Solids,	Duration,	Те	rminal	Extra	pct	
pct	h	pH emf, mV V <sub>2</sub> O <sub>5</sub> U <sub>3</sub>		U <sub>3</sub> 0 <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>	
		COND	A ORE			
50	3	0.7	300	80	88	96
	6	1	340	80	87	97
	18	1.1	360	<b>9</b> 0	87	98
	24	1.1	360	89	87	99
	F	ORT H	ALL ORE			
40	3	0.8	300	68	79	95
	6	.8	330	73	86	90
	18	.8	340	78	74	89
	24	.8	360	84	87	96

<sup>1</sup>Conditions: 100 g of minus 35-mesh sample leached at 75° C using 1,500 lb/st H<sub>2</sub>O<sub>4</sub>.

TABLE 10. - Acid leaching of phosphorite ores and concentrate<sup>1</sup>

H <sub>2</sub> SO <sub>4</sub> ,	Te	rminal	Extra	ction,	pct			
1b/st	pН	emf, mV	V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> O <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>			
		CONDA OR	.E					
1,500	1.3	370	86	88	97			
2,000	•5	350	94	88	98			
2,500	.2	370	90	75	98			
	F	ORT HALL	ORE					
1,000	1.2	350	58	64	81			
1,500	.9	340	78	74	89			
2,000	.4	333	73	66	80			
2,500	.2	340	78	71	93			
CC	NDA (	HIGH-Mg)	MATERI	AL				
1,500	0.9	320	86	84	93			
2,000	•6	335	91	85	96			
2,500	0	370	95	85	98			
	CONDA CONCENTRATE							
1,500	1.3	165	89	76	90			
2,000	•5	260	98	78	99			
2,500	.3	360	98	23	99			
Conditi	one:	100 g o	f min	35	-mach			

Conditions: 100 g of minus 35-mesh sample leached at 75° C for 18 h at 40 pct solids.

# Conda (High-Mg) Material

The best extractions from this material were 95, 85, and 98 pct of the  $V_2O_5$ ,  $U_3O_8$ , and  $P_2O_5$ , respectively (table 10). These results were achieved by leaching the material at 40 pct solids for 18 h at 75° C using 2,500 lb/st  $H_2SO_4$ .

# Conda Concentrate

The best extractions from the Conda concentrate were 98, 78, and 99 pct of the  $V_2O_5$ ,  $U_3O_8$ , and  $P_2O_5$ , respectively (table 10). These results were achieved by leaching the concentrate at 40 pct solids for 18 h at 75° C using 2,000 lb/st  $H_2SO_4$ . This should be similar to the actual mill result.

#### ROAST AND SALT-ROAST LEACHING

Because roast-leach methods have been effective in treating uranium ore in which the uranium could not be extracted by leaching alone, and because salt-roasting has been a viable process used in vanadium extraction (11), tests were performed to determine if vanadium and uranium extractions could be improved by these methods.

In roast-leaching, the samples were roasted for 4 h at 500° or 800° C; in salt-roasting, the samples were first combined with 10 pct NaCl, then roasted for 4 h at 800° C only. After roasting (or salt-roasting), the calcines were leached at 75° C for 18 h with additions of 1,000 to 2,500 lb/st H<sub>2</sub>SO<sub>4</sub>. Calcines from the Conda and Fort Hall ores were leached at 40 and 50 pct solids; calcines from the Conda (high-Mg) ore and Conda concentrate were leached at 40 pct solids.

#### Conda Ore

The following tabulation (from tables 7 and 11) shows that 500 lb/st less acid produced almost identical extraction levels following roasting or salt-roasting than from leaching alone.

	Leaching alone	Rosst- leach	Salt-roast and leach
Conditions:			
Roast temperature°C	NAp	800	800
Leach temperature°C	90	75	75
Acid additionlb/st	2,000	1,500	1,500
Solids contentpct	40	50	40
Extraction, pct:			
V <sub>2</sub> O <sub>5</sub>	97	99	97
U <sub>3</sub> O <sub>6</sub>	87	88	89
P <sub>2</sub> O <sub>3</sub>	99	98	99

NAp Not applicable.

