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MEASUREMENTS OF ¹²⁹I ANDOACTIVE PARTICULATE CONCENTRATIONS IN THE TMI-2 CONTAINMENT ATMOSPHER[®] GURING AND AFTER THE VENTING

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MEASUREMENTS OF ¹²⁹I AND RADIOACTIVE PARTICULATE CONCENTRATIONS IN THE **TMI-2 CONTAINMENT ATMOSPHERE DURING** AND AFTER THE VENTING

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SUMMARY

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Measurements were made of the 129 I and 85 Kr concentrations in containment atmosphere during the purging of the Three Mile Island Unit 2 containment that occurred from June 28 until July 11, 1980. The measurements showed that although the airborne 129 concentration was reduced by a factor of about 20 during the purge, within 15 days it had nearly returned to the prepurge value. Moreover, the initial fraction of organic iodine of 90% was reduced to about 40% during the purge and it returned to 90% in 15 days. It is believed that the equilibrium concentration and species distribution is the result of partitioning of iodine from the sump liquid and by deposition and resuspension of iodine from containment surfaces. The relative contribution to containment atmosphere from sump water and containment is not known. By comparison, the 85 Kr concentration decreased by a factor of 50 000 during the purge and recovered, most likely, by evolution from the sump water, to a level of 1/500 of its initial value after the main purge. Concentrations of 134Cs, 137Cs and 90Sr were also measured in containment atmosphere.

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MEASUREMENTS OF ¹²⁹I AND RADIOACTIVE PARTICULATE CONCENTRATIONS IN THE TMI-2 CONTAINMENT ATMOSPHERE DURING AND AFTER THE VENTING

BACKGROUND

Previous studies of iodine at nuclear power plants suggested that, after escaping the pressure boundary, iodine was deposited and subsequently resuspended before appearing in ventilation exhaust air. 1,2,3 A mathematical model was derived to explain this behavior. 3 It was a two compartment model in which airborne iodine was allowed to deposit on building surfaces and resuspend later. The chemical form of iodine was assumed to remain unchanged. Laboratory tests were carried out to determine values for parameters in the model.⁴ During these tests it was found that surface reactions with the elemental iodine formed organic and other forms of iodine. The model was modified to include the formation and resuspension of nonelemental forms of iodine and was tested with radioiodine measurements in containment atmospheres at operating PWRs.⁵ One of the purposes for measuring ¹²⁹I prior to, during and after the purging of the TMI-2 containment atmosphere was to provide data for the model. Another purpose of the measurements was to document concentrations of other long-lived nuclides, e.g., ¹³⁴Cs, ¹³⁷Cs and ⁹⁰Sr.

A study¹ of the behavior of ¹³¹I in the auxiliary and fuelhandling buildings at TMI, Unit 2 revealed that the most likely source of the activity in these buildings during the period April-September 1979 was evolution from the interior surfaces of the buildings. It is also most likely that the iodine deposited on the surfaces during or soon after the accident. Evaluations done² for the Nuclear Safety Analysis Center (NSAC) show that, relative to ¹³⁷Cs, about half of the ¹³¹I present in the core at the shutdown cannot be accounted for in the reactor coolant, reactor building sump or atmosphere, or the auxiliary and fuel-handling buildings (liquids surfaces, filters, or releases). It is suggested in the study that the "missing" iodine was evolved either from the hot sump water or through flashing of the hot coolant as it was released into containment and that it deposited on surfaces. This continued until an equilibrium was established

between the sump, surfaces and atmosphere. The equilibrium appears to have been established rather quickly. Samples of containment atmosphere² taken on and following March 31, 1979 suggest that the iodine concentration decreased only with the decay half life of 8.04 days; this indicates that the activity level had reached equilibrium as early as three to six days after the accident (and subsequent isolation).

Radioiodine molecular species measurements made⁶ June 26 and July 9, 1979 showed a relatively high concentration of organic 131 I, 86 and 79%, respectively, with the remainder equally split between the elemental and HOI forms. This iodine molecular composition is quite characteristic of iodine that has been "aged" through deposition and resuspension.

MEASUREMENTS

Samplers installed through a containment penetration sampled radioiodine and particulates from July 4 until August 12, 1980. Venting of containment was started June 28, 1980. It ended July 11 but was also done for very short times on August 1 and August 8.

Sampling System

The radioiodine and particulate sampler was installed in what is known as penetration R-626 in the Unit 2 containment building. This penetration is a 25-cm diameter pipe at the "358" level, about 3.1 m (10.5 ft) above the refueling floor. The physical arrangement is as shown in Figure 1.

The sampler pulled the containment atmosphere through a 1-cm diameter, 241-cm long stainless steel tubing that extended about 0.7 m beyond the containment wall. The tubing was connected in series to a flow measuring rotameter, the sampler and a carbon vane air pump. A glove box attached to the R-626 penetration housed the entire system. Sampling was done at a rate of 0.471 liters per second.





The iodine sampler was the SAI radioiodine molecular species sampler whose properties are described in Reference 3. This sampler has five components placed in a series: a particulate filter, a CdI_2 media, an IPH media, and two charcoal cartridges. The cartridges selectively absorb, respectively, particulate, elemental, HOI, and organic forms of iodine. The final charcoal cartridge is used to ensure no break through of iodine in the sampler.

Containment Venting

The main containment venting began June 28 and ended July 11, 1980. The initial venting used the hydrogen control system that vented from the "305'" level inside the steam generator "dee ring." The fast purge that began July 8 used the main venting lines that exhaust from the "347'" level, the refueling floor.

Venting was done cautiously with considerable starting and stopping. Table 1 gives the purge rates as a function of time for the venting. A histogram of these rates for the main venting is shown in Figure 2.

Although the main purge ended on July 11, there were additional short purges on August 1, August 8, August 14 and August 22 whose rates are shown in Table 1. A plot of the purging that includes these periods is shown in Figure 3.

Data Analysis

129 I Analysis

A thin window and thin crystal NaI(T1) counted the radiation emitted in the decay of the ^{129}I activity. Each cartridge was counted nondestructively with the stream entry side toward the detector. Data were acquired using a pulse-height analyzer. In each case there was sufficient activity that this method of direct counting resulted in an analyzable spectrum of ^{129}I radiation.

Date	Time	Flow (cfm)	Date	Time	Flow (cfm)	<u>Date</u>	Time	Flow (cfm)
6/28	0800-0900 1700-1800 1800-2000 2000-2100 2100-2300	100 15 75 96 89		1700 1800 1900 2000 2100	150 210 210 285 80	1	0830 0845 0943 1219 1235	350 240 380 565 200
6/29	1400-1440 1544 1600 1700 1800 1900 2000 2100 2200 2300 2400	105 220 130 90 100 94 102 108 1·10 98 79	7/1	2200 2300 2400 0300 0632 0752 0850 0935 1105 1132 01-1312	150 220 210 180 110 440 250 500 280 190 0	7/4	1334 1440-1445 1445 1649 1933 2133 2231 2332 0030-0050 0050 0725	360 0 215 460 180 470 230 430 0 406 261
6/30	0100 0200 0400	80 90 64		1312 1555 1725 1925 2130	230 340 520 300 200	7/4-5	0827 1645 1730 2020 2120 2305-0319	485 400 50 240 142
	0700 0800 0900 1000 1100 1200 1300 1400 1500 1600	100 130 155 155 155 155 165 160 190 155	7/2 7/3	0015-0532 0532 0928 1200-1715 1715 1932 2132 0223 0321 0	0 258 490 0 407 230 135 530 318	7/5	2303-0319 0319 0623 0924 1042 1920 2120 2220 2330 0031-0110 0110	150 365 430 570 250 332 165 340 0 150
7/6	0232 0555 0623 0723 2040	300 540 238 540 400		0909 0934 1055 1138 1250	2200 4000 4500 2900 4200			

