From the lab to the real world: sources of error in UF6 gas enrichment monitoring

Marcie Lombardi

Follow this and additional works at: http://digitalrepository.unm.edu/ne_etds

Recommended Citation
http://digitalrepository.unm.edu/ne_etds/17

This Dissertation is brought to you for free and open access by the School of Engineering ETDs at UNM Digital Repository. It has been accepted for inclusion in Nuclear Engineering ETDs by an authorized administrator of UNM Digital Repository. For more information, please contact kevco@unm.edu.
Marcie L. Lombardi

Candidate

Chemical and Nuclear Engineering

Department

This dissertation is approved, and it is acceptable in quality and form for publication:

Approved by the Dissertation Committee:

Adam Hecht

Gary Cooper

Plamen Atanassov

Duncan MacArthur
FROM THE LAB TO THE REAL WORLD: SOURCES OF ERROR IN UF₆ GAS ENRICHMENT MONITORING

by

MARCIE L. LOMBARDI

B.S., Physics, Rutgers University, 2002
M.S., Health Physics, Georgetown University, 2006

DISSERTATION

Submitted in Partial Fulfillment of the
Requirements for the Degree of

Doctor of Philosophy
Engineering

The University of New Mexico
Albuquerque, New Mexico

March 2012
DEDICATION

This dissertation is dedicated to my aunt Shari Troy. I wish you could have seen it completed.
ACKNOWLEDGEMENTS

I’d like to thank my dissertation committee at the University of New Mexico, Adam Hecht, Gary Cooper and Plamen Atanassov. I must thank Duncan MacArthur for serving on my committee, but also for the many hours of guidance, editing, and moral support.

I would like to thank Kiril Ianakiev for his great ideas and allowing me to work on his projects, as well as the whole enrichment monitoring team at Los Alamos. I learned so much while working on this project, and got to go on a trip to England!

Brian Rees – thanks for keeping me motivated. I knew I could always count on you for an “are you done yet??” On that note, I would like to thank all of my coworkers and friends at LANL for their support and encouragement along the way.

Finally, and most importantly, I’d like to thank my friends and family for their support and understanding during this stressful time in my life. I thank my parents for always encouraging me. Last but not least, I’d like to give a special thanks to Nicole, my biggest fan.
Safeguarding uranium enrichment facilities is a serious concern for the International Atomic Energy Agency (IAEA). Safeguards methods have changed over the years, most recently switching to an improved safeguards model that calls for new technologies to help keep up with the increasing size and complexity of today’s gas centrifuge enrichment plants (GCEPs). One of the primary goals of the IAEA is to detect the production of uranium at levels greater than those an enrichment facility may have declared. In order to accomplish this goal, new enrichment monitors need to be as accurate as possible.

This dissertation will look at the Advanced Enrichment Monitor (AEM), a new enrichment monitor designed at Los Alamos National Laboratory. Specifically explored are various factors that could potentially contribute to errors in a final enrichment determination delivered by the AEM. There are many factors that can cause errors in the determination of uranium hexafluoride (UF₆) gas enrichment, especially during the period when the enrichment is being measured in an operating GCEP. To measure enrichment using the AEM, a passive 186-keV (kiloelectronvolt) measurement is used to determine the ²³⁵U content in the gas, and a transmission measurement or a gas pressure
reading is used to determine the total uranium content. A transmission spectrum is generated using an x-ray tube and a “notch” filter.

In this dissertation, changes that could occur in the detection efficiency and the transmission errors that could result from variations in pipe-wall thickness will be explored. Additional factors that could contribute to errors in enrichment measurement will also be examined, including changes in the gas pressure, ambient and UF₆ temperature, instrumental errors, and the effects of uranium deposits on the inside of the pipe walls will be considered. The sensitivity of the enrichment calculation to these various parameters will then be evaluated. Previously, UF₆ gas enrichment monitors have required empty pipe measurements to accurately determine the pipe attenuation (the pipe attenuation is typically much larger than the attenuation in the gas). This dissertation reports on a method for determining the thickness of a pipe in a GCEP when obtaining an empty pipe measurement may not be feasible.

This dissertation studies each of the components that may add to the final error in the enrichment measurement, and the factors that were taken into account to mitigate these issues are also detailed and tested. The use of an x-ray generator as a transmission source and the attending stability issues are addressed. Both analytical calculations and experimental measurements have been used. For completeness, some real-world analysis results from the URENCO Capenhurst enrichment plant have been included, where the final enrichment error has remained well below 1% for approximately two months.
# Table of Contents

ABSTRACT ........................................................................................................................ v  
Table of Contents .............................................................................................................. vii  
List of Figures ..................................................................................................................... x  
List of Tables ................................................................................................................... xiv  
List of Abbreviations ........................................................................................................ xv  

1 Introduction ................................................................................................................ 1  
1.1 Safeguards for Uranium Enrichment Facilities ......................................................1  
1.1.1 GCEP Basics ..............................................................................................3  
1.1.2 GCEP Proliferation Concerns .....................................................................5  
1.2 Traditional Enrichment Measurement Methods ..................................................6  
1.3 X-ray Generator as a Transmission Source .........................................................7  
1.4 Description of Dissertation Research ..................................................................8  
1.4.1 Calibration Method for Unknown Pipe Thickness .......................................10  
1.4.2 Sensitivity to Changes in Temperature and Pressure during Measurement .........................................................11  
1.4.3 Field Trial—Some “Real-World” Data .........................................................12  
1.5 Similar Concepts in Industry and Medicine .........................................................14  
1.5.1 Dual X-ray Absorptiometry .........................................................................14  
1.5.2 Two Gamma-ray Wall Thickness Gauge ....................................................15  
1.5.3 Two-Media Method for Attenuation Coefficient Measurement ..................16  
1.6 Outline of Dissertation .....................................................................................17  

2 X-ray Measurement Concepts .................................................................................. 21  
2.1 X-ray Tube Operation with a Bremsstrahlung Notch Filter ................................21  
2.2 Attenuation of X-rays in Aluminum and Uranium Hexafluoride .....................24  
2.3 Calibration Method for Determining Pipe Thickness .......................................25  
2.4 Addressing Potential X-ray Tube Instability ....................................................29  

3 Experimental Implementation .................................................................................. 31  
3.1 Introduction .......................................................................................................31
# Table of Contents

3.2 X-ray Tube and Collimator ................................................................. 33
3.3 Notch Filters ........................................................................................ 35
3.4 Flux Monitor .......................................................................................... 37
3.5 UF₆ Source ............................................................................................. 38
3.6 Detectors/MCAs .................................................................................... 43

4 Analytical Modeling and Calculations ...................................................... 45

4.1 X-ray Tube ............................................................................................. 45
4.1.1 Generated and Transmitted Spectra ............................................. 46
4.1.2 Issues with Analytical Model ......................................................... 48
4.2 UF₆ Properties ....................................................................................... 51
4.3 Effect of Wall Deposits on Measurements ............................................. 52
4.4 Flux Monitor Diode ................................................................................. 55
4.4.1 Diode Responsivity ........................................................................ 56
4.4.2 Measurement Dose Rate ................................................................. 58

5 Experimental Measurements .................................................................... 60

5.1 Flux Monitor Diode Measurements ....................................................... 60
5.1.1 Long-term Irradiation Measurement .............................................. 60
5.1.2 I-V Characterization, Diode Recovery .......................................... 62
5.2 Dual-Energy X-Ray Measurements ....................................................... 65
5.2.1 UF₆ Calibration Source ................................................................. 65
5.2.2 Transmission Measurements .......................................................... 67
5.2.3 Gamma-ray Spectra ....................................................................... 68
5.3 Sensitivity to Changes in Pressure and Temperature during Measurement .74
5.3.1 Pressure ......................................................................................... 74
5.3.2 Temperature Correction: Using the Flux Monitor ......................... 83
5.3.3 Temperature Correction: Simple Method ...................................... 85
5.4 Field Trial—URENCO Capenhurst ......................................................... 89
5.4.1 Variations in UF₆ Gas Pressure during Measurement ................. 91
5.4.2 Background Determination (Including Pipe Deposits)............... 92
5.4.3 Temperature Effects ..................................................................... 93

6 Pipe-wall Thickness Results and Analysis ............................................... 95
<table>
<thead>
<tr>
<th>Section</th>
<th>Title</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.1</td>
<td>Experimental Results</td>
<td>95</td>
</tr>
<tr>
<td>6.2</td>
<td>Calibration Curve</td>
<td>98</td>
</tr>
<tr>
<td>6.3</td>
<td>“Unknown” Pipe Thickness Measurements</td>
<td>100</td>
</tr>
<tr>
<td>6.4</td>
<td>Comparison between Analytical/Experimental Results</td>
<td>102</td>
</tr>
<tr>
<td>7</td>
<td>Error Analysis</td>
<td>106</td>
</tr>
<tr>
<td>7.1</td>
<td>Errors in Determining Pipe-wall Thickness</td>
<td>106</td>
</tr>
<tr>
<td>7.1.1</td>
<td>“Unknown” Pipe Thicknesses Measurement</td>
<td>106</td>
</tr>
<tr>
<td>7.1.2</td>
<td>Measured Ratios</td>
<td>107</td>
</tr>
<tr>
<td>7.1.3</td>
<td>“Unknown” Pipe Thicknesses Calculation</td>
<td>110</td>
</tr>
<tr>
<td>7.2</td>
<td>Enrichment Error from Wall Thickness Measurement Error</td>
<td>111</td>
</tr>
<tr>
<td>7.3</td>
<td>Enrichment Errors over Time (Passive System)</td>
<td>116</td>
</tr>
<tr>
<td>8</td>
<td>Conclusions</td>
<td>119</td>
</tr>
<tr>
<td>9</td>
<td>Future Work</td>
<td>121</td>
</tr>
<tr>
<td>Appendices</td>
<td>Appendix A: Mass Attenuation Coefficients</td>
<td>122</td>
</tr>
<tr>
<td></td>
<td>Appendix B: Equipment Specifications</td>
<td>123</td>
</tr>
<tr>
<td></td>
<td>Appendix C: MCNPX Input File—Diode Flux</td>
<td>126</td>
</tr>
<tr>
<td>References</td>
<td>Appendix C: MCNPX Input File—Diode Flux</td>
<td>131</td>
</tr>
<tr>
<td></td>
<td>References</td>
<td>133</td>
</tr>
</tbody>
</table>
List of Figures

Figure 1-1: Simplified diagram of an enrichment cascade ......................................................... 3
Figure 1-2: Hypothetical cascade arrangement showing the variation in the number of
    centrifuges in each stage .................................................................................................. 4
Figure 2-1: Notch filter concept ............................................................................................. 21
Figure 2-2: Attenuation as a function of energy in UF₆ gas and the aluminum pipe. ....... 25
Figure 2-3: Calculated energy-dependent bremsstrahlung yield of the x-ray tube for
    varying cutoff energies. .................................................................................................. 26
Figure 2-4: Transmitted spectra as calculated from Eqn. (3), with molybdenum and
    palladium notch filter thicknesses of 0.05 cm. .............................................................. 27
Figure 2-5: Ratios of the transmitted spectra as a function of UF₆ gas pressure ................. 28
Figure 2-6: Part of the experimental setup for the diode test ................................................. 30
Figure 3-1: Active AEM setup with LaBr₃ detector ............................................................... 32
Figure 3-2: Passive AEM setup with NaI detectors ............................................................... 33
Figure 3-3: The x-ray tube mounted in the active measurement head ................................. 34
Figure 3-4: Underside of the active head showing the flux monitor diode mounted in
    place .................................................................................................................................. 35
Figure 3-5: Some of the molybdenum and palladium notch filters attached to their
    mounting rings. ................................................................................................................ 36
Figure 3-6: A filter in place in the base of the active head ..................................................... 37
Figure 3-7: Flux monitor diode in its PEEK holder ................................................................. 38
Figure 3-8: Large horizontal UF₆ source with three pipe thicknesses, variable gas
    pressure, and variable enrichment .................................................................................. 40
Figure 3-9: Horizontal UF₆ source showing the three pipe thickness “steps.” ..................... 41
Figure 3-10: Vertical UF₆ source in the environmental chamber ........................................ 42
Figure 3-11: Typical temperature profile used for temperature sensitivity studies. ......... 43
Figure 4-1: Comparison between a calculated and measured spectrum ............................ 46
Figure 4-2: Effect of varying the x-ray tube cutoff voltage, with a fixed beam current... 47
Figure 4-3: Fit of the data points from Table 4–1 that are used to calculate n, the
    empirical anode coefficient of the x-ray tubes .............................................................. 49
Figure 4-4: The effect of different x-ray tube anode materials on the spectrum generated using a 0.1-mm silver notch filter .......................................................... 50

Figure 4-5: UF₆ phase diagram showing (in blue) the temperature and pressure operating range of our experiments ............................................................. 51

Figure 4-6: Comparison of the attenuation of UO₂F₂ and UF₆ ........................................................................................................... 53

Figure 4-7: Difference in attenuation between the pipe with and without deposits ............................................................. 54

Figure 4-8: Calculated spectrum generated by an x-ray tube with a 0.1-mm-thick silver notch filter ......................................................................................... 56

Figure 4-9: Flux monitor diode responsivity as a function of energy .................................................. 57

Figure 4-10: Energies used in the MCNPX input file to calculate dose to the diode ..... 58

Figure 5-1: Long-term irradiation test of the flux monitor diode at 40 kV and 1 mA...... 62

Figure 5-2: I-V characterization of the flux monitor diode after irradiation ............................... 63

Figure 5-3: Observed recovery of the flux monitor diode after the long-term irradiation test .............................................................................................. 64

Figure 5-4: Refurbished horizontal source ................................................................................. 66

Figure 5-5: Redesigned mounting/shielding box, showing ability to mount on different pipe thicknesses ......................................................................................... 67

Figure 5-6: Screen shot of the Lynx software showing a spectrum with UF₆ gas in the pipe and using a molybdenum notch filter ................................................... 69

Figure 5-7: Spectra generated with a molybdenum notch filter, varying the x-ray tube cutoff voltage and beam current ................................................................. 70

Figure 5-8: Spectra taken with a 0.3-mm-thick palladium notch filter ............................................ 72

Figure 5-9: Close-up of the palladium transmission peak .................................................................. 73

Figure 5-10: Spectrum taken with a 0.3-mm molybdenum notch filter in the pipe ................. 74

Figure 5-11: Spectra with and without UF₆ gas) in the pipe ........................................................... 76

Figure 5-12: Performance of the system with varying pressures of UF₆ gas ............................... 77

Figure 5-13: Total counts per second in the vertical source after the UF₆ gas was pumped out ................................................................................................... 79

Figure 5-14: Spectrum showing the ²³⁴Th peaks at 63 and 93 keV after the UF₆ gas was removed ........................................................................................................... 80
Figure 5-15: Series of gas pressure changes, and measured transmission peak net areas as a function of time. ................................................................. 81

Figure 5-16: Transmission and 186-keV count rates vs. pressure, used to determine $I_0$ and B. ................................................................................................. 82

Figure 5-17: Calculated enrichment vs. gas pressure ........................................ 83

Figure 5-18: Transmission peak count rate for a typical temperature profile running between 35 and 15°C ................................................................. 84

Figure 5-19: Fit of transmission rate data, as a function of temperature .......... 85

Figure 5-20: Transmission peak count rate data (red) corrected with the simple temperature correction method ......................................................... 86

Figure 5-21: Transmission net count rates corrected by postprocessing using the simple temperature correction method........................................ 88

Figure 5-22: The AEM passive measurement head as installed at the Capenhurst enrichment plant ........................................................................... 90

Figure 5-23: Schematic of measurement conditions in the unit header pipe during normal plant operation ............................................................. 92

Figure 6-1: Screen shot of a NaIGEM analysis in progress ................................... 96

Figure 6-2: Choosing a notch filter combination .................................................... 97

Figure 6-3: Ratios of transmitted palladium/molybdenum spectra ......................... 98

Figure 6-4: Pipe thickness calibration curve .......................................................... 99

Figure 6-5: Unknown thickness data points plotted against the calibration curve .... 101

Figure 6-6: Calculated ratios of the "worst case" aluminum alloy, compared to the three-step pipe and the unknown pipes .................................................. 104

Figure 7-1: Example of a NaIGEM fit used to determine the net area of the 186-keV peak ................................................................................................. 108

Figure 7-2: The results of a NaIGEM analysis ....................................................... 109

Figure 7-3: Example of enrichment values calculated as described for UF$_6$ gas at 4.5% enrichment and 50 Torr pressure .............................................. 114

Figure 7-4: Calculated values of the maximum allowable instrumentation error vs. energy ......................................................................................... 115

Figure 7-5: Enrichment calculation as a function of transmission peak energy ...... 116
Figure 7-6: Passive AEM enrichment error over almost two months of running at the
URENCO Capenhurst GCEP................................................................. 117
# List of Tables

Table 2-1: Notch Filter Selection Data ............................................................................. 24
Table 3-1: Precise Thickness Measurements for Horizontal Source ................................. 40
Table 3-2: Comparison between LaBr₃ and NaI Scintillators ............................................ 44
Table 4-1: X-ray Generator—Anode Material Coefficients .............................................. 48
Table 5-1: Average Enrichments at Various Pressures and the Effect of Count Rates in the 186-keV Peak Shown by the Standard Deviations ........................................ 78
Table 5-2: Effect of Transmission Rate Error on Final Enrichment Calculation ............... 87
Table 6-1: Pipe Thickness Results .................................................................................. 102
Table 6-2: 6061 Aluminum Alloy Composition ............................................................. 103
Table 6-3: Worst case error calculations ........................................................................ 105
List of Abbreviations

AEM – Advanced Enrichment Monitor
BDMS – Blend-down Monitoring System
CEMO – Continuous Enrichment Monitor
DXA – Dual X-ray Absorptiometry
EURATOM – European Atomic Energy Community
GCEP - Gas Centrifuge Enrichment Plant
HEU – Highly Enriched Uranium
HPGe – High Purity Germanium
HSP – Hexapartite Safeguards Project
IAEA – International Atomic Energy Agency
keV – Kiloelectronvolt
LaBr3 – Lanthanum Bromide
LANL – Los Alamos National Laboratory
LEU – Low Enriched Uranium
LN2 – Liquid Nitrogen
MCA – Multichannel Analyzer
mCi - Millicurie
MCNPX – Monte Carlo N-Particle eXtended (computer code)
Mo – Molybdenum (used for notch filters)
NaI – Sodium Iodide [refers to NaI(Tl), thallium-activated NaI, used as detector]
NIST – National Institute of Standards and Technology
Pd – Palladium (used for notch filters)

PEEK - Polyether Ether Ketone

PLEU – Product Low Enriched Uranium (BDMS terminology)

Rad – Unit of Absorbed Radiation Dose

ROI – Region of Interest

$^{235}\text{U}$ – Uranium 235

$^{238}\text{U}$ – Uranium 238

UF$_6$ – Uranium Hexafluoride (gas in the enrichment measurements)

UO$_2$F$_2$ – Uranyl Fluoride (composition of pipe-wall deposits)

URENCO - Nuclear Fuel Company Operating Enrichment Plants in the United Kingdom, Germany, the Netherlands, and the United States
1 Introduction

1.1 Safeguards for Uranium Enrichment Facilities

The International Atomic Energy Agency (IAEA) has been applying safeguards at gas centrifuge enrichment plants since the 1970s [1]. These safeguards were strengthened in the 1980s by the Hexapartite Safeguards Project (HSP). The HSP included Japan, Australia, the United States, the IAEA, EURATOM, and the countries comprising URENCO (e.g., Germany, the Netherlands, and the United Kingdom). It addressed nuclear material accountancy, the structural features of the facilities (whether they were “safeguards friendly”), and whether access to the cascade halls would be granted [2]. An “inspection-free” approach was considered because of concerns that access to the cascade halls might reveal proprietary operational details.

The HSP working group ultimately chose to allow inspectors limited access to the cascade halls instead of the inspection-free approach. Limited-frequency unannounced access was chosen, which would allow a set number of inspectors to make unannounced visits to the cascade halls, a set number of times per year. The fact that these visits are unannounced is, in itself, a significant deterrent. If it is possible for inspectors to arrive unannounced at any time, plant operators are less likely to deviate from their regular allowable operations for fear of being caught. The inspectors do not need any special equipment but check to see that the facility has not been modified in any way from the declared equipment configuration and operation.

More recently, an improved model safeguards approach has been developed by the IAEA [3]. It requires new techniques for more detailed inspections of the advanced
technologies and increased output from modern centrifuge enrichment plants. The specific safeguards goals include the timely detection of the following:

- The diversion of nuclear material from the declared nuclear material flows and inventories.
- Facility misuse to produce undeclared UF₆ product at the declared product enrichment levels from undeclared feed (excess production).
- Facility misuse to produce UF₆ at enrichments above the declared maximum, in particular HEU.

Because it has become evident to the IAEA that the actions on the above list could be achieved with little or no modification of the equipment in the cascade hall, the Agency desires a technique by which a piece of equipment (the enrichment monitor) could be mounted on a cascade header pipe to continuously monitor the enrichment of the gas being produced. Some of the earlier versions of these enrichment monitors will be discussed in Section 1.2, “Traditional Enrichment Measurement Methods,” after which the Advanced Enrichment Monitor (AEM), our enrichment monitor, will be presented.

In the introduction, the basics of gas centrifuge enrichment plants (GCEPs) will be briefly reviewed, along with proliferation concerns that arise from the operation of the GCEPs. Traditional enrichment measurement methods such as the blend-down monitoring system (BDMS) [4] and continuous enrichment monitor (CEMO) [5] will also discussed. Next, the use of an x-ray generator, instead of a radioisotopic source, for transmission measurements will be discussed and, finally, the purpose and a brief description of the dissertation will be provided.
1.1.1 GCEP Basics

GCEPs provide a common, relatively economical method for enriching uranium to levels suitable for use as fuel in power reactors. In a GCEP, each individual centrifuge is fed UF$_6$ gas and spins at a very high speed to separate the $^{235}$U from the $^{238}$U. Because the $^{238}$U is slightly heavier than the $^{235}$U, it is pushed to the outer walls by the centrifugal force, and the gas extracted from the center of the centrifuge is (very) slightly more enriched than it was when it entered. Since it is only slightly more enriched, these centrifuges are connected in cascades, with different stages for different levels of enrichment. Figure 1.1 below is a simplified schematic of a typical GCEP cascade. A real cascade is much more complex, with many more stages.

![Figure 1-1: Simplified diagram of an enrichment cascade. Each diamond represents a stage, and all of the stages together make up the cascade. Enriched UF$_6$ gas is fed upwards to the next stage, while depleted gas is fed back into the lower stage.](image-url)
Each stage in the picture above, represented by a diamond, contains UF₆ gas at a certain enrichment. When the gas is separated by the centrifuge, the portion that is more enriched is passed onto the next higher stage, while the portion that is more depleted is passed down to the previous stage. The enrichment continues through a number of stages, (many more than are pictured here) until the desired level is reached. The depleted gas usually passes through fewer stages than the enriched gas because the tails are usually removed at approximately a 0.3% enrichment, whereas the initial feed, if it is natural uranium, is approximately a 0.73% enrichment. The number of centrifuges in each stage also varies. Figure 1.2 shows the stages in a hypothetical cascade arrangement.

![Figure 1-2: Hypothetical cascade arrangement showing the variation in the number of centrifuges in each stage. Typically the most centrifuges are in the stage where feed is introduced—stage 4 in this case.](image-url)
In this example, natural UF$_6$ feed, 0.73% enriched, would enter into the cascade at stage 4. Each stage feeds the next, and tails are fed downwards and removed after stage 1. The product, typically enriched to between 3% and 6% for low enriched uranium (LEU), is removed after stage 12. The number of cascades and the complexity of the entire facility are a proliferation concern because of the ease with which misuse could occur.

1.1.2 GCEP Proliferation Concerns

GCEPs are proliferation concerns because in addition to producing UF$_6$ at enrichments useful for power production, such plants can be used to enrich the UF$_6$ to much higher levels, such as those needed for nuclear weapons production. Enrichment to a higher level can be accomplished with little to no modification of the process being used. The two most common proliferation scenarios are batch recycling and cascade interconnection with partial reconfiguration [6].

With batch recycling, the product removed from the final stage is fed back into the cascade (into stage 4 in the hypothetical scenario in Fig. 1.2). Highly enriched uranium (HEU) can be produced much more quickly using batch recycling than with the second method, which takes some rerouting of plant piping. However, batch recycling is very wasteful since higher enrichments are being discarded as tails.

With the cascade interconnection, the product from one cascade is fed directly into a second cascade; this interconnection can be repeated as often as desired. The cascade interconnection breakout scenario, though taking longer to configure, is more efficient than batch recycling, and once it is up and running could also produce HEU in a fairly short period of time [7].
These possible breakout scenarios make it clear that enrichment monitoring is a necessary technology, especially online unattended monitoring that would detect suspect activities in a timely manner [8].

1.2 Traditional Enrichment Measurement Methods

Traditional active enrichment measurement methods, such as the CEMO [9], use a radionuclide source such as $^{109}$Cd or $^{57}$Co. These systems rely on a passive measurement of the 186-keV (kiloelectronvolt) gamma ray to measure $^{235}$U content and a transmission measurement to determine the gas density. The ratio of $^{235}$U (measured by the 186-keV counts) to the total uranium gives the enrichment [10]. A fairly low energy source is required so that attenuation in the gas can be measured. A CEMO’s capability is limited to distinguishing between UF$_6$ containing LEU (approximately 4% $^{235}$U) and that containing HEU (above 20% $^{235}$U).

With the CEMO method, an empty pipe calibration needs to be performed periodically in a laboratory, with a pipe of similar composition and thickness to the one being measured in the facility [11]. However, pipe thicknesses may vary significantly between the laboratory calibration source and the pipe in the facility because of the nature of the pipe manufacturing process. A pipe with a 100-mm inner diameter and 4-mm wall thickness typically has a ±0.4-mm thickness tolerance. Depending on the enrichment and pressure of the gas in the pipe, this variation could easily cause the measured enrichment error to fall outside of the acceptable range. It should be noted that the calibration error caused by differences in the wall thickness between the calibration and facility pipe has been previously analyzed in detail [12], [13].
Since the CEMO was intended as a go/no-go indicator—simply to inform the IAEA whether a cascade was producing LEU as intended or had reached HEU levels—the error in enrichment introduced by performing the calibration on a different empty pipe was acceptable. However, since we are now trying to achieve much higher precision on our enrichment determination (to within 1%), a more accurate way of calibrating our instrument is required. A CEMO is designed to be installed on the individual header pipe of a single cascade, rather than on the product unit header, making it very hard to monitor a whole plant. To monitor the entire plant, a separate unit would need to be installed on each of the cascades, as discussed above in Section 1.1.2.

The BDMS contains an enrichment monitor to perform unattended measurements during the blending down of Russian HEU [14]. This system was developed under the 1993 HEU Purchase Agreement between the Russian Federation and the United States, which specifies the blending down of 500 metric tons of HEU into reactor grade uranium. Once the HEU is blended down to LEU, it is purchased by the United States for use in power reactors. In this way, Russia has a financial incentive to blend down its surplus weapons-grade material, making the deal mutually beneficial [15]. The BDMS system monitors this process under the agreement, verifying the enrichment and mass flow rate in the three legs of the stream: HEU, LEU, and PLEU (product LEU). In an operating enrichment facility, it may not be feasible to directly measure an empty pipe in order to calibrate for pipe attenuation, as is the case with the BDMS system [16].

1.3 X-ray Generator as a Transmission Source

Using an x-ray tube as a transmission source for UF₆ gas enrichment monitoring eliminates the costly replacement of the traditional gamma-ray source as it decays. A
\(^{109}\)Cd source has a half-life of 463 days. A way to compensate for this relatively short half-life is to start by installing a “hot” source, in order to maintain its useful activity for as long as possible. An attenuator is used to reduce the intensity of the source when it is first installed. Periodically the amount of attenuation is reduced in order to maintain a reasonable intensity. Each time the attenuator is replaced, the system needs to be recalibrated. Typically the source itself must be replaced every two to four years.

An x-ray tube does not need to be replaced as frequently because the expected lifetime of the tube is tens of years. In the operation of the transmission-based AEM, the x-ray tube is run at a very low power, compared to its rated capacity, in order to extend this lifetime. In addition, for system maintenance, the tube can be turned off so no source handling is required. However, the output of the x-ray tube can vary due to a number of factors, such as temperature changes, tube degradation, etc. An in-beam silicon flux monitor diode is used to correct for any instabilities in the output of the tube.

The x-ray tube is operated with a notch filter, the material of which is selected to transform the bremsstrahlung output of the tube into a spectrum with a sharp energy peak, determined by the K-edge of the filter. Thus the user is able to select the transmission peak energy as well as the beam intensity, giving much more flexibility with the x-ray tube than with a radioisotope as a transmission source.

1.4 Description of Dissertation Research

The AEM is designed to measure the enrichment of gaseous UF\(_6\) in cascade header pipes. The enrichment is determined by measuring the ratio of the \(^{235}\)U to the total amount of uranium present. A passive measurement of the 186-keV gamma ray
determines the amount of $^{235}$U, and a transmission measurement determines the total amount of uranium in the gas. The enrichment is calculated by the following formula:

$$E(t) = K_{cal} * \frac{R(t) - B}{I_0 - I(t)} ,$$  \hspace{1cm} (1)$$

where $K_{cal}$ is a calibration constant, $R(t)$ is the 186-keV count rate as a function of time, $B$ represents the 186-keV counts coming from background (including pipe deposits), $I_0$ is the empty pipe transmission rate, and $I(t)$ is the measured transmission rate with gas in the pipe, as a function of time. The numerator determines the amount of $^{235}$U, and the denominator determines total uranium by measuring the attenuation by the gas. The ratio of the two, multiplied by a calibration constant, gives the enrichment as a function of time. To perform this transmission measurement, we use an x-ray tube with a notch filter. The filter material determines the energy of the transmission peak spectrum that is generated. This system is designed to be installed in a facility and run in an unattended operation mode, with data being sent back to a central location.

There are several issues with the AEM operation that may lead to potential sources of error in the enrichment determination. In order to address these issues, this dissertation explores real-world error sources in enrichment measurement, including dynamic variations in operational parameters. Topics include cascade header pipe-wall thickness concerns, x-ray tube instabilities, and notch filter material selection. Further, this dissertation studies the sensitivity of the enrichment measurement to changes in pressure and temperature during measurement. The end goal of this dissertation is to study the contributing factors that lead to errors in enrichment monitoring and to find possible ways to mitigate (wholly or partially) these errors.
1.4.1 Calibration Method for Unknown Pipe Thickness

Based on the IAEA’s requirements, it can be argued that continuous, unattended monitoring of the GCEP is desired. In order to do this, however, the monitor must be calibrated for the specific measurement location before it can be run in unattended mode. Therefore, a method of determining the pipe-wall thickness in this location, while the UF$_6$ gas is present in the pipe, is required. The gas pressure may not be known, so this calibration must be independent of the amount of gas in the pipe. Once the pipe attenuation is known, the gas pressure can be determined with another transmission measurement. Since attenuation in the aluminum pipe is much greater than in the gas, small differences in pipe thicknesses from facility to facility, or even from pipe-to-pipe, would greatly affect the UF$_6$ gas density results if this method were not used.

The following is the simplified formula used in a conventional determination of the enrichment of the UF$_6$ gas [17]:

$$E = K \cdot \frac{I_{186}}{\ln \left( \frac{I}{I_0} \right)} \%,$$

(2)

where $I_{186}$ is the intensity of the 186-keV peak obtained using a passive measurement, and $I$ and $I_0$ are obtained by transmission measurements, with and without attenuation by the UF$_6$ gas. $K$ is a calibration constant. Traditionally, an empty pipe measurement was needed to determine the attenuation by the pipe without any gas present ($I_0$). Another option would be to use a facility declaration of the gas pressure. However, because the purpose of enrichment monitoring may be to detect facilities that are trying to hide improper use, facility declarations cannot be assumed trustworthy. Therefore, this
dissertation proposes a two-energy x-ray transmission method for pipe thickness determination in those cases where empty-pipe measurements are not feasible.

Two transmission measurements of the header pipe, at energies with closely matched attenuation in the UF₆ gas, are performed. Looking at the ratio of these two will enable us to determine the attenuation in the aluminum pipe, since the attenuation in the gas will cancel out [18]. This cancellation is possible because the selected transmission energies are around the uranium L-edge region. While the attenuation at these two transmission energies in the UF₆ gas is nearly equal, attenuation in the aluminum pipe wall at these two energies differs by a factor of about 60. The large effect of the aluminum pipe attenuation on the transmitted spectrum can lead to a large measurement error if the pipe thickness is not determined accurately. A comprehensive error propagation analysis is detailed in Chapter 7, “Error Analysis,” determining the precision needed in the pipe thickness measurement in order to obtain an accurate enrichment measurement.

1.4.2 Sensitivity to Changes in Temperature and Pressure during Measurement

In an operating GCEP, the UF₆ gas pressure is constantly changing. One reason is because there are variations in the pumping power used (our AEM is placed after the compressors that send the gas to the fill stations). Another factor that affects gas pressure is the number of cylinders being filled at any one time. When a chilled, empty cylinder is attached at a fill station, there is a sharp drop in the gas pressure in the header pipe. The number of cylinders that are attached for filling at any one time causes variations in the rate of pressure change in the pipe. Data that illustrate this phenomenon are presented in Section 5.4, “Field Trial—URENCO Capenhurst.”
The temperature in our measurement location also has the possibility of constantly changing, for three reasons. First, the gas is heated by the compressor as work is done on it. Second, there may also be some cooling effect as the chilled cylinder to be filled is attached. Third, many GCEPs are not temperature controlled in the area where the centrifuges/pumping stations are, so daily ambient, and therefore pipe, temperature fluctuations will be seen.

The sensitivity of the enrichment measurement to pressure and temperature has been studied in detail. Not only do we have operational data from a GCEP, but we also have a number of laboratory UF₆ sources with variable gas pressure. One of these sources was small enough to fit into an environmental chamber to do temperature sensitivity measurements. Tests were run in the environmental chamber with the whole system inside the chamber. This simulated operation in an actual facility, where all of the components are subject to temperature variations. We are able to study temperature effects on the x-ray tube, the UF₆ gas, the NaI (sodium iodide) detector, and the various electronics used.

1.4.3 Field Trial—Some “Real-World” Data

In August 2011, a team from Los Alamos National Laboratory (LANL) composed of Kiril Ianakiev, Duncan MacArthur, and Marcie Lombardi (the author) traveled to the URENCO Capenhurst plant in the UK for a field test of our AEM system on a real cascade header pipe. We installed our passive enrichment monitoring system, which uses a passive 186-keV measurement plus a facility-supplied gas pressure reading to determine the UF₆ enrichment. We performed a test fit of the newly designed active enrichment monitoring head on the pipe to demonstrate how it would be mounted during
The plant representatives’ main concern was the weight of the system because it is clamped directly onto the pipe. Our main concern was that we had a tight enough fit, so there would be no shifting at all during operation, which would introduce additional geometrical errors. URENCO personnel determined that the weight and attachment mechanism was acceptable. This system is being returned to LANL for further testing (mostly for mechanical stability) and to complete the electronics package fabrication.

During the August visit, two ½” thick by 3” in diameter NaI detectors, in a “face-to-face” orientation, were installed. This configuration was intended to increase the counting efficiency as well as to provide a backup in case one detector failed. Tungsten composite shielding was used around the pipe and the detectors to avoid measuring background radiation. The system is supported by a table, and Neoprene sheets were placed between the pipe and the shielding pieces. The data acquisition will run over a period of about one year in unattended mode.

We also installed four temperature sensors near the AEM. The output of these sensors is also logged by our data collection system. We are currently monitoring the ambient air, the aluminum product pipe very close to the monitor, the product pipe just after the compressor (upstream of the monitor), and on the steel bellows immediately following the compressor.

Another visit to the Capenhurst GCEP is planned for April of 2012 to install the completed, active AEM and update the passive system software to allow real-time enrichment determination.
1.5 Similar Concepts in Industry and Medicine

A literature review of concepts similar to the two-energy pipe thickness method found the following measurement methods are currently used either in industry or medicine: (1) Dual x-ray absorptiometry is a medical procedure using two transmission measurements to determine patient bone density. (2) The two gamma-ray wall thickness measurement is a method for determining cylinder thickness from the emissions of the radioactive material contained in the cylinders. (3) The two-media method uses repeated transmission measurements of an object, placed in different media. The goal is to determine the linear attenuation coefficient of a sample of any shape. All of these concepts have similar aspects to the two-energy wall thickness method but do not use the idea of “canceling out” attenuation in a media because of its similar attenuation properties at different energies.