TABLE 11. - Roast leaching and salt-roast leaching of Conda ore 1

Solids,	Roast	H <sub>2</sub> SO <sub>4</sub> ,	Te	rminal	Extra	ction,	pct
pct	temp, °C	lb/st	pН	pH emf, mV		U <sub>3</sub> O <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>
		ROA	ST LE	ACH			
40	500	1,000	1.9	360	83	75	70
	800	1,000	1.5	340	80	77	61
	500	1,500	1	360	89	88	94
	800	1,500	1.1	400	97	88	98
	500	2,000	.3	360	94	89	96
	800	2,000	• 4	352	97	88	99
50	800	1,000	1.8	305	80	78	60
	500	1,500	1.1	270	94	87	82
	800	1,500	.9	290	99	88	98
	800	2,000	0	391	93	77	85
	SALT-ROA	ST LEACH	USIN		ct NaC	1	
40	800	1,000	1.7	690	83	78	62
	800	1,500	1.1	700	97	89	99
	800	2,000	0	800	91	87	94
50	800	1,000	1.5	696	66	77	63
	800	1,500	.8	760	97	88	96
	800	2,000	0	773	95	75	96

<sup>1</sup>Conditions: Charges roasted for 4 h and calcines leached at 75° C for 18 h.

# Fort Hall Ore

Roasting with or without salt increased  $V_2O_5$  extractions (tables 7 and 12), as shown in the following tabulation:

	Leaching	Roast-	leach	Salt-roast		
	alone	(2)	(3)	and leach		
	(1)			(4)	(5)	
Conditions:						
Roast temperature°C	NAp	500	800	800	800	
Leach temperature°C	90	75	75	75	75	
Acid additionlb/st	1,500	1,500	1,500	1,500	2,000	
Solids contentpct	40	50	40	40	50	
Extraction, pct:	[					
V205	89	97	96	98	99	
U 3 0 g	89	88	91	90	78	
P205	98	97	99	99	99	

As can be seen, salt-roasting produced slightly higher  $V_2O_5$  values, but surprisingly, leaching the salt-roasted calcine lowered the  $U_3O_8$  extraction considerably at the higher acid and solids content. (See columns 4 and 5 above.) Values are nearly as high after roasting without

salt at only  $500^{\circ}$  C and 50 pct solids. Note that roasting enabled leaching the ore at 50 pct solids.

# Conda (High-Mg) Material

Nearly identical results (tables 10 and 13) were achieved following roasting and salt-roasting at  $800^{\circ}$  C as with leaching alone, but with lower acid additions, as shown in the following tabulation:

	Leaching	Roast-	Salt-roast
	alone	leach	and leach
Conditions:			
Roast temperature°C	NAp	800	800
Leach temperature°C	75	75	75
Acid additionlb/st	2,500	1,500	2,000
Solids contentpct	40	40	40
Extraction, pct:			
V <sub>2</sub> O <sub>5</sub>	95	97	96
U <sub>3</sub> O <sub>8</sub>	85	84	85
P205	98	97	98

TABLE 12.	- Roast leachi	ing and salt-roast	leaching
of Fort	Hall ore <sup>1</sup>		

Solids,	Roast	H <sub>2</sub> SO <sub>4</sub> ,	Te	rminal	Extra	ction,	pct
pct	temp, °C	lb/st	pH emf, mV		V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> O <sub>8</sub>	P <sub>2</sub> O <sub>5</sub>
	<u> </u>	ROA	ST LE	ACH			
40	500	1,000	1.4	350	85	89	83
	800	1,000	1.3	500	94	80	82
	500	1,500	.7	360	88	89	92
	800	1,500	.9	460	96	91	99
	500	2,000	.9	380	96	88	99
	800	2,000	.4	470	95	90	98
50	800	1,000	1.5	485	93	81	86
	500	1,500	.7	310	97	88	97
	800	1,500	.5	640	97	88	99
	500	2,000	.9	380	96	89	99
	800	2,000	.2	410	94	67	70
	SALT-ROA	ST LEACH	USIN	G 10 wt p	ct NaC	1	
40	800	1,000	1.4	665	92	79	80
	800	1,500	.8	700	98	90	99
	800	2,000	.3	760	97	89	99
50	800	1,000	1.2	770	88	81	81
	800	1,500	•5	730	98	87	97
	800	2,000	.1	780	99	78	99
	SALT-ROA 800 800 800 800 800	T LEACH 1,000 1,500 2,000 1,000 1,500	USIN 1.4 .8 .3 1.2 .5	G 10 wt p 665 700 760 770 730 780	ct NaC 92 98 97 88 98	79 90 89 81 87 78	80 99 99 81

