Test of low radioactive molecular sieves for radon filtration in SF$_6$ gas-based rare-event physics experiments

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ABSTRACT: Type 5Å molecular sieves (MS) have been demonstrated to remove radon from SF$_6$ gas. This is important for ultra-sensitive SF$_6$ gas-based directional dark matter and related rare-event physics experiments, as radon can provide a source of unwanted background events. Unfortunately, commercially available sieves intrinsically emanate radon at levels not suitable for ultra-sensitive physics experiments. A method to produce a low radioactive MS has been developed in Nihon University (NU). In this work, we explore the feasibility of the NU-developed 5Å type MS for use in such experiments. A comparison with a commercially available Sigma-Aldrich 5Å type MS was made. The comparison was done by calculating a parameter indicating the amount of radon intrinsically emanated by the MS per unit radon captured from SF$_6$ gas. The measurements were made using a specially adapted DURRIDGE RAD7 radon detector. The NU-developed 5Å MS emanated radon up to 61±9% less per radon captured (2.1±0.1)$\times 10^{-3}$, compared to the commercial Sigma-Aldrich MS (5.4±0.4)$\times 10^{-3}$, making it a better candidate for use in a radon filtration setup for future ultra-sensitive SF$_6$ gas-based experiments.

KEYWORDS: Gas systems and purification; Radon emanation; Low background experiments; RAD7 radon detector

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1 Introduction

Minimisation of radon contamination is essential in ultra-sensitive gas rare-event physics experiments including directional dark matter experiments [1, 2]. The decay of radon gas in these experiments can produce unwanted background events, able to mimic genuine signals. Radon contamination comes primarily from detector materials. These materials can contain a trace amount of $^{238}\text{U}$ due to inevitable material contamination. The radon gas is produced in the $^{238}\text{U}$ decay chain. Many gas rare-event physics experiments manage radon contamination with continuous flow and disposal of the target gas.

The gas SF$_6$ has been identified as a target gas for use in future directional dark matter searches [3]. However, SF$_6$ is the most potent greenhouse gas, making the method of continuous flow and disposal problematic [4]. Furthermore, climate change initiatives, such as the EU F-Gas directive, heavily regulate the use of SF$_6$ and aim to reduce total use fivefold by 2030, compared with levels in 2014 [5]. This restriction creates a hurdle for ultra-sensitive SF$_6$-based rare-event physics experiments, particularly for large-scale plans such as CYGNUS-1000 which utilises a 1000m$^3$ SF$_6$ gas directional dark matter detector [2]. Therefore, an alternative method to minimise radon contamination must be implemented for future SF$_6$ based experiments, where the gas is reused and recycled.

The demonstration of radon removal from SF$_6$ gas was a significant advance towards a radon filtration system [6]. In principle, the SF$_6$ gas can be continuously recirculated and reused by filtration with the 5Å type molecular sieve, reducing the total amount of SF$_6$ used. Unfortunately, commercial molecular sieves intrinsically emanate radon at levels unsuitable for ultra-sensitive rare-event physics experiments. Commercially available sieves are primarily used in the petroleum industry, where having low radioactive content is not essential, so the manufacturer does not need
to screen materials for radioactive content in production. Therefore, it can be assumed that any commercially manufactured molecular sieves will not meet the required low radioactivity level.

Recently, a method to produce low radioactive molecular sieves was developed in Nihon University. A 5Å type Nihon University developed molecular sieve may provide a suitable candidate for use in a radon filtration setup for ultra-sensitive SF$_6$ gas based rare-event physics experiments. In this paper, we investigate the feasibility of a 5Å type NU-developed molecular sieve for use in a radon filtration setup for ultra-sensitive SF$_6$ gas based rare-event physics experiments. This was done by measuring the radon emanated from the NU-developed molecular sieve and its radon capture efficiency from SF$_6$. These measurements were compared with the commercial, Sigma-Aldrich manufactured, molecular sieve used in the original radon filtration demonstration in 2017 [6].

