



# Analyzing existing and planned nuclear facilities as potential sources for unusual xenon radionuclides ( $^{127}\text{Xe}$ , $^{129\text{m}}\text{Xe}$ , $^{125}\text{Xe}$ , $^{122}\text{Xe}$ )

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*Changing the World's Energy Future*

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# **Analyzing existing and planned nuclear facilities as potential sources for unusual xenon radionuclides (127Xe, 129mXe, 125Xe, 122Xe)**

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# An Analysis of Potential Sources of Non-fission Product Xenon Radionuclides ( $^{127}\text{Xe}$ , $^{129\text{m}}\text{Xe}$ , $^{125}\text{Xe}$ , $^{122}\text{Xe}$ )

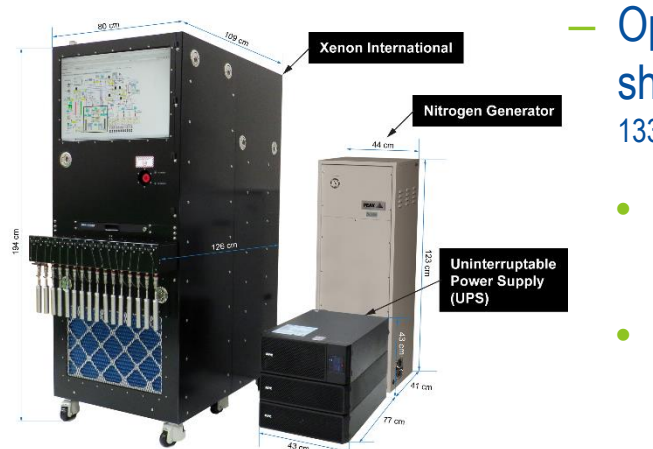
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## Quality Control Standards at Idaho National Laboratory

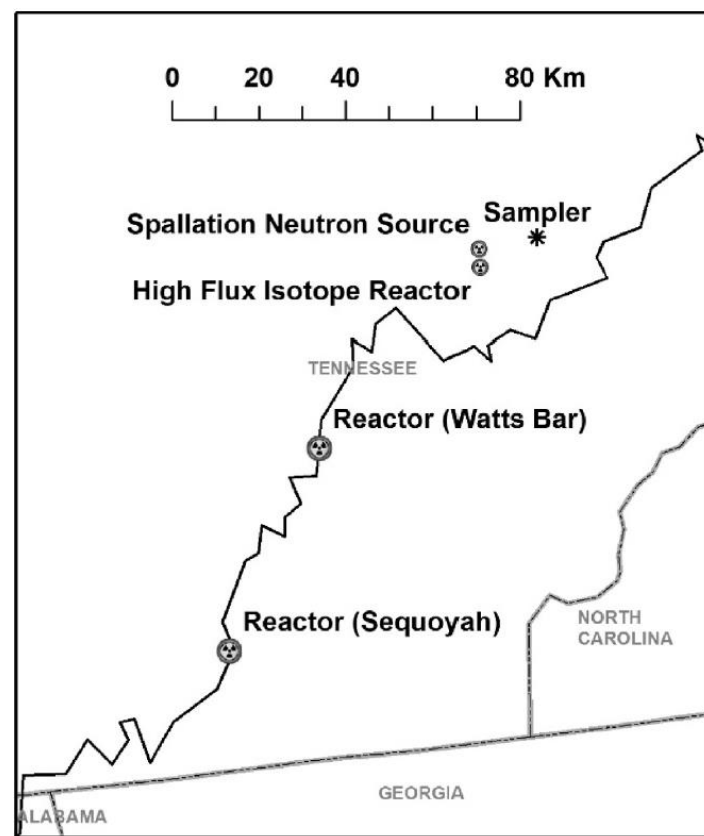
- INL Noble Gas Laboratory (NGL) produces Quality Control standards for laboratories that operate xenon radionuclide monitoring systems.
- Current standards provided to these labs consist of fission product xenon radionuclides ( $^{131\text{m}}\text{Xe}$ ,  $^{133}\text{Xe}$ ,  $^{133\text{m}}\text{Xe}$ , and  $^{135}\text{Xe}$ )
- INL NGL's mission is to provide high quality standards that ensure high accuracy results being reported
  - INL NGL research can demonstrate potential interferences to quantification of fission product xenon radionuclides
  - Through research, INL NGL may demonstrate the need for calibration with other xenon radionuclides not currently in use
    - Potential needs for calibration standards containing non-fission products such as  $^{127}\text{Xe}$ , or  $^{129\text{m}}\text{Xe}$  may exist

