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TITLE: A STUDY OF IN-LINE PLUTONIUM ISOTOPIC ANALYSIS FOR GASEOUS PLUTONIUM HEXAFLUORIDE

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A STUDY OF IN-LINE PLUTONIUM ISOTOPIC ANALYSIS FOR GASEOUS PLUTONIUM HEXAFLUORIDE

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ABSTRACT

In-line plutonium isotopic analysis of gaseous plutonium hexafluoride (PuF_6) is very important for process control and special nuclear material accountability in any plutonium-isotope-separation process that requires a gaseous phase. Although much effort had been devoted to analyse arbitrary plutonium samples, no isotopic analysis had been done on gaseous PuF6 samples. We have initiated a study on the use of a high-resolution, gammaray spectroscopy technique to analyze gaseous plutonium hexafluoride. For the first time, PuF6 gas samples with pressures varying from 0.15 to 31 torr, which were directly fed into a gas cell from a process flow loop, were measured. The isotopic results of these measurements agree very wel) with chose of mass spectrometry measurements of solid PuF4. The precision of a 10-min measurement of a 10-torr reactor-grade PuF6 is 1.5% for 238pu, 0.22% for 239pu, 0.87% for 240pu, and 17.5% for ²⁴¹Pu.

INTRODUCTION

The Los Alamos Special Isotope Separation (SIS) Facility is designed to demonstrate the first large-scale separation of plutonium isotopes by using the molecular laser isotope-separation (MLIS) process to produce special isotopes and to convert plutonium scrap and waste. The MLIS process¹ separates specific plutonium isotopes from gaseous plutonium hexafluoride (PuF_6) using two types of lasers. The PuF6 gas, prepared from reacting plutonium tetrafluoride (PuF4) with flucrine, is mixed with an inert carrier gas. The mixture is cooled through a supersonic nosale to lower its energy. The first laser irradiates the gas and is tuned to excite a specific isotope in the PuFd molecules. Another leser then dissociates the excited PuF6 to form PuF4 in a solid form, which is collected on a series of filters.

In-line isotopic analysis for gaseous Pur_6 is very important for process development, process control, and special nuclear material accountability in any isotope-separation process that requires plutonium in a gaseous phase. Although much effort has been devoted to analyse arbitrary (solid and solution) samples, no isotopic analysis has been done previously on gaseous PuF_6 samples. We have initiated a study of an in-line nondestructive technique to measure PuF_6 gas from the MLIS process flow loop. In this paper we report on the first analysis of plutonium isotopic compositions in gaseous PuF_6 .

MEASUREMENT METHOD

The measurement method is based on high-resolution, low-energy gamma-ray spectroscopy techniques similar to those described in Refs. 2-4. In general, the isotopic ratio N(m)/N(n) of two isotopes m and n can be determined by measuring their selected gamma rays a and b, respectively.

$$\frac{N(m)}{N(n)} = \frac{R(a)}{I(b)} = \frac{I(b)}{I(a)} = \frac{T_{y}(m)}{T_{y}(n)} = \frac{c(b)}{c(a)}$$
(1)

where

- R = measured count rate of gamma rays,
- I = absolute branching intensity of gamma
- rays, T_{χ} = half-life of isotope, and
- E = relative efficiency of selected gamma rays, including detector intrinsic efficiency, counting geometry, and attenuation.

In this work, the isotopic ratios of 238pu/239pu, 240pu/239pu, and 241pu/239pu are determined by analysing the gamma-ray ratios 43.48 keV/51.63 keV, 45.23 keV/51.63 keV, and 148.6 keV/129.3 keV, respectively. The 238pu, 239pu, 240pu, and 241pu compositions in the sample can then be determined by combining isotopic ratios and correcting for the 242pu content, which is predicted by isotope correlation techniques.⁵ All gamma-ray peak areas are calculated by using a channel-by-channel summation method with a linear straight-line background subtraction. Minor interferences in the full-energy peaks are taken into account in the assay equations.

For each spectrum, the gamma-ray relative efficiencies are determined by using the quotient of the measured peak areas and their known specific activities of the select d^{239} Pu gamma rays in the sample. A simple linear ln c vs ln E (gamma-ray energy) interpolation between measured efficiency points at 38.66 and 51.63 keV is used to calculate the relative efficiencies at 43.48 and 45.23 keV. A similar interpolation between 51.63 and 68.72 keV is used to calculate the relative efficiency efficiency at 59.54 keV. The measured efficiency points at 129.3, 144.2 171.3, 195.7, and 203.5 keV are fit to a quadratic to determine the relative efficiency at 148.6 keV.