2 Molecular sieve

Molecular sieves are chemical structures with specific pore sizes. Molecules with a critical diameter equal or smaller than the pore size will diffuse into the pores and be captured. Whereas, molecules with diameters larger than the pore size will not be captured. Instead, the larger molecules will pass between molecular sieve gaps, as shown in figure 1.

![Figure 1](image1.png)

**Figure 1.** Illustration of the separation of smaller radon atom from larger SF$_6$ molecules using molecular sieves. The arrows represent the direction of flow of the gas inside the molecular sieve vessel.

The sizes of the pores are controlled by using different size cations in the ion exchange processes during production. In the 5Å type molecular sieve, the cation used is calcium. The chemical formula for the 5Å type is shown in equation 2.1 [7].

$$0.80\text{CaO} \cdot 0.20 \cdot \text{Na}_2\text{O} \cdot \text{Al}_2\text{O}_3 \cdot 2.0\text{SiO}_2 \cdot x\text{H}_2\text{O}.$$  \hspace{1cm} (2.1)

The source of radioactive contamination in molecular sieves is from raw materials used in production, mainly the metallic components. In commercial production, the screening of raw materials for radioactive content is not necessary. H.Ogawa et al., from Nihon University in collaboration with Union showa K.K, has developed a method of producing low radioactive molecular sieves by extensive material selection and screening of raw materials used. The 5Å type NU-developed MS was made by exchanging calcium ions with the low-radioactive 4Å type MS mentioned in [8]. In figure 2, the 5Å type MS samples used in our comparison are shown. On the left is the
NU-developed, in the form of white irregular granules with approximate size of 1-2 cm. On the right is the MS manufactured by Sigma-Aldrich in uniform beige beads with 8-12 mm diameter.

Figure 2. Images of the 5Å type molecular sieves tested. The NU-developed MS is on the left and Sigma-Aldrich MS on the right.

When choosing a suitable molecular sieve for a radon filtration system, it is important to understand how radon behaves in a sealed chamber. A gas-based detector is fundamentally a sealed gas chamber with various materials which can emanate radon. Once the chamber is sealed, the radon concentration inside the vessel increases continuously until a steady-state is reached. This is when the production rate of radon is equal to the decay rate of radon, termed secular equilibrium.

The steady-state radon concentration can be considered as the maximum radon activity in the chamber, applying a molecular sieve filter reduces this maximum. The determining mechanism now is the equilibration between the amount of radon captured by the MS and the amount of radon intrinsically emanated by the MS.

To determine the suitability of a MS candidate, both the amount of radon intrinsically emanated from the sieve and its radon capturing efficiency must be measured. The ideal MS having a minimal intrinsic emanation whilst having an optimal capturing efficiency.

3 Radon emanation test

Radon is a gaseous element produced in both the $^{238}\text{U}$ and $^{232}\text{Th}$ decay chains. Most materials contain a trace amount of $^{238}\text{U}$ and $^{232}\text{Th}$ due to inevitable material contamination, which is exacerbated by their extremely long half-lives. When radon is produced within a material, due to its gaseous nature, it can escape and emanate into its surroundings. The radon isotope $^{222}\text{Rn}$, produced from the $^{238}\text{U}$ decay chain, is the most problematic isotope for contamination because $^{222}\text{Rn}$ has a half-life of 3.8 days, meaning it has the ample time to reach the material surface and travel within the detector before decaying. In contrast, the second most abundant radon isotope, $^{220}\text{Rn}$, produced from the $^{232}\text{Th}$ decay chain, has a half-life of 55 seconds. Hence, tests reported here focused on measurements from emanation of the $^{222}\text{Rn}$ isotope.
Experimental Setup The radon emanation test for the molecular sieves were performed using the experimental setup shown in figure 3. The setup consist of a 3.5L stainless steel emanation chamber in a loop with two DURRIDGE RAD7 radon detectors. The RAD7 collects radon by electrostatic precipitation, measuring alpha decays from radon progeny such as $^{218}\text{Po}$ and $^{214}\text{Po}$ [9]. The loop includes a tee connection where either an EDWARDS vacuum scroll pump or an input of low-humidity low-activity nitrogen gas was connected. The MS of interest was enclosed inside the **Molecular Sieve Container** with meshed O-rings to stop any small fragments escaping from the vessel during evacuation.

Method The emanation testing procedure started with two rounds of 1-hour gas evacuations using the EDWARDS pump. The first evacuation was to get the emanation chamber into sub-torr pressures, inducing outgassing from materials. Since the initial outgassing was expected to be dominated by the emanation chamber, the gas was evacuated again after an outgassing period of 48 hours. After the second gas evacuation, the MS was left to emanate into the vacuum for seven days. On completion of the emanation period, the chamber was backfilled with nitrogen gas. It was essential to use low activity-low humidity nitrogen gas to backfill the chamber. Low activity nitrogen gas was required to minimise the introduction of any ‘new’ radon to the system. Low-humid gas was required because increased humidity can suppress radon counts measured by the RAD7. Low humidity-low activity gas was achieved by flowing the nitrogen through a desiccant, which removes water molecules, and a charcoal filter cold trap, which removes any radon introduced by the nitrogen tank. Once the chamber was backfilled to 760 torr, the RAD7 detectors were purged with nitrogen for 5 minutes to remove any potential residue inside the detector. Finally, the RAD7 with its internal pump on, recorded radon concentration every four hours for a 48-hour measurement period.

Data Analysis The total radon emanating from the molecular sieves and setup is equal to the radon activity at secular equilibrium. For $^{222}\text{Rn}$ to reach secular equilibrium, an emanation time in the order of a month is required. However, in our molecular sieves emanation testing, the samples
were only left to emanate for seven days. To compensate for the shorter emanation time, equation 3.1, which describes the accumulation of radon in a closed system as a function of emanation time was fitted to the measured data.

\[
A_m = A_s \times \left(1 - \exp\left(-\frac{t}{\tau}\right)\right),
\]

(3.1)

where \(A_m\) is the measured radon activity, \(A_s\) is the radon activity at secular equilibrium, \(t\) is the emanation time and \(\tau\) is the lifetime of \(^{222}\text{Rn}\). Before the raw concentration data was fitted to equation 3.1, radon concentration (Bq/m\(^3\)) was converted to radon activity (Bq). This was achieved by multiplying the radon concentration by the total volume in the emanation measurement loop. The total volume is given by equation 3.2. Where \(V_{EC}\) is the volume of the emanation chamber (3.5 L), \(V_R7\) is the volume of the RAD7 (0.9 L) and \(V_{MS}\) is the volume of the molecular sieve container including the approximate volume occupied by the molecular sieve sample.

\[
V_T = V_{EC} + 2 \times V_{R7} - V_{MS},
\]

(3.2)

A correction to ensure there was ample radon mixing time in the measurement loop for radon to distribute evenly, was also applied. This was done by disregarding the first cycle of the RAD7 data. The nominal flow rate of the RAD7 pump was 1.0 LPM. Therefore, one measurement cycle corresponds to approximately six full system recirculation for mixing.

The total radon emanating from the molecular sieves was calculated by extrapolating the radon activity at secular equilibrium, \(A_s\), from the equation 3.1 fit, and subtracting it with the background radon emanation of the setup.

**Results**

The radon concentration as a function of time is shown in figure 4 (NU-developed MS on the left and Sigma-Aldrich MS on the right). Equation 3.1 non-linear regression fit are shown as dashed red lines. The extrapolated radon emanation from this fit are shown in table 1. Note that background emanation of the setup with an empty molecular sieve container was measured to be 4.69±1.1 mBq and subsequently subtracted from the results. Note that the relative humidity correction applied in section 4 was not required as the humidity was below 15% throughout.

