

IRDFF-II: A New Neutron Metrology Library

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High quality nuclear data is the most fundamental underpinning for all neutron metrology applications. This paper describes the release of version II of the International Reactor Dosimetry and Fusion File (IRDFF-II) that contains a consistent set of nuclear data for fission and fusion neutron metrology applications up to 60 MeV neutron energy. The library is intended to support: a) applications in research reactors; b) safety and regulatory applications in the nuclear power generation in commercial fission reactors; and c) material damage studies in support of the research and development of advanced fusion concepts. The paper describes the contents of the library, documents the thorough verification process used in its preparation, and provides an extensive set of validation data gathered from a wide range of neutron benchmark fields. The new IRDFF-II library includes 119 metrology reactions, four cover material reactions to support self-shielding corrections, five metrology metrics used by the dosimetry community, and cumulative fission products yields for seven fission products in three different neutron energy regions. In support of characterizing the measurement of the residual nuclei from the dosimetry reactions and the fission product decay modes, the present document lists the recommended decay data, particle emission energies and probabilities for 68 activation products. It also includes neutron spectral characterization data for 29 neutron benchmark fields for the validation of the library contents. Additional six reference fields were assessed (four from plutonium critical assemblies, two measured fields for thermal-neutron induced fission on ²³³U and ²³⁹Pu targets) but not used for validation due to systematic discrepancies in C/E reaction rate values or lack of reaction-rate experimental data. Another ten analytical functions are included that can be useful for calculating average cross sections, average energy, thermal spectrum average cross sections and resonance integrals. The IRDFF-II library and comprehensive documentation is available online at www-nds.iaea.org/IRDFF/. Evaluated cross sections can be compared with experimental data and other evaluations at www-nds.iaea.org/exfor/endl.htm. The new library is expected to become the international reference in neutron metrology for multiple applications.

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B. $^{235}\text{U}_{(n_{th},f)}$ PFNS	72	Well-validated nuclear data evaluation techniques are needed to produce a modern nuclear data evaluated library like IRDFE-II. An overview of nuclear data evaluation methodology can be found in Ref. [3]; recent progress has been reviewed in Refs. [4–6]. Nowadays the generalized least-square method is the most common evaluation method, <i>e.g.</i> , see [7–11]. A similar least-squares approach [12–14] that is based on Padé approximants [15–17] has also been extensively employed by K. Zolotarev to produce many of the new dosimetry evaluations for the IRDFE-II library.	
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A. $^{252}\text{Cf}_{(s.f.)}$	91	The development of the IRDFE-II dosimetry library has greatly benefited from the availability of the comprehensive TENDL libraries [24, 25] ¹ , especially to extend the energy range of the existing evaluations for neutron-induced reactions on medium- and heavy-mass targets.	
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¹ The TENDL-2015.s60 file of the TENDL 2015 library contains exclusive representations of reactions and associated covariances up to 60 MeV. This file was extensively used for the extension of IRDFE-II evaluations up to 60 MeV.

A general purpose neutron evaluated nuclear data file (ENDF) uses the evaluation techniques briefly discussed above to combine model calculations with experimental data to address the assembly of a complete and consistent set of neutron-induced reactions for a given target isotope. In this process an evaluator may need to make some compromises in the recommended nuclear data in order to satisfy consistency constraints on the nuclear data between different reaction channels, *e.g.*, the total cross section must be exactly equal to the sum of the various reaction channel cross sections, while matching as much as possible the available experimental data.

In contrast to a general nuclear data evaluation, an entry in a metrology library is reaction-specific. The evaluation can start from a nuclear model calculation, but relies more heavily on experimental data to obtain recommended energy-dependent cross sections, including full covariance information for that particular reaction channel so that an accurate assessment of the uncertainty in reaction rates can be made for a given neutron spectrum field. Whereas information on the consistency of the evaluated data with integral measurements gathered in various reference neutron fields is not directly taken into account in the actual nuclear data evaluation process, it does influence the decisions to declare a reaction as “recommended” for use in metrology applications.

This paper presents a consistent set of nuclear data intended for neutron metrology that support: a) applications in research reactors; b) safety and regulatory applications in the nuclear power generation in commercial fission reactors; and c) material damage studies in research and development of advanced fusion concepts.