<sup>1</sup>Conditions: Charges roasted for 4 h and calcines leached at 75° C for 18 h.

Salt-roasting and leaching with 1,500 lb/st  $H_2SO_4$  produced even higher  $V_2O_5$  extraction (of 98 pct), but  $P_2O_5$  extractions were much lower (78 pct) (table 13).

# Conda Concentrate

with regard to lowering acid requirements, as shown in tables 10 and 13.

	Leaching	Roast-	Salt-roast	
	alone	leach	and leach	
Conditions:				
Roast temperature°C	NAp	800	800	
Leach temperature°C	75	75	75	
Acid additionlb/st	2,000	2,000	2,000	
Solids contentpct	40	40	40	
Extraction, pct:				
V <sub>2</sub> 0 <sub>5</sub>	98	98	98	
ปรู้ 0คู่	78	89	89	
P205	99	98	97	

TABLE 13. - Roast leaching and salt-roast leaching of Conda (high-Mg) material and Conda concentrate1

NaCl in roast,	H <sub>2</sub> SO <sub>4</sub> ,	Te	rminal	Extra	pct	
wt pct	1b/st	pН	emf, mV	V <sub>2</sub> 0 <sub>5</sub>	U <sub>3</sub> O <sub>8</sub>	P <sub>2</sub> 0 <sub>5</sub>
	CONDA (H	IGH-M	g) MATERI	AL		
0	1,500	1.2	440	97	84	97
	2,000	.4	390	95	86	94
	2,500	0	410	94	72	85
10	1,000	1.5	618	61	65	39
	1,500	1	700	98	87	78
	2,000	.1	760	96	85	98
	CONDA	CONC	ENTRATE			
0	1,500	0.8	310	92	89	95
	2,000	.5	340	98	89	98
	2,500	.1	440	98	88	93
10	1,500	.9	715	98	81	96
	2,000	.2	780	98	89	97
	2,500	0	820	98	88	99

<sup>1</sup>Conditions: Charges roasted at 800° C for 4 h and calcines leached for 18 h at 75° C at 40 pct solids.

#### LEACHING UNDER PRESSURE

Leaching tests were performed under pressure to determine if improved vanadium and uranium extraction could be achieved without affecting phosphate extraction and to determine the conditions required for optimum extractions.

Conda ore, Fort Hall ore, Conda (high-Mg) ore, and Conda concentrate samples were pressure leached in a 2-L, glass-lined Parr<sup>5</sup> pressure vessel. The 50-g charges were leached for 3 h at 20 pct solids under an oxygen atmosphere. The

total pressure, temperature, and acid addition levels were varied. Combinations of temperature and pressure were  $90^{\circ}$  C, 50 psi;  $140^{\circ}$  C, 90 psi; and  $200^{\circ}$  C, 260 psi. (Oxygen overpressure was 50 psi.) Acid additions were varied from 1,000 to 2,500 lb/st  $\rm H_2\,SO_4$ . Test results are given in table 14.

Compared with other methods, leaching under pressure generally produced lower extractions; however, this may be due to the shortened leach time for these tests (3 h compared with 18 h for all previous test series). For two of the samples, fairly comparable results were produced at lower acid addition levels than for the 18-h leach tests—Conda ore and Conda concentrate.

<sup>&</sup>lt;sup>5</sup>Reference to specific products does not imply endorsement by the Bureau of Mines.