![Figure 4](image-url)

*Figure 4.* Plot of radon concentration as a function of emanation time. NU-developed MS on the left and Sigma-Aldrich MS on the right.
Table 1. Results of the MS intrinsic radon emanation test.

| Molecular Sieve   | $^{222}$Rn Emanated (mBq) | Amount of MS Used (g) | $^{222}$Rn Emanated per kg (mBq/kg) |
|-------------------|---------------------------|----------------------|------------------------------------|
| NU-developed      | 6.95±1.64                 | 70                   | 99±23                              |
| Sigma-Aldrich     | 45.2±3.2                  | 86                   | 525±37                             |

It can be seen from table 1 that the NU-developed MS has significantly lower intrinsic radon emanation per unit mass by about a factor of 5. This result makes a promising case for the NU-developed MS to be used in radon filtration setups for ultra-low background SF$_6$-based physics experiments. However, to make a complete comparison, the radon capture efficiency of the MS must also be considered.

4 Radon filtration test

Previously it was found that radon atoms have a critical diameter between 4 and 5Å, while SF$_6$ molecules have a critical diameter between 5 to 10Å [6]. This makes it possible to remove radon atoms from SF$_6$ with molecular sieves with 5Å sized pores. In the radon filtration test, the amount of $^{222}$Rn captured by the molecular sieves from radon-contaminated SF$_6$ is measured.

Experimental Setup The radon filtration tests were performed using the experimental setup shown in figure 5. The setup consists of the molecular sieve filter in a loop with a RAD7 radon detector and a gas reservoir. Also, included in the loop were connections to a passive PYLON 5.4kBq radon source; an SF$_6$ gas input; an EDWARDS vacuum pump and an MS filter bypass. The setup used several Swagelok valves to direct the gas flow either via the molecular sieve filter or MS bypass. Note that the RAD7 has a pump inside with a nominal flow of 1.0 litre per minute.

Figure 5. Schematic of the setup used for the radon filtration tests.
**Method**  The filtration testing procedure started with the evacuation of the setup and the introduction of low humidity SF$_6$ gas to 760 torr. The gas was then contaminated with radon by opening the valve to the PYLON radon source, which was left to diffuse for 15 minutes. The valves were initially configured so that the gas flow direction was through the MS bypass. With the pump on, the RAD7 radon detector measurement started, recording every hour. After about 24 hours, the molecular sieve filter was engaged by opening the valves at both ends of the filter and closing the MS bypass valve. The RAD7 radon detector continued to measure for at least a further 20 hours.

**Data Analysis**  To calculate the amount of radon captured by the MS, the data was split into *MS off* and *MS on* data sets, which correspond to the measured data when the gas flow was directed via the MS bypass and MS filter, respectively. Both data sets were converted from radon concentration (Bq/m$^3$) to radon activity (Bq), by multiplying the radon concentration with the total volume in the radon filtration setup. The total volume was approximated to be the sum of the gas reservoir (6L) and the volume of the RAD7 (0.9L).

Two corrections were applied to the radon activity data before calculating the amount of radon captured. The first correction accounts for relative humidity (RH) in the chamber. A high concentration of water molecules in the chamber can suppress radon counts in the RAD7 detector. To compensate for lost radon counts, at RH higher than 15%, equation 4.1 provided by DURRIDGE was applied to correct for high humidity. In this equation, $A_m$ is the radon activity measured, $A_c$ is the corrected radon activity and $RH$ is the associated relative humidity with the radon measurement.

$$A_c = A_m \times \frac{100}{116.67 - 1.1 \times RH}.$$  

The second correction applied ensures there was ample time for the radon from the PYLON source to distribute evenly throughout the system. Therefore, the first 4 hours of the data were disregarded. Using the corrected data, the radon reduction due to the application of the MS was determined by fitting the *MS off* and *MS on* data sets to decay equation 4.2. Note that the *MS on* fit was optimised by only including data after a new equilibrium was reached.

$$A(t) = A_0 \exp (-\lambda t).$$

$A(t)$ is the radon activity at time $t$, $A_0$ is the initial radon activity and $\lambda$ is the radon decay constant. The initial radon activity, $A_0$, for both *MS off* and *MS on* data sets were extrapolated, and the effective radon reduction was determined by calculating the difference between the extrapolated $A_0$ values.