II. OVERVIEW OF THE IRDFF-II LIBRARY

The new IRDFF-II library addresses incident neutron energies from 0 to 60 MeV. The library entries, enumerated in Table 1, include 119 metrology reactions, four cover cross sections used to support shielding corrections, five metrology metrics used by the dosimetry community, seven cumulative fission products yields and corresponding decay data. In support of characterizing the measurement of the residual nuclei from the dosimetry reactions and the fission product decay modes, the library includes decay data, gamma and beta particle emission energies and probabilities for 68 residual nuclei. The library also includes neutron spectral characterization data for 45 neutron benchmark fields that were used to support the validation of the library contents. The following Secs. discuss these entries of the dosimetry library in more detail.

A. Neutron Metrology Reactions

There are several requirements for a given reaction to be accepted for use in metrology applications and included in the IRDFF-II library. These requirements include:

1. The existence of a high-fidelity nuclear data evaluation for the reaction channel that is documented and stored in the library;
2. A residual particle from the reaction that is measurable with a technique in use at multiple laboratories, *e.g.*, gamma activation analysis, beta spectroscopy, mass spectrometry;
3. The existence of uncertainty data for the cross section that must come directly from the cross section evaluation process, captured in the form of a covariance matrix, and stored in the library;
4. A reaction that is of interest to the commercial or research reactor community or to the fusion community (including metrology at high energy accelerators used to support fusion material damage studies, *e.g.*, IFMIF [26]);
5. Evidence that the nuclear data evaluation took into consideration the available body of experimental data.

Table 1 summarizes the 119 metrology reactions that meet these requirements and are included in the IRDFF-II library. The list includes all of the reactions that were present in the earlier IRDF-2002 [1] and IRDFF v1.05 [2] libraries. The differences between library versions are indicated: column 8 indicates whether IRDFF-v1.05 was updated (new) or not (old) compared to IRDF-2002. Likewise, column 9 indicates whether IRDFF-II was updated compared to IRDFF-v1.05. As seen from the table, several of the dosimetry reactions in the IRDFF-II were updated and many reactions were added, such as $^{27}\text{Al}(n,2n)^{26g}\text{Al}$, $^{28}\text{Si}(n,p)^{28}\text{Al}$, $^{29}\text{Si}(n,X)^{28}\text{Al}$, $^{113}\text{In}(n,\gamma)^{114m}\text{In}$, $^{238}\text{U}(n,2n)^{237}\text{U}$, and $^{209}\text{Bi}(n,xn)^{(210-x)}\text{Bi}$. The extensions were introduced to support specialized dosimetry applications.

High resolution atomic mass spectroscopy is often used to measure alpha production in materials such as ^{10}B and ^6Li . Measured alpha particles can originate from reactions other than the (n,α) channel, hence the updated IRDFF-II library also includes reactions for the total alpha production cross section, *i.e.*, $^{10}\text{B}(n,X)^4\text{He}$ and $^6\text{Li}(n,X)^4\text{He}$ reactions. Since boron and lithium are often used with a variety of enrichments in dosimetry applications, nuclear data for the $^{11}\text{B}(n,X)^4\text{He}$ and $^7\text{Li}(n,X)^4\text{He}$ reactions, as well as the composite elemental-based $^{\text{nat}}\text{B}(n,X)^4\text{He}$ and $^{\text{nat}}\text{Li}(n,X)^4\text{He}$ reactions, are included in the library. The availability of the separate ^{11}B and ^7Li alpha-production cross sections enables users to apply the IRDFF-II cross sections to variable enrichment dosimeters that might be employed at various metrology laboratories.

Previous input from users [27] of the IRDFF-1.05 library [2], which was extended to address the energy range up to 60 MeV, indicated that natural elemental dosimeters, rather than isotopic ones, were usually used in dosimetry applications. Since, at the higher incident neutron energies on natural elemental dosimeters, several contributing reaction channels can produce the same residual nucleus that is used as a dosimetry monitor, the IRDFF-II library has also been extended to explicitly address the composite reactions that produce commonly used residual nuclei.

TABLE 1: IRDF-II nuclear data contents and evaluation sources. (r) denotes renormalization. Major sources of data are K.I. Zolotarev evaluations and EAF-2010 [28], JENDL-4 [29], TENDL [24, 25], ENDF/B-VIII.0 [30], and JEFF-3.3 [31] libraries.

