NEUTRON ACTIVATION COOL-DOWN OF THE TOKAMAK FUSION TEST REACTOR

H. W. Kugel,^a G. Ascione,^a C. Tilson Jr.,^a and A. Kumar^b

^aPrinceton Plasma Physics Laboratory Princeton University P. O. Box 451, Princeton NJ 08543 ^bSchool of Engineering and Applied Science, University of California, Los Angeles, CA 90095-159710

ABSTRACT

TFTR final operations and post-shutdown neutron activation measurements were made. Ionization chambers were used to follow TFTR activation during operations and after shutdown. Gamma-rav energy spectroscopy measurements were performed to characterize TFTR activation at accessible vessel-bays, and on sample hardware removed from structures at various distances from the vessel. The results demonstrate long-lived activations from common, commercially available materials used in the fabrication and field engineering of TFTR. The measurements allow characterization of residual TFTR neutron activation, the projection of residual activation decay, and benchmarking of low activation issues simulations.

I. INTRODUCTION

The Tokamak Fusion Test Reactor (TFTR) began high power deuterium-tritium (D-T) fueled operations in December 1993, completed operations in April 1997, and is presently in a shutdown, safe-keeping mode.¹ The achieved performance includes central ion temperatures up to 44 keV, electron temperatures up to 13 keV, fusion power yields up to 10.7 MW, and a D-T fusion energy yield of 1.55 GJ, corresponding to a total D-T neutron vield of more than 5.52×10^{20} . There was also about 0.15 GJ of D-D fusion energy yield. These operations involved about 21,670 high power discharges, including about 1000 D-T discharges of which about 750 D-T discharges involved at least one neutral beam injecting tritium, and about 250 discharges into which tritium gas was puffed into the tokamak with all neutral beams injecting deuterium. This work involved the safe handling and processing about 100g of tritium. This resulted in manageable long-term neutron activation of the vessel and surrounding hardware. In this paper, we report the results of post-shutdown measurements made for the purpose of characterizing residual TFTR neutron activation, projecting residual activation decay, exploring low activation issues, and investigating options for future radioactive waste management.

II. EXPERIMENTAL CONDITIONS

Extensive simulations of TFTR radioactivation have been performed previously using estimates of the Test Cell material density distribution.^{2, 3} The materials used in the construction of the TFTR facility are known but due to the evolution of the experimental requirements the facility grew in complexity, and hence, the exact material density distribution has been difficult to quantify. However, during D-T operations, the configuration remained relatively unchanged to meeting maintenance, reliability, and performance requirements. Fig. 1 shows a partial schematic plan view of the TFTR concrete Test Cell showing the relative locations of the vessel, neutral beam injection systems, the vacuum system pump ducts, and the activation measurements. The characteristics of the Test Cell shielding and radiation field have been described previously.⁴



Fig. 1 Partial schematic plan view of the TFTR Test Cell showing the location of the activation measurements relative to the vessel, and other components.

III. ACTIVATION MEASUREMENTS

A. Ionization Chamber Measurements



Fig. 2. The dose rate at Vessel Bay-C, 1.2 m from lower vessel (Fig.1, Location 1) as measured at 6:30 am each morning with Ionization Chamber.

Activation measurements were performed with four gamma-ray Argon Ionization Chamber detectors. Three detectors were located near the vessel (1.2 m outboard of vessel at R = 4.9 m, and 2 m below mid-plane) and the fourth at a distance at R = 24.5 m at about 1 m below midplane. These measurements were performed automatically every 15 minutes and computer archived. During experimental operations, TFTR operated in the pulse mode with pulse lengths 5 to 10 sec with a dwell time between pulses about 8 to 15 minutes, from about 0800 hrs to 2300 hrs, 5 days per week. Fig. 2 shows Ionization Chamber measurements of the TFTR activation gamma dose rate at Bay C, about 1.2 m from the lower vessel (Fig.1, Location 1), as measured at 0630 hrs each morning, about 7.5 hrs after the end of operations on the previous day. The activity shown in Fig. 2 exhibits a large increase after each day of D-T operation which then rapidly decayed to a long-term level higher than prior to the sequence of D-T discharges. A similar behavior was exhibited in the activations measured at the other TFTR locations but with differences attributed to different local materials and shielding. During D-T operations, all of the ionization chamber locations exhibited a burst of activation during D-T operations which decayed to an activation level higher than before the D-T burst (about 1-10 mSv/hr after 7 hr). Typical contact dose rates at a vessel flange 2 weeks after D-T operations were about 1 mSv/hr. The midplane contact dose rates about 400 days after shutdown were about 0.3 mSv/hr (30 mR/hr). The measured dose rates exhibited complex decay rates due to many contributing short-lived (hours to days), intermediate (months), and long-lived (years) activities.² The vessel activation decayed with a

effective fast 3.2 hr decay half-life, followed by about an 18 hr decay half-life, and finally a longer decay half-lives in the range from 230 d and longer. At shorter times, the observed decay half-lives indicate the importance of ⁵⁶Mn (2.6 hr) from the ⁵⁶Fe(n,p)⁵⁶Mn reaction, and ²⁴Na(15 hr) from the 27 Al(n, α) 24 Na reaction due to iron and aluminum structural materials, and other short-lived high intensity activations due to unforeseen impurities in certain construction materials. The longer lived activity results from Fe-Cr-Ni related radioisotopes, such as ⁵⁴Mn, and 60 Co generated by various reaction channels and the 58 Co from activation of structural alloys containing Ni such as, SS-304, SS-316, Nitronic, and Inconels.² The long-lived contributions became increasingly observable as the shorter-lived activities decayed and thereby improved the signal-to-background ratio for sensitive gamma-ray energy spectroscopy measurements.

B. Gamma-ray Energy Spectroscopy Measurements

Gamma-ray energy spectroscopy measurements described here were performed at selected Test Cell locations about 326 days after shutdown, to identify the long-term isotopic contributions to the observed decay activities. The measurements were performed with a high purity Ge, high efficiency (41% efficiency relative to NaI at 1333 keV), portable gamma-ray detector connected to a PC-based signal processing and data analysis system. The results exhibited considerable dependence on location due to local variations in material density and isotopic composition, as was found with the Argon Ionization Chamber measurements described above.

Fig. 3 shows the gamma-ray energy spectrum measured at the Northwest Entrance to the Test Cell, at vessel Bay-R, at major radius R = 22.4 m (Fig.1, Location 2) for 301 s with a detection dead-time of 0.96%. The location background dose was 0.4 µSv/hr (40 µrem/hr). This background is significantly less than a previously reported measurement at the same location with a similar detector system performed 14 hrs after a day of intense D-T operation when the activity was 7 μ Sv/hr and the system dead-time was 7% using 1.1 cm wide Pb slits in front of the detector. In the case of the present measurements, the dominant activities shown in Fig. 3 are 54 Mn (γ -ray energy = 834.8 keV), ⁶⁰Co (1173.2 keV, 1332.5 keV), and ⁴⁰K (1460.8 keV). The ⁵⁴Mn (half life = 312 d) activity is attributed to the ⁵⁵Mn(n, 2n)⁵⁴Mn and ⁵⁴Fe(n, p)⁵⁴Mn reactions from the activation of structural alloys (e.g., SS-304,-316,-321, Nitronic-33, and Inconels-600,-718,-750). The 60 Co (5.26 yr) activity is generated through various reaction channels in structural materials. The ⁴⁰K (1.27 x 10^9 yr) activity is attributed to natural background radiation from the concrete in the Test Cell shielding structure. An



Fig. 3. Gamma-ray energy spectrum measured the NW entrance to the Test Cell at R = 22.4 m (Fig. 1, Location 2).



Fig. 4. Gamma-ray energy spectrum measured at Vessel Bay-R/Q at R = 10.7 m (Fig.1, Location 3).



Fig. 5. Gamma-ray energy spectrum measured at Vessel Bay-B at R = 9.8 m (Fig.1, Location 4).



Fig. 6. Gamma-ray energy spectrum measured at Vessel Bay-K at R = 15.3 m (Fig.1, Location 8).



Fig. 7. Gamma-ray energy spectrum measured at the RF antenna and feedthrough assembly after removal from in-vessel service at Bay-L (Fig.1, Location 9).

indication of ⁵⁷Co (122.1 keV) and ⁵⁸Co (810.8 keV) were also observed, but were more evident at other measurement locatons (refer below).

Fig. 4 shows the gamma-ray energy spectrum measured at almost the same toroidal angle, passing between the NW Pump-Duct and a Neutral Beam Injection system, at vessel Bay-R/Q but closer to the vessel at R=10.7m (Fig.1, Location 3) for 309 s with detection dead-time of 3.1%. The location background dose rate was 2 μ Sv/hr (200 μ rem/hr). The dominant activities include those shown in Fig.3, and in addition, the ⁵⁷Co (122.1 keV) and ⁵⁸Co(810.8 keV). The ⁵⁷Co (270.9 d) activity is attributed to the ⁵⁸Ni(n, d)⁵⁷Co and ⁵⁸Ni(n, n'p)⁵⁷Co reactions, and the ⁵⁸Co (70.8 d) activity to the ⁵⁸Ni(n, p)⁵⁸Co reaction from nearby nickel alloys.

Fig. 5 shows the gamma-ray energy spectrum measured at vessel Bay-B at R = 9.8 m (Fig.1, Location 4) for 305 s with a detection dead-time of 4.1%. This location was between the NE Pump-Duct and the Li Pellet Injection system. The background dose rate was 2.5 μ Sv/hr (250)

µrem/hr). As in the case of Fig. 4, at this similarly close location, the dominant activities include those shown in Fig. 4. In addition, the spectrum exhibits a weak indication of ¹²⁴Sb (1690 keV) (not seen in the Fig.4 semi-log plot). After intense D-T operations, short-lived ¹²²Sb (2.75 d) was observed, and is attributed to components of a protective tape used to wrap cable installations. The weak ¹²⁴Sb activity may have a similar origin, or may be contributed by components which occur in lead and brazing alloys. A similar spectrum with weaker intensity was measured at vessel Bay-B at R = 15.5 m (Fig.1, Location 5).

Gamma-spectra energy were measured at Vessel Bay-F, at R= 8.8 m (Fig.1, Location 6) and R = 14 m (Fig.1, Location 7). The location background dose rates were 2 μ Sv/hr (200 μ rem/hr) and 1.2 μ Sv/hr (120 μ rem/hr), respectively; the Ge gamma-ray detection dead-times were 3.6% and 1.5%, respectively. This region is flanked by the NE Pump-Duct on one side, and is part of the toroidal arc from Bays-A to -F containing a high concentration of diagnostic hardware of various materials. The measured gamma-ray energy spectra exhibited the same activities identified in Fig. 5 but in different intensities. The results are summarized in Table 1.

Fig. 6 shows the gamma-ray energy spectrum measured at Vessel Bay-K on the south side of TFTR at R = 15.3 m (Fig.1, location 8) for 303 s with a detection dead-time of 1.2%. This location was between a Neutral Beam Injection system and an array of RF wave-guides running radially toward the vessel mid-plane. The location background dose rate was 1.1 μ Sv/hr (110 μ rem/hr). The dominant activities include those exhibited at Bays-B and -F, and in addition ⁶⁵Zn (1115.4 keV). The ⁶⁵Zr (245 d) activity may have resulted from the ⁶⁴Zn(n, γ)⁶⁵Zn (natural has 4.6% ⁶⁴Zn), and the ⁶⁶Zn(n, 2n)⁶⁵Zn (natural Zn has 27.9% ⁶⁶Zn) reactions from Zn containing copper alloys in the RF antenna wave-guide structures.

Fig. 7 shows the gamma-ray energy spectrum measured for 322 s with a detection dead-time of 7% at a distance of about 1 m at the RF antenna and feedthrough assembly that had been removed from its port location at Vessel Bay-L and placed within a tritium containment tent in the NE corner of the Test Cell at R = 20 m (Fig.1, Location 9). The location background dose rate was 5 μ Sv/hr (500 μ rem/hr). This background rate was significantly higher than comparable major radius locations in the Test Cell and is attributed predominantly to the activation of the RF antenna assembly. As was observed at locations near similar construction materials, the dominant

Location (Refer to Fig.1)	Isotope, γ-ray energy (keV)	Net Activity (CPM)	Location (Refer to Fig.1)	Isotope, γ-ray energy (keV)	Net Activity (CPM)
NW Entrance Bay-R R=22.4m, (2)	Mn-54, (835) Co-60, (1173) Co-60, (1332) K-40, (1461)	116 258 253 14.4	Bay-F R=8.8m (6)	Co-57, (122) Co-58, (811) Mn-54, (835) Co-60, (1173) Co-60, (1332) K-40, (1461) Sb-124, (1690)	792 228 1899 2582 2487 16.7 9.0
Bay-R/Q R=10.7m (3)	Co-57, (122) Co-58, (811) Mn-54, (835) Co-60, (1173) Co-60, (1322) K-40, (1461)	1056 1451 1578 2638 2467 11.8	Bay-F R=14m (7)	Co-57, (122) Co-58, (811) Mn-54, (835) Co-60, (1173) Co-60, (1332) K-40, (1461) Sb-124, (1690)	252 82 632 995 932 12.4 1.2
Bay-B R=9.8m (4)	Co-57, (122) Co-58, (811) Mn-54, (835) Co-60, (1173) Co-60, (1322) K-40, (1461) Sb-124, (1690)	1095 287 2308 3048 2867 8.2 11.2	Bay-K R=15.3m (8)	Co-57, (122) Co-58, (811) Mn-54, (835) Zn-65, (1115) Co-60, (1173) Co-60, (1332) K-40, (1461)	196 54 523 76 921 866 14.2
Bay-B R=15.5m (5)	Co-57, (122) Co-58, (811) Mn-54, (835) Co-60, (1173) Co-60, (1332) K-40, (1461) Sb-124, (1690)	382 134 865 989 972 16.5 0.4	At Removed RF Antenna (9)	Co-57, (122) Co-57, (136) Co-58, (811) Ag-110m, (658) Ag-110m, (885) Ag-110m, (1384) Mn-54, (835) Co-60, (1173) Co-60, (1332) K-40, (1461)	12551 2186 5144 504 349 159 78 9724 1944 1842 18.7

Table 1. Summary of the gamma-ray energy spectroscopy measurements shown in Fig.s 3-7. All measurements were made with the same detector 326 days after shutdown. Refer to text for discussion of net activity in detector counts per minute (CPM).

activities were 54 Mn (834.8 keV), 60 Co (1173.2 keV, 1332.5 keV), 40 K (1460.8 keV), 57 Co (122.1 keV) and 58 Co(810.8 keV), and 124 Sb (1690.0 keV), and in addition, 110m Ag (657.8 keV, 884.7 keV, 937.5 keV, 1384.3 keV). The 110m Ag (249.9d) activity is attributed to the

 109 Ag(n, γ) 110m Ag reaction from extensive Ag electrical contact coatings on antenna parts. Table 1 summarizes the gamma-ray energy spectroscopy measurements discussed above. All measurements were made with the same detector. Listed are the observed isotopes at each location

Table 2. Summary of activity measurements on samples removed from TFTR and counted about 390 d after shutdown. Shown are the sample description, mass (g), approximate major radius R(m), shortest distance from sample to Vessel Top (VT) or shortest distance from sample to Vessel Midplane (VM), the toroidal angle ϕ (degree), dominant activities, and activation per unit mass (μ Ci/g).

Sample	Mass	R(m)	d(m)	φ	Isotope	e Activity	Sample	Mass	R(m)	d(m)	φ	Isotope	Activity
Description	(g)	~ /		т	I	uCi/g	Description	(g)	~ /		Т		uCi/g
#980061 SS bolt from glow probe	57.56	3.9	2.13 from VT	270	Mn-54 Co-57 Co-58 Co-60	4.25E-5 1.84E-5 3.38E-6 2.39E-4	#980092 brass bus joint bolt	175	3.9	0.61 from VT	180	Mn-54 Co-60	1.29E-5 5.94E-4
#980062 brass bus joint bolt	174.00	3.9	0.61 from VT	234	Mn-54 Co-60	2.20E-5 8.59E-4	#980093 SS bus joint washer	21.9	3.9	0.61 from VT	180	Mn-54 Co-57 Co-58 Co-60	3.45E-4 7.74E-3 7.02E-4 5.78E-3
#980063 Cu /SS/plastic ground strap	12.4	3.9	0.20 from VT	126	Co-60 Zn-65 Sb-124	3.04E-3 1.45E-4 7.68E-5	#980094 Cu tritium ground strap	69.37	3.9	1.82 from VT	9	Co-60 Sb-124	8.89E-5 2.30E-6
#980064 SS support bolt	7.39	3.9	0.10 from VT	126	Mn-54 Co-57 Co-58 Co-60	3.39E-3 3.14E-3 3.61E-4 5.24E-3	#980095 Cu ground strap	66.41	3.9	1.82 from VT	63	Co-60 Sb-124	9.35E-5 2.17E-6
#980071 brass bus joint bolt	174.9	3.9	0.46 from VT	0	Mn-54 Co-60	1.11E-4 5.09E-4	#980100 brass screws	3.11	3.9	0.10 from VT	126	Co-60 Zn-65	2.08E-3 7.93E-3
#980072 SS bus joint washer	21.99	3.9	0.46 from VT	0	Mn-54 Co-57 Co-58 Co-60	3.04E-4 6.91E-3 6.48E-4 5.47E-3	#980101 SS support nut	3.11	3.9	0.10 from VT	126	Mn-54 Co-57 Co-58 Co-60	4.58E-3 3.27E-3 3.57E-4 4.85E-3
#980073 brass/plastic Swagelock fitting	12.63	4.7	0.91 from VM	0	Co-60 Zn-65	2.12E-4 9.63E-4	#980102 brass bus joint bolt	175	3.9	0.46 from VT	126	Mn-54 Co-60	1.01E-4 4.76E-4
#980074 SS/plastic Swagelock fitting	11.88	4.6	0.76 from VM	0	Mn-54 Co-57 Co-58 Co-60	1.14E-3 8.13E-4 1.09E-4 2.15E-3	#98103 SS bus joint washer	21.9	3.9	0.46 from VT	126	Mn-54 Co-57 Co-58 Co-60	2.79E-4 5.70E-3 6.14E-4 6.24E-3
#980075 support plate	163.5	5.32	1.52 from VM	0	Co-60	2.50E-5	#980104 brass bus joint bolt	175	3.9	1.52 from VT	288	Mn-54 Co-60	1.10E-4 6.11E-4
#980076 Cu/plastic ground strap	3.9	3.9	2.01 from VM	270	Co-60	1.42E-4	#980105 SS bus joint washer	22	3.9	1.52 from VT	288	Mn-54 Co-57 Co-58 Co-60	3.65E-4 8.19E-3 7.40E-4 6.19E-3
#980082 brass bus joint bolt	128.6	3.9	0.81 from VT	0	Mn-54 Co-57 Co-58 Co-60	1.83E-4 9.07E-5 1.76E-5 2.13E-3	#980124 SS support bolt	21.3	5.0	1.22 from VM	0	Mn-54 Co-57 Co-58 Co-60	3.45E-4 2.59E-4 3.37E-5 8.81E-4
#980083 SS bus joint washer	31.26	3.9	0.81 from VT	0	Mn-54 Co-57 Co-58 Co-60	3.57E-4 2.06E-4 2.04E-5 1.12E-3	#980126 SS insulator bolt	32.17	5.0	1.22 from VM	72	Mn-54 Co-57 Co-58 Co-60	2.17E-4 1.41E-4 1.72E-5 4.44E-4
#980090 SS support bracket	42.1	3.9	0.46 from VT	126	Mn-54 Co-57 Co-58 Co-60	2.37E-3 1.40E-3 1.52E-4 2.80E-3	#980127 SS insulator washer	7	5.0	1.22 from VM	72	Mn-54 Co-57 Co-58 Co-60	3.54E-4 4.06E-4 2.63E-5 1.31E-3
#980091 SS support bolt	18.77	3.9	0.46 from VT	126	Mn-54 Co-57 Co-58 Co-60	2.05E-3 2.04E-3 2.33E-4 4.34E-3							

and the measured detector counts per minute uncorrected for detector absolute efficiency. There are a number of difficult issues involved in attempting to convert the measured detector count rate for a particular isotopic gamma-ray energy to yield the absolute activity of the source in the TFTR Test Cell. In particular, these are extended sources of gamma-rays with varying geometries, gamma-ray absorption, attenuation, shielding, scattering, and angle of incidence on the detector. The available absolute efficiency calibration procedures usually involve small point or disk calibration sources on a known backing, placed in front of a detector to give an accurately defined solid angle with minimal scattering and background effects. The complexity of the TFTR experimental configuration prevents this approach in a manner that would yield sufficient accuracy for benchmarking purposes.

In view of these difficulties, an alternate approach was pursued to obtain an activity calibration of the global TFTR activation decay with sufficient accuracy to provide benchmarking for the present measurements and future simulations. Small activated TFTR parts, fabricated from representative TFTR materials were removed from well characterized vessel and near-vessel locations. The activations of these samples were measured in the PPPL Radiological and Environmental Measurements Laboratory about 390 d after shutdown using high resolution, high efficiency, low background, gamma-ray spectroscopy systems with efficiency calibrations traceable to NIST standards. The results of these measurements are summarized in Table 2. Shown are the sample description, exposure location, mass, dominant activities and activation per unit mass. Future work will use these and the results of similar measurements in progress to benchmark simulations of the TFTR cool-down for testing low activation reactor design codes

III. SUMMARY AND CONCLUSIONS

The measurements have characterized residual TFTR neutron activation. The results indicate that, in general, the observed gamma activities are similar to those expected from previous simulation work, and can be used for several applications, including accurate determinations of low activation long-lived cooling, and benchmarking advanced simulations of D-T fusion reactor neutron activation.

ACKNOWLEDGMENTS

The technical contributions of Keith Chase and Joseph Winston are gratefully acknowledged. This work was supported by US DOE Contract No. DE-AC02-76CH03073.

REFERENCES

1. R. J. Hawryluk, *et al.*, "Fusion Plasma Experiments on TFTR: A 20 Year Retrospective", Phys. Plasmas, **5**(5) 1577 (1998).

2. L.-P. Ku and J. G. Kolibal, "Radioactivation Characteristics for the Tokamak Fusion Test Reactor", Nuc. Technol./Fus.,4 (11), 586 (1983).

3. L. P. Ku and S. L. Liew, "Global Dose Rate in TFTR Due to Neutron Induced Residual Radioactivities During DT Operation", Proceedings of the 15th IEEE/NPSS Symposium on Fusion Engineering, p 313, October 11-15, 1993, Hyannis, MA, and references therein.

4. H. W. Kugel *et al.*, "Measurements of TFTR D-T Radiation Shielding Efficiency", Fus. Eng. & Design, **28**, 534 (1995).