TABLE 1. PURGE RATES OF TMI-2 CONTAINMENT FROM JUNE 28 TO AUGUST 23, 1980

TABLE 1. (Continued)

<u>Date</u>	Time	Flow (cfm)	Date	Time	Flow (cfm)	Date	Time	Flow (cfm)
7/7	0030-0105 0105 0225 0425 0445 0525 0721 0810 2041 2133 2320	0 170 89 354 445 300 153 321 540 400 260 560	7/10 7/11	1330 1442 1755 1846 0148-1531 1531 1634 2045 2145 2235 0100-0118 C118 0933	4350 2100 1500 18500 19000 18500 13000 16250 18500 0 18500 0			
7/8	0037-0052 0052 0622-1200 1200-1300 1600-1700 1800 1850 1850 1900-2000	0 560 0 1000 ~1500 1200 500 1750 1000	8/1 8/8 8/14	1300 1400 1520 0900 1000 1100 1200 1230 1000	0 7600 0 4000 8000 9600 9800 0 8000			
7/8-9	2000-0058 0058 0430-0444 0444 0520 0620-0700 0700 0730	0 560 0 1000 1800 0 1000 1600	8/22	1100-1500 1600-2100 2100-2300 1000 1100 1700	9000 0 20000 5000 20000 0			

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Figure 2. Histogram of TMI-2 containment purge rates during the period June 28 to July 12, 1980.



Figure 3. Histogram of TMI-2 containment purge rates during the period June 28 to August 23, 1980.

Efficiency calibration relied on the known intensities of the x-ray and 662-keV transitions from 137 Cs. The charcoal was removed from an unused cartridge, wetted with a solution containing about 1 µCi of 137 Cs, dried, mixed thoroughly, and loaded into a plastic cup. The filled cup was capped and sealed. This cup was analyzed for total Cs activity with a calibrated Ge(Li) detector (using the 662-keV gamma ray). We then calculated the emission rate of the K x-ray from the sample. Next, the sample in the cup was counted in a reproducible geometry using the thin window NaI(T1) detector and the x-ray counting efficiency for this geometry was determined to be 1.90%. Three cartridges that had been counted nondestructively were taken apart and the charcoal mixed thoroughly and loaded in plastic cups identical to those for the 137 Cs secondary standard. These cups were then counted on the thin window NaI(T1). The efficiency for the nondestructive geometry was thus determined and used for all of the analyses.

The principal assumption in the use of this calibration number was that the distribution of ¹²⁹I through the cartridges was the same for all cartridges. Roughly, the same volume of air was pulled through each cartridge in the sampling. The cartridges have shown reproducible activity distributions in past analyses. Hence, the assumption is considered as valid at least within the other sampling uncertainties.

Particulate Analysis

Analysis of the particulate filters for gamma-ray emitting nuclides used Ge(Li) spectrometers. The analysis revealed the presence of only the two cesium nuclides 134 Cs and 137 Cs. The filters were also analyzed nondestructively for 90 Sr activity by beta pulse-height spectrometry, and destructively for both 89 Sr and 90 Sr using the standard radiostrontium procedures at our laboratory.

Results

Table 2 gives the analysis results in terms of microcuries of ¹²⁹I per cubic centimeter of containment air for each sampling period. The table lists the values obtained for each component of the sampler as well as the