ID No.	Reaction	Label	MAT	Energy Interval	Evaluation source	Eval. Date	Consistency	
							v2002 to v-1.05 ¹	v-1.05 to IRDF-II ²
1	^{nat} Li(n,X) ³ H	LiH3	300	0–60 MeV	IRDF-II Li6H3,Li7H3	2019	–	–
2	^{nat} Li(n,X) ⁴ He	LiHe4	300	0–60 MeV	IRDF-II Li6He4,Li7He4	2019	–	–
3	⁶ Li(n,t) ⁴ He	Li6t	325	0–1 MeV	IAEA STD 2017 [32]	2018	new	new
			325	1–2.75 MeV	ENDF/B-VIII.0	2018	new	new
			325	2.75–60 MeV	EAF-2010(r)	2010	new	new
4	⁶ Li(n,X) ³ H	Li6H3	325	0–60 MeV	IRDF-II Li6t	2019	–	–
5	⁶ Li(n,X) ⁴ He	Li6He4	325	0–20 MeV	MT24–ENDF/B-VIII.0	2018	–	–
			325	20–60 MeV	EAF-2010(r) (n,2n α)	2010	–	–
			325	0–20 MeV	MT32–ENDF/B-VIII.0 (n,n')	2018	–	–
			325	20–60 MeV	EAF-2010(r) (n,nd)	2010	–	–
			325	0–60 MeV	IRDF-II Li6t	2019	–	–
6	⁷ Li(n,X) ⁴ He	Li7He4	328	0–60 MeV	MT24–ENDF/B-VIII.0	2018	–	–
			328	20–60 MeV	EAF-2010(r) (n,2n α)	2010	–	–
			328	0–20 MeV	MT25–ENDF/B-VIII.0	2018	–	–
			328	20–60 MeV	EAF-2010(r) (n,np)	2010	–	–
			328	0–20 MeV	MT33=ENDF/B-VIII.0 (n,n')	2018	–	–
			328	20–60 MeV	EAF-2010(r) (n,n α)	2010	–	–
			328	0–20 MeV	MT102–ENDF/B-VIII.0	2018	–	–
			325	20–60 MeV	EAF-2010(r) (n, γ)	2010	–	–
7	⁷ Li(n,X) ³ H	Li7H3	328	0–20 MeV	MT33–ENDF/B-VIII.0 (n,n')	2018	–	–
			328	20–60 MeV	EAF-2010(r) (n,n α)	2010	–	–
8	^{nat} B(n,X) ³ H	BH3	500	0–60 MeV	IRDF-II B10H3,B11H3	2019	–	–
9	^{nat} B(n,X) ⁴ He	BHe4	500	0–60 MeV	IRDF-II B10He4,B11He4	2019	–	–
10	¹⁰ B(n, α) ⁷ Li	B10a	525	0–1 MeV	MT107–IAEA STD 2017 [32]	2018	new	new
			525	1–60 MeV	EAF-2010(r) (n, α)	2010	new	new
11	¹⁰ B(n,X) ⁴ He	B10He4	525	0–20 MeV	MT22–ENDF/B-VIII.0 (n,n')	2018	–	–
			525	20–60 MeV	EAF-2010(r) (n, α)	2010	–	–
			525	0–20 MeV	MT35–ENDF/B-VIII.0 (n,n')	2018	–	–
			525	20–60 MeV	EAF-2010(r) (n,d2 α)	2010	–	–
			525	0–20 MeV	MT106–EAF-2010 (n, ³ He)	2010	–	–
			525	0–60 MeV	IRDF-II B10a	2019	–	–
12	¹⁰ B(n,X) ³ H	B10H3	525	0–20 MeV	MT113–ENDF/B-VIII.0 MT700	2018	–	–
			525	20–60 MeV	EAF-2010(r) (n,t)	2010	–	–
			525	E_{th} –60 MeV	MT155–EAF-2010 (n,t α)	2010	–	–
13	¹¹ B(n,X) ³ H	B11H3	528	0–20 MeV	MT105–ENDF/B-VIII.0	2018	–	–
			528	20–60 MeV	EAF-2010(r) (n,t)	2010	–	–
			528	0–60 MeV	MT33–EAF-2010 (n,nt)	2010	–	–
14	¹¹ B(n,X) ⁴ He	B11He4	528	0–60 MeV	MT11–EAF-2010 (n,2nd)	2010	–	–
			528	0–20 MeV	MT22–ENDF/B-VIII.0(r) (n,n α)	2018	–	–
			528	20–60 MeV	EAF-2010(r) (n,n α)	2010	–	–
			528	0–60 MeV	MT24–EAF-2010	2010	–	–
			528	0–60 MeV	MT33–EAF-2010 (n,nt)	2010	–	–
			528	0–20 MeV	MT107–ENDF/B-VIII.0 (n, α)	2018	–	–
			528	20–60 MeV	EAF-2010(r) (n, α)	2010	–	–
			528	0–60 MeV	MT117–EAF-2010 (n,d α)	2010	–	–
15	¹⁹ F(n,2n) ¹⁸ F	F192	925	E_{th} –30 MeV	RRDF-2002	2002	old	old
				30–60 MeV	TENDL-2010(r)	2010	–	old
16	²³ Na(n,2n) ²² Na	Na232	1125	E_{th} –60 MeV	INDC(NDS)-0705	2017	old	new
17	²³ Na(n, γ) ²⁴ Na	Na23g	1125	0–20 MeV	INDC(NDS)-0705	2017	–	new
				20–60 MeV	TENDL-2010(r)	2012	–	new