The measurement system consists of a highresolution hyperpure germanium planar detector and associated electronics, a Canberra Series 90 multichannel analyzer (MCA) with a 16-k channel analog-to-digital converter, and a Digital Equipment (DEC) Micro-11 computer and peripherals. The MCA is controlled by the computer, which has 128-k 16-bit words of memory and is a processor for data acquisition and analysis. A two-point digital stabilizer locked to the 51.63- and 129.3-keV gamma rays from ²³⁹Pu is used to maintain the energy calibration. A two-point stabilizer locked to the 51.63- and 129.3-keV gamma rays from $^{\rm 239}{\rm Pu}$ is used to maintain the energy calibration. The data acquisition and analysis program is written in FORTRAN under DEC's RT-11 V-5.02 operating system in the extended memory environment.

The detector has dimensions of 1000 mm^2 by 13 mm and a resolution (full width at half maximum) of 560 eV at 122 keV. It is located outside the glove box directly under the gas sample cell, which is separated from the detector by a 3.2-mm thick by 17.3-cm-diam polycallonate window. The cell is a right circular eluminum cylinder (~76 cm² by 6.4 cm) with a 1.6.mm-thick window facing the detector. It is installed in a sample chamber that provides a minimum of 5 cm of lead shielding to prevent the detection of gamma reys from extraneous plutonium in the vicinity. Two valves control the flow of PuF₆ gas through the cell.

RESULTS AND DISCUSSION

To study the sensitivity in measuring a gaseous sample using this low-energy gamma-ray technique, we measured PuFs feed gaseous samples with plutonium pressures varying from 0.15 to 31 torr that were directly fed into the gas cell from the process flow loop. Plutonium isotopic distributions of a typical reactor-grade PuF6 feed gaseous samples are listed in Table I. In Fig. 1, the estimated precision (10) of plutonium isotopes for 10-min measurements are plotted as a function of PuF6 partial pressure in torr (bottom horisontal scale) and as a function of plutonium mass in milligrams (top horizontal scale). The estimated precisions (solid circles for 238 Pu, open triangles for 239 Pu, solid squares for 240 Pu, and open circles for 241Pu) are calculated from counting statistics, including uncertainties from

TABLE I

ISOTOPIC DISTRIBUTIONS (in wt%) OF PuF6 FEED GASEOUS SAMPLE

Isotope	Weight Percent
238 _{Pu}	0.059
239 _{Pu}	87.32
240 _{Pu}	11.50
241 _{Pu}	0.916
242 _{Pu}	0.202

relative efficiencies and background run. Typically, within a 10-min count time, the precisions for a 10-torr PuF₆ sample are 1.5% for 238 Pu, 0.22% for 239 Pu, 0.87% for 240 Pu, and 17.5% for 241 Pu. The larger uncertainty of 241 Pu is due to the low quantity in the samples and the larger uncertainty of relative efficiency at 148.6 keV, which is determined by the efficiency points at lower gamma-ray intensities from 129.3 keV to 203.5 keV.

A typical relative efficiency curve for a 135-mg PuF₆ gaseous sample is compared to that from a 25-mg PuF₄ solid sample in Fig. 2. For gamma-ray energy above 100 keV, the relative efficiency curves are similar. However, the relative efficiency of the PuF₄ solid sample decreases steeply as energy decreases at lower gamma-ray energies. This shows that the effect of sample self-attenuation on the PuF₆ gas is very small as compared to that of the solid PuF₄.

In addition to sample mass, PuF6 partial pressure, isotopic distribution, and Am/Pu ratio, the precision obtained from gamma-ray techniques is also affected by count time. Figure 3 shows that the precision (10) obtained for a 85-mg (~11 torr) PuF6 sample for count time from 1 min to 240 min. The curves indicate estimated precision and data points indicate measured precision. The measured procisions are obtained from 15 repeated runs. The measured precisions appear to be better than the estimated precisions. This is because the same background data have been used for 15 repeated runs. In this case, ev ry run of the 15 subtracts a constant background By using different background runs for every measurement, especially when background is high, one may expect that the measured precision agrees well with estimated precision. Background is slowly increased from decomposed PuF4 deposited on the inner surface of the aluminum gas sample cell. For a 85-mg PuFg sample in the gas cell, we demonstrate that one can expect better than 1% precision within 20 s for 230 Pu, 7 min for 240 Pu, and 25 min for 238pu. The rapid plutonium isotopic analysis for a PuF6 gas development sample is very valuable for process development and process control.