1.5.1 Dual X-ray Absorptiometry

Dual photon (x-ray) absorptiometry is a medical procedure that uses transmission measurements at two different energies. It could use a radionuclide such as Gd-153, which has peaks at 41 and 100 keV, or an x-ray generator. In the techniques using x-rays, it is better known as dual x-ray absorptiometry, or DXA. The bone mineral content of an area such as the lumbar vertebrae can be determined by using the different attenuation coefficients of the bone and soft tissue. Once soft tissue absorption has been subtracted out, bone mineral density can be determined using the absorption of each beam by the bone [18].

In this technique, which is most similar to our two-energy thickness determination method, the x-ray tube is operated at a constant output voltage, and K-edge filtering is
used to produce spectra with peaks at two different energies. The transmission of the patient is then determined, for each pixel, using spectroscopy and energy windowing. This is similar to our method of using regions of interest (ROIs), one for each transmission peak. Norland uses a samarium filter that has a K-edge at 45 keV, and the x-ray tube is operated at 80 kVp [20]. A peak with a maximum energy of 45 keV is generated, and the higher energy transmission uses the bremsstrahlung above 40.4 keV, up to 80 keV. With the GE Lunar Bone Densitometer, a cerium filter is used, resulting in peaks at 38 and 70 keV, using the same principal [21].

1.5.2 Two Gamma-ray Wall Thickness Gauge

The two-gamma-ray wall thickness gauge was developed to account for variations in the wall thicknesses of UF₆ type 5-A containers [22]. It uses the 144- and 205-keV lines from $^{235}$U to determine a thickness correction factor for the walls.

Initially, $^{75}$Se was used as an external source. Count rates in two ROIs were determined with and without an absorber (using slab geometries). With this method, if the initial intensities of the gamma rays are equal, the difference in the attenuation of the two through an absorber indicates thickness. Ratios in attenuation are used to determine the thickness. This was used as a proof of principle experiment only.

An internal two gamma-ray transmission method has also been used for instances where measuring unattenuated gamma-ray count rates is not possible, such as measuring a sealed source in a container. The ratio between the transmission of two gamma rays with known intensities can be used to determine the container thickness. In this case, detection efficiency, containment attenuation, and matrix self-attenuation must be corrected for. Multiple gamma-ray measurements were also explored. For example, if the
ratio of each of two gamma-ray lines is taken with a third, then two thickness measurements can be obtained, providing greater accuracy.

An enrichment measurement technique is discussed, where the $^{235}\text{U}$ 186-keV count rate is measured with a high-purity germanium (HPGe) detector. In this method, a standard of known enrichment is needed. To account for minor wall thickness fluctuations between the standard and cylinders being measured, a two-gamma-ray method is used and a correction factor is determined, relating the standard to the unknown thickness. Results are compared using an ultrasonic gauge to measure the wall thicknesses.

The main conclusion of the wall thickness work is that ultrasonic thickness determination requires about $\frac{1}{4}$ the count time of the two gamma-ray measurements to achieve the same accuracy. Uncertainties in the ultrasonic method were also much lower.

1.5.3 Two-Media Method for Attenuation Coefficient Measurement

The two-media method [23] is an experimental procedure for determining the linear attenuation coefficient of a sample. It is useful when a sample is irregularly shaped and it is not easy to determine the sample thickness. A 100-mCi $^{241}\text{Am}$ radioisotopic source is used for transmission measurements with a 2” × 2” NaI detector. The sample is immersed in two different media with known linear attenuation coefficients. Two separate transmission measurements are performed, one with the sample in each medium. The entire setup is placed in an acrylic box, and the source is collimated to transmit straight through the sample and into the detector.

By performing two measurements in media with known attenuation coefficients, all distances cancel out because these do not change between the two measurements. The
absolute linear attenuation coefficient of the sample is determined by looking at the ratio of the two measurements.

This method is similar to the two-energy transmission method that is used for determining GCEP pipe thickness because it uses the ratio of two transmission measurements to determine an unknown parameter. However, in the wall thickness measurement, the L-edge region of the UF₆ gas must be used to cancel out gas attenuation between the two measurements. The two-media method is too basic for our purposes, where we have two unknowns: the pipe thickness and the UF₆ gas density.

1.6 Outline of Dissertation

This dissertation is structured into eight chapters. In this first chapter, enrichment monitoring fundamentals and AEM requirements are explored, plus a review of some of the concepts in industrial or medical applications that are similar to the two energy thickness determination method is presented. In Chapter 2, the use of the x-ray tube for transmission measurements is discussed. The notch filters are explained in more detail, including the methodology for choosing which notch filters to use and why they work for this application. In addition, the attenuation properties of UF₆ and aluminum are discussed, and the basis for the two-energy method for thickness determination is examined. This leads into an explanation of the two-energy method of calibration. A solution for any instabilities in the output of the x-ray tube is then presented, and the use and testing of the flux monitor diode are discussed.

In Chapter 3, the setup and equipment used in this dissertation are described. In addition, this chapter includes a general diagram of a typical measurement setup and
explains each of the individual components used. These components include discussion of the following:

1. the x-ray tube used for transmission measurements and pipe-wall thickness determination,

2. the notch filters used to generate a transmission peak with the x-ray tube,

3. the silicon diode used as a flux monitor to correct for any instability in the output of the x-ray tube,

4. the laboratory UF$_6$ sources used for testing, and

5. the different detectors and multichannel analyzers (MCAs) that were used.

Chapter 4 discusses analytical modeling and calculations. The general equation used to calculate the output of the x-ray tube is presented, from the original bremsstrahlung to the final spectrum using the K-edge notch filters. This discussion includes attenuation by the pipe, gas, and notch filters themselves. The calculations used to select notch filter thicknesses to test experimentally are shown. Some issues with the analytical model are also presented. The properties of the UF$_6$ gas that might affect our measurements and the temperature and gas pressure ranges that we operate in, as related to these properties, are presented. Furthermore, the effect of wall deposits, or holdup, on both the active and passive measurements are discussed, and calculations that show why the possibility of deposits will not negatively impact the measurements are given. Finally, the equations used to calculate the radiation hardness of the flux monitor diode specific to our application are provided. The dose that was received by the diode was calculated with an energy responsivity curve and an MCNPX (Monte Carlo N-Particle eXtended computer code) calculation.
Chapter 5 presents all of the measurements performed, both in the laboratories at LANL and in the URENCO Capenhurst plant. The chapter begins by explaining the testing performed on the flux monitor diode to establish whether the diode would hold up to long-term use in a facility. It then describes the experimental procedure used for the pipe-thickness measurements, discussing both the laboratory UF₆ sources and the transmission measurements performed. Finally, this chapter discusses experimental measurements to determine the sensitivity of our enrichment measurements to changes in gas pressure and temperature and concludes with a discussion of the field trial in Capenhurst (with the passive system only).

Chapter 6 presents the results of the pipe-wall thickness experiments. The calibration curve that was measured and used to determine three “unknown” pipe thicknesses is given. The experimental results of the determination of the three pipe thicknesses are presented, and the initial analytical results are compared with the experimental results.

Chapter 7 presents a detailed error analysis, concentrating on the factors that contribute to the error in our final enrichment determination. The error in measurement parameters and how these propagate into error in the pipe-wall thickness determination are discussed, as well as the manner in which this contributes to enrichment errors. In this chapter, enrichment errors over time are also examined, using some data from the Capenhurst plant that have been acquired continuously over a period of months.

Chapter 8 presents the conclusions and the main findings of this dissertation. These include the less than 2% accuracy that can be achieved when determining pipe-wall thickness when an empty pipe calibration is not possible. This type of calibration is
the only option for calibration, and therefore, enrichment determination in an unfriendly country where facility declarations (such as gas pressure) cannot be trusted. Also summarized are the real-world analysis results from the URENCO Capenhurst plant, accounting for variations in temperature and pressure and achieving a final enrichment error of less than 1% during almost two months of unattended operation.
2 X-ray Measurement Concepts

This chapter details the operation of the x-ray generator as it is used in the AEM. The concept of a notch filter is presented and explained. The attenuation of x-rays in aluminum and UF₆ are discussed, and from these the calibration method for determining pipe thickness in the presence of gas is detailed. Finally, ideas that address potential x-ray tube instabilities are presented.

2.1 X-ray Tube Operation with a Bremsstrahlung Notch Filter

In the operation of the active AEM, thin targets are used as notch filters to transform the bremsstrahlung spectrum produced by the x-ray tube into a more useful spectrum with a sharp peak [24]. The maximum energy of this peak is determined by the K-edge of the filter material. Figure 2.1 demonstrates the basic concept of a notch filter, showing the bremsstrahlung output of the x-ray tube being transformed into a sharply-peaked spectrum because of the attenuation by the notch filter selected.

Figure 2-1: Notch filter concept demonstrating the transformation of the bremsstrahlung spectrum from the x-ray tube into a useful transmission peak spectrum, with the maximum energy of the peak determined by the K-edge of the notch filter.
The x-ray tube is operated so that the energy it emits is slightly higher than the K-absorption edge of the filter. A sharp peaked spectrum is emitted because the filter absorbs radiation above the K-edge energy that corresponds to the binding energy of the electrons in the K-shell of the atoms in the filter material.

One advantage of using an x-ray tube with notch filters is that it allows greater flexibility in selecting transmission energies. Traditionally, when measurements were performed with an isotopic source, the 22-keV silver x-ray from a decaying $^{109}$Cd source was used to measure attenuation in the UF$_6$ gas. There are not many choices available that have both an optimum energy and a long enough half-life to be useful. With the notch filter method, a wide range of transmission peak energies is available. However, there is a trade-off between the attenuation in the gas and attenuation in the pipe, which is large for such low energies. For the transmission measurement that determines the UF$_6$ gas density, the goal is to try to maximize the attenuation in the gas and minimize the attenuation in the pipe. In order to do so, the AEM uses the highest energy possible that will still give acceptable attenuation results in the gas. A silver filter is often used for this purpose in normal operation of the AEM. For the pipe thickness measurement, however, it is more important that the attenuation in the gas can be canceled out for two subsequent transmission measurements, using the two-energy technique described previously. Because this is a one-time measurement to characterize the pipe before the enrichment measurements are performed, longer count times are acceptable, allowing lower energies to be used. Because it is feasible to go to (slightly) lower energies than the silver notch filter that has a K-edge at 25.5 keV, two filters can be chosen that have transmission peaks at equal attenuation in the UF$_6$ gas but different in the aluminum pipe. This
technique takes advantage of the fact that in the uranium L-edge region there are multiple energies with equal attenuation coefficients.

Table 2–1 shows a number of options for x-ray transmission notch filter materials compared with two traditionally used radioisotopes. This table compares some of the data previously presented [12] with an additional material, molybdenum. The table includes K-edge energies of the various materials and attenuations in the 5-mm wall thickness aluminum pipe. Also shown are attenuations in 10 cm of UF₆ gas at 50 Torr (typical of a downstream pipe header, where the AEM will be placed) and at 5 Torr (typical of an upstream header before a pump). The K-edge of zirconium (18 keV) was also considered because of its similar attenuation in UF₆ to using a ruthenium notch filter, but the attenuation in the aluminum pipe would have been unfeasibly large. It is important to note the similar attenuation of molybdenum and palladium in the UF₆ gas, both at 5 and 50 Torr. Because of these properties, molybdenum and palladium were selected as the two notch filter target materials for the pipe thickness determination. All further work on the two-energy pipe thickness method focuses on these two materials.

A silver filter was being considered for the final unattended transmission measurements used to determine the enrichment [25]. Another possibility could have been to use a palladium filter for the enrichment transmission measurement as well as one of the two filters for the pipe attenuation determination. With less than a 1-keV difference in the silver and palladium K-edges, this would have increased the absorption in the gas by 5% at 50 Torr, but would have almost doubled the count time because of the higher attenuation in the aluminum pipe. This may or may not be an acceptable compromise.
### Table 2-1: Notch Filter Selection Data

<table>
<thead>
<tr>
<th>X-ray Notch Filter</th>
<th>Isotopic Source</th>
<th>X-ray Notch Filter</th>
<th>Isotopic Source</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mo</td>
<td>Ru</td>
<td>Cd-109</td>
</tr>
<tr>
<td>Peak Energy (keV)</td>
<td>20.00</td>
<td>22.10</td>
<td>22.16</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>10.28</td>
<td>12.44</td>
<td>N/A</td>
</tr>
<tr>
<td>Attenuation Factor in Al</td>
<td>12,322</td>
<td>1086</td>
<td>1029</td>
</tr>
<tr>
<td>Attenuation Factor in UF6 at 50 Torr</td>
<td>1.63</td>
<td>1.86</td>
<td>1.86</td>
</tr>
<tr>
<td>Attenuation Factor in UF6 at 5 Torr</td>
<td>1.05</td>
<td>1.06</td>
<td>1.06</td>
</tr>
</tbody>
</table>

#### 2.2 Attenuation of X-rays in Aluminum and Uranium Hexafluoride

Figure 2.2 demonstrates that the attenuation in an aluminum pipe drops much more steeply as a function of energy than the attenuation in UF₆ gas. Because the K-edges of molybdenum and palladium are located in the L-edge region of uranium, transmission spectra using these notch filters will have equal attenuation in the UF₆ gas. Because of this large difference in attenuation in the pipe, an error in the pipe thickness has a very large effect on the calculated enrichment. Attenuation coefficients, from the National Institute of Standards and Technology (NIST) XCOM database, are plotted in Appendix A.
2.3 Calibration Method for Determining Pipe Thickness

The following analytical formula is used to determine the energy dependence of the transmitted spectra:

\[
I_0(E) = k \left( \frac{E_c}{E} - 1 \right)^n \exp[-\mu_{\text{Mo}}(E) \cdot \rho_{\text{filter}} \cdot d_{\text{filter}}] \cdot \exp[-\mu_{\text{Al}}(E) \cdot \rho_{\text{Al}} \cdot d_{\text{Al}}] \cdot \exp[-\mu_{\text{UF}_6}(E) \cdot \rho_{\text{UF}_6} \cdot d_{\text{UF}_6}],
\]

where \( k \) is a scaling constant, \( E_c \) is the cutoff energy (determined by the high voltage), \( n \) is an empirical coefficient depending on the anode material, and \( \mu(E), \rho, \) and \( d \) are the
mass attenuation coefficient, density, and thickness, respectively. This equation includes the energy-dependent bremsstrahlung yield of the x-ray tube [26] (shown in Fig. 2.3 for varying cutoff energies) multiplied by the exponential attenuation in the notch filters, as well as the attenuation of the aluminum pipe and the UF₆ gas. The three exponential terms apply to the filter, the aluminum pipe, and the UF₆ gas. Finally, \( d_{UF₆} \) is the equivalent thickness for UF₆ pressures in the pipe. Attenuation by the flux monitor diode is considered negligible compared to the rest of the system. Because the ratio of two transmission measurements is used to determine pipe thickness, any small amount of attenuation by the flux monitor would cancel out since it is used in both measurements.

Figure 2-3: Calculated energy-dependent bremsstrahlung yield of the x-ray tube for varying cutoff energies.

![Graph showing calculated energy-dependent bremsstrahlung yield for different x-ray tube voltages.]

Figure 2.4 illustrates the spectra calculated by Eqn. (3) on a normalized scale. The ratio of these transmitted spectra is used to determine the pipe-wall thickness, since the gas attenuation factors cancel out. This is because the AEM operates in the L-edge region of uranium. The average energy of the peaks can be adjusted by varying the target
thicknesses. Choosing a thicker filter causes the transmission peak to be narrower because more of the low-energy portion of the peak is attenuated.

![Figure 2-4: Transmitted spectra as calculated from Eqn. (3), with molybdenum and palladium notch filter thicknesses of 0.05 cm.](image)

Looking at the ratio of the two transmitted spectra (one with each filter), the gas attenuation factors cancel each other out. Since the filter thicknesses and the operating voltage of the x-ray tube are chosen by the user, the attenuation in the aluminum can be solved for, thereby determining the effective thickness of the pipe. To perform this calculation, a preselected ROI for each spectrum is used. Figure 2.5 demonstrates two examples of ratios of these ROIs as a function of gas pressure.
Figure 2-5: Ratios of the transmitted spectra as a function of UF₆ gas pressure. The aluminum pipe wall was 5 mm thick. Since the "0.5/0.5" curve has a positive slope and the "0.3/0.3" curve has a negative slope, there will be a combination of notch filter thicknesses that would have a slope of zero for the measured ratios vs. pressure, indicating no sensitivity to gas pressure. The factor of 50 to 60 is mostly from the difference in attenuation in the aluminum pipe at the two transmission energies.

A slope of zero on Fig. 2.5 would indicate a combination of molybdenum and palladium target thicknesses that would generate results independent of the UF₆ gas pressure. Because the lines for the two thicknesses used have opposite slopes, there is an optimum thickness somewhere in-between the two shown. From these calculations, it was determined that filter thicknesses between 0.3 and 0.5 mm of each material would be tested. For implementation in a facility, it is necessary to determine the range of target thicknesses that would yield an acceptably low sensitivity to a change in the pressure of the gas. Once the correct notch filters are selected, a user could go into a facility “blind”
and calibrate the AEM system on an operational system, without knowing the amount of UF₆ gas in the pipe.

### 2.4 Addressing Potential X-ray Tube Instability

An International Radiation Detectors Inc. (IRD Inc.) silicon flux monitor diode [27] was explored for the measurement of the output of an x-ray tube used for active transmission measurements on the pipe containing UF₆ gas. The measured flux in the diode can be used to correct for any instabilities in the x-ray tube or the high voltage power supply. Temperature sensitivity and radiation hardness tests were performed to determine the suitability of these diodes for use in the active implementation of the AEM. Although these diodes have been extensively tested for radiation hardness in the ultraviolet range, the enrichment monitor is operated in the 10- to 40-keV x-ray region. Radiation hardness testing over this energy range was performed using the energy spectrum that would pass through the diode during normal operation. Figure 2.6 shows the experimental setup used for this testing. The inset on the upper right is a picture of the diode. Temperature sensitivity measurements were also performed with the diodes.
Figure 2-6: Part of the experimental setup for the diode test, showing the x-ray tube and an inset (upper right) of the diode. The x-rays are directed into a steel "collimation" plate with the diode at the far side. The shielded cable connects to the Keithley picoammeter for the readout of the diode current.
3 Experimental Implementation

The general experimental setup that is described in the following sections was used as the basis for both the analytical calculations detailed in Chapter 4 and the experiments performed to test the active AEM. A diagram of the passive AEM is also included here to help distinguish the two systems.

3.1 Introduction

Figure 3.1 is a drawing of the active AEM, showing the x-ray tube and power supply, notch filter, collimator, in-beam flux monitor (to correct for instabilities in the x-ray output), pipe, and lanthanum bromide (LaBr$_3$) detector with an MCA. The wall thickness tests were performed with a LaBr$_3$ detector, and the temperature and gas pressure sensitivity tests were performed using a NaI detector.
Figure 3-1: Active AEM setup with LaBr₃ detector. This is a typical experimental setup for the active system and is also the basis for analytical calculations described in Chapter 4.

The passive AEM, as installed in Capenhurst, does not use a transmission measurement to determine the total amount of uranium. Rather, it uses a facility gas pressure reading, as shown in Fig. 3.2. The detectors are positioned in a “face to face” configuration, each with its own MCA. This gives the operator both a redundant system, in case of one detector or MCA failure, plus a way to check the system health by comparing one side to the other.
Figure 3-2: Passive AEM setup with NaI detectors. This is the way the passive system is currently configured, collecting data in Capenhurst. A pressure reading gives the total amount of uranium in the UF₆ gas, and 186-keV counts give the amount of $^{235}$U.

### 3.2 X-ray Tube and Collimator

The x-ray generator used in the AEM is a Varian model VF-50J industrial tube (see Appendix B for specifications) [28], with either a tungsten, silver, or palladium anode. Of these, the tungsten anode tube is the most efficient. A greater amount of incident radiation is converted to x-rays, rather than heat, due to its higher atomic number [29]. The tube has a beryllium window and is powered by an XRM series Spellman high voltage supply (described in Appendix B [30]). The operating current of the tube is about 150 $\mu$A, and the operating voltage is in the range of 35 kV. This x-ray tube was chosen because it is capable of operating at beam currents much higher than needed for those used in the AEM, suggesting a long operational lifetime. The x-ray tube is embedded in the active measurement head (a steel fixture that gets mounted to the tungsten box around the pipe), as shown in Fig. 3.3. An interlock switch, also shown in the figure of the active
head, is used to guarantee that the tube cannot operate if it is not affixed to the pipe. This picture shows a small fan mounted on the back of the x-ray tube for cooling. Figure 3.4 shows the underside of the active measurement head. Two dowel pins ensure that the system is mounted on the pipe in the same orientation each time a filter is changed. A temperature sensor is also visible in the background, attached to the body of the x-ray tube. Additional temperature sensors can be monitored with the data collection software.

Figure 3-3: The x-ray tube mounted in the active measurement head, protected by an interlock switch. Also shown is a fan used for cooling, mounted above the x-ray tube.
Figure 3-4: Underside of the active head showing the flux monitor diode mounted in place as well as the dowel pins that are used to fix the orientation of the x-ray tube when notch filters are changed.

3.3 **Notch Filters**

Temperature sensitivity tests were performed using a silver notch filter, which gives a transmission peak with the maximum energy at 25.5 keV. This filter was affixed to the active head using an aluminum ring, placed directly in front of the x-ray tube. The molybdenum and palladium notch filters for the pipe thickness experiment were fabricated by cutting the 0.1-mm-thick sheets into 1.25” squares and attaching these squares to aluminum rings that could be mounted directly above the flux monitor. This is shown in Fig. 3.5. Filter thicknesses of 0.3 mm, 0.4 mm, and 0.5 mm were used for each material; these were fabricated by stacking the required number of thicknesses of each material on the aluminum rings.
Figure 3-5: Some of the molybdenum and palladium notch filters attached to their mounting rings.

The flux monitor and notch filter are mounted on the opposite side of the steel fixture from the x-ray tube, as seen in Fig. 3.6. A hole collimator through the middle ensures all of the beam hits the target and is directed through the flux monitor. The positioning of the filter on the side of the collimator farthest from the x-ray tube allows changing of the filters without affecting the position of the x-ray tube. This stability allows spectra to be compared directly, without worrying about any geometrical errors that may have been introduced when removing and reattaching the x-ray tube.
3.4 Flux Monitor

The beam intensity of the x-ray tube is monitored with a silicon junction p-n photodiode, model AXUV100GX, developed by IRD Inc. [31] Photodiode specifications are given in Appendix B. This diode has a silicon thickness of 104 microns and a thin (3 to 7 nm) silicon dioxide junction with a passivating, protective entrance window. A polyether ether ketone (PEEK) insulator, shown in Fig. 3.7, was machined to hold the diode in place and screw into the active head through the notch filter holder. The collimated beam passes through the notch filter, and all of the resulting radiation then passes through the diode. The diode is operated in the photovoltaic mode (with no bias voltage), and current is measured with a Keithley picoammeter [32]. This current is also read by the AEM software and can be used to correct for any instabilities (or temperature changes) with postprocessing of the data.
Figure 3-7: Flux monitor diode in its PEEK holder. A plus sign was drawn on one side of the PEEK material to ensure the diode would be remounted correctly in case it had to be removed between tests.

3.5 \textit{UF}_6 \textbf{Source}

A number of sealed, gaseous UF$_6$ laboratory sources were built for experimental testing. These sources were fabricated, evacuated, leak tested, and passivated by introducing a small amount of depleted UF$_6$ gas into the pipe and then evacuating it. This was performed by the Materials Science Group, MST-6, at LANL.

Figure 3.8 shows a large UF$_6$ source that has a pipe with three calibrated wall thicknesses. This is the source that the wall-thickness measurements were performed on. A standard 6061 aluminum pipe was used to make this source, with the outer diameter machined down to the three thicknesses shown in Table 3–1. The thicknesses at each step were averaged for use in analytical calculations. A close-up of the source, showing the three steps, is shown in Fig. 3.9. Precise thickness measurements were performed on the pipe at two locations for each step before it was attached to the rest of the system. These
data give us an idea of the error in the pipe thickness, which our density measurements are very sensitive to.

This source allows for variable UF₆ gas pressure (up to ~50 Torr), as well as different enrichment levels. The manifold on the left end of the source has four connections for bottles of UF₆. To increase gas pressure in the pipe, the valve to the bottle of the desired enrichment is opened. A high precision Baratron pressure gauge is used to monitor the pressure in the pipe, and the valve is closed when the desired pressure is reached. To lower gas pressure in the pipe, liquid nitrogen (LN₂) cooling is used. A small vessel, filled with LN₂, is placed around the bottle that the UF₆ is returned to. The valves are then opened and the gas is cryogenically pumped back into the bottle.
Figure 3-8: Large horizontal UF₆ source with three pipe thicknesses, variable gas pressure, and variable enrichment. The active system is mounted on the pipe with the x-ray tube on the top pointing downwards; the LaBr₃ detector is mounted on the bottom in its tungsten composite shielding.

<table>
<thead>
<tr>
<th>Table 3-1: Precise Thickness Measurements for Horizontal Source</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Step 1</strong></td>
</tr>
<tr>
<td>Point 1</td>
</tr>
<tr>
<td>Point 2</td>
</tr>
<tr>
<td><strong>Step 2</strong></td>
</tr>
<tr>
<td>Point 1</td>
</tr>
<tr>
<td>Point 2</td>
</tr>
<tr>
<td><strong>Step 3</strong></td>
</tr>
<tr>
<td>Point 1</td>
</tr>
<tr>
<td>Point 2</td>
</tr>
</tbody>
</table>
Figure 3-9: Horizontal UF$_6$ source showing the three pipe thickness “steps.” The thicknesses of the three steps are given in Table 3–1.

Another UF$_6$ source used is a smaller vertical source that fits into an environmental chamber. This setup is shown in Fig. 3.10. This source only has a connection for one bottle of UF$_6$; 3.3% enriched is currently attached and was used for the temperature/pressure sensitivity measurements. An aluminum heat sink designed to clamp directly onto the x-ray tube is pictured.
Figure 3-10: Vertical UF$_6$ source in the environmental chamber. This source has an enrichment of 3.3%. The active AEM used for temperature and pressure sensitivity tests is mounted on the pipe. A heat sink around the x-ray tube is visible.

The environmental chamber is typically run at temperatures between 15°C and 45°C when performing tests with UF$_6$. These measurements are discussed in more detail in Section 4.2, “UF$_6$ Properties.” The typical temperature profile used, with 8-hour hold times at each temperature, is shown in Fig. 3.11. Temperature ramp rates are held at less than 0.5°C per hour, and the system is held at each temperature for 8 hours to ensure data are collected once the system has been allowed to reach equilibrium.
Figure 3-11: Typical temperature profile used for temperature sensitivity studies. Temperature ramp rates are kept less than 0.5°C per hour, and the system is held at each temperature for 8 hours to ensure data are collected once the system has been allowed to reach equilibrium.

### 3.6 Detectors/MCAs

For the two-energy pipe thickness measurements, a planar ½” thick by 3” diameter LaBr$_3$ spectrometer was used. This detector, because of its shorter time constant [33], is capable of handling the higher dead times anticipated when performing attenuation measurements comparing spectra with molybdenum and palladium filters. The data were acquired with a Canberra Lynx digital signal analyzer.

In order to accurately compare spectra taken with different notch filters, the same x-ray tube high voltage and beam current were used with both filters. Because of the low count rates seen in the transmission peak with the molybdenum filters, tube settings were optimized to achieve a reasonable count rate. Therefore, somewhat higher count rates were seen with the palladium filter using the same settings. The LaBr$_3$ detector was able to handle all observed count rates. Spectra were collected for 30, 60, or 90 minutes in order to get sufficient counts in the transmission peak net areas.
For the temperature and pressure sensitivity measurements, NaI(Tl) detectors were used. These are the most commonly used detectors in enrichment monitoring because of their low cost, good efficiency, and sufficient resolution for the task. Table 3–2 compares some of the significant characteristics of NaI and LaBr₃ detectors [34].

**Table 3-2: Comparison between LaBr₃ and NaI Scintillators**

<table>
<thead>
<tr>
<th></th>
<th>Density (g/cm³)</th>
<th>Resolution (%) 122 keV</th>
<th>Resolution (%) 662 keV</th>
<th>Light yield (photons/keV)</th>
<th>Decay time (ns)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaI:Tl</td>
<td>3.67</td>
<td>8.0</td>
<td>6.5</td>
<td>39</td>
<td>16</td>
</tr>
<tr>
<td>LaBr₃:Ce</td>
<td>5.1</td>
<td>6.0</td>
<td>2.8</td>
<td>65</td>
<td>250</td>
</tr>
</tbody>
</table>

An ORTEC DigiDART MCA was used with the NaI detector for the temperature and gas pressure sensitivity measurements [35].
4 Analytical Modeling and Calculations

Analytical modeling and calculations were completed before any experimental measurements were performed. Typically these consisted of one-dimensional linear attenuation calculations performed in Excel, using slab geometries for the materials. The formulas used for these calculations are presented in this chapter, along with plots of the various calculated spectra. The UF₆ attenuation was calculated using 50 Torr of gas with a density of 0.001 g/cm³, and the aluminum attenuation was through 1 cm of pipe (both walls of a 0.5-cm pipe). Properties of the UF₆ gas were also explored to determine the range of temperatures in which to operate to keep the UF₆ in gas form. Calculations were performed to determine if wall deposits would affect the pipe thickness measurement. Finally, MCNPX calculations were performed to determine the dose to the flux monitor diode during the long-term radiation hardness test, described in Chapter 5.

4.1 X-ray Tube

Analytical calculations were performed to estimate the output of the x-ray tube using different operating parameters. This allowed the appropriate range of thicknesses to be selected for experimental testing of the notch filters and also helped determine the final settings for the x-ray tube. Calculations were compared with experimental results, as shown in Fig. 4.1. A 0.4-mm-thick palladium notch filter with a K-edge at 24.35 keV was used. The spectrum was collected with the AEM positioned on Step 2 of the pipe, which has a thickness of 0.5 cm.
4.1.1 Generated and Transmitted Spectra

The most noticeable difference between the measured and calculated spectra is the width of the peaks, which is affected by the detector resolution in the actual data. Also evident in the measured spectrum is the lanthanum x-ray peak that comes from internal radioactivity of the LaBr₃ detector, which does not show up in calculations that only display radiation incident on the detector.
35 kV and beam current of 150 nA for more than two years, with very few interruptions. These operational interruptions were not because of AEM or x-ray tube failure but were typically power outages or procedural shutdowns for safety reasons, such as for interlock checks, lasting days at a time.

X-ray tube voltage and beam current settings needed to be determined for the pipe thickness determination testing. The goal was to have a high intensity peak but not have too much bremsstrahlung above the peak. The intention was also to use the same settings with both the molybdenum and the palladium notch filters in order to directly compare the ratios of the transmitted spectra without having to correct for any differences. The first calculations were performed using a palladium notch filter to observe the effect of changing the tube voltage on the spectrum. The results of this calculation are shown in Fig. 4.2.

Figure 4-2: Effect of varying the x-ray tube cutoff voltage, with a fixed beam current. A palladium notch filter was used for these calculations.
The calculations showed that while increasing the cutoff voltage of the tube does increase the intensity of the peak, it does not change its width. However, there is a tradeoff because making the cutoff voltage too high allows more and more of the bremsstrahlung above the K-edge of the notch filters to leak through, as was shown in the above figure. For this reason, a cutoff voltage of 30 kV was chosen, and the beam current was increased from the typical setting of 100 µA to 160 µA. Increasing the beam current simply increases the flux from the tube without changing the shape of the spectrum.

### 4.1.2 Issues with Analytical Model

The following analytical formula was used to determine the transmitted spectra, as detailed in equation 3 from Section 2.3:

\[
I_0(E) = k \cdot \left( \frac{E_c}{E} - 1 \right)^n \cdot \exp[-\mu_{\text{fiber}}(E) \cdot \rho_{\text{fiber}} \cdot d_{\text{fiber}}] \cdot \exp[-\mu_{\text{Al}}(E) \cdot \rho_{\text{Al}} \cdot d_{\text{Al}}] \cdot \exp[-\mu_{\text{UF}_6}(E) \cdot \rho_{\text{UF}_6} \cdot d_{\text{UF}_6}]
\]

The empirical coefficient describing the anode material \( n \) was determined by extrapolation from the data in McCall’s paper (shown in Table 4–1), which came from Jakschik [26].

#### Table 4-1: X-ray Generator—Anode Material Coefficients

<table>
<thead>
<tr>
<th>Element</th>
<th>Z</th>
<th>( n )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beryllium</td>
<td>4</td>
<td>1.28</td>
</tr>
<tr>
<td>Aluminum</td>
<td>13</td>
<td>1.23</td>
</tr>
<tr>
<td>Copper</td>
<td>29</td>
<td>1.08</td>
</tr>
<tr>
<td>Silver</td>
<td>47</td>
<td>0.96</td>
</tr>
<tr>
<td>Gold</td>
<td>79</td>
<td>0.91</td>
</tr>
</tbody>
</table>
One type of x-ray tube had a silver anode. Two others had palladium and tungsten anodes, for which $n$ needed to be determined. The fit of the data is shown in Fig. 4.3, and from this the anode coefficients of palladium and tungsten, 0.97 and 0.9, respectively, can be determined.

![Graph showing the fit of data points from Table 4–1 used to calculate $n$, the empirical anode coefficient of the x-ray tubes.](image)

Figure 4-3: Fit of the data points from Table 4–1 that are used to calculate $n$, the empirical anode coefficient of the x-ray tubes.

Figure 4.4 shows the effect of different x-ray tube anode materials on the spectrum generated using a 0.1-mm silver notch filter. While a silver anode coefficient may have been a suitable substitute for calculations using a palladium anode, this figure shows that silver is not as good an approximation for a tungsten anode.
Figure 4-4: The effect of different x-ray tube anode materials on the spectrum generated using a 0.1-mm silver notch filter. This figure shows that although silver may be a good approximation for a palladium anode, it cannot substitute in calculations for a tungsten anode.

Also, note that $I(E)$ is the calculation of the beam that hits the detector, but it is not necessarily what the detector measures. Since the two transmission peaks generated using the notch filters are both below 25 keV, detector efficiency in either the LaBr$_3$ or the NaI should not cause any differences between the two measured spectra. However, the resolution of the detector will affect the shape of the spectrum measured. For this reason, calculated spectra appear to have a sharp drop in intensity at the K-edge of the notch filter used, whereas measured spectra are broadened from the detector resolution.
4.2 \textit{UF}_6\textit{ Properties}

A phase diagram of UF$_6$ is presented in Fig. 4.5 [36]. All of the temperature sensitivity testing was performed in the range shown in blue on the diagram. Since the operating gas pressures ranged from 0 to 50 Torr in our laboratory sealed UF$_6$ sources, the operating temperature range selected was between 15°C and 45°C. The lower temperature limit of 15°C kept the gas from freezing and solidifying in the pipe at higher pressures.

Figure 4-5: UF$_6$ phase diagram showing (in blue) the temperature and pressure operating range of our experiments. Tests were performed with UF$_6$ in the gas phase only. The only area that approached the freezing of the gas was at 15°C at 50 Torr, so the source temperature was always kept at or above 15°C.
4.3 **Effect of Wall Deposits on Measurements**

Calculations were also performed to determine whether there was a measureable effect of pipe-wall deposits on the aluminum pipe thickness results. A UF$_6$ source was built that was plated with thin deposits of uranyl fluoride (UO$_2$F$_2$) for testing a multidetector deposits characterization system [37]. The physical characteristics of the deposits were characterized by x-ray photoelectron spectroscopy and Rutherford backscattering spectrometry and were used as input for the following calculations.

First the attenuation of the deposits was calculated using the NIST photon cross section database [38]. The result was compared to the UF$_6$ attenuation, calculated from the same database. These data are plotted in Fig. 4.6. The UF$_6$ attenuation was plotted as a red dashed line to show that the attenuations are exactly equal in the energy range between 10 and 25 keV. This is because the uranium L-edge region dominates the attenuation in both materials: the deposits and the gas. Because of these material properties, the method of factoring out UF$_6$ gas attenuation at the two energies should also apply to the deposits.
Figure 4-6: Comparison of the attenuation of UO$_2$F$_2$ and UF$_6$. It appears that uranium dominates the attenuation factor at the energies in which the AEM operates. For this reason, the presence of wall deposits inside the pipe should not affect the wall thickness measurement, as any attenuation due to deposits will cancel out just as the gas attenuation does with the two-energy transmission method.

Next, calculations were performed to determine whether there would be measureable attenuation of an x-ray transmission measurement by the deposits. Using our laboratory source as a model (0.5-µm-thick deposit, density of 6.45 g/cm$^3$), the calculations show that there would only be about a 0.017% difference in the net peak area of a transmission measurement, both with and without the deposit (i.e., the deposits described above would only cause a 0.017% increase in the attenuation). This was done using a 0.4-mm palladium filter. Figure 4.7 shows the difference in attenuation between the two scenarios, with the ROI for the palladium peak shaded in blue.
The effect was even smaller with a molybdenum filter because those generated peaks had lower net count rates due to the higher pipe attenuation. Since the statistical uncertainty in the net area with the palladium filter was about 0.08% (for a 1,800-s measurement), there was no significant measurable attenuation by the deposits. Moreover, once the one-time thickness measurement had been completed, we switch to another filter with a higher transmission energy, such as silver, for the unattended enrichment monitoring. A transmission peak generated by a silver notch filter would be even less attenuated by uranium deposits than the palladium, since the peak would be at a higher energy (K-edge at 25.5 keV).
On the other hand, if there is a known enrichment for calibrating the system, such as from a mass spectrometer measurement, deposits in the pipe can be corrected for. Since the deposits should not change significantly over the monitoring time (as long as the pipe is not brand new when monitoring is begun), this correction should remain valid. Even if the deposits are a different enrichment than the UF₆ gas, the two-energy pipe thickness determination method should work because attenuation by the uranium is what cancels, and it is independent of enrichment. The intent of the AEM is not to quantify the deposits but rather to determine the portion of the 186-keV count rate that comes from deposits (of any enrichment) to enable a correct enrichment determination over time.

### 4.4 Flux Monitor Diode

The following analytical formula is used to describe the energy spectrum seen by the flux monitor diode:

\[
I = k \cdot \left( \frac{E_c}{E} - 1 \right)^n \exp \left[ -\mu_{Ag}(E) \cdot \rho_{Ag} \cdot d_{Ag} \right].
\]  

(4)

The intensity of the energy spectrum I is the bremsstrahlung yield of the tube [26] multiplied by the attenuation in the silver notch filter. The terms \( \mu_{Ag}, \rho_{Ag}, \) and \( d_{Ag} \) are the mass attenuation coefficient, density, and thickness of the silver notch filter, respectively. The cutoff voltage of the tube, \( E_c \), was set to 40 kV. This is a typical voltage setting that could be used in operation in a GCEP. The beam current can then be increased to raise the flux at the diode to simulate very long operating times. Figure 4.8 shows the calculated spectrum that is generated by the silver attenuated x-ray tube beam. A 0.1-mm-thick silver notch filter was used for the hardness testing to reduce the
attenuation from that of the typical 0.5-mm-thick silver filter, in order to have as much
dose to the diode as possible. Although the 0.1-mm-thick silver notch filter used is
thinner than what would be used in normal operation by a factor of between 2 and 5, it
still gives us the typical transmission spectrum to which the diode would be subjected—
just a slightly broader peak.

![Normalized Spectrum](image)

Figure 4-8: Calculated spectrum generated by an x-ray tube with a
0.1-mm-thick silver notch filter.

### 4.4.1 Diode Responsivity

The theoretical responsivity of the flux monitor diode, as a function of x-ray
energy for the 104-micron silicon thickness, was used to determine the absorbed dose in
the diode over the long-term irradiation. Responsivity is defined as a measure of the
amount of output current produced by the diode for an incident radiant power. The IRD
Inc. website ([www.ird-inc.com](http://www.ird-inc.com)) compares experimental responsivity results for a
calibrated diode to this theoretical formula, which shows excellent agreement [31]. The
responsivity curve shown in Fig. 4.9 was used to determine the photon flux at the detector.

![Diagram of flux monitor diode responsivity as a function of energy.](image)

Figure 4-9: Flux monitor diode responsivity as a function of energy. Responsivity data taken from the IRD website [30]. This plot was used to calculate the energy-dependent absorbed dose to the diode over the long-term irradiation test.

Equation 4 relates the responsivity to the photon flux:

$$ R = \frac{I_{ex}}{\phi \cdot E}, $$

where $R$ is the responsivity in amperes per watt, $I_{ex}$ is the average measured current in the diode during irradiation, $\phi$ is the flux in photons/second (assumed to be constant here), and $E$ is the approximate irradiation energy in keV. If we use an average energy of 22.3 keV for $E$ (the actual spectrum is discussed in Section 4.3), then Fig. 4.9 gives the
responsivity. $I_{ex}$ was measured in our experiment. By substituting these values and solving for $\phi$, a flux of $\sim1.4 \times 10^{10}$ photons/s was calculated.

### 4.4.2 Measurement Dose Rate

The flux determined above, $1.4 \times 10^{10}$ photons/s, was used to calculate a dose rate and total dose. The absorbed dose in the diode during the long-term (30 day) irradiation was evaluated to be about 40 rad, or 55 mrad/hr. The dose and dose rate calculations were performed using the MCNPX code [39], using an F6 (energy deposition) tally with a beam of photons directed at the silicon diode. Figure 4.10 is a plot of the input energies of this beam of photons (used to approximate the spectrum shown in Fig. 4.8), and the entire MCNPX input file is given in Appendix C.

![Figure 4-10: Energies used in the MCNPX input file to calculate dose to the diode. This is an approximation of the spectrum shown in Fig. 4.8.](image-url)
The spectrum counts are in arbitrary units; MCNPX normalizes the input intensities before running. These MCNPX calculations, using the energy responsivity curve provided by the diode manufacturer, determined that the absorbed dose in the diode during the long-term irradiation was about 40 rad, or 55 mrad/hr. Section 5.1, “Flux Monitor Diode Measurements,” will show the results of the long-term diode test, demonstrating that the diode continued to operate in the desired manner after receiving a dose of 40 rad.
5 Experimental Measurements

Experimental measurements form the basis of this dissertation, as the purpose is to operate the AEM in the field as accurately as possible. First, radiation hardness measurements were performed on the flux monitor diode to test whether it could withstand the long-term x-ray radiation used in the AEM. Next the dual-energy method was evaluated. These dual-energy measurements comprised the largest portion of the experimental work in this dissertation, as close to 100 individual transmission measurements were performed to optimize the notch filter ratio and x-ray tube settings. Temperature and pressure sensitivity tests were performed with both the active and passive AEM systems. Finally, some details of initial measurements from the URENCO Capenhurst field trial have been presented.

5.1 Flux Monitor Diode Measurements

A long-term measurement was performed at a high flux, simulating more than 80 years of operation. No significant degradation was seen during this time. Fluctuations were found to be within the 0.1% operationally acceptable error range. After irradiation, an I-V characterization showed a temporary irradiation effect that decayed over time. This effect was small because the diode was operated without external bias.

5.1.1 Long-term Irradiation Measurement

The diode was irradiated at a tube voltage of 40 kV and a beam current of 1 mA. With the 0.1-mm-thick silver filter in place, an average current of 3,491 nA was induced in the diode. The average diode current was measured with a Keithley picoammeter,
Model 6485. A readout from the board of the x-ray tube high voltage power supply provided temperature data. Figure 5.1 shows the temperature-corrected diode current over the 30-day period of irradiation. A temperature correction factor (change in diode output current per degree centigrade) had been obtained previously for the diode being characterized. This factor was obtained by placing the diode in an environmental chamber and using a temperature profile with a range of temperatures between 5°C and 45°C, similar to the one shown in Figure 3.11. Each temperature was maintained for at least one hour to allow the diode to reach equilibrium, and a slow ramp rate of between 4 and 5 degrees per hour was used. High voltage also is plotted to demonstrate that most of the major fluctuations in the temperature-corrected data can be attributed to changes in the high voltage, which is also affected by temperature.

The narrow drops in the current are unexplained instrumental artifacts; however, even with these, the temperature-corrected current falls within the acceptable 0.1% error range. This operation of the diode at a flux of 1,000 times that of normal operation for 30 days is equivalent to approximately 80 years of normal operation.
Figure 5-1: Long-term irradiation test of the flux monitor diode at 40 kV and 1 mA. Temperature-corrected diode data are shown, plus high voltage, to help explain the remaining fluctuations.

5.1.2 I-V Characterization, Diode Recovery

An I-V characterization of the diode was performed following the long-term irradiation. Figure 5.2 shows the measured curves over a 13-day period. The diode was held at 25°C for the first 7 days following irradiation, and the decay of charge trapping was recorded. At this point the diode was heated to 45°C. The final two measurements shown in Fig. 5.3 were taken with the diode back at 25°C. Additional measurements showed no significant change with time.
Figure 5-2: I-V characterization of the flux monitor diode after irradiation. The first measurement (blue diamonds—greatest effect) was taken 30 minutes after stopping the irradiation. The last (orange x’s—lowest effect) was taken 311 hours after irradiation ceased. Notice the decay of charge trapping over time, after irradiation was stopped.

Prolonged ionizing radiation can cause charge trapping (ionizing damage) in the oxide layer of the diode and at the oxide/silicon interface [40]. After the diode was irradiated, a fast (short-term) recovery was observed, followed by a slow (long-term) recovery. This recovery is shown in Fig. 5.3 for three different voltages on the I-V curve. By increasing the temperature, it is possible to free the charge stored in the oxide. Little sensitivity to this effect is expected during actual operation because the diodes in the AEM are operated without external bias (very close to 0 V).
Figure 5-3: Observed recovery of the flux monitor diode after the long-term irradiation test for three voltages on the I-V curve shown in Fig. 5.2. Short- and long-term recovery of the diode from charge trapping is shown.

No permanent damage to the diode appears to have been done by any of the long-term measurements. Small fluctuations of the measured current were seen, but these can be explained by temperature changes in the room and fluctuations in the x-ray tube high voltage. After an equivalent of 80 years of standard operation, the diode still continued to function as needed for use in the active AEM.

No permanent damage done to the silicon diode was seen when testing with x-ray energies that are useful for performing transmission-based enrichment measurements in a GCEP. It appears that a temporary charge-trapping effect may exist that was caused by irradiation of the diodes with x-rays in the 10- to 40-keV region. However, the diode received an absorbed dose of about 40 rad with no observed degradation in its operation. Therefore, these flux monitors will be useful tools for stabilizing x-ray tubes during the long durations of possibly unattended monitoring in a facility.
5.2 *Dual-Energy X-Ray Measurements*

The following sections describe the dual-energy transmission measurements that were performed. The UF$_6$ calibration source that was built for the laboratory testing of this method is detailed, and some of the generated spectra are presented.

5.2.1 *UF$_6$ Calibration Source*

A horizontal source (Fig. 5.4) incorporating a new UF$_6$ manifold with thicker piping was rebuilt for laboratory testing. This $\frac{1}{2}$” piping makes it easier to pump down the source with liquid nitrogen to lower the gas pressure in the pipe. With the previous $\frac{1}{4}$” piping, it was necessary to heat the manifold while pumping down so that UF$_6$ wouldn’t freeze in the small openings and block the flow. The source currently has two bottles of UF$_6$ attached: depleted uranium and 4.5% enriched. While this source was being refurbished, an older tungsten mounting fixture was redesigned (Fig. 5.5) to attach to the pipe with the three different thicknesses. To do so, the bottom half was made with a circular curvature slightly larger than the largest pipe to be measured. The top half was designed with 45 degree angles, ensuring three-point contact on all pipe thicknesses.
Figure 5-4: Refurbished horizontal source. Shown are the new ½” diameter pipes and the two bottles of UF₆ (foreground, bottom). The bottle on the right is empty.
5.2.2 Transmission Measurements

A number of measurements were performed to test the technique for determining attenuation, including the following variables:

- Pipe thickness
- Notch filter thickness
- UF₆ gas pressure

While the source was disassembled it was cleaned, and precise wall-thickness measurements were performed. Thicknesses were determined at two points on each step of the pipe. The measurements were performed at the ends of each step. The data, shown in Table 3–1, give us a better idea of the error in the pipe thickness, which our density measurements are very sensitive to.
More than 90 measurements were taken to acquire the spectra needed to optimize the combination of molybdenum and palladium notch filters. Both molybdenum and palladium filters of 0.3-mm, 0.4-mm, and 0.5-mm thicknesses were used. First, empty pipe measurements were taken and spectra were collected with each of the six filters on each step of the pipe. The UF₆ (4.5% enriched) gas pressure was then increased in steps up to 50 Torr, which is the limit of the UF₆ contained in this closed system. Spectra were collected for either 1,800 or 3,600 seconds, depending on the count rate in the transmission peak. The transmission of the peak generated by the molybdenum notch filter was much lower than the peak with palladium because there was much higher attenuation in the aluminum pipe for the 20-keV molybdenum peak than for the 24.3-keV palladium peak.

### 5.2.3 Gamma-ray Spectra

A ½” thick by 3” diameter LaBr₃ spectrometer was used for the pipe thickness measurements. This detector is capable of handling the higher dead times that were anticipated when performing attenuation measurements comparing spectra with molybdenum and palladium filters. The MCA was a Canberra Lynx digital signal analyzer, to further help with the possible high dead times that might have been encountered. A screen shot of Canberra’s Lynx data acquisition interface is shown in Fig. 5.6.
Figure 5-6: Screen shot of the Lynx software showing a spectrum with UF₆ gas in the pipe and using a molybdenum notch filter. The preset time of this spectrum is visible at 3,600 s.

To accurately compare spectra taken with different notch filters, it was a requirement to use the same x-ray tube high voltage and beam current. Because of the low count rates seen in the transmission peak when using the molybdenum filters, the tube settings for molybdenum were optimized without adversely affecting the palladium spectra. The effect on the spectrum of changing the x-ray tube high voltage was examined. Data were taken with the high voltage at both 30 kV and 28 kV and a beam current of 100 µA. As long as the cutoff voltage of the tube was kept above the K-edge of the notch filter, the width of the peak was not affected.

Figure 5.7 shows spectra taken with a 0.3-mm molybdenum notch filter. PeakEasy software [41], which is capable of reading in many different file formats, was used to manipulate and plot all spectral data. Shown is the 20-keV peak from the molybdenum filter as well as the x-rays between 32 and 37 keV that are from the $^{138}$La component of the LaBr₃ detector. In the figure, the bremsstrahlung evident in the black
spectrum (30 kV, 100 µA) is cut off in the blue spectrum that was taken at 25 kV and 160 µA. The peak generated by the molybdenum filter is of much lower intensity than the one observed when palladium is used. This is due to attenuation in the aluminum pipe, as well as the self-attenuation of the molybdenum. The molybdenum has a K-edge at a low energy of 20 keV; thus the generated peak is easily attenuated. In order to compensate for this lower count rate, the beam current of the x-ray tube and cutoff voltage was optimized while staying within the operating power of the tube.

Figure 5-7: Spectra generated with a molybdenum notch filter, varying the x-ray tube cutoff voltage and beam current. Higher cutoff voltage allows bremsstrahlung above the transmission peak through, whereas higher beam current gives a more intense transmission peak.

The maximum operating power of the VF-50 x-ray tube is 50 watts. During normal AEM operation, the power was kept well below this level, but that was for long-
term measurements. Measurement times for the pipe-wall thickness experiment are either 30 or 60 minutes, depending on the count rate in the transmission peak. The final x-ray tube settings selected for the measurements were 30 kV and 160 $\mu$A. Thus the molybdenum peak was measurable (good enough statistics to analyze with a 3,600-s count time) in most cases, and the palladium peak still had a reasonable dead time—less than a 3% maximum—which the LaBr$_3$ detector and Lynx system are easily able to handle.

A few selected spectra are shown below. First, in Figs. 5.8 and 5.9, spectra taken with a 0.3-mm-thick palladium notch filter are compared. In those figures, the black line was taken with 40 Torr of UF$_6$ gas in the pipe, and the blue was an empty pipe measurement. Figure 5.8 has ROIs highlighted in red, showing the transmission peak at 24.3 keV and the 185.7-keV $^{235}$U peak. A log scale was used to make all peaks visible. Figure 5.9 is a close-up of the transmission peak generated with the palladium notch filter, showing the effect of attenuation by 40 Torr of UF$_6$. A linear scale was used to show the difference in spectra.
Figure 5-8: Spectra taken with a 0.3-mm-thick palladium notch filter. Black: 40 Torr of UF₆. Blue: empty pipe. ROIs show the transmission peak at 24.3 keV and the 185.7-keV $^{235}$U peak. Spectra are plotted on a log scale.
Figure 5-9: Close-up of the palladium transmission peak. Black: transmission through 40 Torr of UF₆. Blue: transmission through the empty pipe. Data are displayed on a linear scale to emphasize the difference in peak areas.

Figure 5.10 shows a spectrum taken using a 0.3-mm molybdenum notch filter with 40 Torr of UF₆ in the pipe. ROIs (in red) show the molybdenum transmission peak at 20 keV and the 185.7-keV $^{235}\text{U}$ peak. In this spectrum there is bremsstrahlung visible between the molybdenum peak and the $^{138}\text{La}$ x-ray peak. This is because a 30-kV cutoff voltage on the x-ray tube is used.
Figure 5-10: Spectrum taken with a 0.3-mm molybdenum notch filter. Red ROIs show the transmission peak at 20.0 keV and the 185.7-keV $^{235}\text{U}$ peak. A linear scale was used.

5.3 Sensitivity to Changes in Pressure and Temperature during Measurement

Sensitivity of the AEM to changes in gas pressure and temperature are explored in this section. Two methods were tested for correcting changes in the output of the x-ray tube due to variations in the ambient and system temperature, both with and without the flux monitor.

5.3.1 Pressure

The system was calibrated by first removing all gas from the sealed pipe for the UF$_6$ source and then measuring the intensity $I_0$ of the transmitted x-ray beam unattenuated by any gas (averaged over eight days) and the background $B$ under the
186-keV peak from $^{235}$U. The method for extracting $B$ and $I_0$ from data is explained later in this section. Next, the pipe was filled with 3.3% enriched UF$_6$ to a pressure of 60 Torr to determine the transmitted intensity $I$ with gas and the 186-keV count $R$ averaged over five days. By knowing these values, the calibration constant $K$ can be determined in the following formula:

$$E(t) = K \times \frac{R(t) - B}{\ln \left( \frac{I}{I_0} \right)}$$

(6)

After calibration, this formula gives the enrichment for subsequent measurements. The stability of the system with 60 Torr of UF$_6$ gas in the pipe was examined. The stability of the system without gas in the pipe was also investigated. A comparison of two example spectra, one with 60 Torr of gas in the pipe and one after the pipe was pumped down, is shown in Fig. 5.11. There is no measurable net count rate in the ROI set for measuring $^{235}$U around 186 keV for the empty pipe. The 186-keV count rate, corrected for background or deposits, is $R(t) - B$ in Eqn. 6, the piece of the equation that determines the portion of the uranium that is $^{235}$U. The absence of a 186-keV peak in the blue (empty pipe) spectrum verifies that all of the gas has been removed from the pipe and there is no significant deposit of $^{235}$U. In the empty pipe case, both $R(t)$ and $B$ are zero. In a GCEP deposits in the pipe, or other $^{235}$U nearby such as cylinders being moved around, would contribute 186 keV counts to $B$. In this case, however, $B$ was determined to be zero by performing the empty pipe measurement, and is used for all subsequent enrichment calculations for this experiment. There is also a noticeable difference in the palladium transmission peak at 24.35 keV, with a higher intensity peak for the blue (empty pipe).
spectrum than for the black one with gas in the pipe. This affects the denominator of Eqn. 6, which determines the total amount of UF₆ gas.

![Figure 5-11: Spectra with UF₆ gas (black) and without UF₆ gas (blue) in the pipe. The red coloring shows the ROI for the 186-keV peak.](image)

The accuracy of the enrichment determination with varying pressures of UF₆ gas was investigated. The gas pressure in the source within the environmental chamber was stepped down from 60 to 20 Torr, 10 Torr at a time. Each pressure was held for a day, allowing multiple spectra to be collected at each step. The transmission count rates, using a silver notch filter with a 25.5-keV transmission peak maximum energy and the 186-keV count rates, were measured, and the enrichment was calculated from these data. The raw data are shown in Fig. 5.12, and the calculated average enrichment at each pressure is shown in Table 5–1. Table 5–1 shows that the average enrichment did not deviate much.
from the known value of 3.3% used in the calibration. Except for the data at 10 Torr, where count rates in the 186-keV peak were very low, the standard deviation from the “known” enrichment at each pressure was less than 1% and improved as the pressure increased. This figure, which is a screen capture of the trend analysis section of the AEM software, plots parameters such as transmission peak net count rate (top), 186-keV net area count rate (middle), and enrichment (bottom) over time.

Figure 5-12: Performance of the system with varying pressures of UF₆ gas. Left to right: 20, 30, 40, 50 and 60 Torr. The top plot shows that the net area of the 25.5-keV silver transmission peak decreases with increasing gas pressure (once background has been subtracted). The middle plot is of the 186-keV peak. The bottom plot shows the calculated enrichment. Red points are individual data points collected when each spectrum was saved, and the green lines are a trend line of the data.
The trend analysis portion of the AEM software (from which the screen shot above was taken) allows user-chosen parameters to be displayed over time. In this case, the parameter that was changed over time was the pressure, raised in steps from 20 to 60 Torr, 10 Torr at a time. The top plot shows that the net area of the transmission peak decreases with increasing gas pressure (once background has been subtracted). In the plot beneath it, the 186-keV peak increases as the gas pressure is raised. The bottom plot shows that the calculated enrichment holds steady over all of the changes in the gas pressure. The enrichment plot also shows that the statistics of the enrichment measurement improve with increasing gas pressure. This improvement is due to the lower count rates in the 186-keV peak at low pressures.

Table 5-1: Average Enrichments at Various Pressures and the Effect of Count Rates in the 186-keV Peak Shown by the Standard Deviations

<table>
<thead>
<tr>
<th>Pressure (Torr)</th>
<th>Average Enrichment (%)</th>
<th>Standard Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>3.279 ± 0.097</td>
<td>2.95%</td>
</tr>
<tr>
<td>20</td>
<td>3.319 ± 0.029</td>
<td>0.86%</td>
</tr>
<tr>
<td>30</td>
<td>3.329 ± 0.027</td>
<td>0.80%</td>
</tr>
<tr>
<td>40</td>
<td>3.324 ± 0.023</td>
<td>0.69%</td>
</tr>
<tr>
<td>50</td>
<td>3.318 ± 0.020</td>
<td>0.60%</td>
</tr>
<tr>
<td>60</td>
<td>3.322 ± 0.018</td>
<td>0.55%</td>
</tr>
</tbody>
</table>

A significant effect occurred when the gas was removed from the pipe after it had been in the pipe for several weeks. As shown in Fig. 5.13, the total count rate in the spectrum continued to decrease after the gas had been removed. Investigation identified the cause as the decay of $^{234}$Th (half-life 24 days), which is formed by the decay of $^{238}$U.
and deposited on the inside wall of the pipe. When the UF₆ is removed, the $^{234}\text{Th}$ remains and continues to decay. Figure 5.14 shows the $^{234}\text{Th}$ spectrum, which has significant peaks at 63 and 93 keV. These peaks should not interfere with our enrichment measurement because they are in-between the transmission and 186-keV peaks.

Figure 5-13: Total counts per second in the vertical source after the UF₆ gas was pumped out. This represents the decay of the slight amount of deposit on the inside of the pipe wall.
The decay of $^{238}$U to $^{234}$Th in the deposits and its subsequent decay is an interesting phenomenon discovered when the pressure sensitivity measurements were performed. However, it should have no effect on the enrichment measurement because it is in a different region of the spectrum than either the transmission or the 186-keV peak. Laboratory testing has shown that pressure changes in the pipe (for the active AEM system) are handled well. Accurate enrichment monitoring is maintained over the range of UF$_6$ gas pressures that are expected in a working GCEP. Low count rates in the 186-keV peak force larger statistical errors in the enrichment measurements at low pressures, but this can be compensated for by using longer measurement times or averaging over more than one measurement.
Another example of using changes in pressure to determine the gas enrichment was performed with a source that did contain deposits on the walls. The active system was held at 25°C, and put through a series of pressure changes. This was done to simulate the gas pressure changes in a GCEP, and to show how these changes in pressure can be used to calculate $B$ and $I_0$, to determine the enrichment. Figure 5.15 shows the changing of the gas pressure, as well as $I(t)$, the transmission peak net count rate as determined by NaIGEM.

From the data taken at different gas pressures, plots of $I(t)$ and $R(t)$ (transmission count rate and 186-keV count rate) were created, shown in Figure 5.16. These data were fit to determine $I_0$ and $B$, the empty pipe transmission count rate and the portion of the 186-keV count rate that comes from deposits in the pipe. The fits of the data indicate a value of 4286.1 counts per second as the empty pipe transmission rate, and 21.008 as the background, when the gas pressure in the pipe is zero.
Figure 5-16: Transmission and 186-keV count rates vs. pressure, used to determine $I_0$ and $B$.

$B$ and $I_0$, determined from Figure 5.20, were then used to calculate the average enrichment at the various gas pressures using Equation 6: 

$$E(t) = K \frac{R(t) - B}{\ln \left( \frac{I(t)}{I_0} \right)}.$$ 

$B$ is calculated by fitting a line to the 186-keV count rate vs. pressure, and extrapolating to zero pressure. This determines the 186-keV count rate when there is no gas in the pipe, therefore coming from the deposits. $I_0$ is determined in the same way, by fitting an exponential to the transmission count rate vs. gas pressure to determine what the transmission rate would be for an empty pipe. Figure 5.17 shows the calculated enrichment over the range of pressure tested.
Figure 5-17: Calculated enrichment vs. gas pressure. Error in enrichment is driven by the statistical uncertainty in the 186-keV peak, which is highest at low pressures.

This calculated enrichment is constant at 3.3%, within the error at each pressure. The error in enrichment determination is mainly driven by statistical error in the count rate of the 186-keV peak, which is worse with decreasing pressure.

5.3.2 Temperature Correction: Using the Flux Monitor

An accurate temperature measurement of the gas is not always possible. Because of this measurement problem, postprocessing the transmission count rate data using the diode flux monitor data to correct for temperature effects was explored. Again, while keeping the gas pressure fixed, a temperature profile was run in the chamber that ranged
from 15 to 35°C. Over this range, the transmission peak count rate data changed by plus or minus 0.8%. Figure 5.18 shows the uncorrected data (relative change, in percent) and also the corrected data.

![Graph showing relative change in transmission peak count rate](image)

Figure 5-18: Transmission peak count rate for a typical temperature profile running between 35 and 15°C. The blue squares are corrected with the flux monitor diode data. (Relative change in percent is plotted.)

Although the diode-corrected data reduce the temperature effects of the x-ray tube in the count rate, they actually overcorrect the data. This is because the diode itself is slightly sensitive to temperature changes. Diode correction does reduce the error in the net count rate of the transmission peak from 0.8% to 0.3%.
5.3.3 Temperature Correction: Simple Method

A simple temperature correction for the transmission data on the active system has also been tested. This correction was determined by taking empty pipe measurements at 15, 25, and 35°C. Using these data, if the temperature of the system (including the UF₆ gas in the pipe) is known, the transmission data can be corrected for temperature effects.

First, empty pipe transmission count rates were measured. Each spectrum was collected over an hour, and the average count rate was recorded. X-ray tube high voltage and beam current were held constant, and the ramp rate of the temperature change was kept very low (under 1°C per hour). The flux monitor was placed in the collimated beam of the x-ray tube, mounted just after the notch filter. Transmission count rates at each temperature hold were averaged, and a line was fitted to the data at each temperature, as shown in Fig. 5.19. This fit was used to correct the raw transmission count rate data as it changed with temperature.

![Figure 5-19: Fit of transmission rate data, as a function of temperature, used to correct raw transmission rate data in the simple temperature correction method.](image)

*y = 1.3211x + 1946.6
R² = 0.9984*
A plot showing the flux monitor diode correction and the result of the simple temperature correction method is shown in Fig. 5.20. The temperature correction method shows less than a 0.2% relative change in the transmission peak over a range of temperatures from 15 to 35°C.

Table 5–2 relates these errors in transmission peak count rates to the final errors in enrichment determination. Because the natural log of the transmission rates is used to calculate enrichment, errors in the transmission rate have a higher effect on enrichment.
results. The table details one of the spectra collected as an example. It shows the raw data behind Fig. 5.20 above, when the environmental chamber was being held at 5°C.

**Table 5-2: Effect of Transmission Rate Error on Final Enrichment Calculation**

<table>
<thead>
<tr>
<th>Transmission Rate</th>
<th>Deviation in Transmission Rate (from baseline)</th>
<th>Calculated Enrichment (%)</th>
<th>Enrichment Error</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Raw Data (Uncorrected)</strong></td>
<td>1965.572</td>
<td>−0.65%</td>
<td>3.272</td>
</tr>
<tr>
<td><strong>Flux Monitor Corrected</strong></td>
<td>1982.619</td>
<td>0.21%</td>
<td>3.342</td>
</tr>
<tr>
<td><strong>Simple Temperature Correction</strong></td>
<td>1979.012</td>
<td>0.03%</td>
<td>3.327</td>
</tr>
</tbody>
</table>

The temperature change from 25°C to 5°C caused a 0.65% deviation in the transmission peak count rate, which caused the enrichment to be underreported by 1.60%. Using the flux monitor improves but overcorrects the raw transmission peak count rate. It shows a 0.21% deviation from the baseline transmission rate at this point, which causes the enrichment to be over-reported by 0.52%. Finally, using the simple temperature correction method, the transmission peak count rate is corrected to within 0.03% of the baseline, resulting in only a 0.07% error in the enrichment determination.

The next test was to check that this temperature correction could be used for data taken at other UF₆ gas pressures. Spectra were collected at UF₆ gas pressures of 20 and 40 Torr, and the data were postprocessed with the correction factor that had been determined at 0 Torr. Figure 5.21 shows the results, with the relative change in the transmission rate still within 0.25% over the 20-degree range.
This simple temperature correction is less complicated than using the flux monitor diode, which has its own temperature effects. It was verified that the simple temperature correction method works for pressures other than those in empty pipes by postprocessing data at 20 and 40 Torr of UF$_6$ gas pressure. Corrected results are in the range of 0 to 0.25% relative change in the transmission count rate per degree Celsius. This method provides a quick and easy way to correct for temperature changes, independent of gas pressure, as long as temperature data are available for the system. Although this method works on the laboratory UF$_6$ source in an environmental chamber, it will not be so straightforward in a GCEP, where the actual gas temperature is unknown and difficult to measure.
5.4 *Field Trial—URENCO Capenhurst*

A passive AEM, partially shown in Fig. 5.22, was installed on a cascade product header pipe at the URENCO Capenhurst enrichment plant. It is currently running in unattended mode, continuously collecting and saving data to a local computer. These data include gamma-ray spectral information as well as temperature information, as was described in Section 1.4.3, “Field Trial—Some Real World Data.” This picture shows the two inward-facing NaI detectors mounted on the pipe. The various grey colored pieces are the tungsten composite shielding designed specifically for this application by Tungsten Heavy Powder Inc. [42].

Above the passive system is the mounting for the active system, which was dry-fitted to the pipe as a demonstration for our Capenhurst collaborators. The x-ray tube will be mounted on the left side (not visible), and a third NaI detector will be mounted on the right side in the shielding ring. Also visible in the photo is the IOtech interface, which is used to read temperatures into our AEM software. The support table and brackets were designed by LANL and shipped to Capenhurst ahead of the installation trip. On the lower level of the table, not shown in the picture, are two ORTEC digiDART MCAs.
Figure 5-22: The AEM passive measurement head as installed at the Capenhurst enrichment plant. Two NaI detectors are mounted face-to-face across the unit header pipe, surrounded by tungsten composite shielding. The shielding for the active system is mounted above the passive detectors, without a detector or x-ray tube.

The LANL data collection software is set to collect spectra every 10 minutes. The 10 minute time was selected in order to have enough statistics in the spectrum (counts in the 186-keV peak) while still being able to capture any effects in the plant that occur on a short time scale. Each time a spectrum is saved, temperature information, averaged over the 10 minute period, is also saved. Our collaborators at Capenhurst periodically retrieve these data and send them to us for analysis, along with the corresponding gas pressure information.

Much of the data and processes that we are analyzing are plant-specific operational details and are therefore proprietary information. For this reason actual data
acquired are not shown here, but important findings that might affect our enrichment measurements are discussed.

5.4.1 Variations in UF₆ Gas Pressure during Measurement

Since the AEM is collecting data from the plant over a long period of time, cylinder fill cycles can be observed, along with all of the various parameters that change over these cycles. Figure 5.23 shows the basic concept of how some of these parameters change, relative to each other. For example, when a cylinder is attached at a fill station, the gas pressure at our measurement location drops, and the $^{235}$U count rate also drops. If the active system was in use, transmission through the gas would increase because of the lower gas pressure in the pipe. The cooled cylinder has the highest pumping power when it is first attached to be filled, and at this point the gas temperature also drops.
Figure 5-23: Schematic of measurement conditions in the unit header pipe during normal plant operation. When a new cylinder is connected to be filled, the gas pressure drops, the gas temperature and the $^{235}$U count rate drop, and transmission through the gas increases. The dashed lines represent B, the contribution to the $^{235}$U count rate from background (and deposits), and $I_0$, the empty pipe transmission scenario.

The idea of using pressure transients, or rapid drops in the UF$_6$ gas pressure in the pipe at the measurement location when a new fill cylinder is attached, was first presented at the European Safeguards Research and Development Association’s 2010 meeting [43]. These rapid pressure changes can be used to calibrate the passive portion of the AEM, since the enrichment cannot change in such a short time period.

5.4.2 Background Determination (Including Pipe Deposits)

By measuring the 186-keV count rate before and after a pressure transient, we can determine B, the background (which includes deposits) in Eqn. 7.

$$E(t) = K \ast (R(t) - B) \frac{T(t)}{p(t)}$$  \hspace{1cm} (7)
This is similar to the transmission-based enrichment formula given in Eqn. 5, but instead of using an attenuation term in the denominator to determine the total uranium content, it uses temperature corrected \((T(t))\) gas pressure \((p(t))\) to determine the total amount of gas in the pipe. This calculation is possible because the enrichment in a GCEP cascade cannot change over such a short period of time and can be treated as a constant over the pressure transient. Therefore, since the measurement system operates with declared operator pressure readings, one can interpolate down to a zero pressure and determine the portion of the 186-keV signal that is coming from background or the background plus pipe deposits. This is a fairly straightforward calculation, since it is assumed that the background \(B\) is also a constant over the pressure transient time. In the same way, with the active AEM, we could use this method to solve for \(I_0\), the empty pipe transmission count rate in Eqn. 6 from Chapter 5, “Experimental Measurements.”

Another method that has been even more accurate for determining background is to plot the calculated enrichment errors over time vs. the pressure at which the spectrum was collected. Then \(B\) can be optimized so that the plot of the enrichment error vs. pressure has a slope of zero—ensuring that the background determination is independent of gas pressure and, therefore, as accurate as possible.

### 5.4.3 Temperature Effects

The greatest challenge for calibrating the passive system, however, is the gas temperature, which not directly measurable. While there are temperature sensors on the aluminum pipe, the steel bellows after the compressor (just upstream of the AEM) and measuring ambient temperature, there is no direct measurement of the gas temperature. Furthermore, the temperature is the one variable that cannot be treated as a constant
during a pressure transient. For analysis of the current data being taken with the AEM in Capenhurst, a temperature correction factor is used that is derived from a combination of the measured temperatures. Once the active system is installed next to the passive system, the transmission measurement results can be used as an additional piece of information to help correct for gas temperature changes.
6  **Pipe-wall Thickness Results and Analysis**

This chapter presents the results and analysis of the pipe-wall thickness experiments. First the measured transmission ratios are shown to be independent of gas pressure, which allows this method to be used for an unknown gas pressure or in a facility where the AEM operator does not have access to facility pressure declarations. Next a calibration curve has been created, using a standard pipe of three known thicknesses as the basis for determining unknown pipe thicknesses. Finally, a brief discussion is presented, comparing the earlier analytical results to the experimental ones.

6.1  **Experimental Results**

A series of measurements was taken for the pipe thickness experiment, varying a number of factors.

1. Two different notch filters were used at each measurement setting: molybdenum and palladium.
2. Three thicknesses of each filter material were used: 0.3, 0.4, and 0.5 mm.
3. The transmission measurements were performed on three pipe thicknesses, as shown in Fig. 2.6.
4. The UF$_6$ gas pressure in the pipe was increased in steps up to 50 Torr.

After the data were collected, they were analyzed with a version of Ray Gunnink’s NaIGEM software [44], which he modified so it could be used to determine the net areas of both the 186-keV peak from the UF$_6$ and the transmission peak at either 20.0 keV (molybdenum filter) or 24.35 keV (palladium filter). Figure 6.1 shows a screen shot of a NaIGEM analysis in progress.
Figure 6-1: Screen shot of a NaIGEM analysis in progress. Fitting of the 24.3-keV transmission peak using the palladium notch filter is shown for the LaBr₃ detector.

With the matrix of measurements described above, ranges of filter thicknesses were explored to find the optimum combination that was most independent of gas pressure. Figure 6.2 shows the ratios of the net number of counts for the two transmitted spectral lines for a number of the best notch filter combinations, including 0.3-mm palladium/0.3-mm molybdenum, 0.4-mm palladium/0.3-mm molybdenum, and 0.5-mm palladium/0.3-mm molybdenum. These were combinations that had the lowest sensitivity to changes in the gas pressure. From these, the red data points, which were taken with a combination of 0.4-mm palladium/0.3-mm molybdenum filters, were selected as the best. Figure 6.3 is a plot of the ratios using this filter combination for different pipe thicknesses.
Figure 6-2: Choosing a notch filter combination. Ratios of the palladium (24.35 keV) to molybdenum (20 keV) transmission peaks for different notch filter thickness combinations on each step of the horizontal UF$_6$ source.

Figure 6.3 shows the ratios of transmitted palladium/molybdenum spectra, as a function of UF$_6$ gas pressure, for three pipe thicknesses. These data were taken with a
0.4-mm-thick palladium filter and a 0.3-mm-thick molybdenum filter. The ratio of these transmitted spectra vs. the gas pressure has slopes closest to zero over the range of pipe thicknesses tested. The slopes of these lines are very close to zero in all three cases, showing that these ratios are largely independent of gas pressure.

![Figure 6-3: Ratios of transmitted palladium/molybdenum spectra, as a function of UF₆ gas pressure, for three pipe thicknesses. The notch filter combination of 0.4-mm palladium/0.3-mm of molybdenum was chosen (the red points from Fig. 6.2).](image)

**6.2 Calibration Curve**

A calibration curve was determined (Fig. 6.4) with the transmission ratios shown above, allowing the measurement of a transmission ratio on an unknown aluminum pipe thickness to determine its thickness [45]. To determine the thickness of an unknown pipe,
two measurements are made on the pipe: one with a 0.3-mm molybdenum filter and one with a 0.4-mm palladium filter. The thickness of the pipe can then be determined from the ratio of the peaks generated. This measurement can be performed for any gas pressure that falls within the expected range of a working header pipe in a GCEP.

![Figure 6-4: Pipe thickness calibration curve. With a measured transmission ratio we can use this curve to determine an unknown aluminum pipe thickness.](image)

\[ y = 0.1181 \ln(x) + 0.0635 \]
\[ R^2 = 0.999 \]

The error in the ratios used for the calibration curve is too small to be seen except for the point that represents the thickest pipe. This error is driven by the statistical uncertainty in the net area of the peaks and is larger for the low count rates observed with thicker pipe segments.
6.3 “Unknown” Pipe Thickness Measurements

Next the two-step transmission measurement through three “unknown” pipe thicknesses was performed. Again, standard 6061 aluminum ¼”-thick pipes were used. Two sections of this pipe were machined down to thicknesses that were within the expected range of the header pipe thickness in a GCEP. A third pipe section that had a thickness out of our expected operating range was measured as well. Using the calibration curve that was achieved with the three-step source, $d_{\text{calc}}$, the calculated thickness of each pipe was determined. Once the transmission measurements had been performed and spectra had been collected, each pipe was cut at the transmission location for precise thickness measurements. The pipes were measured with a tube micrometer at two locations—the “entry” and “exit” spots of the beam—and averaged. Figure 6.5 shows the measured thickness of each pipe segment as a function of the ratio for each of the unknown pipe thicknesses. Error in the measured ratio comes from error in the calculation of the transmission peak net areas, as determined by the NaIGEM software. The error bars are too small to see on all but the ratio for the thickest section of pipe. This error was driven by the statistical uncertainty in the molybdenum transmission peak, which has the highest attenuation in the pipe.
Figure 6-5: Unknown thickness data points plotted against the calibration curve. Most of the error bars are smaller than the data points. True caliper-measured values for the pipe thicknesses were used for plotting the unknown pipes. In thicker pipes, statistics in the transmission peak will drive the error in the thickness determination when using the two-energy method.

Table 6–1 shows the ratio measured for each pipe thickness, the measured and calculated pipe thicknesses, and the percent difference in calculation using the two-step method. All three were calculated within 2% accuracy. The thickest pipe segment is out of the range of pipe thicknesses that we should encounter in a GCEP header pipe. This method loses accuracy for increasing pipe thicknesses because of the low transmission count rates seen in the detector in these cases. While the method still works, the error in the net area of the transmission peak gets too large because of the statistics. This is why the thickest unknown pipe ratio is the only one with visible error bars. Table 6-1 also
shows the effect of the pipe thickness error when determining the relative enrichment error, and the error in a 5% enriched gas, as an example.

### Table 6-1: Pipe Thickness Results

<table>
<thead>
<tr>
<th>Pd/Mo Ratio</th>
<th>(d_{\text{meas}}) (cm)</th>
<th>(d_{\text{calc}}) (cm)</th>
<th>Difference (%)</th>
<th>Enrichment Error</th>
<th>Enrichment (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>33.933 ± 0.132</td>
<td>0.472 ± 0.0018</td>
<td>0.480 ± 0.0001</td>
<td>1.7</td>
<td>8.3%</td>
<td>5.0±0.4</td>
</tr>
<tr>
<td>74.725 ± 0.750</td>
<td>0.561 ± 0.0024</td>
<td>0.573 ± 0.0089</td>
<td>2.0</td>
<td>11.6%</td>
<td>5.0±0.6</td>
</tr>
<tr>
<td>131.315 ± 2.843</td>
<td>0.632 ± 0.0016</td>
<td>0.640 ± 0.0073</td>
<td>1.2</td>
<td>7.8%</td>
<td>5.0±0.4</td>
</tr>
</tbody>
</table>

There appears to be a systematic error to the data, since all of our measurements fall below the calibration curve. This may be because unknown pipes, while being of the same alloy as the one used to create the calibration curve, may have had slightly different amounts of components such as copper, magnesium, silicon, etc. This may introduce a small error when performing real measurements in an enrichment plant, as will be determined by the results of the Capenhurst active AEM installation trip. However, there are only small variations in alloy materials allowed in these aluminum pipes, as shown in Table 6–2.

### 6.4 Comparison between Analytical/Experimental Results

In all of the analytical calculations performed, aluminum was used for the pipe material in a slab geometry. This was an approximation—the actual measurements were done with an aluminum alloy in a pipe geometry. The pipe used for the three-step source was a 6061 alloy with a nominal composition of 0.15% copper, 0.8% magnesium, and 0.4% silicon, with the remainder being aluminum [46]. This could be one reason why the
analytical notch filter thickness determination was slightly off but still close enough to give a good range of thicknesses of molybdenum and palladium to test. Table 6–2 shows the alloy composition of 6061 aluminum, detailing the ranges of materials that are accepted [47].

![Table 6-2: 6061 Aluminum Alloy Composition](image)

<table>
<thead>
<tr>
<th>Material</th>
<th>Amount (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aluminum</td>
<td>Balance</td>
</tr>
<tr>
<td>Chromium</td>
<td>0.04–0.35</td>
</tr>
<tr>
<td>Copper</td>
<td>0.15–0.4</td>
</tr>
<tr>
<td>Iron</td>
<td>0–0.7</td>
</tr>
<tr>
<td>Magnesium</td>
<td>0.8–1.2</td>
</tr>
<tr>
<td>Manganese</td>
<td>0.15 max</td>
</tr>
<tr>
<td>Other (Each)</td>
<td>0.15 max</td>
</tr>
<tr>
<td></td>
<td>0.05 max</td>
</tr>
<tr>
<td>Silicon</td>
<td>0.4–0.8</td>
</tr>
<tr>
<td>Titanium</td>
<td>0.15 max</td>
</tr>
<tr>
<td>Zinc</td>
<td>0.25 max</td>
</tr>
</tbody>
</table>

Calculations were performed to compare spectra generated using 0.3-mm molybdenum and 0.4-m palladium notch filters, for a 1-cm-thick aluminum pipe and a 1-cm-thick aluminum alloy pipe. Assuming a “worst-case scenario,” that the three-step pipe used to generate the calibration curve was pure aluminum, another curve was fit to compare it to using the high end of all of the alloy materials. This is shown in Figure 6.6, along with the calculated ratios for those three pipe thicknesses, using the density and attenuation coefficients calculated for this aluminum alloy.
Figure 6-6: Calculated ratios of the "worst case" aluminum alloy, compared to the three-step pipe and the unknown pipes.

Table 6-3 shows what the calculated thickness error would be when using the calibration curve generated with the three step source on pipes made of the worst case 6061 aluminum alloy. While the relative enrichment errors are high, the effect on the final calculated enrichment (shown again here with an example of a 5% enriched gas) are acceptable. If the goal of this system is to determine whether a product is low or highly enriched, the method will definitely work.
As for the error introduced by using a slab geometry instead of a pipe, an MCNP calculation could be performed using the correct geometry. However, this is not a concern since the basis of this work is experimental. Calculations were accurate enough to help choose experimental settings.
7  **Error Analysis**

The following formula, the general formula for error propagation, is used as the basis of this error analysis section:

\[ q = f(x_1, x_2, x_3, x_4), \quad \text{and} \]

\[ \delta q = \sqrt{\left(\frac{\partial q}{\partial x_1}\right)^2 + \left(\frac{\partial q}{\partial x_2}\right)^2 + \left(\frac{\partial q}{\partial x_3}\right)^2 + \left(\frac{\partial q}{\partial x_4}\right)^2}. \]  

7.1 **Errors in Determining Pipe-wall Thickness**

The first section details the calculated errors in the determination of the pipe-wall thickness. This includes physical measurement errors, errors in the transmission peak net areas due to statistical uncertainties, and a comparison of the effect of the wall thickness vs. instrument errors on the final enrichment determination.

7.1.1 **“Unknown” Pipe Thicknesses Measurement**

After the transmission measurements were completed, the empty pipes of “unknown thicknesses” were cut so their thicknesses at the measurement location (directly where the beam passed through) could be measured. Each pipe thickness was measured in two locations across from each other on the pipe. Ten measurements were performed in each spot with a pipe micrometer. The standard deviation was calculated at each location (of 10 measurements), and the average thickness of each pipe section was calculated.

The average thickness of each pipe section was calculated with the following equation:
\[ \text{Avg}_1 = \frac{\text{Avg}_1 + \text{Avg}_2}{2}, \]  

(10)

and the error in the average thickness is

\[ \Delta \text{Avg}_1 = \frac{\sqrt{\text{Avg}_1^2 + \text{Avg}_2^2}}{2}. \]  

(11)

In this way, the unknown pipe thicknesses were measured to the values listed as \( d_{\text{meas}} \) in Table 6–1.

### 7.1.2 Measured Ratios

The errors in the ratios of the transmission peaks using molybdenum and palladium filters were derived from the reported errors in net peak areas, as determined by the NaIGEM software. NaIGEM reports an error for each net area that it fits and calculates. Figure 7.1 shows an example of a NaIGEM fit of the 186-keV region of a spectrum taken with the LaBr\(_3\) detector with 50 Torr of UF\(_6\) gas in the horizontal source. Background is shown by the light blue line, and contributions from other \(^{235}\text{U}\) peaks at 144, 163, and 205 keV are taken into account (shown by the green line) when fitting the 186-keV peak.
Figure 7-1: Example of a NaIGEM fit used to determine the net area of the 186-keV peak. There is 50 Torr of UF$_6$ gas in the pipe, and the measurement was performed on Step 2, the middle (0.5 cm) thickness.

Figure 7.2 shows the results of a NaIGEM analysis, giving the live time, counts (and error) in the palladium transmission peak net area, and the counts (and error) in the 186-keV peak. Live time was needed because the net count rates in each peak were used for analysis instead of net counts, since longer measurement times were used when count rates were lower. Thirty-minute count times were typical for measurements using the palladium notch filter, but 60-minute count times were used in most cases with the molybdenum notch filter because it was at a lower energy and had higher attenuation in the aluminum pipe.
Figure 7-2: The results of a NaIGEM analysis, giving the live time, counts (and error) in the palladium transmission peak net area, and the counts (and error) in the 186-keV peak.

The ratios use counts per second as a normalization, so percent error was calculated as in Eqn. 11.

\[
\text{% Error} = \frac{\text{Uncertainty (NaIGEM)}}{\text{Net Area}}.
\]  \hspace{1cm} (12)

This percent error was then multiplied by the count rate (cps) where

\[
cps = \frac{\text{Net Area}}{\text{Live Time}}.
\]  \hspace{1cm} (13)

is used to get the percent error in the count rate.

Finally, the calculated error in the measured ratios is

\[
\Delta \text{Ratio} = \frac{Pd}{No} \times \left(\frac{\Delta No}{No}\right)^2 + \left(\frac{\Delta Pd}{Pd}\right)^2,
\]  \hspace{1cm} (14)
where palladium and molybdenum are the net areas of the transmission peaks with each of the notch filters, and ΔMo and ΔPd are the uncertainties in the net areas of each peak. This was used to determine the error in the ratios of the unknown pipe thicknesses in Table 6–1.

### 7.1.3 “Unknown” Pipe Thicknesses Calculation

To calculate the thicknesses of the unknown pipes, the calibration curve shown in Fig. 6.4 was used. This curve is a fit to the measured ratios of the palladium to molybdenum transmission peaks for each pipe segment, with errors described above. The fit is given by the following equation:

\[
d_{\text{calc}} = 0.1181 \times \ln(\text{Ratio}) + 0.0635
\]

To find the error in \(d_{\text{calc}}\), the calculated pipe thicknesses, the above fit for \(d_{\text{calc}}\) was used to determine the error in \(d_{\text{calc}}\) for each pipe thickness. Using Eqn. 9, it follows that

\[
\delta q = \sqrt{\left(\frac{\partial q}{\partial x_1}\right)^2 \delta x_1^2}
\]

where \(q = d_{\text{calc}}\), and \(x_1 = \text{Ratio}\).

Therefore,

\[
\delta d_{\text{calc}} = \sqrt{\left(\left(\frac{\delta (0.1181 \times \ln(\text{Ratio}) + 0.0635))}{\delta \text{Ratio}}\right) \times \delta \text{Ratio}\right)^2}.
\]

Because

\[
\frac{\partial}{\partial x} (\ln x) = \frac{dx}{x}
\]
we find that

$$\delta d_{\text{calc}} = \sqrt{0.1181 \times \left( \frac{\delta \text{Ratio}}{\text{Ratio}} \right) \times \delta \text{Ratio}^2}. \quad (19)$$

which because we already solved for $\Delta \text{Ratio}$ above, simplifies to

$$\delta d_{\text{calc}} = 0.1181 \times \left( \frac{\Delta \text{Ratio}^2}{\text{Ratio}} \right). \quad (20)$$

### 7.2 Enrichment Error from Wall Thickness Measurement Error

The following error analysis examines the effect of a measured 2% error in wall thickness on the feasibility of keeping the enrichment measurement within the 5% relative error limit set by mass balance requirements. In actuality, we strive for much better than this 5% relative error. The data from Capenhurst with the passive system alone demonstrate that we are able to achieve less than a 1% error in the enrichment determination if we are able to use declared gas pressures and calibrate with mass spectrometer data, which are already taken periodically for process control. However, situations where facility declarations are not available or cannot be trusted must be explored. For this reason, the effect of wall thickness measurement error on the enrichment error is studied.

The previously reported CEMO calibration is based on an initial laboratory calibration with the same type of pipe as the one in the plant [11]. While the composition of the pipe material is kept within very tight tolerances, the pipe geometry may differ significantly due to the extrusion manufacturing process. Because the attenuation in the pipe is much higher than attenuation in the UF$_6$ gas, any difference in the pipe thickness between the calibration pipe and header pipe in the plant could lead to a significant
calibration error. A calibration error caused by an error in the wall thickness can be
determined using an analysis similar to the one used to determine the transmission error
reported previously by Ianakiev et al. [13]. This error is compared to the difference in the
thickness between the facility and calibration pipes, as explored in the Ianakiev et al. This
analysis is described below.

If the same composition in pipe material between the calibration and the facility
pipe is assumed, the variable $d_{w0}$ is used for the wall thickness of the calibration pipe,
along with $I_0$ and the variable $d_e$ for the thickness of the header pipe. Similar to the way it
was expressed in Eqn. (2) in “New Generation Enrichment Monitoring Technology for
Gas Centrifuge Enrichment Plants,” [13] the transmission ratio $I/I_0$ can be expressed as
follows:

$$
\frac{I}{I_0} = \frac{I_S(t)}{I_S(t_0)} \cdot \frac{\epsilon}{\epsilon_0} \cdot \frac{e^{-\mu_e \rho_e d_e}}{e^{-\mu_w \rho_w d_{w0}}} \cdot \frac{e^{-\mu_{UF6} \rho_{UF6} d_{UF6}}}{e^{-\epsilon}}
$$

(21)

where $I_S(t)$ and $I_S(t_0)$ are the intensities of the transmission source for the
measurement and calibration times, respectively; $\epsilon$ and $\epsilon_0$ are the detection efficiencies
for the measurement and calibration, respectively, incorporating all contributing factors
(geometry, NaI(Tl) intrinsic efficiency, MCA dead time, etc.); $\mu_w, \rho_w,$ and $d_{w0}$ are the
attenuation parameters of the calibration pipe; $\mu_w, \rho_w,$ and $d_w$ are the attenuation
parameters of the header pipe; and $\mu_{UF6}, \rho_{UF6},$ and $d_{UF6}$ are the attenuation parameters
of the UF$_6$ gas. The attenuation of the vacuum is shown as $e^{-\epsilon}$. 