TABLE	14.	-	Pressure	leaching	of	phosphorite	ores
and	cond	cer	itrate <sup>1</sup>				

H <sub>2</sub> SO <sub>4</sub> ,	Temp,	Total pressure,	Te	rminal	Extra	ction,	pct			
lb/st	°C	psi	pН	emf, mV	V <sub>2</sub> O <sub>5</sub>	U308	P <sub>2</sub> O <sub>5</sub>			
	CONDA ORE									
1,500	90	50	0.8	400	86	87	92			
	140	90	.9	450	92	88	92			
	200	260	.7	610	89	86	82			
2,000	90	50	.4	390	90	88	94			
	140	<b>9</b> 0	.3	415	93	87	94			
	200	260	•2	550	75	73	80			
		FORT HALL	ORE							
1,500	90	50	0.9	390	78	88	92			
	140	<b>9</b> 0	8.	350	78	88	93			
	200	260	.8	375	88	78	90			
2,000	<b>9</b> 0	50	.6	380	81	88	93			
	140	90	1	390	84	89	96			
	200	260	.3	370	71	14	84			
		CONDA (HIGH-Mg		ERIAL						
2,000	90	50	0.5	420	88	84	90			
	140	<b>9</b> 0	.4	470	87	73	93			
	200	260	.4	630	91	66	91			
2,500	<b>9</b> 0	50	.8	420	90	84	99			
	140	<b>9</b> 0	.2	440	91	71	99			
	200	260	•5	390	91	56	94			
		CONDA CONCE	NTRAT							
1,500	90	50	1.2	470	91	77	89			
	140	<b>9</b> 0	1.1	553	94	79	89			
	200	260	1.2	618	94	88	89			
2,000	90	50	.4	404	87	88	92			
	140	90	.6	322	60	61	61			
	200	260	.3	612	95	79	91			

<sup>1</sup>Conditions: Samples leached for 3 h at 20 pct solids under 49 to 50 psi  $0_2$ .

# Conda Ore

Preferred leaching conditions were  $140^{\circ}$  C, 90 psi, and 1,500 lb/st  $\rm H_2SO_4$ ; this resulted in extractions of 92, 88, and 92 pct, respectively of the  $\rm V_2O_5$ ,  $\rm U_3O_8$ , and  $\rm P_2O_5$  (table 14). Acid requirement was 500 lb/st less than with leaching alone.

#### Fort Hall Ore

Highest  $V_2O_5$  extraction of 88 pct resulted from using the following conditions: 200° C, 260 psi, and 1,500 lb/st  $H_2SO_4$  (90 and 78 pct  $P_2O_5$  and  $U_3O_8$ , respectively); but much higher  $U_3O_8$  and  $P_2O_5$  extractions were obtained by

leaching at  $140^{\circ}$  C, 90 psi, and acid additions of 2,000 lb/st. (See table 14.) The extractions were 84, 89, and 96 pct of the  $V_2O_5$ ,  $U_3O_8$ , and  $P_2O_5$ , respectively. These extractions were higher than those from leaching alone, but lower than those from roasting plus leaching.

# Conda (High-Mg) Material

Best extractions required 2,500 lb/st  $\rm H_2SO_4$ , which is no better than from leaching alone or from roasting followed by leaching. This acid addition level at 90° C at 50 psi enabled extractions of 90, 84, and 99 pct, respectively of  $\rm V_2O_5$ ,  $\rm U_3O_8$ , and  $\rm P_2O_5$ .

# Conda Concentrate

Preferred leaching conditions were 200° C, 260 psi, and 1,500 lb/st H<sub>2</sub>SO<sub>5</sub>.

This resulted in extractions of 94, 88, and 89 pct, respectively of the  $V_2O_5$ ,  $U_3O_8$ , and  $P_2O_5$  (table 14).

#### SOLVENT EXTRACTION TESTS

Subsequently, tests were performed to investigate methods to obtain separate products of vanadium and uranium from the various leach solutions. Table 15 summarizes the results of these tests, which involved adjusting the solution emf prior to contacting the pregnant solution with extractants.