**Results**  The radon activity as a function of time is shown in figure 6 (NU-developed MS on the left and Sigma-Aldrich MS on the right). The *MS off* and *MS on* non-linear regression fits are shown in red and green dashed lines, respectively. The highlighted green area represents that the molecular sieve filter was engaged. The blue points represents the relative humidity in the setup. The implementation of the molecular sieves, resulted in a drop in relative humidity, as expected. In addition, it is clear that the effective radon activity also decreases, results shown in table 2. The NU-developed MS demonstrates the removal of radon from SF$_6$. However, at a lower radon capture efficiency ($35\pm2$ Bq kg$^{-1}$) compared to the Sigma-Aldrich MS ($97\pm1$ Bq kg$^{-1}$).
Figure 6. Plot of the change in radon concentration observed due to application of molecular sieve. NU-developed MS on the left and Sigma-Aldrich MS on the right.

Table 2. Results of the radon filtration test.

| Molecular Sieve | $^{222}$Rn Captured (Bq) | Amount of MS Used (g) | $^{222}$Rn Captured per kg (Bq kg$^{-1}$) |
|-----------------|--------------------------|----------------------|------------------------------------------|
| NU-developed    | 2.39±0.10                | 68                   | 35±2                                      |
| Sigma-Aldrich   | 9.67±0.12                | 100                  | 97±1                                      |

5 Molecular sieve feasibility

To provide a complete comparison of the molecular sieve candidates, the results from the emanation and filter tests were combined. A parameter indicating the amount of radon emanated by the MS per radon captured is shown in table 3. This parameter was used as the figure of merit for the suitability of the MS for use in radon filtration setups for ultra-low background SF$_6$-based physics experiments, with a value that needs to be minimised. The NU-developed 5Å MS emanated radon 48±15% less per radon captured, compared to the commercial Sigma-Aldrich MS. This result shows that the NU-developed MS is a better candidate.

Table 3. Calculated comparison parameter, indicating the amount of radon emanated by the MS per radon captured by the MS from SF$_6$.

| Molecular Sieve | $^{222}$Rn Emanated per $^{222}$Rn captured ($\times 10^{-3}$) |
|-----------------|---------------------------------------------------------------|
| NU-developed    | 2.8±0.7                                                       |
| Sigma-Aldrich   | 5.4±0.4                                                       |

To determine if the NU-developed MS activity levels are acceptable for ultra-low background SF$_6$ gas-based physics experiments, let us consider a practical example. The DRIFT (Directional Recoil Identification From Tracks) Experiment is a gas-based direction dark matter experiment with
chamber dimensions of 1.5 x 1.5 x 1.5 m³. The radon activity due to material contamination in the experiment was measured to be 372±66 mBq [1]. The NU-developed MS granules was measured to capture 35±2 Bq kg⁻¹. Therefore, it is theoretically possible to use about 1g of the granules to reduce the radon activity in the detector chamber. Not only would this be an engineering challenge but using 1g to capture a limited amount of radon atoms in such a large volume would make the timescales unrealistic. Since radon capture is a probabilistic kinetic process, using more sieves would reduce the timescale required to reach radon adsorption equilibrium. Therefore, depending on the radioactive tolerances of the experiment, it is recommended an amount considerably more than required of molecular sieve should be used. For instance, using 1 kg of NU-developed MS granules is more than sufficient to capture all the radon (372±66 mBq) in the 3.4 m³ DRIFT-II experiment chamber, while only contributing 99±23 mBq.

6 Macroscopic geometry optimisation

The radon reduction result for the NU-developed MS is significantly less than the Sigma-Aldrich MS result, seen in table 2. Optimising the radon filtration for NU-developed MS could possibly improve the radon emanated per radon captured parameter. The filtration discrepancy may be attributed to the difference in their geometries and surface area to volume ratio, seen in figure 2. In a bid to improve the radon capturing capabilities of the NU-developed MS, its surface area to volume ratio was increased by crushing it from its irregular granular form into a fine powder with a mortar and pestle, as shown in figure 7. The radon filtration test, discussed in section 4, was repeated with the NU-developed MS in its powder form. It is worth noting that the significant increase in surface area to volume ratio has implications that there will be an increase in the amount of radon emanated from the MS, as the path from inside the material to the surface is reduced. Therefore, the emanation test, discussed in section 3, was also repeated with the powder form.

![Figure 7](image_url). Images of the NU-developed MS before and after crushing it into a fine powder.

The results for the powder form of the NU-developed MS test are shown in table 4, with the corresponding filtration and emanation plots in figure 8. The radon reduction efficiency has significantly improved from 35±2 to 330±3 Bq kg⁻¹. As expected, the intrinsic radon emanation has significantly worsen from 99±23 to 680±30 mBq. However, the radon emanated per radon captured parameter has remained the same within errors. Overall, crushing the MS into powder did not improve suitability for use in radon filtration setups for ultra-low background as demonstrated by our parameter.
Figure 8. Plot of the change in radon concentration observed while using the powdered NU-developed MS (left) and a plot of radon concentration as a function of emanation time (right) in the powdered NU-developed MS emanation test.

Table 4. Radon filtration, intrinsic MS emanation and comparison parameter results for the NU-developed MS in granule and powdered form. Note that 85g of NU-developed MS were used in the powdered tests.

| NU-developed MS | $^{222}$Rn Captured per kg (Bq kg$^{-1}$) | $^{222}$Rn Emanated per kg (mBq kg$^{-1}$) | $^{222}$Rn Emanated per $^{222}$Rn Captured ($\times 10^{-3}$) |
|----------------|------------------------------------------|------------------------------------------|------------------------------------------|
| Granules       | 35±2                                     | 99±23                                    | 2.8±0.7                                  |
| Powder         | 330±3                                    | 680±30                                   | 2.1±0.1                                  |

7 Conclusions

In this work, a method to determine the suitability of a molecular sieve candidate for use in ultra-sensitive SF$_6$ gas-based rare-event physics experiments was proposed. A parameter indicating the amount of radon intrinsically emanated per unit radon captured by the sieve from SF$_6$ was experimentally determined. A low radioactive MS candidate developed by Nihon University in collaboration with Union Showa K.K was tested and compared to a commercially manufactured Sigma-Aldrich MS. It was found that the NU-developed MS emanated 48±15% less per radon captured (2.8±0.7)$\times 10^{-3}$, compared to the commercial Sigma-Aldrich MS (5.4±0.4)$\times 10^{-3}$. An attempt to improve this result was made by crushing the MS into a powder, increasing its surface area to volume ratio. The powder form significantly increased the radon capture efficiency. However, the intrinsic radon emanation also worsened. Overall, there was no change in the emanation per radon captured parameter for the NU-developed MS powder (2.1±0.1)$\times 10^{-3}$. The total reduction in our parameter with the powdered NU-developed MS compared to the Sigma-Aldrich MS was 61±9%, which is within errors of the original granulated NU-developed MS. In practice, when constructing a radon-filtration setup, it is recommended an amount of MS considerably more than required be used. This is because radon capture is a probabilistic kinetic process, using more sieves will reduce the timescale required to reach radon adsorption equilibrium within the gas detector. The NU-developed MS provides a suitable candidate for use in a radon filtration setup for future ultra-sensitive SF$_6$ gas-based experiments. However, efforts towards minimising the emanation per radon
captured parameter must continue so that the total amount of MS allowed by the radioactive budget of an experiment is maximised.

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