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TITLE MEASUREMENTS OF URANIUM HOLDUP IN AN OPFRATING GASEOUS DIFFUSION ENRICHMENT PLANT

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MEASUREMENTS OF URANIUM HOLDUP IN AN OFERATING GASEOUS DIFFUSION ENRICHMENT PLANT\*

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# ABSTRACT

Holdup of nuclear material in process equipment is one of the major sources of uncertainty in materials balances, particularly for highthroughput facilities with large equipment and extensive piping, such as gaseous diffusion uranicm-enrichment plants. Locating and measuring the holiup while the plant is operating is a challenging problem because or background from the process material and the neighboring equipment. This paper reports NDA measurements performed at the Goodyear Atomic Gaseous Diffusion Plant, Portsmouth, Ohio, on enrichment equipment at the higher enrichment end (>102  $^{235}$ U isotopic abundance) of the cascade. Both neutron and gamma-ray measurements were made to locate anomalously large deposits in converters and compressors and, within the limitations of the techniques, to quantify the amount of the deposit.

#### I. FACILITY DESCRIPTION

All the measurements were made in the 326 building at the high-enrichment end of the cancade. In this building, the two sizes of equipment are designated Type 27 (lower enrichments) ard Type 25 (higher enrichments). The cancade is organized by individual stages with 12 stages per cell, 20 cells per unit, and 9 units in the 326 building. A cell is identified by an equipment size nomber (25 or 27), a unit number, and then the individual cell, for example, 27-3-5.

Each cell, No m wide and 30 m long, is enclosed by walts and reading made of thin metal and thermal insulation. Fach entrichment stage contains a converter, cooler, compressor, and connecting piping. The nondestructive assay 9. Harbarger, J. Hicks, G. Timmons, D. Shissler, R. Tayloe, and S. Jones Goodyear Atomic Corporation Fiketon, OH 45661

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(NDA) measurements were made from the roof of the cell housing.

Within a cell, the stages are numbers 1-12; number 1 is the input side and number 12 is the output. The unit structure is shown in Fig. 1. A stage comprises piping, a converter containing the barrier material, a cooler, and a compressor. The stages are inverconnected in a complicated cluster arrangement with . Tee stages per cluster. Figure 2 is a drawing of two typical interconnected cells. It is important to understand the details of the equipment layout because it strongly influences the chuice of measurement location and data interpretation. For example, note how the converters are displaced from the center line, alternating from one stage to the next. The choice of measurement location must take this into account.

#### 11. GAMMA-RAY MEASUREMENTS

A. Uranium-235 Detection with 1.27-cm-Thick, 5-cm-Diam Nal Detectors

Hose of the gamma-ra survey Tata were col-lected with two portable 1.27-cu-thick, 5-cmdiam Nal detectors coupled to Evenine SAM-11 (two-coannel) electronics parkages. This thick-Here of Nallis nearly optimum for the detection of  $^{235}$ U 186-keV gamma rays in the presence of higher energy back tound gamma rays from the daughters of  $^{238}$  U. Each Nal detector is bound in a 0.64-cm-thick lead collimator with a handle and an additional 0.95-cm-thick lead shield over the region containing the Nal crystal. The collimator diameter is 6 cm, and the distance from the front of the collimator to the front of the detector was set at 5 cm, corresponding to a distance to the "effective" detection position of 6 cm. The single-channel analyzers were set ar every windows of 161-211 keV and 237-287 keV to measure simultaneously 186-keV gamma tave from 2350 and a count above this energy for ba kground subtraction.

<sup>&</sup>quot;Work supported by the US Department of Energy, Office of Safeguards and Security.



Fig. 2. Locations of stage components in two cells.

This technique and the cell overhead geometry used for this exercise are very similar to those used several years ago by Los Alamos to evaluate holdup in the high-enrichment portion of the shutdown cascade at K-25 Oak Ridge Gaseous Diffusion Plant.<sup>1</sup> Except for the collimator, the Nal thickness, the background subtraction capability, and the temperature stabilization of the SAM-II, the method is also similar to the RASCAL survey instrument recently developed and implemented by Goodyear Atomic Corporation. The advantages of detector collimation are higher detection sensitivity and the capability of measuring holdup in individual components of an enrichment scage. The principal disadvantage of the collimator and shield Assemblies used in the present work is their weight. They are too heavy for routine surveys that are performed manually by production permonnel.