		Concentr (µi/cm ³	Percentage of Total on Each Cartridge				
Period (1980)	Cadmium Iodide	Iodophenol	Charcoal		<u>Cd1</u> ²	<u>IPH</u>	<u>Charcoal</u>
7/4-7/6	7.69(-13)	5.40(-12)	2.14(-11)	2.76(-11)	2.1	19.1	78.8
7/6-7/8	5.69(-13)	2.99(-12)	1.10(-11)	1.46(-11)	3.9	20.6	75.5
7/8-7/9	3.39(-13)	1.60(-12)	2.85(-12)	4.80(-12)	7.1	33.3	59.5
7/10-7/12	2.89(-13)	7.84(-13)	7.43(-13)	1.82(-12)	15.9	43.2	40.9
7/12-7/15	4.55(-13)	2.56(-12)	4.18(-12)	7.21(-12)	6.3	35.7	58.0
7/15-7/18	5.69(-13)	3.19(-12)	1.20(-11)	1.58(-11)	3.6	20.2	76.2
7/21-7/22	1.18(-12)	3.09(-12)	2.30(-11)	2.72(-11)	4.3	11.4	84.3
7/23	1.27(-12)	2.94(-12)	2.25(-11)	2.67(-11)	4.7	11.0	84.3
7/25-7/26	1.24(-12)	2.56(-12)	2.53(-11)	2.91(-11)	4.2	8.8	87.0
7/30	1.12(-12)	3.25(-12)	3.99(-11)	4.44(-11)	2.5	7.3	90.2
8/2-8/3	9.28(-13)	2.06(-12)	2.82(-11)	3.12(-11)	3.0	6.6	90.4
8/5-8/6	7.91(-13)	3.01(-12)	2.95(-11)	3.33(-11)	2.4	9.0	88.6
8/7-8/8	9.87(-13)	2.58(-12)	3.51(-11)	3.88(-11)	2.6	6.7	90.8
8/9-8/10	7.47(-13)	2.62(-12)	2.10(-11)	2.44(-11)	3.1	10.8	86.2
8/12	5.10(-13)	2.53(-12)	2.72(-11)	3.03(-11)	1.7	8.3	90.0

TABLE 2.129I CONCENTRATIONS FOR TMI-2 CONTAINMENT DURING VENTINGS.SAMPLES TAKEN FROM R-626 PENETRATION

total concentration. The table also contains columns that list the percentage of the total activity on each cartridge. Figure 4 shows a plot of the \approx ¹²⁹I concentrations as a function of the sampling date.

Table 3 lists the measured concentrations of 134 Cs, 137 Cs and 90 Sr. Activities of 89 Sr were found to be below the minimum detectable limit. Figure 5 shows a graph of these concentrations.

For comparisons, Table 4 gives concentrations of 85 Kr in samples of containment air during the same time period as the 129 I sampling program. These data were obtained from the TMI staff. The table gives the location where the sample was pulled. There were qually two samples taken. The results are given for both samples. Figure 6 shows a plot of the 85 Kr concentrations as a function of the sampling date.

DISCUSSION

The concentration of ${}^{129}I$ in the TMI-2 containment atmosphere before the purge was measured⁷ to be about $7.5 \pm 2.0 \times 10^{-11} \,\mu\text{Ci/cm}^3$. This value is comparable to the expected value of $6.7 \times 10^{-11} \,\mu\text{Ci/cm}^3$ that is based on earlier measurements of ${}^{131}I$ and relative isotope inventories of the iodine at the time of the accident. This agreement suggests that equilibrium of airborne iodine with the sump and building surfaces has been continuous since early in the accident.

Containment venting reduced the 85 Kr concentration by about a factor of 50 000. When the main purge was terminated, the activity rose nearly a factor of 100 to what appeared to be a new equilibrium level. The recovery was most probably the result of 85 Kr coming out of the sump water until a new equilibrium concentration was established. Ninety percent recovery was in about 17 days. Subsequent short purges continued to reduce the equilibrium concentrations of 85 Kr.

Indine activity levels showed similar but significantly different behavior during and after the main venting. It is unfortunate that our measurements of $^{129}\mathrm{I}$ did not begin at or before the venting began. The



Figure 4. Plots of the ¹²⁹I concentrations as a function of time during the venting of the TMI-2 containment. The different curves show the concentrations of the different molecular forms of airborne iodine.