In the first step,  $U_3O_8$  values were extracted by contacting the pregnant solution with  $0.5\underline{M}$  D2EHPA (di(2-ethlhexyl) phosphoric acid) plus  $0.125\underline{M}$  TOPO (trioctylphosphinic oxide), using a kerosene diluent. Contact was made at 25° C

for 5 min at an aqueous-to-organic (A/0)ratio of 4:1 following adjustment of the emf on all aqueous samples. Depending upon the aqueous material, the emf adjustment involved oxidation, using NaClO3, or reduction via addition of iron filings. The emf must be adjusted to the value given in table 15 or coextraction of V205 will occur. The only pregnant solution that would extract only U308 over a range of emf values was the pregfrom leaching nant solution produced Conda ore alone. The range was 500±20 mV.

TABLE 15. - Solvent extraction recovery of vanadium and uranium from phosphorite ore leach solutions

Method used	Feed solution,		Firs	t contact	;	Second contact	
to prepare	g	/L	(D2EHP	A plus To	)PO)	(OPAP p	lus IDA)
filtrate	V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> O <sub>8</sub>	Solution,	Extracti	ion, pct	Solution,	Extraction,
			emf, mV	V <sub>2</sub> O <sub>5</sub>	U <sub>3</sub> O <sub>8</sub>	emf, mV	pct V <sub>2</sub> O <sub>5</sub>
			CONDA O	RE			
Leach only	0.50	0.022	500	0	98	108	95
Roast-leach	.49	.027	491	0	99	182	96
Salt-roast and							
leach	.57	.030	490	0	97	180	87
Pressure-leach	.32	.016	510	0	94	-60	99
			FORT HALI	ORE			
Leach only	0.48	0.026	482	0	97	205	96
Roast-leach	.71	.027	488	0	96	200	89
Salt-roast and							
leach	.81	.025	491	0	96	212	89
Pressure-leach	.32	.016	515	0	94	182	99
		CONE	A (HIGH-Mg)	MATERIA	Ĺ		
Leach only	0.30	0.017	500	3	94	120	96
Roast-leach	.36	.018	478	6	94	142	97
Salt-roast and							
leach	.32	.017	480	0	94	-160	79
Pressure-leach	.16	.007	510	13	86	-30	91
			CONDA CONCE	ENTRATE			
Leach only	0.29	0.017	500	0	94	70	96
Roast-leach	.28	.031	480	0	97	200	97
Salt-roast and							
leach	.39	.032	480	0	97	202	90
Pressure-leach	.20	.017	510	0	94	190	98

The second step, which extracted  $V_2O_5$  values, involved contacting the raffinate from the first contact with  $0.3\underline{M}$  OPAP (octylphenyl acid phosphate) plus 5 vol pct IDA (isodecanol), using a kerosene diluent. Contact was made for 30 min at 25° C at an A/O ratio of 1:1, following adjustment of the emf downward. The emf values are given in table 15, but in most cases  $V_2O_5$  could be extracted when the emf was less than 212 mV.

This method generally worked for all sample solutions, but small amounts of  $V_2O_5$  were extracted during the uranium separation step for three of the four Conda (high-Mg) material solutions (table 15). Typical single contact results (leach only for Conda ore) extracted 98 pct of the  $U_3O_8$  and none of the  $V_2O_5$  in the first step, and 95 pct of the  $V_2O_5$  during the second step. In all cases the  $P_2O_5$  remained in the raffinate.

#### SUMMARY AND CONCLUSIONS

Extractions of 91 to 99 pct of the  $V_2O_5$ , 78 to 89 pct of the  $U_3O_8$ , and 89 to 99 pct of the  $P_2O_5$  were achieved by processing Conda, Conda (high-Mg) material, and Conda concentrate samples using the following methods: leaching only; roasting with or without salt, followed by leaching; and leaching under pressure. Metal extractions required 1,500 to 2,000 1b/st H<sub>2</sub>SO<sub>4</sub> for all samples except the Conda (high-Mg) material, which required 2,000 to 2,500 lb/st. All tests involved leaching for 18 h except the pressure test, for which time was reduced to 3 h. Roasting was done for 4 h, with or without salt.

