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Souza, E.B.D., Park, B.J., Adhikari, G. et al. (48 more authors) (2020) Study of cosmogenic radionuclides in the COSINE-100 NaI(TI) detectors. Astroparticle Physics, 115. 102390. ISSN 0927-6505

https://doi.org/10.1016/j.astropartphys.2019.102390

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# Study of cosmogenic radionuclides in the COSINE-100 NaI(Tl) detectors

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by looking for nuclei recoiling from dark matter-nucleus scattering [1, 2] and report null results. One notable exception is the DAMA/LIBRA experiment that has consistently reported the observation of an annual event-rate

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Preprint submitted to Astroparticle Physics

for three different configurations of the detector, it remains controversial because it is in conflict with the bounds from other direct detection experiments using different target materials [5, 6, 7, 8, 9, 10] and indirect searches [11]. However, since these conflicts depend on the details of the models for dark matter-nucleus scattering [12] and the properties of the galactic dark matter halo [13, 14, 15], a conclusive statement about the DAMA/LIBRA signal can only be made by conducting an independent experiment using the same NaI(Tl) target material. This is the

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Figure 1: (a) The COSINE-100 detector. From outside inward, the four shielding layers include: 3 cm thick plastic scintillator panels (green), 20 cm of lead (khaki), a 3 cm thick copper box (orange), and liquid scintillator (light blue). (b) A side view of the detector geometry used in the Geant4 simulations.

prime motivation of COSINE-100 and a number of other NaI(Tl)-crystal-based experiments [16, 17, 18, 19, 20]

COSINE-100 is a dark matter direct detection experiment [21, 22] that uses a 106 kg array of eight lowbackground NaI(Tl) crystals situated in a 2000 liter liquid scintillator veto counter. The experiment is located 700 m underground at the Yangyang Underground Laboratory (Y2L), where it has been operating since September 2016. The search for an annual modulation signal requires a complete understanding of background sources and their time dependence. To accomplish this, a complete simulation that accurately models the background energy spectra measured in the detector is required [23]. In addition to backgrounds from long-lived radioactive contaminations in the crystal bulk and surfaces, we have to deal with time-dependent backgrounds from short-lived cosmogenically activated radionuclides. These are isotopes that are created by interactions of cosmic rays with stable nuclides in the detector material. In COSINE-100, almost all of the cosmogenic isotopes come from cosmic ray interactions with either Na or I nuclei.

This paper is organized as follows. The COSINE-100 detector is described in section 2. In section 3, the cosmogenic isotopes that are produced in NaI(Tl) are listed and the determination of the activity levels at the time of their initial deployment underground at Y2L is described. The use of these initial activity levels to infer production rates for cosmogenic isotopes at sea level and their comparison with ACTIVIA and MENDL-2 calculations [24, 25] and with experimental data are discussed in section 4. The fitted activities of <sup>3</sup>H and <sup>129</sup>I from the background modeling are evaluated in section 5 and conclusions are provided in section 6.

## 2. The COSINE-100 experimental setup

The experimental setup of COSINE-100, shown in Fig. 1(a), is described in detail in Ref. [21]. Eight NaI(Tl) crystals, arranged in two layers, are located in the middle of a four-layer shielding structure. From outside inward, this comprises plastic scintillator panels, a leadbrick castle, a copper box, and a tank of scintillating liquid. The eight encapsulated NaI(Tl) crystal assemblies and their support table are immersed in the scintillating liquid that serves both as an active veto and a passive shield. The eight NaI(Tl) crystals were grown from powder provided by Alpha Spectra (AS). Two crystals (Crystal-5 and Crystal-8) are not considered in this paper because their low light yields result in poorer energy resolution and because of their relatively high background contamination levels, especially at low energies.

Since the detailed cosmic ray exposure history of each crystal is unknown, we estimated the time period for each crystal's exposure, listed in Table 1, from the time between the powder production by Alpha Spectra at Grand Junction, Colorado, to the date delivered to Y2L. We considered that the preparation of the NaI powder precedes the crystal growth date by 2 months. It includes 30 days as the duration of transportation to Y2L. Since Crystal-3 has a complicated exposure history, having been repaired once before deployment at Y2L, we can only be certain that the corresponding period is more than 9 months. The radioactivity cooling time for each crystal between delivery to Y2L and the start of data-taking is also listed in Table 1.