Period (1980)	134 _{Cs}	¹³⁷ Cs	⁹⁰ Sr	⁸⁹ Sr
7/4-7/6	5.87(-12)	4.05(-11)	7.0(-12)	
7/6-7/8	2.84(-12)	1.80(-11)	6.4(-12)	
7/8-7/9	1.74(-11)	1.21(-10)	2.6(-11)	<1.1(-12)
7/10-7/12	1.95(-11)	1.40(-10)	4.67(-11)	<1.6(-12)
7/12-7/15	2.80(-11)	2.02(-10)	6.5(-11)	
7/15-7/18	246(-11)	1 78(-10)	7 94(-11)	
7/21-7/22	9,80(-11)	7.30(-10)	2.5(-10)	
7/23	2.76(-11)	1.96(-10)	7.4(-11)	
7/25-7/26	5.22(-11)	3.75(-10)	1.32(-10)	
7/30	1.096(-11)	7.62(-11)	2.6(-11)	
8/2-8/3	1 (19(-11))	7 75(-11)	2.8(-11)	
8/5-8/6	8 40(-12)	5 89(-11)	2.0(-11)	
8/5-8/0	1,314(-11)	9.74(-11)	2.12(-11)	
0/7-0/0	1.314(-11)	9.74(-11)	3.4(-11)	•
8/9-8/10	1.50/(-11)	1.123(-10)	3.8(-11)	
8/12	2.03(-11)	1.49(-10)	5.3(-11)	

TABLE 3. PARTICULATE CONCENTRATIONS FOR TMI-2 CONTAINMENT DURING VENTINGS. SAMPLES TAKEN FROM R-626 PENETRATION (Values in $\mu \text{Ci}/\text{cm}^3$)



Figure 5. Flots of the particulate concentrations measured in TMI-2 containment atmosphere during the period July 4 to August 12, 1980.

Date 1980	HP-R-227 354' Sample 1	HP-R-227 354' Sample 2	HP-R-227 469' Sample 1	HP-R-227 469' Sample 2	HP-R-229X	HP-R-219	R-626
6/27 6/29 6/30 7/1 7/2 7/3 7/4 7/5	1.02 1.01	0.998	0.96 0.95 0.92 0.88 0.72 0.61 0.461 0.358	0.88 0.89 0.71 0.61 0.468 0.372			
7/6 7/7	0.342 0.215	0.328 0.203	0.259	0.263	0.12	0.154	0.216
7/8 7/9 7/10	0.0072 1.7(-4) 1.5(-4) 2.0(-4)	0.0083 1.7(-4) 1.9(-4)	0.134 0.0094	0.135 0.0080	0.197		
7/11 7/15	2.0(1)	1.0(4)	3.6(-5) 6.4(-5) 6.4(-4)	3.5(-5) 5.8(-5) 6.2(-4)			
7/16 7/17 7/22 7/23 7/24 7/25 7/26 7/27 7/28 7/29 7/30 7/31 8/1 8/1 8/4 8/6 8/8	6.9(-4) ^a	6.7(-4) ^a	$\begin{array}{c} 6.3(-4) \\ 7.5(-4) \\ 1.3(-3) \\ 1.4(-3) \\ 1.5(-3) \\ 1.6(-3) \\ 1.6(-3) \\ 1.6(-3) \\ 1.7(-3) \\ 1.7(-3) \\ 1.7(-3) \\ 1.9(-3) \\ 2.3(-3) \\ 3.4(-3) \\ 3.5(-3) \\ 2.0(-3) \\ 2.5(-3) \\ 2.4(-3) \\ 2.4(-3) \\ 1.4(-3) \end{array}$	7.1(-4) 7.7(-4) 1.4(-3) 1.4(-3) 1.5(-3) 1.6(-3) 1.6(-3) 1.7(-3) 1.8(-3) 1.7(-3) 2.0(-3) 3.3(-3) 3.4(-3) 2.4(-3) 2.3(-3) 2.3(-3) 2.4(-3) 1.4(-3)		·	

TABLE 4. KRYPTON-85 CONCENTRATION FOR TMI-2 CONTAINMENT DURING VENTING (Values of μ Ci/cm³)

a. 315 ft.