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Fig. 1. Estimated precisions (10) of plutonium isotopes for 10-min measurements are plotted as a function of PuF6 partial pressure in torr (bottom horizontal scale) and as a function of plutonium mass in milligrams (top horizontal scale). The solid circles are for 238 Fu, the open triangles are for 239 Pu, the solid squares are for 240 Pu, and the open circles are for 241 Pu.



Fig. 2. Typical relative efficiency curves for PuF_6 gaseous and PuF_4 solid samples.



Fig. 3. The precision (10) of platonium isotopes on a 85-mg (~11 torr) PuF_6 sample as a function of count time (min). The curves indicate estimated precisions and data points indicate measured precisions. The measured precisions are obtained from 15 repeated runs. The solid curve and solid circles are for ²³⁸Pu, the dotted curve and open triangles are for ²³⁹Pu, the dashed curve and solid squares are for ²⁴⁰Pu. and the dashed- dotted curves and open circles are for ²⁴¹Pu.

To evaluate accuracy, three samples with different isotopic compositions and PuF₆ gas pressures were decomposed from PuF₆ gas to PuF₄ solid form and then sent to the Analytical Chemistry Group for mass spectrometry analysis. The uncertaintic: represent the estimated precision of gamma-ray spectroscopy. The average ratios of gamma spectroscopy to mass spectrometry are 0.999 for 238 pu/239 pu, 0.9945 for 240 pu/239 pu, and 1.0143 for 241 Pu/239 Pu, as shown in Table II.A. The average ratios of gamma spectroscopy to mass spectrometry are 1.0041 for 238 Pu, 1.0012 for 239 pu, C.9913 for 240 Pu, and 1.0071 for 241 Pu, as shown in Table II.B. These results show negligible bias when compared with mass spectrometry results.

CONCLUSION

In summary, we have demonstrated the first isotopic analysis of gaseous PuF₅ by using a nondestructive gamma-ray spect oscopy technique. The repid and accurate in-line isotopic analysis of PuF₅ in a process flow loop provides important information on process development, process control, and nuclear safeguards for any plutoniumisotope-separation process that requires plutonium in a gaseous phase. 1

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TABLE II

COMPARISON OF PLUTONIUM ISOTOPIC ANALYSIS BY GAMMA-RAY SPECIROSCOPY WITH MASS SPECTROMETRY

A. Plutonium Isotopic Ratios

	D. D	<u>Ratio: Gamma</u>	Spectroscopy/Mass	Spectrometry
Sample	Pu Partial Pressure (torr)	238 _{Pu/} 239 _{Pu}	240 _{Pu} / ²³⁹ Pu	²⁴¹ Pu/ ²³⁹ Pu
1	13.8	1.0046 ± 0.19%	0.9975 ± 0.10%	1.0265 ± 2.1%
2	3.1	1.0037 ± 0.68%	0.9895 ± 0.35%	1.0040 ± 6.1 %
3	5.9	0.9887 ± 0.37%	0.9965 ± 0.2%	1.0123 ± 3.4%
Average		0.9990	0.9945	1.0143
Std Dev		±0.0089	±0.0044	±0.0114

B. Plutonium Isotopes

Pu Pa	Pu Partial	Ratio: Gamma Spectroscopy/Mass Spectrometry				
Sample	Pressure mple (torr)	238 _{Pu}	239 _{Pu}	240 _{Pu}	241 _{Pu}	
1	13.8	1.0102 ± 0.19%	1.0008 ± 0.03%	0.9939 ± 0.11%	1.0188 ± 2.1%	
2	3.1	1.0102 ± 0.2%	1.0018 ± 0.08%	0.9875 ± 0.37%	0.9974 ± 6.1%	
3	5.9	0.9918 ± 0.37%	1.0011 ± 0.04%	0.9924 ± 0.2%	1.0051 ± 3.4%	
Average		1.0041	1.0012	0.9913	1.0071	
Std Dev		±0.0106	±0.0005	±0 0033	±0.0108	

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