¹ Column (v2002 to v-1.05) marks any change in IRDF-II-v1.05 compared to IRDF-2002 as a reference up to 20 MeV.² Column (v-1.05 to IRDF-II) marks any change in IRDF-II compared to IRDF-II-v1.05 as a reference.

Consistency legend: new = new evaluation; old = unchanged; – = not present in reference.

TABLE 1: (continued). IRDF-II nuclear data contents and evaluation sources. (r) denotes renormalization.

ID No.	Reaction	Label	MAT	Energy Interval	Evaluation source	Eval. Date	Consistency	
							v2002 to v-1.05 ¹	v-1.05 to IRDF-II ²
18	^{nat} Mg(n,X) ²⁴ Na	MgNa24	1200	Up to 60 MeV Minor channels	IRDF-II Mg24p TENDL-2015(r)	2019 2019	–	–
19	²⁴ Mg(n,p) ²⁴ Na	Mg24p	1225	E_{th} -21 MeV 21-60 MeV	INDC(NDS)-0526 TENDL-2010(r)	2008	new	old
20	²⁷ Al(n,p) ²⁷ Mg	Al27p	1325	E_{th} -22 MeV 22-60 MeV	INDC(CCP)-0438 TENDL-2010(r)	2004	new	old
21	²⁷ Al(n, α) ²⁴ Na	Al27a	1325	E_{th} -30 MeV 30-60 MeV	INDC(NDS)-0546 TENDL-2010(r)	2009	old	old
22	²⁷ Al(n,X) ²⁴ Na	Al27Na24	1325	Up to 60 MeV Minor channels	IRDF-II Al27a TENDL-2015	2019	–	–
23	²⁷ Al(n,2n) ^{26g} Al	Al272g	1325	E_{th} -60 MeV	INDC(NDS)-0705	2017	–	–
24	²⁸ Si(n,p) ²⁸ Al	Si28p	1425	E_{th} -21 MeV 21-60 MeV	INDC(NDS)-0668 TENDL-2015(r)	2014	–	new
25	²⁹ Si(n,X) ²⁸ Al	Si29Al28	1428	E_{th} -60 MeV Minor channels	IRDF-II Si29p TENDL-2015(r)	2019	–	–
26	^{nat} Si(n,X) ²⁸ Al	SiAl28	1400	E_{th} -60 MeV	IRDF-II Si-28,29	2019	–	–
27	³¹ P(n,p) ³¹ Si	P31p	1525	E_{th} -21 MeV 21-60 MeV	INDC(NDS)-0668 TENDL-2015(r)	2014	new	old
28	^{nat} S(n,X) ³² P	SP32	1600	E_{th} -60 MeV	IRDF-II S32p	2019	–	–
29	³² S(n,p) ³² P	S32p	1625	E_{th} -21 MeV	INDC(NDS)-0526 TENDL-2013(r)	2008	new	old
30	⁴⁵ Sc(n, γ) ⁴⁶ Sc	Sc45g	2125	E_{th} -21 MeV 21-60 MeV	IRDF-2002 TENDL-2010(r)	1991	old	old
31	⁴⁶ Ti(n,2n) ⁴⁵ Ti	Ti462	2225	E_{th} -20 MeV 20-60 MeV	INDC(CCP)-0360 TENDL-2010(r)	1993	old	old
32	⁴⁶ Ti(n,p) ⁴⁶ Sc	Ti46p	2225	E_{th} -20 MeV 20-60 MeV	IRDF-2002 TENDL-2010(r)	2005	old	old
33	⁴⁷ Ti(n,p) ⁴⁷ Sc	Ti47p	2228	E_{th} -20 MeV 20-60 MeV	RRDF-2002 TENDL-2011(r)	2002	new	old
34	⁴⁸ Ti(n,p) ⁴⁸ Sc	Ti48p	2231	E_{th} -20 MeV 20-60 MeV	IRDF-2002 TENDL-2010(r)	2005	old	old
35	^{nat} Ti(n,X) ⁴⁵ Ti	TiTi45	2200	E_{th} -60 MeV Minor channels	IRDF-II Ti462 TENDL-2015	2019	–	–
36	^{nat} Ti(n,X) ⁴⁶ Sc	TiSc46	2200	E_{th} -60 MeV Minor channels	IRDF-II Ti46p TENDL-2015	2019	–	–
37	^{nat} Ti(n,X) ⁴⁷ Sc	TiSc47	2200	E_{th} -60 MeV Minor channels	IRDF-II Ti47p TENDL-2015	2019	–	–
38	^{nat} Ti(n,X) ⁴⁸ Sc	TiSc48	2200	E_{th} -60 MeV Minor channels	IRDF-II Ti48p TENDL-2015	2019	–	–
39	⁵¹ V(n, α) ⁴⁸ Sc	V51a	2328	E_{th} -20 MeV 20-60 MeV	IRDF-2002 TENDL-2010(r)	2005	old	old
40	⁵¹ V(n,X) ⁴⁸ Sc	V51Sc48	2328	E_{th} -60 MeV Minor channels	IRDF-II V51a TENDL-2015	2019	–	–
41	^{nat} Cr(n,X) ⁵¹ Cr	CrCr51	2400	E_{th} -60 MeV Minor channels	IRDF-II TENDL-2015	2019	–	–
42	⁵⁵ Mn(n,2n) ⁵⁴ Mn	Mn552	2525	E_{th} -60 MeV	K.I.Zolotarev	2019	new	new

¹ Column (v2002 to v-1.05) marks any change in IRDF-II-v1.05 compared to IRDF-2002 as a reference up to 20 MeV.² Column (v-1.05 to IRDF-II) marks any change in IRDF-II compared to IRDF-II-v1.05 as a reference.

Consistency legend: new = new evaluation; old = unchanged; – = not present in reference.

TABLE 1: (continued). IRDF-II nuclear data contents and evaluation sources. (r) denotes renormalization.

ID No.	Reaction	Label	MAT	Energy Interval	Evaluation source	Eval. Date	Consistency	
							v2002 to v-1.05 ¹	v-1.05 to IRDF-II ²
43	⁵⁵ Mn(n,γ) ⁵⁶ Mn	Mn55g	2525	E_{th} -60 MeV	ENDF/B-VII.1 [33, 34]	2011	new	old
44	^{nat} Fe(n,X) ⁵¹ Cr	FeCr51	2600	Up to 60 MeV Minor channels	IRDF-II Fe54a TENDL-2015	2019	–	–
45	^{nat} Fe(n,X) ⁵⁴ Mn	FeMn54	2600	Up to 60 MeV Minor channels	IRDF-II Fe54p TENDL-2015	2019	–	–
46	^{nat} Fe(n,X) ⁵⁶ Mn	FeMn56	2600	Up to 60 MeV Minor channels	IRDF-II Fe56p TENDL-2015	2019	–	–
47	^{nat} Fe(n,X) ⁵³ Fe	FeFe53	2600	Up to 60 MeV Minor channels	IRDF-II Fe542 TENDL-2015	2019	–	–
48	⁵⁴ Fe(n,2n) ⁵³ Fe	Fe542	2625	E_{th} -20 MeV 20-60 MeV	INDC(CCP)-0360 TENDL-2010(r)	1993	old	old
49	⁵⁴ Fe(n,p) ⁵⁴ Mn	Fe54p	2625	E_{th} -20 MeV 20-60 MeV	INDC(NDS)-0657 TENDL-2013(r)	2013	new	old
50	⁵⁴ Fe(n,α) ⁵¹ Cr	Fe54a	2625	E_{th} -20 MeV 20-60 MeV	IRDF-2002 TENDL-2013(r)	2005	old	old
51	⁵⁶ Fe(n,p) ⁵⁶ Mn	Fe56p	2631	E_{th} -20 MeV 20-60 MeV	INDC(CCP)-0438 TENDL-2011(r)	2004	old	old
52	⁵⁸ Fe(n,γ) ⁵⁹ Fe	Fe58g	2637	E_{th} -20 MeV 20-60 MeV	JEFF-3.1 [35] TENDL-2011(r)	2011	new	old
53	⁵⁹ Co(n,2n) ⁵⁸ Co	Co592	2725	E_{th} -20 MeV 20-60 MeV	INDC(NDS)-0546 TENDL-2010(r)	2009	new	old
54	⁵⁹ Co(n,3n) ⁵⁷ Co	Co593	2725	E_{th} -20 MeV 20-60 MeV	INDC(NDS)-0584 TENDL-2010(r)	2010	–	new
55	⁵⁹ Co(n,γ) ⁶⁰ Co	Co59g	2725	E_{th} -20 MeV 20-60 MeV	IRDF-2002 TENDL-2010(r)	2005	old	old
56	⁵⁹ Co(n,p) ⁵⁹ Fe	Co59p	2725	E_{th} -20 MeV 20-60 MeV	INDC(NDS)-0546 TENDL-2010(r)	2009	new	old
57	⁵⁹ Co(n,α) ⁵⁶ Mn	Co59a	2725	E_{th} -20 MeV 20-60 MeV	IRDF-2002 TENDL-2010(r)	2005	old	old
58	⁵⁹ Co(n,X) ⁵⁶ Mn	Co59Mn56	2725	Up to 60 MeV Minor channels	IRDF-II Fe542 TENDL-2015	2019	–	–
59	^{nat} Ni(n,X) ⁵⁷ Ni	NiNi57	2800	Up to 60 MeV Minor channels	IRDF-II Ni582 TENDL-2015	2019	–	–
60	^{nat} Ni(n,X) ⁵⁸ Co	NiCo58	2800	Up to 60 MeV Minor channels	IRDF-II Ni58p TENDL-2015	2019	–	–
61	^{nat} Ni(n,X) ⁶⁰ Co	NiCo60	2800	Up to 60 MeV Minor channels	IRDF-II Ni60p TENDL-2015	2019	–	–
62	⁵⁸ Ni(n,2n) ⁵⁷ Ni	Ni582	2825	E_{th} -20 MeV 20-60 MeV	INDC(NDS)-0657 TENDL-2013(r)	2013	new	old
63	⁵⁸ Ni(n,p) ⁵⁸ Co	Ni58p	2825	0-20 MeV 20-60 MeV	K.I.Zolotarev (RRDF-2002) TENDL-2013(r)	2002	old	old
64	⁶⁰ Ni(n,p) ⁶⁰ Co	Ni60p	2831	E_{th} -21 MeV 21-60 MeV	INDC(NDS)-0526 TENDL-2011(r)	2007	new	old
65	^{nat} Cu(n,X) ⁶⁰ Co	CuCo60	2900	Up to 60 MeV Minor channels	IRDF-II Cu63a TENDL-2015	2019	–	–
66	^{nat} Cu(n,X) ⁶² Cu	CuCu62	2900	Up to 60 MeV Minor channels	IRDF-II Cu632 TENDL-2015	2019	–	–

¹ Column (v2002 to v-1.05) marks any change in IRDF-II v1.05 compared to IRDF-2002 as a reference up to 20 MeV.² Column (v-1.05 to IRDF-II) marks any change in IRDF-II compared to IRDF-II v1.05 as a reference.

Consistency legend: new = new evaluation; old = unchanged; – = not present in reference.