The overhead surveys were performed with the detectors placed on the cell rooftops looking downward at selected components. Given the distance of 213 cm from the rooftop to the merdian place of the converters and most of the UF6 gas transfer components, the detector views a 175-cm-diam circle in the median plane and, in the case of a line source, a 155-cm effective length. The effective source dimensions were derived from the measured angular response of the detectors.

Figure 3 shows a plan view of the areas and components viewed by the 1.27- by 5-cm collimated NaI detectors in the overhead cell surveys. For each stage, measurements were performed at three positions centered above the circles and normal to the converter axis. The measured response was from the materials located within the circles superimposed on the cell layout drawings. Note that signals from adjacent stages are blocked by the collimator.

These two detectors were calibrated by two independent methods. The first method, pointsource calibration, was made by first measuring the counting rate with a thin  $^{235}$ U foil standard (0.395 g  $^{235}$ U) at 36.5 cm from the effective detection position in the NaI. This calibration was then scaled to the distance used in



Fig. 3. Gamma-ray det ctor viewing areas of a stage.



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Fig. 1. Layout of the enrichment cascade at the Goodyear Atomic Gascous Diffusion Plant.

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#### TABLE I

235 <sub>U</sub>	HOLDUP	IN	CALIBRATION	CELL
		(gr	ams)	

Stage	Converter
1	116
2	22
3	26
4	15
5	20
6	47
7	45
8	35
9	77
10	139
11	117
12	255

<sup>4</sup>Quantities derived using calibration constant and assuming uniform distribution of material in converter.

example of direct holdup measurement results, in this case for the evacuated calibration cell. The quantities shown for stages 1-9 can be attributed essentially all to solid deposits, whereas those for the higher stages also comprise significant amounts of residual gas-phase 2350. Based on stages 1-9 data, the average holdup in a single entire converter is 45 g 2350 and the average for an inlet region is 4 g 2350. The minimum detectable amounts of the 2350 in an entire converter and in an inlet region, that is, the detection sensitivities for counting times of 30 s are 10 and 3 g, respectively.

Figure 4 shows an example of the peak stripping procedure applied to the converter data measured with a collimated 1.27- by 5-cm Nai detector over a cell that was operating at the time of the measurement. Significant depositu are evident in converters of stages 4 and 11. A variance parameter, 0, was used to identify anomalously large 235U quantifies above the baseline composite of gas-phase 235U and smaller amounts of uniform holdup. Signs is defined as the square root of the sum of the squares of the per cent variance of the UF6



Fig. 4. Grams 2350 (from gamma-ray measure-ments) vs stage for three viewing areas.

gas phase in the converters in the calibration cell, (7.7%), and the standard deviation (%) of a 30-s count of a cell converter that showed no significant anomaly. The detection limit of 20 permits us to identify anomalies that lie outside the stage-to-stage UF6 gas-phase variations in a normally operating cell.

Table II shows the difference between the baseline count and the counts for stages 1, 2, 8, and 10 of another cell, together with the conversion to  $g^{235}U$ . The first-pass results for  $^{235}U$  were so large that an iteration to account for the self-attenuation of these deposits was needed. The numbers in the last column were obtained assuming that the uranium was uniforally deposited within each converter and using data taken over the center of the converter. The largest strenuation correction was 1.09.

Without independent information about the operating parameters of the cell, it is non-possible to subtract the gas-phase 2320 to obtain the smaller, more uniformly distributed

the survey measurements, 219 cm, by the appropriate  $1/r^2$  ratio. Next, the angular response of the detector was measured so that the effective area of the component viewed by the collimated detector could be determined for twodimensional analysis, or for one-dimensional analysis, the effective length of the source could be determined. The angular or lateral response was determined by measuring the counting rate of a large-mass <sup>235</sup>U sample as a function of the perpendicular distance from the collimator line of sight at a point 169 cm from the detector. The attenuation of the cell roof cover hatch relative to that at zero lateral displacement was included in the final lateral response function used to reduce the overhead survey data.

The next step in applying the point-source calibration was calculating the attenuation of the 186-keV gamma rays by the components being viewed. For the purpose of calculating attenuation corrections, most UF<sub>6</sub> gas transfer components, except for the converter, the central (heat exchanger) region of the cooler, and the compressor, can be considered as simple pipes with 0.95-cm walls. A detailed point-source calibration was applied only to the overhead measurement of converters, a case that is amenable for quantitative assay based on 186-keV gamma-ray detection any important because converters frequently collect significant deposits of uranium solijs.

The second calibration method used was the gas-phase technique, which is based on measuring the components of stages in a c-11 operating normally with  $UF_6$  and repeating measurements with the  $UF_6$  displaced from the cell. Figure 3 shows the viewing circles and enclosed components in the median horizontal plane of a generic stage. The centers of the circles correspond to the three gamma-ray weasured to the three gamma-ray weasured to the cell roof. The difference between the observed counts will and without UF<sub>6</sub> can the amount of  $235_{\rm U}$ respond to the three gamma-ray measurement posibe attributed directly to the amount of in the detector viewing area, providing that the gamma-ray attenuations of the various components in the field of view are the same. This condition is met in the case of the measurements made directly above the converter centers. At the other extreme, the heat exchanger regions of the coolers are very dense and transmit "ssentially no gamma rays to the detector. Thus, the cooler region is omitted entitely as a component in the measurement calibration of the converter input region. Finally, the gas-phase calibration requires that UF6 masses be obtained from stage temperature and pressure measurements and purity and isotopic analyses of samples. The 235Umasses in the components encompassed in the three measurement viewing areas used in the surveys were provided by Goodyear Atomic Corporation.

The point-source and gas-phase calibration methods agreed within 17%. The gas-phase value was used in the subsequent analyses.

B. Overhead Measurements with the 1.27-cm-Thick Detectors These surveys were conducted with the

SAM-II set for automatic background subtraction in the 186-keV window from Compton tails of the high-energy gamma rays from <sup>238</sup>U daughters, cosmic rays, and the environment. Separate channel counts were taken at least once during the survey of a cell to obtain a background count for estimation of statistical precision. A typical net count and a background count over a converter in an operating cell in Unit 27-3 were 2400 and 100 counts/min, respectively. The counting time was 30 s for each measurement point. The stability of each instrument was checked at least once during a cell survey by counting a  $^{235}$ U foil in fixed geometry. The measurement point above the converter was 2.4 m from the cell housing wall on the side of the compressor for that stage, and the measurement points for the converter inlet and outlet regions were 1.5 w from one side or the other of the cell walls, depending on the stage and cell number. Because of this topography, measurement points were designated as left, center, and right, relative to the direction of increasing stage number; the correspondence with the actual component regions was made as a part of the data analysis.

The calibration constants obtained from the UF<sub>6</sub> gas-phase measurements performed on the calibration cell can be used to convert observed counting rates to g 235U. If the quantity of gas-phase 235U in the stage components is obtained from pressure and temperature measurements and analyses of gas samples, this portion of the measured 235U can be subtracted to yield 235U holdup. Alternatively, the baseling in a particular equipment zone as a function of cell stage can be subtracted to find the amounts of large, localized deposits, which correspond to peaks in the data. Of course, the latter method, referred to as "peak stripping" or baseline subtraction, will not give the amount of 235U that hes built up uniformly in a cell. This uniform holdup combined with the in-process UF<sub>6</sub> is the source of the baseline course.

For shutdown cells, the collimated NaI detector provides a direct measurement of the holdup in its viewing field. Table I gives an

# TABLE II

# PEAK-STRIPPING EVALUATION OF ANOMALOUS 235U DEPOSITS IN CONVERTERS OF OPERATING CELL

Stage (Converter)	Counts (Baseline per 30 s)	Net Counts/min	First-Pass 235 g U	Attenuation Corrected 235 U
1	2375	4750	1185	1292
2	528	1056	263	268
8	2176	4352	1085	1172
10	417	834	208	212
			Total	2944

baseline holdup in the cell converters. For this example of peak stripping, the minimum amount of  $^{235}$ U that can be detected in a single operating converter, that is, the detection sensitivity, is 90 g. These numbers also correspond to the uncertainties that we assign to the measu ements, neglecting, of course, errors from nonuniform converter distributions.

At an example of data reduction using the gas-phase inventory and enrichment obtained from Goodyear Atomic Corporation operations and analytical chemistry laboratory, we show in Table III results for measurements of converters in cell C. This cell had no anomalies. Clearly, there is a negative bias of at least 14 g per stage in the unlikely event that the converters in this cell have no holdup. A more reasonable assumption is that this cell has a baseline average uranium holdup about the same as that of the calibration cell, which is  $\sim 20$  g  $^{235}$ U per converter.

Possible sources of the negative bias of ∿34 g <sup>235</sup>U (20 g + 14 g) per converter are (1) changes in cell gas-phase inventory and enrichment in the time interval between the temperature, pressure, and enrichment measurements and the NDA measurements; (2) a 5% error in the gasphase calibration; and (3) instrument drift. The last of these is ruled out because of the stability of the SAM-II unit and the record of the measurement control data with the standard  $^{235}$ U foil. On the other hand, the chrichment of the calibration cell changed by 25% in 1 day during the measurement campaign. Even larger negative numbers for converters, except one with a large deposit, were obtained when the gasphase subtraction method was applied to data for cell E. Clearly, taking the difference between two large numbers can yield meaningful results

# TABLE III

# HOLDUP IN CELL C CONVERTERS DERIVED FROM OVERHEAD MEASUREMENTS AND KNOWN GAS-PHASE INVENTORY A" ENRICHMENT

Stage	Measured Total 235 <sub>U (g)</sub>	Stage Average 235 U (g) in Converter	Net 235 Holdup
1	473	447	25
2	407	447	- 40
3	415	447	- 32
4	367	447	- 80
5	415	447	- 33
6	438	447	- 9
7	470	447	23
8	424	447	- 23
9	472	447	25
10	413	447	- 42
11	427	. 447	- 20
12	484	447	37
	Mean = 433	Total =	-169
	σ <b>= 3</b> 5		

only if the recuracies of these numbers are very good, that is, a few per cent uncertaincy.

As a consistency check, we used the  $UF_6$  gas subtraction method to evaluate the holdup in the calibration cell. These results,

together with the results of the cirect measurement with the  $UF_6$  displaced, are given in Table IV. Although the differences between the by-difference holdup values and those measured directly for individual stages fluctuate considerably as expecied, the totals are in excellent agreement. This may be a consequence of measuring the process parameters and enrichment simultaneously with the NDA measurements.

C. Conclusions and Recommendations for Gamma-Ray Measurements of Holdup

(i) The most valuable and probably the only overhead measurement needed for early detection of localized holdur was that made with the collimated 1.27- by 5-cm NaI 186-keV detector positioned on the cell roof directly over the converter center. Results of this study support the assumption that converters function as endicient filters for collection of solids. No new information was gained from measurements with this detector over the converter inlet and outlet components, and the amount of holdup in these components was consistently much less than that in the converter (except in cell A where the sudden multiple leaks occurred).

# TABLE IV

# 235U HOLDUP IN CELL G DERIVED FROM OVERHEAD MEASUREMENTS AND KNOWN GAS-PHASE INVENTORY AND ENRICHMENT

	By-	Directly
	Difference	Measured
	Holdup	Holdup
Stage	235 <sub>U (g)</sub>	235 <sub>U (g)</sub>
1	163	116
2	43	22
3	16	26
4	40	15
5	6	20
6	66	47
7	34	45
8	4	35 -
9	27	77
Totals	402	403

- (2) The contents of cooler heat exchangers could not be determined by gamma-ray measurements because they are opaque to the gamma rays used in this study.
- (3) The collimated 186-keV gamma-ray detector calibrated by the gas-phase technique can measure absolute holdup to an accuracy usually better than ±20%. It also sharply isolates the <sup>235</sup>U signal from individual stage components. Figures 5 and 6 compare gamma-ray measurements with and without collimation. The minimum amount of <sup>235</sup>U that can be detected in a single operating converter in *e* counting time







Fig. 6. Gamma-ray counts ve stage comparing collimated and uncollimated detectors.

of 12 s is 100 g if the data for a cell are analyzed by peak stripping. This sensitivity for detection of anomalies is limited by variations in UF6 gas-phase loadings of the converters within a cell. For a shutdown cell, the sensitivity for detecting <sup>235</sup>U in a converter is 10 g in 12 s.

- (4) Converter holdup in an operating cell can also be derived from gamma-ray measurements by subtracting the signal from the UF6 gas phase, providing the gas-phase <sup>235</sup>U is determined from careful and simultaneous measurements of stage process parameters and the cell enrichment.
- (5) On the basis of gamma-ray measurements of shutdown cells, converter holdup can be classified as normal if less than 100 g  $^{235}$ U and anomalous if greater. Data for 46 shutdown converters having less than 100 g  $^{235}$ U indicate that the nominal holdup in a size 27 converter is on the average of 100 g uranium.
- (6) Inasmuch as the average anomalous deposit in a converter was approximately 1000 g 235U for the cells measured, the collimated 186-keV gamma-ray detector can be used for detection of holdup from new leaks long before they reach this level.
- (7) The collimated 186-keV detector showed promise for detection of holdup in compressors either from the floor or several feet above the compressor.

# III. NEUTRON MEASUREMENTS

A. Description of the Neutron Measurements The neutron emission from UF6 comes primarily from the  $F^{19}(\alpha, n)$  reaction driven by alpha particles from  $2^{34}U$ . Thus, if the  $2^{34}U$ isotopic percentage is known, it is possible to calibrate a detector geometry for measuring the uranium in the cascade. The detector-geometry arrangement chosen to measure holdup in the Goodyear Atomic Corporation equipment was a large neutron area monitor, suspended over a cell with the active detector area looking down on a stage. Figure 7 shows the detector with polyethlene moderator and <sup>3</sup>He gas tubes. The viewing angle of such a detector is quite wide, particularly when compared with the collimated gamma detectors. By making two measurements, positioned as shown in Fig. 8, along the axis

Fig. 7. Large neutron area monitor used for holdup measurements.



Fig. 8. Counting geometry (1/2 maximum cone) for neutron measurements of converter-cooler components.

of a converter, an almost equal weighting is given to uranium neutrons emitted by material in a stage, except for that in the compressor. A third measurement was made to determine the quantity in the compressor (Fig. 9). The side viewing angle also includes neutrons emitted from adjacent stages. Therefore, to measure the quantity in a single stage, the data must be unfolded, subtracting the neutron rate from other stages. Such an unfolding procedure har been developed and applied to the data. The calibration of this detector geometry was performed by measuring each stage of a specific cell with and without UF6 gas present, just as for the gamma detectors. Knowing the amount of UF6 gas from plant data, the calibration constant, neutron count rate per kilogram of  $23^4$ U was calculated. because the emphasis of





Fig. 9. Counting geometry (1/2 maximum cone) for neutron measurements of compressor-cooler components.

the present measurements is on material held up in cascade equipment, the calibration constant must be corrected for the different neutron emission rates of UF6 and U02F2. The chemical form of holdup material,  $U0_2F_2$ , has an emission that is only 0.44 times that of Uf6. To determine the amount of uranium holdup in an operating stage, one first subtracts the neutron rate for the known gas-phare inventory and applies ... the corrected  $U0_2F_2$  calibration constant. This procedure assumes that the amount and isotopic composition of the stage gas-phase inveniory and the isotopic composition of the holdup material are known. Holdup isotopic composition is subject to large uncertointies because it is not known when the deposits were formed.

The neutron detector geometry wa calibraned as described. A cell was measured with  $UF_6$ , then shut down and pumped out. By subtraching the neutron counts with and without gas present, the counts due to only the gas can be determined. These data were then unfolded. Table V summarizes the data to this point. For this cell, the gas-phase inventory was calculated from process information and a sample taken for isotopic analysis.

The neutron data, while involving a series of analysis steps, hold promise of giving a quantitative estimate of large holdup deposits located by the gamma survey measurement. The following steps summarize the data analysis.

- 1. Add right and left measurements.
  - Measurement positions allow approximately equal weighting to be given to the various locations in the stage except for the compressor. Compressor measurements are analyzed separately.

•

TABLE V						
COUNT	DATA FOR CALIBRATION OF CONVERTER PLUS COOLER Count/100 a = Laf. + Right - Background					

Stage	Cas + Holdup	Joldup	Gas (Calculated) Inventory	Unfolde Gas	rd
1	445	94	351	211	
2	607	87	520	290	
3	682	72	610	359	
4	638	67	571	249	
5	716	82	634	353	
6	702	68	634	358	
7	651	81	576	262	
b	671	89	582	295	
9	704	87	617	373	
10	640	121	519	217	
11	633	120	513	326	
12	481	144	337	198	
		1112	5824 Te	3491	

Note: (Gas)<sub>Calc</sub> = (Gas > holdup)<sub>Meas</sub> = (Holdup)<sub>Meas</sub>

- 2. Subtract background.
  - Background, measured by looking down at the end of a unit with only one cell on one side and doubling it, was consistently 100 counts/100 s ± 30 counts/100 s.
- Unfold resultant stage net count rates next to give the contribution from a single stage.
  - This requires solving a set of 12 coupled but linear equations.

 $C_1$  is the measured quantity and  $S_1$  is the desired neutron source:

 $c_{i} = s_{i} + \alpha(s_{(i+1)} + s_{(i-1)})$ 

+  $\beta$ [S(1+2) + S(1-2) + ...]

- $\alpha$ ,  $\beta$  need to be determined. From the detector angular distribution and the stage geometry, the values used in this analysic are  $\alpha = 0.36$ ,  $\beta = 0.10$ .
- Separate the single-stage count rates into a component from gas-phase inventory and a second component from uranium holdup.
  - Gas-phase inventory is estimated from process parameters and assay values. Then using the calibration

constant counts/time/kg 234U, the counts from gas-phase inventory are calculated and subtracted from the total.

- The uranium holdup is determined from the counts remaining but using the calibration factor for U02F2.
- The assay (isotopic composition) of the holdup material will probably be different from the gas phase and not known This will contribute to the uncertainty in the holdup determination.

# B. Results

1. Holdup Determinations for Converter-Cooler Combination. Figures 10-12 show the stage-by-stage unfolded neutron counts for eight cells. These results correlate exactly with the gamma measurements. The deposits in coolers also contributed to the neutron count, whereas for the gamma count they did not. Thus, neutron detection samples more of the stage components. As for the gamma method, it is only possible to calculate a total holdup number for those cells not on-line or if the gas-phase inventory is known.

# Cell A

This cell had 'uptures in three expansion joints and was off-line when mersured. After measuring each stage, converter 5 was removed from the cell, and the cell was remeasured. Converter 5 was also measured in a location isolated from the cascade equipment.

Сe	1	l AHei	fore and	after	converter	5	WAB	removed

	Messured Kilograms	of Uranium
Stage	Before	After
1	1.5	1.6
2	7.1	7.1
3	6.6	6.4
4	12.9	13.3
5	27.1	25.2
6	33.5	25.8
7	10.5	13.2
8	2.7	2.6
9	5.9	5.6
10	i <b>.1</b>	1.2
11	0.6	0.6
12	3.0	2.9
	115	105.5
	$\Delta = 7 \text{ kg U}$	•

These measurements indicate a total cell holdup of 105.5 kg U after removal of converter 5 and that converter 5 contained 7 kg U.



Fig. 10. Neutron counts vs stage.



Fig. 11. Neutron counce vs stage.

#### Converter 5

Unfolding was not required because these measurements were made sfter the converter was removed and isolated from the cell. The counts were 147/100 s corresponding to 9.2 kg U.

This is compared with 7 kg from the previous difference value. The isolated measurement should be more accurate than the difference measurement made on the cell with and without converter 5.

#### Cell B

Off-line--drawing a negative pressure, that is, no UF6 gas. Holdup was 19 kg U.



# Fig. 12. Neutron counts vs stage.

Cell C

Operating cell--so an estimated count rate from the gas-phase must be subtracted.

	Estimated				
	Total	Gas-Phase	Difference		
Stage	Counts	Counts	(Holdup)		
1	209	163	46		
2	219	163	56		
3	281	163	118		
4	180	163	27		
5	206	163	43		
6	252	163	89		
7	223	163	60		
8	255	162	82		
9	186	163	23		
10	223	163	60		
11	306	163	6		
12	306	153	143		
			753		

#### Equivalent to 39 kg U

NOTE: If the background had been 30 counts/ 100 s higher, then the holdup would be reduced to 20.2 kg U, showing that the difference approach can be very sensitive to the background value.

2. Analysis of Compressor Data. The neutron data analysis procedure for the overhead compressor measurements is similar to that for the converter portion of the stage. The unfolding procedure is simpler because the compressors are farther apart. To the first approximation, the present analysis used  $\alpha = 0.15$  and  $\beta = 0$ . Figure 9 shows that the detector viewed the compressor, some piping, and most of the next up-stream stage cooler within the half-maximum ef-

ficiency cone. This overlap makes interpretation of the data difficult because it is not possible to separate the compressor and cooler contributions.

Table VI summarizes the data and indicates in average holdup per compressor-cooler of 1.4 kg U for the cell measured. For cell G thus stage-by-stage holdup is as follows:

Stage	Con	pressor Holdup U (kg)
1		1.5
2		1.7
2		1.5
5		1.5
4		1.4
5		0.4
0		0.8
7		0.8
8		0.9
9		1.6
10		1.3
11		1.7
12		1.7
	Total	15.3

3. Summary of Neutron Measurements. Table VII con, ites the results of the neutron measurements with amounts of uranium recovered for three cells and one converter. Adding the converter and compressor values given higher values than the amounts recovered. This is

## TABLE VI

# COMPRESSOR-COOLER HOLDUP

Cell	Total Count B	Gas-Phase Inventory	Δ	Uranium ( <u>+</u> E)
٨	988	a. <del>.</del> .	988	71
B	410		410	10
С	1418	960	458	27.4
		(1080)	(338)	(20.3)
E	1932	1572	360	10.8
		(1769)	(163)	(4.9)
r	2063	1464	599	20.2
		(1647)*	(416)	(14)
Ċ	625		625	15.4
		AV/stage (exc	luding A)	= 1.4 kg

"Attempt to include cooler in gas-phase counts.

# TABLE VII

# COMPARISON OF NEUTRON MEASUREMENTS VS RECOVERY

<u>Cell</u>	Neutron Measurements (kg U)			
	Converter	Compressor	Total	Recovery (kg U)
٨	105	71	176	120
B	19	10	29	25
С	39	27	66	
E	18	11	39	
F	12	20	32	28
C	7-15	15	22-30	
Stage	5 converter		9.2	6.9

"Total" counts the cooler twice; therefore, total values should be high; how high depends on the amount of material in the cooler.

expected because some components, for example, coolers, are weighted too much by the simple addition of results. As more experience is gained, a more suitable data analysis can be applied.

> C. Conclusions and Recommendations for Neutron Measurement's

The purpose of NDA holdup measurements is to identify cells with problems by detecting deposits, and in some cases, observing the growth of deposits from one month to the next. Ideally, the measurements should localize the deposit within a cell to a given stage and within the stage to a specific component, for example, converter, cooler, or compressor. In addition to locating the deposit, the measurement should quantify the amount within a known uncertainty.

Using the present neutron measurements, the deposit can only be localized to within one or two stages because of the relatively wide detector viewing angls. The detector looks at all components in a stage but cannot effectively indicate whether the material is in the converter, compressor, or upstream cooler. The quantitative comparisons of neutron results for a cell with Goodyear Atomic Corporation results (Table VII) are in reasonable agreement (±50%) but at present are based on only four cases. The factors affecting the measurement uncertainty are understood but more information is needed to determine the range of these factors.

To interpret the data, the UF<sub>6</sub> gas-phase inventory and isotopic assay must be known. The detector background, holdup isotopic assay, and holdup chemical composition directly affect the quantitative rusults. Therefore, effort should be made to improve the knowledge of these factors.

The principal disadvantages to routine use of the neutron measurements are the time and effort required to collect the data and the uncertainties in certain key information. The advantage to the neutrons is that they "see" the entire stage with approximately equal weighting, which is particularly important for criticality mafety measurements.

#### REFERENCE

1. R. B. Walton, "The Fearibility of Nondestructive Assay Measurements in Uranium Enrichment Plants," Los Alamos scientific Laboratory report LA-7212-MS (April 1978).