 $V_2O_5$  extraction from Fort Hall ore was (1) 78 pct when the ore was leached at 75° C for 18 h, (2) 78 to 88 pct when leached under pressure for 3 h at 90° to 200° C with an oxygen overpressure of about 50 psi, and (3) 97 to 98 pct when roasted without salt at 500° or 800° C or roasted with salt at 800° C for 4 h, followed by leaching for 18 h.  $U_3O_8$  and  $P_2O_5$  extractions were 74 to 91 pct and

89 to 99 pct, respectively. The required acid addition for this ore was  $1,500 \text{ lb/st H}_2 \text{ SO}_4$ .

Preliminary solvent extraction tests showed that vanadium and uranium products could be separated from these leach solu-Tests conducted at ambient temperature showed that when the emf of the leach solutions was adjusted to approximately 500 mV, an average of 95 pct of the  $U_3O_8$  and none of the  $V_2O_5$  was extracted using D2EHPA plus TOPO. Adjusting the raffinate emf to less than 200 mV and contacting with OPAP plus IDA enabled extraction of 79 pct or more of the  $V_2O_5$ after one contact. All of the phosphate values were contained in the final raffinate, and therefore were available for H<sub>3</sub>PO<sub>4</sub> production.

Work needs to be done on a continuous basis to determine the effect of impurity buildup on metal recovery. In addition, an economic evaluation must be conducted to determine which leaching procedures are cost effective.

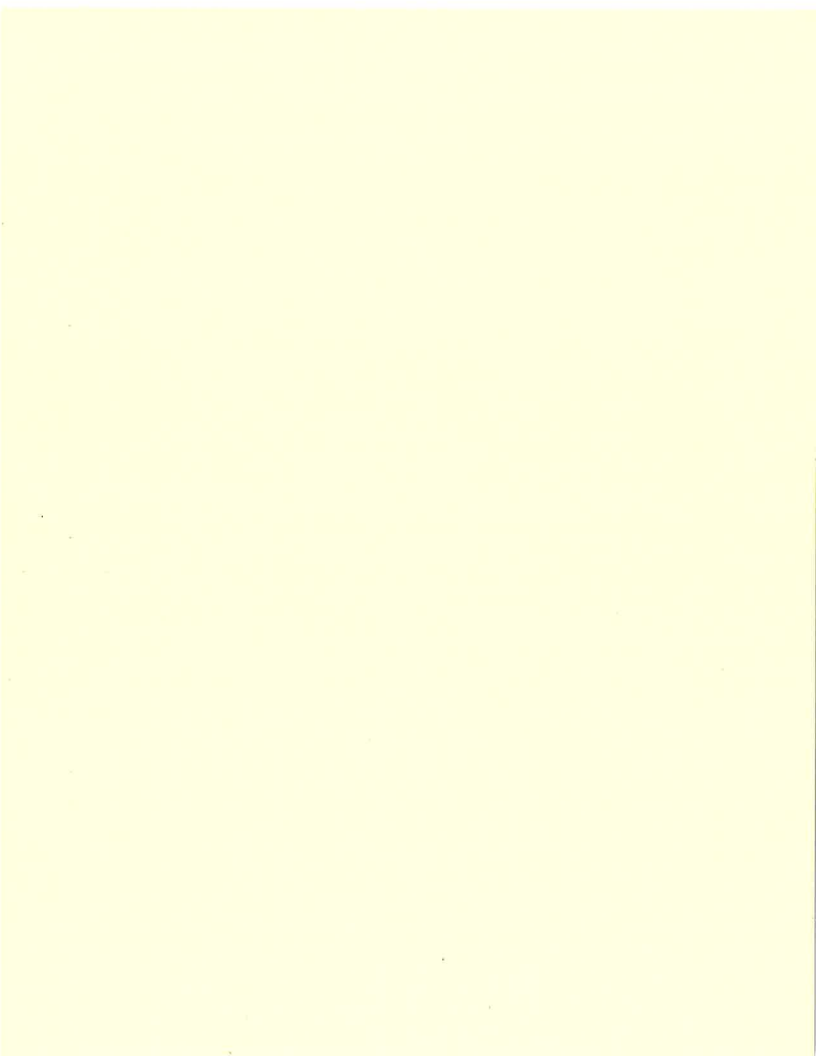
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