## Background: Detection of Non-fission Product Radioxenons

- Testing of a new continuous noble gas monitoring system (Xenon International, Teledyne Brown Engineering) showed some unusual radionuclide detections<sup>1</sup>
- System was operated in proximity (~20 km) of Oak Ridge National Laboratory
  - Operators noted that a large number of samples showed all 4 radionuclides of interest<sup>1</sup> and more  $^{133\text{m}}\text{Xe}$  than  $^{133}\text{Xe}$
  - Samples showing all 4 radionuclides are relatively uncommon when analyzing the atmosphere
  - $^{133\text{m}}\text{Xe}$  is almost never more abundant than  $^{133}\text{Xe}$

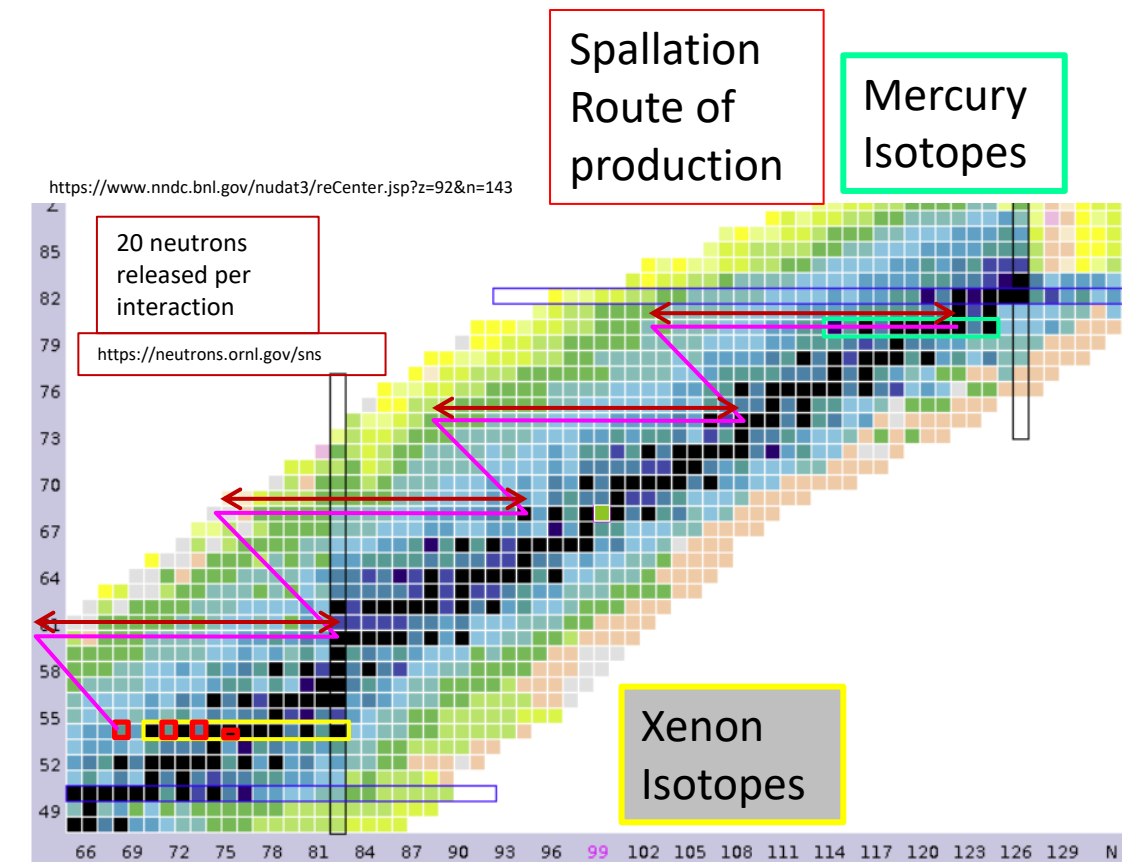


<https://www.tbc.com/en-us/what-we-do/markets/energy/rmg/New-Unit-2023-2>  
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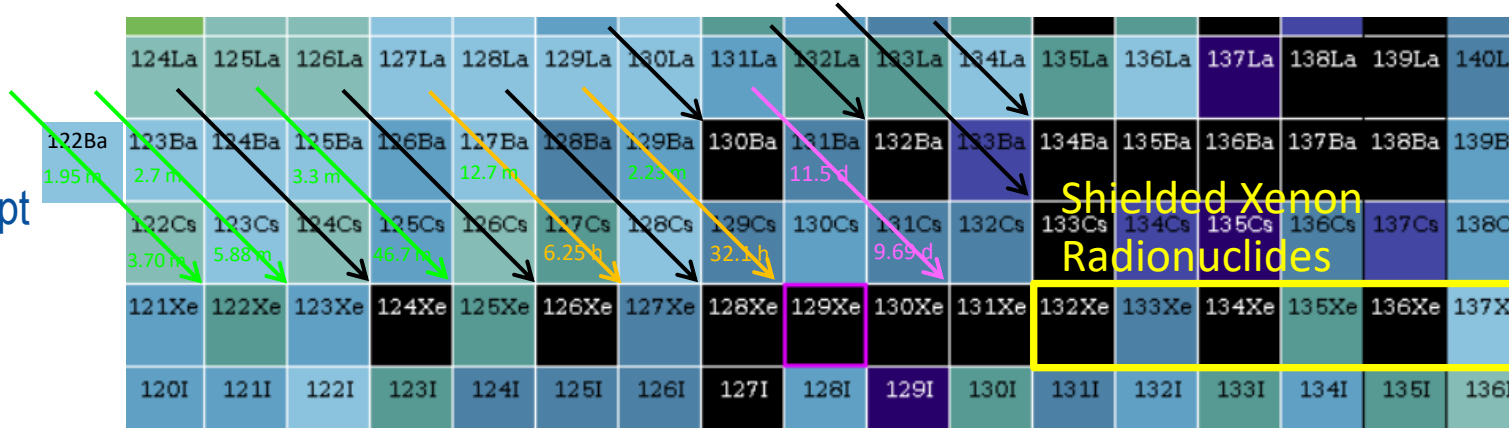
## Production Routes for Non-fission production Radionuclides: SNS

- The path to create  $^{122}\text{Xe}$  from the mercury target (shown in pink zig zag above) requires 4 spallation interactions to reduce mass of mercury to xenon
  - $^{202}\text{Hg}$  (Rel. Abund. = 29.86%) – 4 interactions x (20 neutrons / interaction) =  $^{122}\text{Xe}$



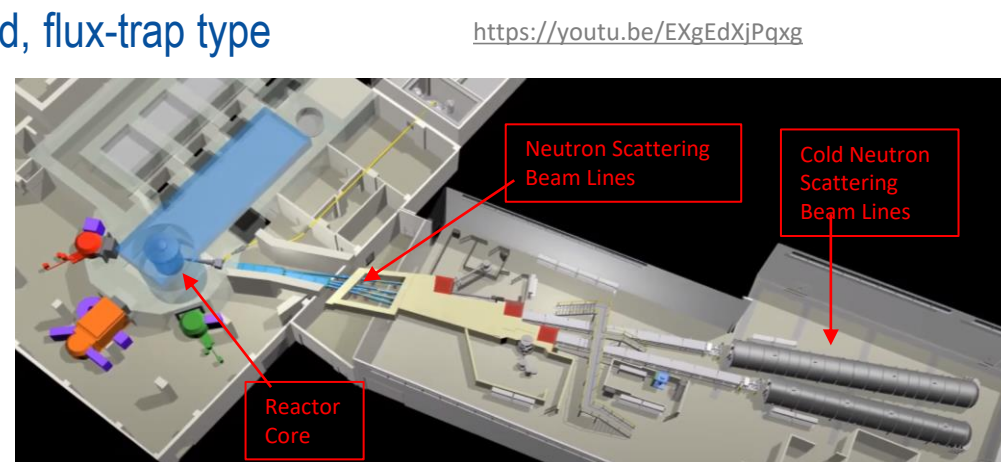
## Radionuclides Shielded from Production Route at SNS

- Spallation Sources produce radionuclides through decay cascades from above
- All fission product xenon radionuclides are shielded except  $^{131\text{m}}\text{Xe}$
- $^{131\text{m}}\text{Xe}$  formation is impeded by relatively long-lived  $^{131}\text{Ba}$  ( $T_{1/2} = 11.5$  d) and  $^{131}\text{Cs}$  ( $T_{1/2} = 9.69$  d)

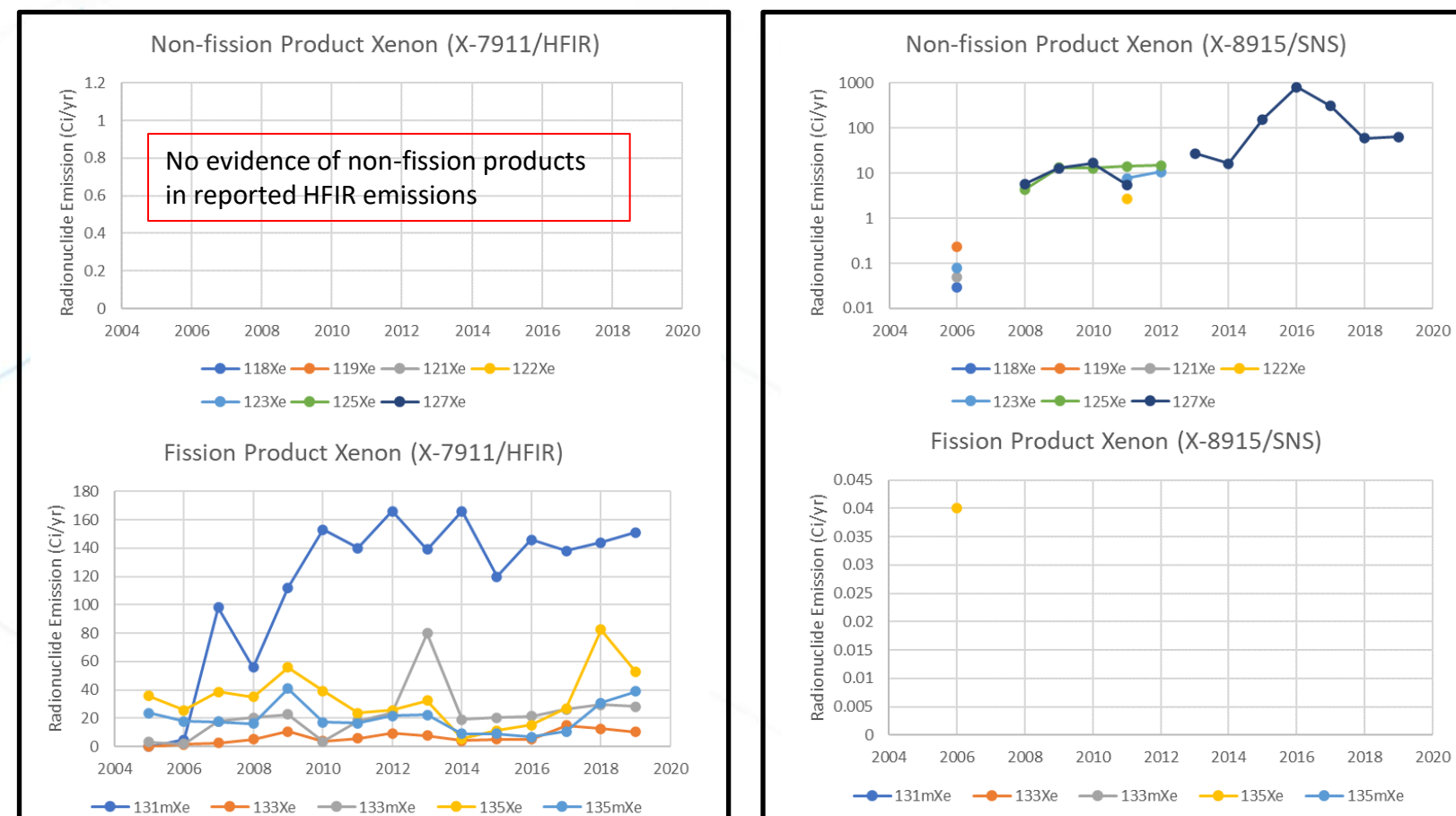


## Potential source#2: High Flux Isotope Reactor (HFIR)

- HFIR: "Beryllium-reflected, light-water cooled and -moderated, flux-trap type reactor" (<https://neutrons.ornl.gov/hfir/parameters>)
- Isotope Production ( $^{252}\text{Cf}$ , others)<sup>4</sup>
- $^{252}\text{Cf}$  Starting Materials =  $^{242}\text{Pu}$ ,  $^{243}\text{Am}$ ,  $^{244}\text{Cm}$
- Neutron Scattering experiments
- [Too many types to list here]



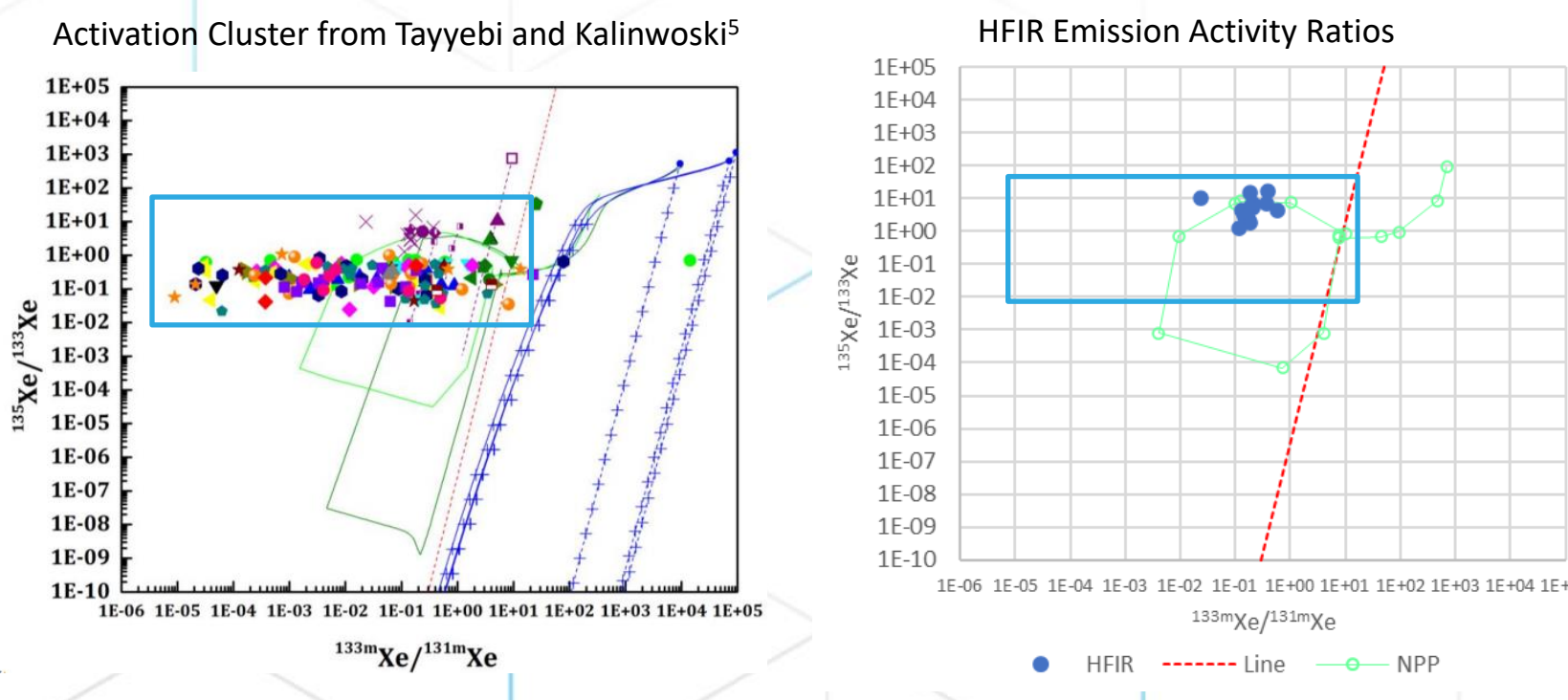
## Reported Emissions from HFIR and SNS



Radionuclide emission data from: Oak Ridge Reservation Annual Site Environmental Reports: years 2005 to 2019

## HFIR emission Activity Ratios align with Activation Cluster

- Isotope Ratio region identified as Activation Cluster by Tayyebi and Kalinowski<sup>5</sup>
- HFIR reported emissions lie within Activation Cluster
- Also lies within the aged fission gas from nuclear power plant region



## Analysis of Another High Flux Reactor: INL Advanced Test Reactor

- Four stack emission grab samples (5 min assay) from 8/10/2023 were averaged to obtain observed daily emission activities
- $^{135}\text{Xe}$  to  $^{41}\text{Ar}$  ratio =  $0.010 \pm 0.002$

## Recreation of $^{135}\text{Xe}$ : $^{41}\text{Ar}$ ratio by Air Activation Modeling (ATR Neutron Flux)

- Using measured neutron fluxes from ATR, the activation of air was modeled
  - $^{135}\text{Xe}$  to  $^{41}\text{Ar}$  ratio (Modeled, no decay) =  $1.30\text{E-}07$
  - $^{135}\text{Xe}$  to  $^{41}\text{Ar}$  ratio (Stack Emission, Observed) =  $0.010 \pm 0.002$
- The ratios can be reconciled by assuming decay (residence time in ATR building before exiting the stack)
  - $^{135}\text{Xe}$  to  $^{41}\text{Ar}$  ratio (Modeled,  $36.7 \pm 1.5$  hr decay) =  $0.010$
  - Other Activation Product emitted in this scenario →

Radionuclide	Classification	Stack Emission (No decay) Ci/day	Stack Emission (36.7 hr Decay) Ci/day
$^{41}\text{Ar}$	Pure Activation	10	10
$^{125}\text{Xe}$	Pure Activation	1.4E-05	5 ± 2
$^{127}\text{Xe}$	Pure Activation	4.6E-09	0.008 ± 0.003
$^{131\text{m}}\text{Xe}$	Activation or Fission	1.4E-07	0.14 ± 0.16
$^{139}\text{Xe}$	Activation or Fission	6.9e-07	0.6 ± 0.1
$^{133\text{m}}\text{Xe}$	Activation or Fission	1.6e-07	0.11 ± 0.01
$^{135}\text{Xe}$	Activation or Fission	1.3e-06	0.09 ± 0.02

If stack emission  $^{135}\text{Xe}$  is from purely activation of air, the quantity of  $^{125}\text{Xe}$  would be easy to detect, but it is not observed/ reported in stack emissions assay.

## $^{125}\text{Xe}$ as an Indicator of Air-Activation non-Fission Products

- Modeling of air activation products (this work) shows that  $^{125}\text{Xe}$  is the largest radioxenon activity when air is activated, next followed by fission product radioxenon ( $^{131\text{m}}\text{Xe}$ ,  $^{133}\text{Xe}$ ,  $^{133\text{m}}\text{Xe}$ , and  $^{135}\text{Xe}$ )
- $^{127}\text{Xe}$  is produced 3 orders of magnitude lower activity than  $^{125}\text{Xe}$  (see plots)
  - $^{125}\text{Xe}$  should be relatively easy to detect in stack emission gamma quantification systems
  - If  $^{125}\text{Xe}$  is found to be present,  $^{127}\text{Xe}$  will likely not be detectable, but is likely present at 3 orders of magnitude lower activity

Radionuclide	Half-life	Classification	Ci/day
$^{41}\text{Ar}$	109.61 min	Activation	$10.0 \pm 0.4$
$^{133}\text{Xe}$	5.2475 day	Activation or Fission	$0.08 \pm 0.04$
$^{135}\text{Xe}$	9.14 hr	Activation or Fission	$0.10 \pm 0.02$
$^{135\text{m}}\text{Xe}$	15.29 min	Activation or Fission	$0.10 \pm 0.02$
$^{139}\text{Xe}$	14.14 min	Activation or Fission	$0.26 \pm 0.04$
$^{87}\text{Kr}$	76.3 min	Activation or Fission	$0.04 \pm 0.01$
$^{88}\text{Kr}$	2.825 hr	Activation or Fission	$0.04 \pm 0.01$

## ATR emissions dominated by Fission Products

- The existence of extremely short-lived (<1 hr) radionuclides in the stack emissions indicates that 36.7 hour decay time for activation products is not feasible
- Significant fraction of fission  $^{135}\text{Xe}$  explains the  $^{135}\text{Xe}$ : $^{41}\text{Ar}$  ratio better than activation and decay

$^{127}\text{Xe}$  emission is negligible compared to fission product radioxenon emitted from high flux reactors (both ATR and HFIR)

## Analysis of Neutron Spallation Sources World-wide

- ORNL SNS is in Knoxville, TN, USA
- Japan operates a similar spallation source to the ORNL SNS (liquid mercury target)
- Solid target (W, Pb) spallation sources are sealed and emit no significant spallation products

## Solid vs. Liquid Spallation Targets

- Liquid targets are continuously purged of neutron spallation products
  - Both Japanese and American treatment systems are similar and developed by shared experience<sup>7</sup>
  - Mercury off-gas treatment systems prioritize separation and emission of spallation products while preventing mercury emissions
- Solid targets (W, Pb, Ta) appear to be operated without any attempts to remove accumulated neutron spallation products
  - Because no attempts are made to remove contaminants while targets are in service, containments are designed with the goal of retaining all spallation products

## Conclusions

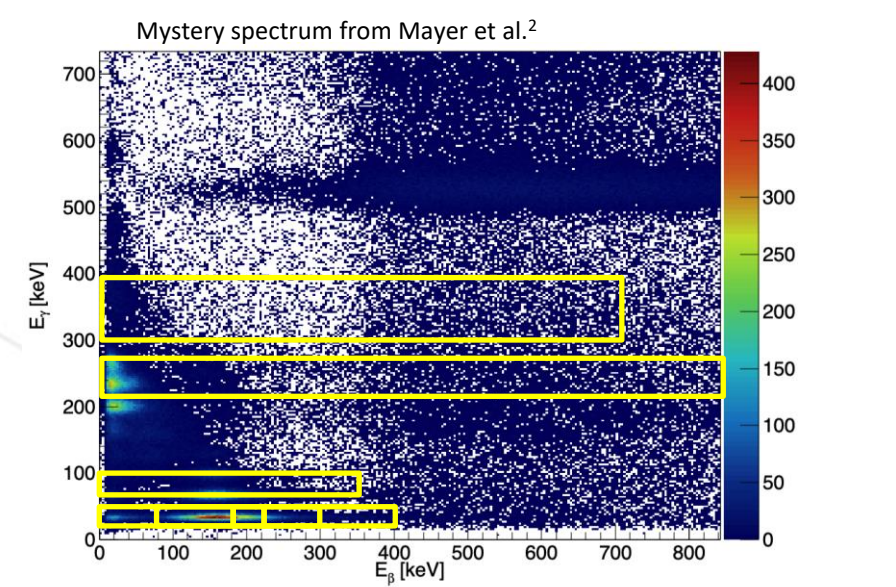
- Liquid mercury target spallation neutron sources are the most likely source of detectable  $^{127}\text{Xe}$ 
  - Proton rich neutron spallation products are produced in the absence of fission product xenon radionuclides
  - Facilities using liquid mercury targets are constantly removing neutron spallation products to prolong the life of mercury target containers
- Solid target spallation neutron sources also produce proton rich neutron spallation products but are more likely to retain produced radionuclides
- Neutron activation of air in nuclear power plants and research reactors produces significantly more  $^{125}\text{Xe}$  than  $^{127}\text{Xe}$ 
  - If gamma spectrum of stack-emissions doesn't show signs of  $^{125}\text{Xe}$ , the emissions are likely dominated by fission gas, and won't be a strong source of  $^{127}\text{Xe}$

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## Unusual Spectra Observed

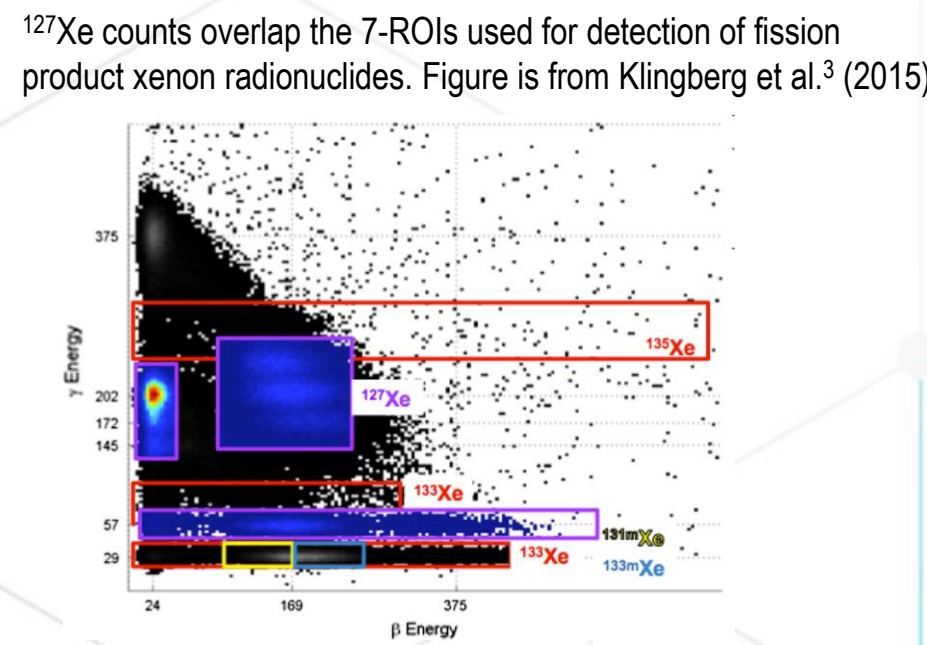
- Testing of a new continuous noble gas monitoring A spectra was published<sup>2</sup> of the unusual count pattern observed
- Although the automatic quantification algorithm showed hits for all 4 radionuclides of interest, it appears that no  $^{135}\text{Xe}$  or  $^{133}\text{Xe}$  is present



ROIs for  $^{131\text{m}}\text{Xe}$ ,  $^{133}\text{Xe}$ ,  $^{133\text{m}}\text{Xe}$ , and  $^{135}\text{Xe}$  are approximated in this plot for comparison. The pattern formed by the observed beta-gamma spectrum do not fit inside the defined ROIs.

## Theorized Radionuclides Responsible for Unusual $\beta/\gamma$ Coincidence Spectra

- Various xenon radionuclides are theorized to contribute to the spectrum recorded during testing of Xenon International
- $^{122}\text{Xe}$ ,  $^{125}\text{Xe}$ ,  $^{127}\text{Xe}$ , and  $^{129\text{m}}\text{Xe}$ , are potentially radionuclides that could create this type of spectrum
- The spectrum of  $^{127}\text{Xe}$  is shown on the right<sup>3</sup> and has counts in the spectra that overlaps with some of the 7 ROIs for determining the 4 radionuclides of interest



## Theorized Radionuclides Responsible for Unusual $\beta/\gamma$ Coincidence Spectra

- Coincidence counts observed during assay of  $^{127}\text{Xe}$  by beta-gamma coincidence counting shows overlap with  $^{135}\text{Xe}$ ,  $^{133}\text{Xe}$ ,  $^{133\text{m}}\text{Xe}$  and  $^{131\text{m}}\text{Xe}$  ROIs
- Because  $^{127}\text{Xe}$  was identified in an environmental sample, it is important to understand the potential for sources of  $^{127}\text{Xe}$
- Operators of systems benefit from understanding the potential of having  $^{127}\text{Xe}$  present in samples
- Understanding types of facilities that have the potential to emit  $^{127}\text{Xe}$  will put operators in mind to watch for  $^{127}\text{Xe}$  interference if the potential exists

## Potential Source#1: Spallation Neutron Source (SNS)

- Power Source = 1.4 MW pulsed (60 Hz) beam (proton) accelerator
- Target Material = Liquid Mercury
- Neutron production Mechanism: High energy protons collide with mercury target atoms to knock out neutrons
- 19 instruments for neutron scattering experiments
- Mercury off-gas treatment system removes radioxenon by-products
  - In operation since 2006