TABLE 1: (continued). IRDF-II nuclear data contents and evaluation sources. (r) denotes renormalization.

ID No.	Reaction	Label	MAT	Energy Interval	Evaluation source	Eval. Date	Consistency	
							v2002 to v-1.05 ¹	v-1.05 to IRDF-II ²
67	^{nat} Cu(n,X) ⁶⁴ Cu	CuCu64	2900	Up to 60 MeV Minor channels	IRDF-II Cu63g TENDL-2015	2019	–	–
68	⁶³ Cu(n,2n) ⁶² Cu	Cu632	2925	E_{th} -20 MeV 20–60 MeV	INDC(NDS)-0526 TENDL-2011(r)	2007	new	old
69	⁶³ Cu(n, γ) ⁶⁴ Cu	Cu63g	2925	E_{th} -20 MeV 20–60 MeV	IRDF-2002 TENDL-2011(r)	2005	old	old
70	⁶³ Cu(n, α) ⁶⁰ Co	Cu63a	2925	E_{th} -20 MeV 20–60 MeV	IRDF-2002 TENDL-2011(r)	2005	old	old
71	⁶⁵ Cu(n,2n) ⁶⁴ Cu	Cu652	2931	E_{th} -20 MeV 20–60 MeV	INDC(NDS)-0526 TENDL-2011(r)	2007	new	old
72	^{nat} Zn(n,X) ⁶⁴ Cu	ZnCu64	3000	Up to 60 MeV Minor channels	IRDF-II Zn64p TENDL-2015	2019	–	new
73	^{nat} Zn(n,X) ⁶⁷ Cu	ZnCu67	3000	Up to 60 MeV Minor channels	IRDF-II Zn67p TENDL-2015	2019	–	new
74	⁶⁴ Zn(n,p) ⁶⁴ Cu	Zn64p	3025	0–20 MeV 20–60 MeV	INDC(NDS)-0526 TENDL-2011(r)	2008	new	old
75	⁶⁷ Zn(n,p) ⁶⁷ Cu	Zn67p	3034	E_{th} -20 MeV 20–60 MeV	INDC(NDS)-0526 TENDL-2011(r)	2008	new	old
76	⁶⁸ Zn(n,X) ⁶⁷ Cu	Zn68Cu67	3034	E_{th} -60 MeV	INDC(NDS)-0796	2018	–	new
77	⁷⁵ As(n,2n) ⁷⁴ As	As752	3325	E_{th} -20 MeV 20–60 MeV	IRDF-2002 TENDL-2011(r)	2005	old	old
78	⁸⁹ Y(n,2n) ⁸⁸ Y	Y892	3925	E_{th} -40 MeV 40–60 MeV	INDC(NDS)-0584 TENDL-2010(r)	2010	new	old
79	^{nat} Zr(n,X) ⁸⁹ Zr	ZrZr89	4000	Up to 60 MeV Minor channels	IRDF-II Zr902 TENDL-2015	2019	–	new
80	⁹⁰ Zr(n,2n) ⁸⁹ Zr	Zr902	4025	E_{th} -20 MeV 20–60 MeV	INDC(NDS)-0546 TENDL-2010(r)	2009	new	old
81	⁹³ Nb(n,2n) ^{92m} Nb	Nb932m	4125	E_{th} -40 MeV 40–60 MeV	INDC(NDS)-0584 TENDL-2010(r)	2010	new	old
82	⁹³ Nb(n,n') ^{93m} Nb	Nb93nm	4125	E_{th} -20 MeV 20–60 MeV	IRDF-2002 TENDL-2010(r)	2005	old	old
83	⁹³ Nb(n, γ) ⁹⁴ Nb	Nb93g	4125	E_{th} -20 MeV 20–60 MeV	INDC(NDS)-0657 TENDL-2013(r)	2013	new	old
84	⁹³ Nb(n, γ) ^{94m} Nb	Nb93gm	4125	E_{th} -20 MeV 20–60 MeV	INDC(NDS)-0657 TENDL-2013(r)	2013	new	old
85	^{nat} Mo(n,X) ^{92m} Nb	MoNb92m	4200	Up to 60 MeV Minor channels	IRDF-II Zr902 TENDL-2015	2019	–	new
86	⁹² Mo(n,p) ^{92m} Nb	Mo92pm	4225	E_{th} -40 MeV 20–60 MeV	INDC(NDS)-0657 TENDL-2010(r)	2013	new	old
87	¹⁰³ Rh(n,n') ^{103m} Rh	Rh103nm	4525	E_{th} -20 MeV 20–60 MeV	IRDF-2002 TENDL-2010(r)	2005	old	old
88	¹⁰⁹ Ag(n, γ) ^{110m} Ag	Ag109gm	4731	0–60 MeV 20–60 MeV	JENDL-3.2 TENDL-2010(r)	1994	–	old
89	^{nat} In(n,X) ^{114m} In	InIn114m	4900	Up to 60 MeV Minor channels	IRDF-II In113gm TENDL-2015	2019	–	new
90	¹¹³ In(n, γ) ^{114m} In	In113gm	4925	0–20 MeV 20–60 MeV	INDC(NDS)-0668 TENDL-2015(r)	2015	–	old