TABLE 4. (continued)

Date 1980	HP-R-227 354' Sample 1	HP-R-227 354' <u>Sample 2</u>	HP-R-227 469' Sample 1	HP-R-227 469' Sample 2	HP-R-229X	HP-R-219	R-626
8/11			1.5(-3)	1.5(-3)			
8/13			1.6(-3)	1.6(-3)			
8/14			1.6(-3)	1.6(-3)			
			4.6(-4)	4.3(-4)			
8/15			<6.7(-6)	<7.0(-6)			
8/18			1.2(-4)	1.3(-4)			
8/20			5.6(-4)	6.0(-4)			
8/22			5.6(-4)	5.5(-4)			
			4.3(-5)	3.5(-5)			
8/25			1.0(-4)	1.2(-4)			
8/27			1.4(-4)	1.4(-4)			





fact that they did not, results in some problems in comparing our measurements during and after the venting to those taken before the venting. Purging flow rates prior to the installation of the sampler was quite low (~200 cfm). By comparing the relative decrease of 85 Kr activity prior to the beginning of the sampling period, the equilibrium value on the scale of the present measurements is estimated to be about 4 x 10⁻¹¹ µCi/cm³. It thus appears that the equilibrium concentration of total 129 I after the purge is about the same as that before the purge. Furthermore, each molecular specie of 129 I appears to recover to about the same concentration although the rate of recovery seems to be somewhat longer for I₂ and CH₃I than for HOI.

The makeup of the iodine species changed radically during the measurements with the relative concentrations of I_2 and HOI increasing to 16 and 43%, respectively, at the end of the purge and then returning to about 2.5 and 8% after the purging ended. The final equilibrium molecular species distribution is very similar to that of ¹³¹I measured in June and July of 1979, mentioned in Section 1. The significant relative change of species can quantitatively be explained on the basis of partition coefficients for surfaces⁴ for the different forms of iodine. Quantitative explanation would require computations involving the models developed⁵ to describe the transport of iodine in systems. Changes in iodine species distribution were observed⁹ during operations of "kidney filters" in containment at the Rancho Seco plant.

Concentrations of particulate activity was found to increase during the fast purge stage of the venting and to remain relatively high after the venting was terminated. A possible explanation is that the fast purge significantly disturbed the air currents in the vicinity of the sampler inlet, suspending considerable particulate activity into the air and that it remained suspended for a considerable time.

SUGGESTED ADDITIONAL WORK

Emission of iodine into the air from liquids and surfaces contributes largely to airborne iodine concentrations. This has significance in releases from nuclear reactors into confined areas from personnel safety or clean-up considerations.

Two additional tasks are desirable to fully understand the mechanism and implications of the iodine releases.

- o Extend and apply the iodine model to the data to extract the needed resuspension, deposition, and partition coefficients.
- o Measure the deposits of ¹²⁹I on the surfaces in TMI-2 containment to attempt to establish a better estimate of the relative contributions of the water and the surfaces.

The model in its current form does not include deposition of nonelemental forms of iodine. However, the fact that nonelemental forms of ^{129}I reached in equilibrium show that deposition does exist, and the model should be modified.

If the only source of ^{129}I were that deposited on interior surfaces early in the accident, the current model would predict that all the iodine would be in the air as organic and one purge would rid the atmosphere of it. It is possible that the sump water is also a source of ^{129}I . The model should be modified to include iodine partitioning after an accident as a source of iodine. Such a model would be capable of predicting iodine concentrations in containment after incidents such as those at Crystal River and Arkansas 1.

Using only airborne concentration measurements to verify and modify the model of iodine behavior provides only a portion of the evidence. Measurements of ^{129}I deposited on interior surfaces at TMI-2 offer along with air concentration measurements unambiguous evidence to verify or modify the model. Therefore, a program for measuring ^{129}I on surfaces at TMI-2 should be developed and carried out.

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