The six crystals that are considered in this analysis have light yields of about 15 photoelectrons/keV; the energy threshold for an accepted signal from a crystal is 2 keV. Selection criteria that are used to isolate scintillation-light

Table 1: Mass, dimensions, powder, surface exposure, and underground radioactivity cooling times for each one of the analyzed crystals (see text).

Crystal	Mass	Size	Powder type	Exposure time	Radioactivity
	(kg)	$(diameter \times length)$		(years)	cooling time at Y2L
		(inches)			(years)
Crystal-1	8.3	$5.0 \times 7.0$	AS-B	2.17	3
Crystal-2	9.2	$4.2 \times 11.0$	AS-C	0.92	2.75
Crystal-3	9.2	$4.2 \times 11.0$	AS-WSII	> 0.92	1.2
Crystal-4	18.0	$5.0 \times 15.3$	AS-WSII	1.83	0.5
Crystal-6	12.5	$4.8 \times 11.8$	AS-WSIII	0.5	0.6
Crystal-7	12.5	$4.8 \times 11.8$	AS-WSIII	0.5	0.6

Table 2: Cosmogenic radionuclides in the NaI(Tl) crystals identified in other studies and taken into consideration here.

Cosmogenic	Half-life $[27, 28, 29]$	Decay type
isotopes	(days)	& Emissions energy
$^{125}I$	59.4	EC, 35.5+31.7=67.2 keV
$^{121}\mathrm{Te}$	19.17	EC, $4.1-4.7$ and $30.5 \text{ keV}$
$^{121m}$ Te	164.2	EC, $4.1-4.7$ and $30.5 \text{ keV}$
$^{123m}$ Te	119.3	IT, 247  keV
$^{125m}\mathrm{Te}$	57.4	IT, $145 \text{ keV}$
$^{127m}\mathrm{Te}$	106.1	IT, 88  keV
$^{113}$ Sn	115.1	EC, $3.7$ – $4.2$ and $28 \text{ keV}$
$^{109}\mathrm{Cd}$	462	EC, $25.5$ and $88 \text{ keV}$
$^{22}$ Na	950	$\beta^{+}, 511 \text{ and } 1274.6 \text{ keV}$
$^{3}\mathrm{H}$	4494	$\beta^{-}$
$^{129}$ I	$1.57{\times}10^7 { m yr}$	$\beta^{-}$

generated signals from photomultiplier tube noise are described in detail in Ref. [21]. Events that have abovethreshold signals in only one of the crystals and none in any of the other crystals or the liquid scintillator are classified as *single-hit* events. Those with above-threshold signals in more than one crystal and/or the liquid scintillator are classified as *multiple-hit* events.

Monte Carlo simulations based on the Geant4 toolkit [26] are used to better understand the background spectra from the cosmogenic isotopes in the crystals; the geometry used for these simulations is shown in Fig. 1(b).

## 3. Cosmogenic radionuclides and initial activities

Although the eight NaI(Tl) crystals had underground radioactivity cooling times that range from several months to three years, there are still backgrounds from the longlived cosmogenic isotopes that were activated by cosmic rays while they were on the surface.

To understand these backgrounds, we first considered the list of cosmogenic radioactive isotopes that are produced in NaI(Tl) reported in Ref. [30, 31, 32, 33]. In Table 2, we list the contributing cosmogenic isotopes with their half lives and decay modes; short-lived isotopes, for which half lives are less than a year, are <sup>125</sup>I, <sup>121</sup>Te, <sup>121m</sup>Te, <sup>123m</sup>Te, <sup>125m</sup>Te, <sup>127m</sup>Te, and <sup>113</sup>Sn and long-lived isotopes are <sup>109</sup>Cd, <sup>22</sup>Na, <sup>3</sup>H, and <sup>129</sup>I. Since there are no characteristic peaks from the decay of  $^{123m}\text{Te}/^{125m}\text{Te}$  in the low energy below 100 keV their contributions are negligible in all crystals in Table 1 and, thus, they are not further considered in the analysis.

The short-lived  $(T_{1/2} < 1 \text{ y})$  isotopes are not expected to contribute significantly to either Crystal-1 or Crystal-2 because their cooling times are long enough to reduce these activities to a negligible level. However, we expect some backgrounds from the short-lived isotopes in other crystals because their production rates at sea level, as estimated in [30, 31, 32, 33], are high and their cooling times are less than or equal to a year.

Data points in Fig. 2 show the energy spectra for the six considered NaI(Tl) crystals during the first (blue) and last (green) 25 day segments of the dataset taken between October 21, 2016 and July 18, 2018. A significant reduction of peaks from short-lived cosmogenic isotopes in Crystals 4, 6, and 7 for both single- and multiple-hit events is evident, while the differences for Crystals 1 and 2 are small, as expected. To associate the specific peaks with its cosmogenic nuclide, we simulated each isotope in Table 2 as a radioactive contaminant randomly distributed inside the NaI(Tl) crystal bulk. Fig. 3 shows the differences between the initial and final data segments for Crystal-4. The subtracted spectrum is well fitted by the simulated cosmo-



Figure 2: Background spectra for six NaI(Tl) crystals during the first (blue points) and the last (green points) 25 days of the dataset taken from October 21, 2016 to July 18, 2018. The upper plots show single-hit events and the lower ones show multiple-hit events.

genic components that are treated as parameters floating in the fit, thereby validating our selection of the main cosmogenic contributors to the low-energy single-hit distribution. However, the derived weight of each isotope from the fit are not further considered in the analysis. Those two structures at about 12 and 48 keV in Fig. 3 are charac-



Figure 3: Difference between the first and last 25 day spectra of the 1.7 year data for Crystal-4 (in black) together with the fitted spectrum (in red) from several simulated individual contributions (in several colors). The subtracted spectra in Crystals-4, 6, and 7 are similar to each other due to their relatively short cooling times underground. However, the short-lived isotopes are not expected to contribute to either Crystal-1 or Crystal-2 due to their relatively long cooling times underground.

teristic of  $^{210}$ Pb. It decays over time with the half-life of 22.3 yr and there is a little difference between the first and last 25 day spectra of the 1.7 years data.

Four long-lived nuclides, <sup>109</sup>Cd, <sup>22</sup>Na, <sup>3</sup>H, and <sup>129</sup>I have low energy deposits and are, therefore, potentially troublesome. It is essential to understand their background contributions to the low energy spectra regions, especially in the (2–6) keV dark matter signal region of interest (ROI). The beta-decay spectrum of tritium has an endpoint energy of 18 keV and the electron capture decay of <sup>22</sup>Na produces 0.87 keV emissions. The beta decay of <sup>129</sup>I to <sup>129</sup>Xe\* is followed by <sup>129</sup>Xe\* transitioning to the stable <sup>129</sup>Xe isotope via the emission of a 39.6 keV  $\gamma$ -ray. Its spectral feature from this process has a distribution with a peak around ~45 keV. The electron capture decay of <sup>109</sup>Cd contribute peaks at 25.5 and around 3.5 keV.

Because it is impossible to compute the initial activities of the cosmogenic radioisotopes from the production rates without knowing their detailed cosmic ray exposure conditions: i.e., time, location, altitude, etc. [30], we, when possible, extrapolated the time-dependent reduction of characteristic peaks from their decays to determine their activity levels at the time of their initial deployment at Y2L. For the activities of the long-lived <sup>22</sup>Na and <sup>109</sup>Cd isotopes we investigated temporal and spatial correlations of characteristic  $\gamma$ /X-rays peaks produced in their decays. The details of these technique are discussed in the following sections.

Table 3: Initial activity  $A_0$  (mBq/kg) of  $^{125}I$  in each crystal as measured by the decay rate method. This includes the statistical uncertainty.

	Crystal-1	Crystal-2	Crystal-3	Crystal-4	Crystal-6	Crystal-7	
$^{125}I$	-	-	$9.0{\pm}0.9$	$3.4{\pm}0.1$	$5.1 {\pm} 0.2$	$5.4 {\pm} 0.2$	



Figure 4: Average rate in 60-70 keV region of Crystal-4's single-hit spectrum over time. The rate shows a clear decrease, which can be modeled with an exponential decay, for  $^{125}$ I in this case, in addition to a flat background component. The data is divided in bins of 15 days. However, down times between runs can result in some bins having less statistics than others. It is not expected to contribute to either Crystal-1 or Crystal-2 due to their relatively long cooling times underground.

### 3.1. Measurement of decay rates

One way to measure the activities of the cosmogenic isotopes is through their decay rates. This measurement requires a selection of events of the specific decays studied, which can be identified by investigating the main contributions of the decay to our data spectra. Therefore, we first simulate each of the cosmogenically activated isotopes with the COSINE-100 GEANT4 package, studying their generated spectra and selecting the energy regions where they can have a significant contribution in comparison to the flat background.