112
For these calculations, a constant intensity of the transmission source is assumed, so the intensities of the source in each instance cancel each other out. The direct dependence of the enrichment, \(\text{Enrichment} = K \cdot (R - B)/\ln \left(\frac{I}{I_0}\right)\) (Eqn. 6), can be presented as

\[
T \left( \frac{\Delta \varepsilon \cdot \Delta d_w}{\varepsilon_0 \cdot d_{w0}} \right) = \ln \left( \frac{I}{I_0} \right) = -\mu_{UF6} \cdot \rho_{UF6} \cdot d_{UF6} - \mu_w \cdot \rho_w \cdot d_{w0} \cdot \left( \frac{\Delta d_w}{d_{w0}} \right) + \ln \left( \frac{\Delta \varepsilon}{\varepsilon_0} + 1 \right),
\]

where \(\varepsilon = \varepsilon_0 + \Delta \varepsilon\) and \(\frac{\Delta \varepsilon}{\varepsilon_0}\) is the relative change of the detection efficiency, \(\ln \left( \frac{\Delta \varepsilon}{\varepsilon_0} + 1 \right)\) is the corresponding instrumental error of the transmission ratio, and \(d_w = d_{w0} + \Delta d_w\) and \(\frac{\Delta d_w}{d_{w0}}\) are the relative changes in the wall thickness. And finally, \(\mu_w \cdot \rho_w \cdot d_{w0} \cdot \left( \frac{\Delta d_w}{d_{w0}} \right)\) is the transmission error due to different attenuations in the pipes.

The enrichment formula, including the calibration and instrumental errors, is

\[
E \left( \frac{\Delta \varepsilon \cdot \Delta d_w}{\varepsilon_0 \cdot d_{w0}} \right) = \frac{-K \cdot I_{186}}{\mu_{UF6} \cdot \rho_{UF6} \cdot d_{UF6}} - \mu_{UF6} \cdot \rho_{UF6} \cdot d_{UF6} + \mu_{UF6} \cdot \rho_{UF6} \cdot d_{UF6} + \mu_w \cdot \rho_w \cdot d_{w0} \cdot \left( \frac{\Delta d_w}{d_{w0}} \right) \right) - \ln \left( \frac{\Delta \varepsilon}{\varepsilon_0} + 1 \right),
\]

where \(\frac{-K \cdot I_{186}}{\mu_{UF6} \cdot \rho_{UF6} \cdot d_{UF6}}\) is the enrichment for \(\Delta d_w = 0\) and \(\Delta \varepsilon = 0\), and

\[
\frac{\mu_{UF6} \cdot \rho_{UF6} \cdot d_{UF6}}{\mu_{UF6} \cdot \rho_{UF6} \cdot d_{UF6} + \mu_w \cdot \rho_w \cdot d_{w0} \cdot \left( \frac{\Delta d_w}{d_{w0}} \right) - \ln \left( \frac{\Delta \varepsilon}{\varepsilon_0} + 1 \right)}
\]

is the combined error factor [12].
With the measured 2% error in wall thickness, \( \frac{\Delta d}{d_0} \), the maximum instrumental error allowable was calculated in order to keep the enrichment measurement within the 5% (full range) relative error limit set by mass balance requirements. Figure 7.3 shows one of the plots used to calculate this error for a transmission energy of 30 keV. The enrichment measurement error is approximately symmetrical for positive and negative errors in pipe thickness measurement.

Figure 7-3: Example of enrichment values calculated as described for UF\(_6\) gas at 4.5% enrichment and 50 Torr pressure. These values were calculated at a transmission energy of 30 keV. The range of instrumentation errors for varying wall thickness error is shown. Specifically detailed is the area that falls within the 5% relative error limit set by mass balance requirements.
The calculation above was repeated over a range of transmission energies, with the results shown in Fig. 7.4. Note that if the wall thickness error is fixed, as in our case at 2%, the maximum allowable instrumentation error decreases with increasing energy.

Figure 7-4: Calculated values of the maximum allowable instrumentation error vs. energy for the enrichment result, to be correct within 5%. This is for a fixed error in wall thickness of 2%.

Finally, an enrichment calculation as a function of transmission energy is shown in Fig. 7.5, with the instrumentation error fixed at 1.5% and the wall thickness error at 2%. A notch filter such as one made from tin, which has a K-edge at 29.2 keV, would be a good choice using these results. This was calculated using Eqn. 23, shown above. The portion that falls in the 5% (total) relative error, as desired by mass balance requirements, is detailed.
Figure 7-5: Enrichment calculation as a function of transmission peak energy, including the combined error factor described in the error analysis section. The range that falls into the 5% relative error is shown.

7.3 Enrichment Errors over Time (Passive System)

The calculations above, using the combined error formula, show the contributions to the enrichment error from errors in certain static parameters such as the calculated pipe-wall thickness, detection efficiencies, geometric factors, etc. Since the intended use of the AEM is for unattended monitoring over long periods of time between calibrations, effects of the parameters that might affect the enrichment calculations over time is also important.

The data acquired with the passive version of the AEM at URENCO Capenhurst (Section 5.4, “Field Trial—URENCO Capenhurst”), illustrate how factors such as temperature and gas pressure affect the error in the enrichment measurement over time.
We are able to calculate the enrichment to a very high accuracy when changes in temperature and pressure are corrected for. Using facility pressure declarations plus a one-time mass spectrometer measurement for calibration, the gas enrichment over time can be determined. Figure 7.6 shows the enrichment error averaged over 12-hour time segments. If all factors contributing to error are considered, the final error assessment of enrichment made in the experimental verification test run of the passive AEM at the URENCO Capenhurst centrifuge enrichment plant was very promising. The plot below shows these data over a six-week period, with the average enrichment error easily remaining below the desired maximum error of 5% (full scale of the graph). This was based on an initial calibration with a mass spectrometer sample and assumes a constant enrichment over time. Part of the enrichment error shown below may actually be due to fluctuations in enrichment of the product UF₆ at the plant, but actual enrichments cannot be shown.

Figure 7-6: Passive AEM enrichment error over almost two months of running at the URENCO Capenhurst GCEP. These data have been corrected for variations in gas pressure and temperature.
By averaging the data over 12-hour time segments, the UF₆ gas enrichment can be determined to ±1%. This is well within the limit set by mass balance requirements and if improved further may even be useful to the plant operators for process control.
8 Conclusions

This dissertation has described a method for determining the cascade header pipe thickness with an enrichment monitor, using an x-ray source and a LaBr₃ detector, when an empty pipe measurement is not feasible. The various possible x-ray filters and isotopic sources were reviewed, and molybdenum and palladium (with K-edges at 20.0 and 24.35 keV, respectively) were chosen on the basis of initial analytical calculations. These analytical calculations showed that the ratios of transmitted spectra should be completely independent of UF₆ gas pressure if notch filter thicknesses are optimized.

From the measurements performed, it was determined that notch filters made of a combination of 0.4-mm palladium and 0.3-mm molybdenum gave the ratio of transmission spectra that was most independent of UF₆ pressure in the pipe. With these notch filters, a calibration curve was created using our UF₆ source with three pipe thicknesses. This curve allows the pipe thickness to be determined simply by measuring the ratio of two transmission peak measurements with the two-energy thickness measurement method. This one-time measurement could be completed in less than an hour, and once the calibration has been performed for a specific measurement location it is not necessary to perform it ever again.

When the pipe thickness had been determined, the AEM could be switched to an unattended mode of operating after selecting an transmission energy that would maximize transmission through the pipe but still allow measureable attenuation in the gas. At that point, a single transmission measurement is needed to determine the gas density, and thus the enrichment. Typically a silver notch filter, with a K-edge at 25.5
keV, is used for this purpose. This will be the notch filter used in the next text trial at Capenhurst, when the active system is installed.

Major issues with the AEM operation that may lead to potential sources of error in the enrichment determination have been investigated. In order to address these issues, analytical calculations and experiments have been run to study cascade header pipe-wall thickness concerns, x-ray tube instabilities, and notch filter material use. This dissertation briefly looked at the effect of wall deposits on enrichment determination, as well as the sensitivity of the enrichment results to changes in pressure and temperature during measurement. In an unfriendly facility, the two-energy thickness measurement method provides a way to determine the UF₆ gas enrichment without using facility pressure declarations. Alternatively, if facility pressure readings are readily available and can be trusted, the passive AEM has been shown to produce enrichment results, when corrected, to within 1% of the actual values.
9 Future Work

As a direct follow-on to the work that is ongoing in the URENCO Capenhurst plant, a field trial is planned for mid-April of this year to install an active AEM system directly above the passive AEM which is currently installed and collecting data. This will be the first time an active system, using an x-ray generator as a transmission source, has been used in an enrichment monitor in a GCEP. The installation will allow a real analysis of the system performance, because it will be exposed to real temperature fluctuations and actual changes in gas pressure over the cylinder fill cycles.

One important improvement to the system which will be explored once real data are acquired is an attempt at performing a running calibration. Using pressure transients to recalibrate the system will make sure it does not drift over time. These quick (known) drops in gas pressure when a new cylinder is attached can be periodically used to check the calibration of the system, and adjust it if needed.

Another possible future path is to look at the whole system – to monitor not only the GCEP product line, but also to look at the feed and the tails. This could help the operators monitor their complete process to improve efficiency. It could also be used by inspectors in the future as a mass balance check (with information from load cells and other factors) to determine if the input of the system matches the output or whether there might be diversion taking place.
Appendices

Appendix A: Mass Attenuation Coefficients

Appendix B: Equipment Specifications

Appendix C: MCNPX Input File—Diode Flux
Appendix A: Mass Attenuation Coefficients
UF6

\[ \text{(cm}^2/\text{g}) \]

\[ \text{Photon Energy (MeV)} \]

- Total Attenuation with Coherent Scattering
Appendix B: Equipment Specifications
APPLICATION

The VF-50 series x-ray tube is a beryllium window x-ray tube designed for use as a radiation source for x-ray fluorescence systems.

CONSTRUCTION

The beryllium x-ray window is located at the end of the tube and the beam is projected along the longitudinal axis of the tube. The cathode operates at ground potential and the envelope is ceramic with the high voltage section potted to increase the high voltage stand off.

<table>
<thead>
<tr>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Envelope</td>
</tr>
<tr>
<td>Be Window</td>
</tr>
<tr>
<td>Anode</td>
</tr>
<tr>
<td>Standard Target Materials</td>
</tr>
<tr>
<td>Target Angle</td>
</tr>
<tr>
<td>Focal Spot</td>
</tr>
<tr>
<td>Maximum Anode Dissipation with 10 cfm forced air cooling</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Filament Characteristics</td>
</tr>
<tr>
<td>Maximum Anode Potential</td>
</tr>
<tr>
<td>Maximum Tube Current</td>
</tr>
<tr>
<td>Cooling Method</td>
</tr>
<tr>
<td>Weight</td>
</tr>
</tbody>
</table>
Spellman's XRM Series of regulated X-ray power supplies offer output voltages to 65kV and incorporate a filament supply which provides regulated dc current adjustable between 0.4A and 3.5A at 5.5kV. High voltage and filament current can be linearly ramped up. The XRM incorporates local and remote programming, monitoring, safety interlock, short-circuit and overload protection.

**TYPICAL APPLICATIONS**

- Powering grounded cathode X-ray tubes from Kavo, Oxford, Poh, Superior, Vianen and Trufocus.

**OPTIONS**

- AC: AC Filament
- CPC: Constant Power
- BIAS: Bias Supply
- TP(x): Alternate Test Point Scaling

**SPECIFICATIONS**

- **Input:**
  - 424Vdc±10%, 4.25A maximum.
- **Output:**
  - 4 models with positive output polarity and adjustable voltages from zero to maximum voltage and current.
- **Voltage Control:**
  - Local: Internal multi-turn potentiometer to set voltage from 0 to full output voltage.
  - Remote: 0 to +10Vdc proportional from 0 to full output voltage.
- **Resistance Control:**
  - Local: Internal potentiometer to set beam current between 0 and full output.
  - Remote: 0 to +10Vdc proportional from 0 to full output current.
- **DC Filament Supply:**
  - Current: 35A, adjustable.
  - Voltage: 6.5V.
- **Voltage Regulation:**
  - Load: ±0.1% of output voltage no load to full load.
  - Line: ±0.05% for ±10% change in input voltage.
- **Current Regulation:**
  - Load: ±0.5% of output current from 0 to rated voltage.
  - Line: ±0.01% of rated current over specified input range.
- **Rigidity:**
  - 0.25% p-p of output voltage.
- **Temperature Range:**
  - 0°C to 65°C operational
  - -40°C to 85°C ambient
- **Temperature Coefficient:**
  - ±0.01% per °C, voltage or current regulated.
- **Stability:**
  - ±0.05% per 6 hours after 12-hour warm-up.
- **Voltage and Current Monitors:**
  - 0 to ±10Vdc (proportional from 0 to full output).
- **Accuracy:** ±1%

- **Dimensions:**
  - 6.58"H x 2.93"W x 10"D (168mm x 75mm x 254mm).

- **Connectors:**
  - HV Output Connector: Delrin type connector, recessed.
  - HV Input Connector: 10-pin mini-D-type Phoenix connector for power, filament and monitor connectors.

- **Remote Programming:**
  - (SCPI compliant, 17-pin D-sub control interface) Permits remote adjustment of the output voltage and current via an external potentiometer and the internal +10V reference. By adjusting the potentiometer from minimum to maximum, the desired output may be selected.

- **Remote Monitor:**
  - Test points are available at J4 for monitoring voltage and current output. The output polarity is positive from 0 to 10V, equal to 0 to 100% of the output.

- **Regulatory Approvals:**
  - Compliance to 2004/108/EC, the EMC Directive and 2009/47/EC, the Low Voltage Directive.
XRM SELECTION TABLE

<table>
<thead>
<tr>
<th>Voltage</th>
<th>Current (mA)</th>
<th>Model Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>25</td>
<td>2.0</td>
<td>XRM252.0</td>
</tr>
<tr>
<td>50</td>
<td>1.67</td>
<td>XRM501.67</td>
</tr>
<tr>
<td>65</td>
<td>1.00</td>
<td>XRM651.00</td>
</tr>
<tr>
<td>85</td>
<td>0.77</td>
<td>XRM850.77</td>
</tr>
</tbody>
</table>

J2 POWER CONNECTOR—-2 PIN PHOENIX

<table>
<thead>
<tr>
<th>PIN</th>
<th>SIGNAL</th>
<th>PARAMETERS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>L &amp; R Input</td>
<td>+24Vdc @ 1.25Amps Input</td>
</tr>
<tr>
<td>2</td>
<td>L &amp; R Return</td>
<td>Power Return</td>
</tr>
</tbody>
</table>

J3 FILAMENT CONNECTOR—-3 PIN PHOENIX

<table>
<thead>
<tr>
<th>PIN</th>
<th>SIGNAL</th>
<th>PARAMETERS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Filament Out</td>
<td>0 to 3.5 Amps @ 25.5-volt filament output</td>
</tr>
<tr>
<td>2</td>
<td>Filament Return</td>
<td>Filament Return</td>
</tr>
<tr>
<td>3</td>
<td>Spare</td>
<td>NC</td>
</tr>
</tbody>
</table>

J4 MONITOR CONNECTOR—-4 PIN

<table>
<thead>
<tr>
<th>PIN</th>
<th>SIGNAL</th>
<th>PARAMETERS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Monitor Return</td>
<td>Signal Ground</td>
</tr>
<tr>
<td>2</td>
<td>RX Monitor</td>
<td>0 to 10mV @ 0 to 100% of regulated output, Code = 100</td>
</tr>
<tr>
<td>3</td>
<td>mA Monitor</td>
<td>0 to 10mV @ 0 to 100% of regulated output, Code = 100</td>
</tr>
<tr>
<td>4</td>
<td>Status Enable</td>
<td>Connecting ground through 10,000 ohm (2.5 to 5V) to close monitor</td>
</tr>
</tbody>
</table>

J5 CONTROL INTERFACE—-8 PIN MINI D CONNECTOR

<table>
<thead>
<tr>
<th>PIN</th>
<th>SIGNAL</th>
<th>PARAMETERS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>+10Vdc Reference</td>
<td>+10Vdc @ 1mA</td>
</tr>
<tr>
<td>2</td>
<td>Spare</td>
<td>NC</td>
</tr>
<tr>
<td>3</td>
<td>RX Program Input</td>
<td>0 to 10mV @ 0 to 100% of regulated output, Code = 100</td>
</tr>
<tr>
<td>4</td>
<td>TX Program Output</td>
<td>0 to 10mV @ 0 to 100% of regulated output, Code = 100</td>
</tr>
<tr>
<td>5</td>
<td>Spare</td>
<td>NC</td>
</tr>
<tr>
<td>6</td>
<td>mA Program Input</td>
<td>0 to 10mV @ 0 to 100% of regulated output, Code = 100</td>
</tr>
<tr>
<td>7</td>
<td>mA Program Output</td>
<td>0 to 10mV @ 0 to 100% of regulated output, Code = 100</td>
</tr>
<tr>
<td>8</td>
<td>Spare</td>
<td>NC</td>
</tr>
<tr>
<td>9</td>
<td>Ground</td>
<td>Signal Ground</td>
</tr>
</tbody>
</table>
Appendix C: MCNPX Input File—Diode Flux
MCNPX Input File—Diode Flux

c  Flux at Si diode
1   1  -2.329 -1
2  204  -0.001225  -2  #1
3   0     2

1  rpp 1  1.0104  -0.5  0.5  -0.5  0.5  $Si
2  so 2

mode p e
ml  14000.04p       1 $MAT1
m204  7014.60c   -0.755636 $MAT204
   8016.60c   -0.231475 18000.59c  -0.012889
imp:p  1  lr     0       $ 1, 3
imp:e  1  lr     0       $ 1, 3

c  source is photons, left of Si det pointed in the x direction
sdef par=2  erg=d1  pos = .9  0  0  axs= 1 0 0 vec = 1 0 0 dir=1
si1  L 0.0125  .015  .0165  .01725  .018  .019  .02  .021  .022  .023  .024  .025  .0259  .0301  .0328  .0349  .0376  .0397
spl1  d 2.28  24.59  54.92  75.99  93.77  120.36  145.13  167.80  185.96  199.5  209.27  212.27  2.08  7.55  11.01  11.82  8.47  1.47

c  f6:p 1    $Energy Deposition in the Si Detector (MeV/g)
*f6:p 1    $Energy Deposition in the Si Detector (GJ/g)
ctme  300
References


[40] A. Holmes-Seidle and L. Adams, Handbook of Radiation Effects, New York:


