Standard Practice for Determining Neutron Fluence, Fluence Rate, and Spectra by Radioactivation Techniques

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1. Scope

1.1 This practice describes procedures for the determination of neutron fluence rate, fluence, and energy spectra from the radioactivity that is induced in a detector specimen.

1.2 The practice is directed toward the determination of these quantities in connection with radiation effects on materials.

1.3 For application of these techniques to reactor vessel surveillance, see also Test Methods E1005.

1.4 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.

NOTE 1—Detailed methods for individual detectors are given in the following ASTM test methods: E262, E263, E264, E265, E266, E343, E393, E481, E523, E526, E704, E705, and E854.

2. Referenced Documents

2.1 ASTM Standards:

E170 Terminology Relating to Radiation Measurements and Dosimetry

E181 Test Methods for Detector Calibration and Analysis of Radionuclides

E262 Test Method for Determining Thermal Neutron Reaction Rates and Thermal Neutron Fluence Rates by Radioactivation Techniques

E263 Test Method for Measuring Fast-Neutron Reaction Rates by Radioactivation of Iron

E264 Test Method for Measuring Fast-Neutron Reaction Rates by Radioactivation of Nickel

E265 Test Method for Measuring Reaction Rates and Fast-Neutron Fluences by Radioactivation of Sulfur-32

E266 Test Method for Measuring Fast-Neutron Reaction Rates by Radioactivation of Aluminum

E343 Test Method for Measuring Reaction Rates by Analysis of Molybdenum-99 Radioactivity From Fission Dosimeters

E393 Test Method for Measuring Reaction Rates by Analysis of Barium-140 From Fission Dosimeters

E481 Test Method for Measuring Neutron Fluence Rates by Radioactivation of Cobalt and Silver

E523 Test Method for Measuring Fast-Neutron Reaction Rates by Radioactivation of Copper

E526 Test Method for Measuring Fast-Neutron Reaction Rates by Radioactivation of Titanium

E693 Practice for Characterizing Neutron Exposures in Iron and Low Alloy Steels in Terms of Displacements Per Atom (DPA), E 706(ID)

E704 Test Method for Measuring Reaction Rates by Radioactivation of Uranium-238

E705 Test Method for Measuring Reaction Rates by Radioactivation of Neptunium-237

E722 Practice for Characterizing Neutron Fluence Spectra in Terms of an Equivalent Monoenergetic Neutron Fluence for Radiation-Hardness Testing of Electronics


E944 Guide for Application of Neutron Spectrum Adjustment Methods in Reactor Surveillance, E 706 (IIA)

E1005 Test Method for Application and Analysis of Radiometric Monitors for Reactor Vessel Surveillance, E 706(IIIA)

E1018 Guide for Application of ASTM Evaluated Cross Section Data File, Matrix E706 (IIB)


2.2 ISO Standard:


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Guide in the Expression of Uncertainty in Measurement

3. Terminology
3.1 Descriptions of terms relating to dosimetry are found in Terminology E170.

4. Summary of Practice
4.1 A sample containing a known amount of the nuclide to be activated is placed in the neutron field. The sample is removed after a measured period of time and the induced activity is determined.

5. Significance and Use
5.1 Transmutation Processes—The effect on materials of bombardment by neutrons depends on the energy of the neutrons; therefore, it is important that the energy distribution of the neutron fluence, as well as the total fluence, be determined.

6. Counting Apparatus
6.1 A number of instruments are used to determine the disintegration rate of the radioactive product of the neutron-induced reaction. These include the scintillation counters, ionization chambers, proportional counters, Geiger tubes, and solid state detectors. Recommendations of counters for particular applications are given in General Methods E181.

7. Requirements for Activation-Detector Materials
7.1 Considerations concerning the suitability of a material for use as an activation detector are found in Guide E844.

7.2 The amounts of fissionable material needed for fission threshold detectors is given in Test Methods E343, E393, and E854, and Guide E844.

8. Irradiation Procedures
8.1 The irradiations are carried out in two ways depending upon whether the instantaneous fluence rate or the fluence is being determined. For fluence rate, irradiate the detector for a short period at sufficiently low power that handling difficulties and shielding requirements are minimized. Then extrapolate the resulting fluence rate value to the value anticipated for full reactor power. This technique is sometimes used for the fluence mapping of reactors (1,2).

8.2 The determination of fluence is most often required in experiments on radiation effects on materials. Irradiate the detectors for the same duration as the experiment at a position in the reactor where, as closely as possible, they will experience the same fluence, or will bracket the fluence of the position of interest. When feasible, place the detectors in the experiment capsule. In this case, long-term irradiations are often required.

8.3 It is desirable, but not required, that the neutron detector be irradiated during the entire time period considered and that a measurable part of the activity generated during the initial period of irradiation be present in the detector at the end of the irradiation. Therefore, the effective half-life, \( t'_{1/2} = 0.693/\lambda' \), of the reaction product should not be much less than the total elapsed time from the initial exposure to the final shutdown.

8.4 As mentioned in 9.11 and 9.12, the use of cadmium-shielded detectors is convenient in separating contributions to the measured activity from thermal and epithermal neutrons. Also, cadmium-shielding is helpful in reducing activities due to impurities and the loss of the activated nuclide by thermal-neutron absorption. The recommended thicknesses of cadmium is 1 mm. When bare and cadmium-shielded samples are placed in the same vicinity, take care to avoid partial shielding of the bare detectors by the cadmium-shielded ones.

9. Calculation
9.1 The activity of the sample, \( A \), at the end of the exposure period is calculated as follows:

\[
A = \lambda D[(1 - \exp(-\lambda t_i)) \exp(-\Lambda t_w)]
\]

where:
\( \lambda \) = decay constant for the radioactive nuclide,
\( t_i \) = time interval for counting,
\( t_w \) = time elapsed between the end of the irradiation period and the start of the counting period, and
\( D \) = number of disintegrations (net number of counts corrected for background, random and true coincidence losses, efficiency of the counting system, and fraction of the sample counted) in the interval \( t_i \).

9.1.1 If, as is often the case, the counting period is short compared to the half-life \( = 0.693/\lambda \) of the radioactive nuclide, the activity is well approximated as follows:

\[
A = D[t_i \exp(-\Lambda t_w)]
\]

9.2 For irradiations at constant fluence rate, the saturation activity, \( A_s \), is calculated as follows:

\[
A_s = A/(1 - \exp(-\lambda t_i))
\]

where:
\( t_i \) = exposure duration, and
\( \lambda' \) = effective decay constant during the irradiation.

Note 2—The saturation activity corresponds to the number of disintegrations per foil per unit time for the steady-state condition in which the rate of production of the radioactive nuclide is equal to the rate of loss by radioactive decay and transmutation.

9.2.1 The effective decay constant, which may be a function of time, is related to the decay constant as follows:

\[
\lambda' = \lambda + \int_0^\infty \sigma_s(E) \phi(E) \, dE
\]

where:
\( \sigma_s(E) \) = neutron absorption cross section for the product nuclide, and
\( \phi(E) \) = neutron fluence rate per unit energy.

9.2.2 Application of the effective decay constant for irradiations under varying fluence rates is discussed in this section and in the detailed methods for individual detectors.

9.3 The reaction rate is calculated as follows:
\[
R_s = A_s \lambda' / N_A \lambda
\]  
(5)

where:
\(N = \) number of target nuclei in the detector at time of irradiation.

9.3.1 The number of target nuclei can often be assumed to be equal to \(N_0\), the number prior to irradiation.

\[N_{\infty} = N_0 F/m \Lambda M\]  
(6)

where:
\(N_A = \) Avogadro’s number
\(= 6.022 \times 10^{23} \text{ mole}^{-1}\),
\(F = \) atom fraction of the target nuclide in the target element,
\(m = \) mass of target element, \(g\), and
\(M = \) atomic mass of the target element.

9.3.2 Calculations of the isotopic concentration after irradiation is discussed in 9.6.6 and in the detailed methods for individual detectors.

9.4 The neutron fluence rate, \(\phi\), is calculated as follows:

\[
\phi = R / \sigma
\]  
(7)

where:
\(\sigma = \) the spectral weighted neutron activation cross section.

9.4.1 Cross sections should be processed from an appropriate cross-section library that includes covariance data. Guide E1018 provides information and recommendations on how to select the cross section library. The International Reactor Dosimetry File (IRDF-2002) (3) is one good source for cross sections. The SNLRLML cross section compendium (4) provides a processed fine-group representation of recommended dosimetry cross sections and covariance matrices.

9.4.2 If spectral-averaged cross-section or spectrum data are not available, one of the alternative procedures discussed in 9.10 to 9.13 may be used to calculate an approximate neutron fluence rate from the saturation activity.

9.5 The neutron fluence, \(\Phi\), is related to the time varying differential neutron fluence rate \(\phi(E, t)\) by the following expression:

\[
\Phi = \int_0^\infty \int_0^{t_i} \phi (E, t) \, dt \, dE
\]  
(8)

where:
\(t_2 - t_1 = \) duration of the irradiation period.

9.5.1 Long irradiations usually involve operation at various power levels, and changes in isotopic content of the system; under such conditions \(\phi(E, t)\) can show large variations with time.

9.5.2 It is usual to assume, however, that the neutron fluence rate is directly proportional to reactor power; under these conditions, the fluence can be well approximated by:

\[
\Phi = \left( \frac{\phi}{P} \right) \sum_{i=1}^n P_i t_i
\]  
(9)

where:
\(\phi/P = \) average value of the neutron fluence rate, \(\phi\), at a reference power level, \(P\).

9.5.2.1 Alternate methods include measuring the power generation rate in a fraction of the reactor volume adjacent to the volume of interest.

9.6 Transmutation Processes:

9.6.1 The neutron fluence rate spectrum, \(\phi(E)\), can be determined by computer calculations using neutron transport codes, and adjustment techniques using radioactivation data from multiple foil irradiations.

9.6.2 The reaction rate is related to the fluence rate by the following equation:

\[
R_i = \int_0^\infty \sigma(E) \phi(E) \, dE
\]  
(10)

where:
\(\sigma(E) = \) activation cross section at energy \(E\), and
\(\phi(E) = \) differential neutron fluence rate, that is the fluence per unit energy per unit time for neutrons with energies between \(E\) and \(E + dE\).

9.6.3 The number of nuclei, \(N_p\), of a radioactive product nuclide is related to the reaction rate by the following equation:

\[
dN_p/dt = NR_i - N_p \lambda'
\]  
(11)

9.6.4 Solution of Eq 11, for the case where \(R_s\) and \(N\) are constant, yields the following expression for the activity of a foil:

\[
A = N_{\lambda'} = (\lambda / \lambda') N R_i \left(1 - e^{-\lambda' t}\right)
\]  
(12)

9.6.5 The saturation activity of a foil is defined as the activity when \(dN_p/dt = 0\); thus Eq 11 yields the following relationship for the saturation activity:

\[
A_s = (\lambda / \lambda') N R_i
\]  
(13)

9.6.6 The isotopic content of the target nuclide may be reduced during the irradiation by more than one transmutation process and it may be increased by transmutations of other nuclides so that the rate of change of the number of target nuclei with time is described by a number of terms:

\[
dN_p/dt = -N (R_i + \sum_{j=1}^n R_j) + \sum_{j=1}^n N_j R_j
\]  
(14)

where:
\(i = \) discrete transmutation path for removal of the target isotope, and
\(j = \) discrete transmutation reaction whereby the target isotope is produced from isotope \(N_j\) and each of the \(R_i\) and \(R_j\) terms could be calculated from equations similar to Eq 10, using the appropriate cross sections.

9.6.6.1 The \(R_i\) term may predominate and, if \(R_i\) is constant, Eq 14 can be solved as \(N = N_{\infty} \exp (-R_i t)\). The change in the target composition may be negligible and \(N\) may, in that case, be approximated by \(N_{\infty}\).

9.6.7 During irradiation, the effective decay rate is increased by transmutations of the product isotope (see Eq 4).

9.7 Long Term Irradiations: