Preliminary Results of an On-Line, Multi-Spectrometer Fission Product Monitoring System to Support Advanced Gas Reactor Fuel Testing and Qualification in the Advanced Test Reactor at the Idaho National Laboratory

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Preliminary Results of an On-Line, Multi-Spectrometer Fission Product Monitoring System to Support Advanced Gas Reactor Fuel Testing and Qualification in the Advanced Test Reactor at the Idaho National Laboratory

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Abstract—The Advanced Gas Reactor-1 (AGR-1) experiment is the first experiment in a series of eight low-enriched uranium oxycarbide tri-isotropic (TRISO) coated particle fuel (in compact form) experiments scheduled for irradiation in the Advanced Test Reactor (ATR) located at the Idaho National Laboratory (INL). The experiment began irradiation in the ATR with a cycle that reached full power on December 26, 2006 and will continue irradiation for about 2.5 years. Six separate test fuel capsules are being irradiated in an inert sweep gas atmosphere with on-line fission product monitoring of each capsule’s effluent to track fission gas emissions from the fuel during irradiation.

The online fission product monitoring system incorporates 7 HPGe spectrometers and 7 NaI(Tl) gross radiation detectors to monitor and quantify the fission gas releases that are important indicators of fuel performance. Details of the design and operation of this detection system and the preliminary results of the fuel performance measurements are presented in this paper.

I. INTRODUCTION

The US Department of Energy has embarked on a series of tests of coated-particle reactor fuel intended for use as part of a future Advanced Gas Reactor (AGR) design [1]. The AGR-1 experiment is the first in a series of eight fuel tests planned for irradiation in the Advanced Test Reactor (ATR) located at the Idaho National Laboratory (INL). The AGR is based on high-temperature gas-cooled reactor (HTGR) technology. The distinguishing features of HTGRs are the use of helium coolant, a low-power-density ceramic core capable of withstanding very high temperatures, and coated-particle fuel. The AGR fuel consists of low-enriched uranium oxycarbide in tri-isotropic (TRISO) coated particles in a carbon matrix compact as shown in Fig. 1.

One important measure of the fuel performance in these irradiation experiments is quantification of the fission gas release over the course of the 2.5-year (approximately 700 Effective Full Power Days) experimental irradiation cycle, during which the fuel compacts are expected to reach a burnup of 18 atom percent [1].

The AGR-1 experiment includes six (6) individual fuel capsules, each containing twelve (12) fuel compacts arranged in 3 stacks per capsule, held in a graphite spacer, incorporated into a test train as shown in Fig. 2. The experiment is what is termed a “lead” experiment, thus the fueled section of the test train that is inserted into the ATR irradiation port connects through an umbilical cord (termed a lead-out) to systems outside the vessel. In the first experiment (AGR-1) each capsule contains about 51,000 TRISO-coated fuel particles. This experiment is intended to serve as a shakedown test of a multi-capsule design to be used in subsequent irradiations [2].

Fig. 1 TRISO Coated fuel particles (left) are formed into fuel compacts (center) and inserted into graphite block matrix (right). The coatings on the TRISO fuel form (primarily the SiC layer) provides the primary barrier to fission product release.
The AGR-1 test train is located in the east large B position (B-10) of the ATR core. During irradiation, independent gas lines route a mixture of inert gasses through each of the six capsules to sweep released fission products to the Fission Product Monitoring System (FPMS) which monitors each capsule’s effluent in near real time [3]. Fig. 3 shows the AGR-1 experiment flow path. The FPMS consists of seven (7) fission product monitors, one for each capsule plus a spare, each consisting of a heavily-shielded high-purity germanium (HPGe) gamma-ray spectrometer for energy resolution and a sodium iodide [NaI(Tl)] scintillation detector for count rate monitoring to provide indication when a “puff” release occurs [4]. The primary functions of this system are to detect the fission gas release that occurs when a TRISO fuel coating may have failed and to provide the release data to support fuel performance evaluations. The on-line analysis of the HPGe spectra provides a record of the isotopes detected in the effluent gas, the most common being inert fission gases.

II. THE AGR-1 FPM SYSTEM

The AGR-1 experiment began irradiation in the ATR operations cycle that reached full power on December 26, 2006 and will continue irradiation for about 2.5 years. To support this experiment, the FPMS was completely assembled, tested, and calibrated in a laboratory at the INL, and then moved and reassembled in its final location in the ATR reactor basement. Fig. 4 shows the equipment assembled in the reactor cubicle at the ATR.
A. The Fission Product Monitoring Software

The FPM software provides the user interface for the operation of the HPGe spectrometers and the NaI(Tl) gross activity monitors for each of the six fuel capsules and for the seventh reserve spectrometer. This interface provides the capability to modify the operation of any of these peripherals in real time. Typically, under normal operation, computerized data acquisition is continuous without operator intervention. The FPM software allows the operator to display the contents of all seven spectrometers or all seven of the gross activity monitors in near real time as is shown in fig. 5 and fig. 6 respectively. The software also allows the operator to view any single spectrum in either near real time or previously saved to disk as shown in fig. 7. The operator can similarly view the activity in a single gross activity monitor as shown in (fig. 8), where the inset is displaying saved data from a completed acquisition segment.

![Fig. 5 The FPM Software allows the operator to look at all 7 HPGe detectors at a glance.](image)

![Fig. 6 The FPM software allows the operator to look at all 7 NaI(Tl) gross activity monitors at a glance.](image)

Each HPGe spectrometer and NaI(Tl) gross monitor has the ability to be placed off-line without adversely affecting the operation of the other spectrometers. This feature is especially useful when we experience either a detector or electronic failure. The gas lines that transport the fission gas effluent from each capsule to the six FPM detector stations also has the flexibility of being able to be re-routed to the spare FPM detector station while a failed monitor is being repaired. As partially stated above, the normal operation of the monitors will be a continuous loop of collecting and then analyzing the resulting spectrum. The length of the data collection is an operator-modifiable parameter. The time range of the Multi-Channel Scaler (MCS) spectra (showing the NaI(Tl) detector count rates as a function of time) is matched to the sample measurement time used for the corresponding HPGe spectra. All spectra are stored with unique file names that identify the spectrometer as well as the time and date of the measurement.

![Fig. 7 The FPM allows the operator to choose to view the effluent from an individual capsule (in real-time or stored).](image)

![Fig. 8 The FPM Software allows the operator to view the effluent from an individual NaI(Tl) gross activity monitor in near real-time or from disk (in the inset). This feature is especially useful for viewing particle failures in near-real time. The FPMS is extremely sensitive as evidenced by the blip in the inset which was caused by changing the supply gas bottle for the sweep gas.](image)

B. Use of FPM to verify prevention of Cross Flow between the Capsules

Because of thermal expansion differences, the individual capsules of the AGR-1 test train could not be completely sealed. A “slip” fitting was required between each capsule’s bottom closure plate and tubes passing through it. This raised the possibility that under some flow conditions fission gasses from one capsule could leak to the common test train volume (called the “lead-out”) and from there into a different capsule thus creating capsule-to-capsule crosstalk. To ensure no fission products leak from any capsule into the leadout, the
leadout is supplied with helium, termed the leadout flow, at a rate such that helium leaks into each of the capsules precluding crosstalk. To determine the minimum leadout flow rate required, a leadout flow experiment was conducted in early March of 2007 while the AGR-1 test train was being irradiated. Based on pressure drop measurements made before test train installation, three of the capsules were deemed most likely to produce leakage into the leadout and result in crosstalk if the leadout flow was too low. These capsules 1, 2, and 5 were tested in turn as follows [5].

The leadout flow experiment was conducted by setting the inlet flow conditions and holding them constant for an hour to assure equilibration before making each change. Initially, the flow to the leadout and each of the capsule inlets was set to pure He at 30 cm³/min at STP (termed “sccm”). Next, the inlet flow to one of the capsules to be tested was changed to 28 sccm of He and 2 sccm of Ne. Subsequently at 1 hour intervals the He flow to the leadout was decreased by 2 sccm. This was continued, decreasing the He flow to the leadout in a step-like fashion, until the leadout flow rate was at or near zero (fig. 9). After the leadout flow had been held at its minimum for one hour, the Ne flow to the tested capsule was secured, the flow to the leadout and the capsule returned to 30 sccm He, repeating the initial conditions, and the system maintained at that condition for 1 hour. After the 1 hour purge and wait, the process of neon injection into the inlet and sequential decreases in the flow to the leadout was repeated for the next capsule to be tested.

During all of these leadout flow test periods, the AGR FPMS was acquiring sequential 20 minute spectra for the effluent from capsule 5 during the leadout flow test. The activity increases as Ne is introduced into the sweep gas at a fixed flow rate, and then decreases as the leadout flow is reduced hourly, decreasing the flow of He from the leadout to the capsule outlet, thus increasing the transport time from the capsule to the detector as the combined flow from the detector is reduced.

![Graph showing activity of 23Ne](image)

**Fig. 9** Activity of 23Ne (in micro-Ci per 58 cm³) for consecutive 20 minute spectra for the effluent from capsule 5 during the leadout flow test. The activity increases as Ne is introduced into the sweep gas at a fixed flow rate, and then decreases as the leadout flow is reduced hourly, decreasing the flow of He from the leadout to the capsule outlet, thus increasing the transport time from the capsule to the detector as the combined flow from the detector is reduced.

In order to derive fuel performance parameters (release-to-birth ratios) the fission gas data measured at the FPMS must be corrected for decay during the transport time from the constantly-irradiated fuel capsule to the FPMS measurement location. The capsule effluent gas flow rates change according to the desired experimental conditions, therefore it is convenient to define a capsule-to-FPMS transport volume that is a function of the hardware installation. The previously-described leadout flow experiment provided the additional benefit of sequentially decreasing each capsule’s outlet flow rate while keeping the neon in the capsule inlet constant. This sequentially increased the transport time to the FPMS Spectrometers which also increased the decay time for the short lived (37.2 second half-life) activated 23Ne. The relative decrease in the short-lived 23Ne activity, due to decay during transport, was analyzed as a function of the measured outlet flow-rate to derive capsule-specific transport volumes for the three capsules tested (capsules 1, 2, and 5) in the leadout flow experiments [5]. Transport volumes for the remaining three capsules (capsules 3, 4, and 6) were estimated as the variance-weighted average of the tested capsules. The results of this analysis is presented in Table I along with the associated transport times in seconds for a nominal capsule outlet flow of 30 sccm. At nominal outlet flow rates of 30 sccm these volume uncertainties result in 1σ uncertainties in the capsule-to-detector transport times of 3.5 to 4.5 seconds [5].

<table>
<thead>
<tr>
<th>Capsule</th>
<th>Transport Volumea (cm³)</th>
<th>Transport Timeb (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>126 ± 7</td>
<td>252</td>
</tr>
<tr>
<td>2</td>
<td>138 ± 7</td>
<td>276</td>
</tr>
<tr>
<td>3</td>
<td>128 ± 9</td>
<td>256</td>
</tr>
<tr>
<td>4</td>
<td>128 ± 9</td>
<td>256</td>
</tr>
<tr>
<td>5</td>
<td>120 ± 8</td>
<td>240</td>
</tr>
<tr>
<td>6</td>
<td>128 ± 9</td>
<td>256</td>
</tr>
</tbody>
</table>

*a. Volumes for capsules 3, 4, and 6 were determined by computing the variance-weighted average of the values for capsules 1, 2, and 5.  
b. Transport time in seconds at a nominal outlet flow rate of 30 sccm.

### III. DATA COLLECTION AND FUEL PERFORMANCE

#### A. I-135 Released from AGR-1 Test Fuel

Iodine-135 is a fission product radioiodine that decays by beta particle emission with a 6.57 hour half life. It decays to...
excited states in $^{135}$Xe ($T_{1/2} = 9.1$ hours). Among the populated states of $^{135}$Xe is an isomeric state $^{135m}$Xe that decays by isomeric transition to the $^{135}$Xe ground state with a half-life of 15.3 minutes and the emission of a 526.6 keV photon. The 15.3 minute isomeric state is populated through a 16.4% branch of the $^{135}$I decay and is amenable to determination by gamma-ray spectroscopy using the 526.6 keV gamma line[6].

During the AGR-1 irradiation $^{135}$Xe and $^{135m}$Xe concentrations detected at the FPMS are generated both from direct fission yield and from the decay of their $^{135}$I parent. However, by about 2.5 hours (ten 15.3 minute half lives) after the irradiation stops, the only source of $^{135m}$Xe is the decay of its 6.6 hour $^{135}$I parent. Further, decay systematic dictate that the $^{135m}$Xe will be in secular equilibrium with its $^{135}$I parent, its production rate will equal the decay rate of the parent, and its concentration in the flowing stream will appear to decay with the parent half-life [2]. This equilibrium arrangement enables determination of the amount of $^{135}$I released from the fuel particles and deposited in upstream structures from quantification of the $^{135m}$Xe concentrations in the flowing gas stream following reactor shutdown [6]. The ATR operating schedule includes scheduled outages during which the reactor is shut down for a planned amount of time. Calculations of $^{135}$I content from the $^{135m}$Xe activities detected just after shutdown for the first two scheduled outages determined that the activity of released I-135 for each test fuel capsule was less than 0.5 microCuries.

B. AGR-1 Fuel Performance

For intact fuel particles, the TRISO coating provides a barrier to fission product release. However, particles with failed TRISO coatings, either those that fail during irradiation or the tiny percentage of particle that are initially defective, can release fission products directly to the flowing gas stream. Because reactive fission products like the radioiodine’s and cesium’s quickly deposit on test train and piping structures, the only released fission products that move downstream unattenuated are the noble fission gas isotopes of Kr and Xe.

One important measure of the fuel performance in these tests is quantification of the fission gas releases over the duration of each irradiation experiment. These measured released activities support the calculation of isotopic release-to-birth ratios. Release-to-Birth ratios are used to measure fuel performance and are an accepted measurement of fuel quality [7]. The FPMS was used to acquire and analyze the fission gas isotopes of Kr and Xe as is illustrated in fig. 10 and Table II. From spectrometer data we are able to calculate the released fission gas activity. With this information along with calculated birth rates we are able to report the Release-to-Birth (R/B) ratios. We have computed average release-to-birth ratios for four different 10-day periods during the AGR-1 irradiation. Those average release-to-birth ratios for three fission gas nuclides - $^{85}$Kr, $^{88}$Kr, and $^{135}$Xe for test capsule 6 are presented in Fig. 11 as a function of irradiation exposure [effective full power days (EFPD)]. Up to now there has been no indication of a failed TRISO coating.

<table>
<thead>
<tr>
<th>Isotope</th>
<th>Net Micro-Curie/58cc</th>
<th>Relative Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>36KR-85</td>
<td>1.18 x 10^{-3}</td>
<td>12.72%</td>
</tr>
<tr>
<td>36KR-87</td>
<td>4.53 x 10^{-3}</td>
<td>7.11%</td>
</tr>
<tr>
<td>36KR-88</td>
<td>2.66 x 10^{-3}</td>
<td>17.28%</td>
</tr>
<tr>
<td>36KR-89</td>
<td>2.82 x 10^{-2}</td>
<td>1.89%</td>
</tr>
<tr>
<td>36KR-90</td>
<td>7.23 x 10^{-3}</td>
<td>56.03%</td>
</tr>
<tr>
<td>54XE-131</td>
<td>7.58 x 10^{-3}</td>
<td>101.20%</td>
</tr>
<tr>
<td>54XE-133</td>
<td>7.55 x 10^{-4}</td>
<td>59.41%</td>
</tr>
<tr>
<td>54XE-135</td>
<td>5.15 x 10^{-4}</td>
<td>39.52%</td>
</tr>
<tr>
<td>54XE-135</td>
<td>2.53 x 10^{-3}</td>
<td>4.34%</td>
</tr>
<tr>
<td>54XE-137</td>
<td>1.69 x 10^{-2}</td>
<td>2.11%</td>
</tr>
<tr>
<td>54XE-138</td>
<td>1.02 x 10^{-2}</td>
<td>6.61%</td>
</tr>
<tr>
<td>54XE-139</td>
<td>1.21E x 10^{-3}</td>
<td>20.40%</td>
</tr>
</tbody>
</table>

Table II. Concentrations are as measured at the detector uncorrected for decay during transport.
IV. SUMMARY

The Advanced Gas Reactor-1 Fission Product Monitor System has been designed, tested and installed at the Advanced Test Reactor located at the Idaho National Laboratory. Performance and sensitivity of the FPMS has been excellent. It is anticipated that with continued irradiation of the experiment along with continued operation of the FPMS will produce more valuable insights and lessons learned that may be applied to future AGR experiments.

The effluent activities have been analyzed to determine capsule specific fuel performance data such as isotope-specific release-to-birth ratios, and the cumulative release of radiiodines during irradiation. The acquired data indicate excellent fuel performance to date.

REFERENCES