Once the regions where the main contributions from each cosmogenically activated isotope are identified, we look into the decay rates for each of them, integrating the rates over the specific energy ranges. We fit the decay rate over time for each component. Depending on the energy region selected for each cosmogenic, the decay rate can be fitted by a constant and one or more exponential functions, as following:

$$A + B \cdot e^{\frac{-\ln(2)(t-t_0)}{C}} , \qquad (1)$$

where  $t_0$  is the initial time, A is the expected flat background rate, B is the rate in  $t_0$ , and C is the half-life, a constant of the fit. Fig. 4 shows an example of decay rate modeling with the units given in dru (counts/day/kg/keV).



Figure 5: Average activities of  $^{125}$ I during the first 60 days of data. The values measured through the decay fit method, described in the text, are plotted in green, while the ones measured through the background modeling [23] are shown in red.

The amplitude of the exponential (B) can be used to calculate the activity (in Bq/kg) of the cosmogenic isotope at the indicated initial time:

$$A_0 = \frac{\Delta E \times B}{86400 \times f_{\Delta E}} , \qquad (2)$$

where  $f_{\Delta E}$  is the fraction of the events from that cosmogenic depositing energy in the specified integration region, which can be calculated from the simulated spectra.  $\Delta Es$ for each isotope are 60–70 keV in single hit for <sup>125</sup>I, 20– 40 keV in multiple hit for <sup>121m</sup>Te, 80–94 keV in single hit for <sup>127m</sup>Te, and 20–30 keV in single hit for <sup>113</sup>Sn.

## 3.1.1. Iodine $^{125}I$

Since iodine is one of the main components in the crystal, a significant amount of <sup>125</sup>I is activated. However, the half-life of this isotope is short with  $T_{1/2} = 59.4$  days. We define the integration region for this isotope as 60 and 70 keV of the single-hit spectrum, as shown in Fig. 4; it decays to an isomeric state of <sup>125m</sup>Te by electron capture, producing 31.7 keV emissions for K-shell electrons, which is followed by the emission of an 35.5 keV gamma ray from the isomer transition of <sup>125m</sup>Te. Although there is a contribution from <sup>121m</sup>Te in this region, it is very small due to the low activity of this isotope and more importantly, due to the very small fraction of total <sup>121m</sup>Te events that actually deposit energy in this region. Therefore, the contribution of <sup>121m</sup>Te in the 60 to 70 keV region is insignificant,



Figure 6: Average activities of (a)  $^{121m}$ Te, (b)  $^{127m}$ Te, and (c)  $^{113}$ Sn during the first 60 days of data in six crystals. The values measured through the decay fit method, described in the text, are plotted in green, while the ones measured through the background modeling [23] are shown in red.

Table 4: Initial activity  $A_0$  (mBq/kg) of  ${}^{121m}$ Te,  ${}^{127m}$ Te, and  ${}^{113}$ Sn in each crystal as measured by the decay rate method. This includes the statistical uncertainty.

~						
	Crystal-1	Crystal-2	Crystal-3	Crystal-4	Crystal-6	Crystal-7
$^{121m}$ Te	-	-	$0.90{\pm}0.16$	$0.89 {\pm} 0.06$	$0.44{\pm}0.07$	$0.41{\pm}0.07$
$^{127m}$ Te	-	-	$0.87 {\pm} 0.16$	$0.48 {\pm} 0.03$	$0.38 {\pm} 0.04$	$0.35 {\pm} 0.04$
$^{113}Sn$	-	-	$0.16{\pm}0.05$	$0.15{\pm}0.06$	$0.16{\pm}0.01$	$0.12{\pm}0.01$

and the amounts of activated  $^{125}I$  in the crystals during their deployment at Y2L can be found in Table 3.

Fig. 5 shows the difference between the measured activities of  $^{125}I$  for the six crystals analyzed. The different amounts of  $^{125}I$  are related to the cooling time of the crystals before the start of data taking. We also compare the activities calculated in the background fit [23] to those measured through this method.

## 3.1.2. Tellurium <sup>121m</sup> Te, <sup>127m</sup> Te and Tin <sup>113</sup>Sn

- The line chosen to investigate  $^{121m}$ Te is the one between 20 to 40 keV in the multiple-hit spectrum, contributed by 30 keV emissions from the  $^{121m}$ Te decay via electron capture. This line is dominant in that region, unlike the  $^{121m}$ Te lines in the single-hit spectra. The method used is the same as described above.
- As we can see from the simulated spectra,  $^{127m}$ Te has • only one peak at 88.3 keV in the single-hit spectrum, contributed by the de-excitation of  $^{127m}$ Te. However, there are other components that can have significant contributions in that energy region as well, such as  $^{109}\mathrm{Cd}$  and  $^{121m}\mathrm{Te:}$  a 88 keV emission from the isomer transition of  $^{109m}$ Ag and another one at 81.8 keV from the de-excitation of  $^{121m}$ Te. The contribution coming from  $^{121m}$ Te can be calculated based on the measurement from the multiple-hit spectrum, as described above. The contribution from <sup>109</sup>Cd, however, is calculated based on another study, which will be described in section 3.2.2. Both of these are added to the fitting function, allowing the measurement of  $^{127m}$ Te, activity.

• The electron capture decay of <sup>113</sup>Sn produces 28 keV emissions. The method used is the same as described above. The peaks at 25.5 keV and 30 keV contributed by <sup>109</sup>Cd and <sup>121m</sup>Te can be calculated, as described above, and added to the fitting function, allowing the measurement of <sup>113</sup>Sn.

Fig. 6(a), (b) show the differences between the measured activities of  $^{121m}$ Te and  $^{127m}$ Te for the six crystals analyzed. We also compare the activities calculated in the background fit to those measured through this method. As listed in Tables 3 and 4, the initial activities when crystals were deployed at Y2L were derived for  $^{125}$ I,  $^{121m}$ Te,  $^{127m}$ Te, and  $^{113}$ Sn in each crystal.

#### 3.2. Long-lived isotopes

## 3.2.1. Sodium <sup>22</sup>Na

The decays of <sup>22</sup>Na (Q-value = 2.84 MeV) to <sup>22</sup>Ne<sup>\*</sup> proceed via  $\beta^+$  emission (90.3%) or electron capture (9.6%) with 3.75 yr mean lifetime, followed by <sup>22</sup>Ne<sup>\*</sup> transitioning to the stable <sup>22</sup>Ne isotope via the emission of a 1274.6 keV  $\gamma$ -ray with a 5.3 ps mean lifetime. The electron capture of <sup>22</sup>Na from K-shell produces 0.87 keV emissions. As a result, ~10% of the <sup>22</sup>Na decay will simultaneously produce a 1274.6 keV  $\gamma$ -ray and 0.9 keV emissions. In the case of  $\beta^+$  decay, the final-state positron immediately annihilates to two 511 keV  $\gamma$ -rays. If one of the two 511  $\gamma$ -rays escapes the crystal, the remaining energy deposited in the crystal will be substantially greater than 650 keV. Figure 7(a) shows the energy spectrum in another crystal in coincidence with a signal in the (650–1000) keV energy



Figure 7: (a) Energy spectrum in another crystal in coincidence with a signal in the (650-1000) keV energy interval in Crystal-3. In the fitting, it represents the peak at 609 keV from  $^{214}$ Bi in green and the peak at 511 keV from  $^{22}$ Na, activated in Crystal-3, in red. (b) Energy spectrum in another crystal contributed by triple coincidence events. The red line represents the peak at 511 keV from  $^{22}$ Na in Crystal-3. The same plots for all the other crystals are similar to each other.

Table 5: Initial activity  $A_0$  (mBq/kg) of <sup>22</sup>Na as measured by double and triple coincidences in each crystal. This includes the statistical uncertainty.

	Crystal-1	Crystal-2	Crystal-3	Crystal-4	Crystal-6	Crystal-7
double coincidence	$2.59 \pm 0.27$	$1.46 {\pm} 0.27$	$0.99 {\pm} 0.07$	$1.20{\pm}0.09$	$0.73 {\pm} 0.09$	$0.93{\pm}0.08$
triple coincidence	$2.0{\pm}0.4$	$1.52{\pm}0.37$	$0.84{\pm}0.18$	$1.17{\pm}0.19$	$0.65 {\pm} 0.14$	$0.87{\pm}0.23$

interval in Crystal-3, which are called *double coincidence* events. The  $^{22}$ Na  $\beta^+$  decay events show up as the peak at



Figure 8: Average activities of  $^{22}$ Na during the first 60 days of data in six crystals. The values measured through the double/triple coincidence methods, described in the text, are plotted in green/pink, while the ones measured through the background modeling [23] are shown in red.

511 keV (red color).

Since the eight NaI(Tl) crystal assemblies are immersed in the scintillating liquid (LS), as described in section 2, we can also identify  $^{22}$ Na decay events in which the 1274.6 keV gamma-ray converts in the LS in coincidence with two 511 keV signals in two crystals. These are referred to as *triple coincidence* events. Figure 7(b) shows the peak at 511 keV (red color) in another crystal, contributed by triple coincidence events, while a signal is in the (650–1000) keV energy interval in Crystal-3.

We used the time-dependent reduction of the peak at 511 keV, contributed by the double/triple coincidences of the 1.7 year data divided in bins of 60 days, to extrapolate the activity at the indicated initial time with the relation:

$$A_0 = \frac{N}{m \cdot t \cdot \epsilon} , \qquad (3)$$

where N is the number of events,  $\epsilon$  is the detection efficiency obtained from a Monte Carlo simulation, m is the mass of the NaI(Tl) crystal, and t is the time of the measurement.

Figure 8 shows the measured activities of  $^{22}$ Na for the six crystals analyzed by these methods, compared with the activities determined from the global background fit. The initial activities, when the crystals were first deployed at Y2L, are listed in Table 5.

Table 6: Initial activity  $A_0$  (mBq/kg) of  $^{109}$ Cd in each crystal. This includes the statistical uncertainty.

	Crystal-1	Crystal-2	Crystal-3	Crystal-4	Crystal-6	Crystal-7
$^{109}$ Cd (×10 <sup>-2</sup> )	$4.6 \pm 3.0$	$0.9{\pm}1.8$	$7.5 {\pm} 0.9$	$9.2 {\pm} 0.7$	$1.5 {\pm} 0.7$	$1.5 {\pm} 0.5$

## 3.2.2. Cadmium <sup>109</sup>Cd



Figure 9: Distribution of the time difference between 25.5 keV and an 88 keV signals, selected within three Gaussian width of the peaks in Crystal-3, fitted with the sum of two exponential decay functions. The <sup>109</sup>Cd contribution with lifetime 56 s is shown in red. The parameter Const is an amplitude of the exponential decay of <sup>109</sup>Cd. The parameters p0 and p1 are an amplitude and a mean lifetime of the exponential decay of the background contribution. The same plots for all the other crystals are similar to each other.

The cosmogenic isotope <sup>109</sup>Cd decays via electron capture to an isomeric state of  $^{109m}$ Ag, with a prompt energy deposit of 25.5 keV, the binding energy of the Ag K-shell electron. This is followed by an emission of 88 keV from the isomer transition of  $^{109m}$ Ag that has a mean lifetime of 57.4 s. From the time interval distribution between 25.5 keV and 88 keV signals, selected within three Gaussian width of the peaks in the same crystal, we extract the level of <sup>109</sup>Cd from a fit with two exponential decay functions. As discussed in section 3.1.2 there are significant contributions from  $^{127m}$ Te and  $^{121m}$ Te around 88 keV and  $^{113}$ Sn and  $^{121m}$ Te around 25 keV, which dominate the blue curve in Figure 9. The fitted mean lifetime,  $56\pm14$  s, from the exponential curve in red, as can be seen in Figure 9, is consistent with the mean lifetime of 57.4 s from the isomer transition of  $^{109m}$ Ag.

We determined the <sup>109</sup>Cd activity rates in mBq/kg from these measurements: Fig. 10 shows the measured activity levels for the six crystals analyzed through this method, compared with the activities determined from the global background fit. The crystal activity levels when they were first deployed at Y2L are listed in Table 6.



Figure 10: Average activities of  $^{109}$ Cd during the first 60 days of data in six crystals. The values measured through this method, described in the text, are plotted in blue, while the ones measured through the background modeling [23] are shown in red.

## 4. Results and comparisons for production rates

In section 3 we describe the determination of the crystals' cosmogenic isotope activities at the time they were first deployed underground at Y2L. However, since we do not know the details of their previous exposure conditions, such as times, locations, and altitudes, these cannot be directly related to production rates or saturation activity levels. But an attempt to extract sea level production rates has been made from a simplified mathematical model for production and decay of radionuclides.

The production rate R for activation of an isotope can be expressed as

$$R \propto \int \sigma(E) \cdot \Phi(E) \cdot dE \tag{4}$$

where  $\sigma$  is the neutron capture cross section and  $\Phi$  is the cosmic-ray neutron flux. Since cosmic-ray neutron flux  $\Phi$  depends on altitude, location, and time the production rate R at any arbitrary location can be calculated by scaling the reference production rate  $R_s$  at sea level,

$$R = f \cdot R_s \tag{5}$$

The produced nuclide then decays according to the standard decay law, and the net rate of change for the number of existing radioactive nuclei N is by the differential equation

$$\frac{dN}{dt} = -\lambda \cdot N + R , \qquad (6)$$

	<sup>22</sup> Na	<sup>109</sup> Cd	<sup>125</sup> I	$^{121m}\mathrm{Te}$	$^{127m}\mathrm{Te}$	$^{113}$ Sn
Crystal-1	$132.0{\pm}24.0$	$1.7{\pm}1.1$				
Crystal-2	$148.5 {\pm} 44.9$	$0.6{\pm}1.2$				
Crystal-3	$114.5 {\pm} 19.7$	$4.7 {\pm} 0.6$	$280.1 \pm 29.3$	$31.1 {\pm} 5.5$	$26.9 {\pm} 4.8$	$5.1 {\pm} 1.6$
Crystal-4	$81.0{\pm}12.7$	$3.7 {\pm} 0.3$	$104.2 \pm 3.7$	$24.9{\pm}1.6$	$13.5{\pm}0.7$	$4.1{\pm}1.6$
Crystal-6	$144.0 \pm 31.2$	$1.8{\pm}0.8$	$184.7 {\pm} 6.3$	$23.5 \pm 3.5$	$16.3 {\pm} 1.5$	$7.1 {\pm} 0.5$
Crystal-7	$151.0 {\pm} 52.1$	$1.8{\pm}~0.6$	$194.0{\pm}6.3$	$22.3 \pm 3.5$	$15.0{\pm}1.5$	$5.3{\pm}0.5$
ACTIVIA	66	4.8	221	93	93	9
MENDL-2		4.8	208	102		
ANAIS measurement [31, 32, 34]	$45.1 {\pm} 1.9$	$2.38 {\pm} 0.20$	$220{\pm}10$	$23.5{\pm}0.8$	$10.2{\pm}0.4$	$4.53 {\pm} 0.40$
DM-Ice17 measurement [30]			230	25	< 9	16

Table 7: Production rate  $R_s [kg^{-1} d^{-1}]$  at sea level.

where  $\lambda$  is the decay constant:  $\lambda = \frac{\ln 2}{T_{1/2}}$  ( $T_{1/2}$  is the decay half-life). The solution of Eq. 6 is

$$N = \frac{R}{\lambda} (1 - e^{-\lambda t}), \tag{7}$$

and the activity A(t) is related to the number of existing nuclei N by

$$A(t) = \lambda \cdot N \tag{8}$$

$$=R(1-e^{-\lambda t}) \tag{9}$$

When the time t is sufficiently large, then a saturation activity  $A_s$  is reached at a given place,

$$A_s = f \cdot R_s \tag{10}$$

The ANAIS experiment [31] realized that the cosmic-ray neutron flux  $\Phi$  can be scaled from its sea level reference flux, as reported in Ref. [30], to the NaI(Tl) crystal production point in Grand Junction, Colorado (altitude=1400 m), by a factor of f = 3.6. Using the measured  $A_0$  activity levels reported here and the exposure times listed in Table 1, we compute a production rate  $\mathbf{R}_s$  at sea level from the relation,

$$A_0 = R_s [1 + (f(1 - e^{-\lambda t_1}) - 1) \cdot e^{-\lambda t_2}]$$
(11)

where the exposure time  $t_{exp}$ , listed in Table 1, is  $t_{exp} = t_1 + t_2$ . The crystals are exposed at Alpha Spectra for a time  $t_1$  and exposed at sea level for a time  $t_2$ . We considered  $t_2 = 30$  days as transportation duration to Y2L. Table 7 shows the production rate of cosmogenic isotopes in each NaI(Tl) crystal used for the COSINE-100 experiment compared with measurements from ANAIS [31, 32, 34] and DM-Ice17 [30], and calculations using ACTIVIA and MENDL-2; we used v1.3 of ACTIVIA that follows the parameterization of Gordon [35], valid from 1 MeV to 10 GeV, for the neutron flux spectrum and MENDL-2 that contains neutron reaction data up to 100 MeV. The results obtained with the six NaI(Tl) crystals are in reasonable agreement with each other, except for Crystal-3 that has a complicated exposure history;

we considered 1 year exposure time for Crystal-3 for calculation of the production rates listed in Table 7. The production rates of <sup>22</sup>Na in the six crystals are compatible with each other although they are larger than other measurement and calulation. For Crystal-4, since we do not know clearly the month and day of the powder production and the crystal delivery to Y2L it is possible to have about 60 days uncertainty for the exposure and the cooling times respectivley.

## 5. Discussion on Tritium <sup>3</sup>H and Iodine <sup>129</sup>I

It is generally difficult to measure activity levels of longlived cosmogenic isotopes, directly from the data due to their long half-lives. This is especially the case for <sup>3</sup>H, which has no distinguishing  $\gamma$ /X-ray peak that can be exploited. Therefore, we simulated background spectra from <sup>3</sup>H in the six NaI(Tl) crystals and used the extracted spectral shapes in the data fitting, while floating their unknown fractions [23]. We determine the initial activity  $A_0$  of <sup>3</sup>H by using the average activity during the first 60 days of data, obtained from the global background fitting model [23] described above, with results shown in Table 8. From these we computed exposures time  $t_{exp}$  from Eq. (11). We assumed that the production rate of  ${}^{3}H$  at sea level is  $R_s = (83\pm27) \text{ kg}^{-1} \text{d}^{-1}$ , which was reported by ANAIS [33]. The resultant exposure times, listed in Table 8, are in good agreement with the time period during which they were being produced at Alpha Spectra and undergoing delivery to Y2L, as shown in Table 1.

The presence of cosmogenic <sup>129</sup>I was introduced by DAMA/LIBRA with the estimated concentration of <sup>129</sup>I/ <sup>nat</sup>I =  $(1.7\pm0.1)\times10^{-13}$  [36]. It is used as a floating parameter in the global background fitting modeling for the COSINE-100 NaI(Tl) crystals, with resulting values of 1.01, 1.08, 0.75, 0.72, 0.91, and 0.94 mBq/kg for Crystal-1, 2, 3, 4, 6, and 7, respectively. These values agree well with the ANAIS result: 0.96±0.06 mBq/kg [37].

Table 8: Initial activity (A<sub>0</sub>) of <sup>3</sup>H in the NaI(Tl) crystals derived from background fitting and the derived estimated exposure times.

	Crystal-1	Crystal-2	Crystal-3	Crystal-4	Crystal-6	Crystal-7
$A_0 [mBq/kg]$	$0.38 {\pm} 0.04$	$0.20{\pm}0.04$	$0.25 {\pm} 0.04$	$0.26 {\pm} 0.04$	$0.11 {\pm} 0.04$	$0.09 \pm 0.04$
Exposure time [year]	2.19	1.11	1.37	1.44	0.66	0.52

## 6. Conclusion

We have studied background contributions from cosmogenic isotopes activated by cosmic rays in the COSINE-100 detectors. To understand their time-dependent energy spectra we simulated responses to decays of the most abundantly produced cosmogenic isotopes in NaI(Tl) crystals and identified the energy regions where they make strong contributions to the crystals' background spectra. Based on these simulation studies we measured decay rates of the cosmogenic isotopes using the time-dependent decrease of peaks from characteristic decays of these isotopes. We also exploited the correlations of characteristic emissions in terms of time differences between sequential decays of <sup>109</sup>Cd and double- and triple-coincidences for <sup>22</sup>Na-decayinduced multi-gamma final states. From these measurements, we extrapolated the various isotopes' activity levels to the times that they were first deployed underground at Y2L.

With these data we estimated production rates (at sea level) for the cosmogenic isotopes that are relevant for COSINE-100 and compared them with other experimental data and ACTIVIA/MENDL-2 calculations. As listed in Table 7, the results from different approaches are in reasonable agreement with each other. We extracted exposure times using initial <sup>3</sup>H activities determined from the COSINE-100 global background fitting model and found results that are in reasonable agreement with the times reported in Table 2(b). We also quantified the unknown <sup>129</sup>I activity level by including it as a free-floating parameter in the the global background fit model and found consistency with an ANAIS result.

This study has given us a quantitative understanding of the cosmogenic isotopes in the NaI(Tl) crystals used for the COSINE-100 experiment. It provides important constraints on time-dependent backgrounds in our search for a time-dependent modulation that would be a characteristic signal for dark matter interactions [38, 39].

### Acknowledgments

We thank the Korea Hydro and Nuclear Power (KHNP) Company for providing underground laboratory space at Yangyang. This work is supported by: the Institute for Basic Science (IBS) under project code IBS-R016-A1 and NRF-2016R1A2B3008343, Republic of Korea; UIUC campus research board, the Alfred P. Sloan Foundation Fellowship, NSF Grants No. PHY-1151795, PHY-1457995, DGE-1122492, WIPAC, the Wisconsin Alumni Research Foundation, United States; STFC Grant ST/N000277/1 and ST/K001337/1, United Kingdom; and Grant No. 2017/02952-0 FAPESP, CAPES Finance Code 001, Brazil.

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