¹ Column (v2002 to v-1.05) marks any change in IRDF-II-v1.05 compared to IRDF-2002 as a reference up to 20 MeV.² Column (v-1.05 to IRDF-II) marks any change in IRDF-II compared to IRDF-II-v1.05 as a reference.

Consistency legend: new = new evaluation; old = unchanged; – = not present in reference.

TABLE 1: (continued). IRDF-II nuclear data contents and evaluation sources. (r) denotes renormalization.

ID No.	Reaction	Label	MAT	Energy Interval	Evaluation source	Eval. Date	Consistency	
							v2002 to v-1.05 ¹	v-1.05 to IRDF-II ²
91	¹¹³ In(n,n') ^{113m} In	In113nm	4925	E_{th} -20 MeV 20–60 MeV	INDC(NDS)-0657 TENDL-2010(r)	2013	new –	old old
92	¹¹⁵ In(n,2n) ^{114m} In	In1152m	4931	E_{th} -20 MeV 20–60 MeV	INDC(NDS)-0526 TENDL-2010(r)	2008	new new	old old
93	¹¹⁵ In(n,n') ^{115m} In	In115nm	4931	E_{th} -20 MeV 20–60 MeV	IRDF-2002 TENDL-2010(r)	2005	old –	old old
94	¹¹⁵ In(n, γ) ^{116m} In	In115gm	4931	0–20 MeV 20–60 MeV	INDC(NDS)-0657 TENDL-2010(r)	2013	new –	old old
95	¹²⁷ I(n,2n) ¹²⁶ I	I1272	5325	E_{th} -32 MeV 32–60 MeV	INDC(NDS)-0526 TENDL-2010(r)	2008	new new	old old
96	¹³⁹ La(n, γ) ¹⁴⁰ La	La139g	5728	0–20 MeV 20–60 MeV	INDC(CCP)-0431 TENDL-2010(r)	2002	old old	old old
97	¹⁴¹ Pr(n,2n) ¹⁴⁰ Pr	Pr1412	5925	E_{th} -20 MeV 20–60 MeV	IRDF-2002 TENDL-2010(r)	2005	old –	old old
98	¹⁶⁹ Tm(n,2n) ¹⁶⁸ Tm	Tm1692	6925	E_{th} -40 MeV 40–60 MeV	INDC(NDS)-0584 TENDL-2013(r)	2010	new new	old old
99	¹⁶⁹ Tm(n,3n) ¹⁶⁷ Tm	Tm1693	6925	E_{th} -60 MeV	INDC(NDS)-0657	2013	new	old
100	¹⁸¹ Ta(n, γ) ¹⁸² Ta	Ta181g	7328	0–20 MeV 20–60 MeV	JENDL-3.2 TENDL-2010(r)	1994	old –	old old
101	¹⁸⁶ W(n, γ) ¹⁸⁷ W	W186g	7443	0–60 MeV	ENDF/B-VII.1 [33, 36]	2009	new	old
102	¹⁹⁷ Au(n,2n) ¹⁹⁶ Au	Au1972	7925	E_{th} -40 MeV 40–60 MeV	INDC(NDS)-0526 TENDL-2010(r)	2008	new new	old old
103	¹⁹⁷ Au(n, γ) ¹⁹⁸ Au	Au197g	7925	0–60 MeV	IAEA STD 2017 [32]	2018	new	new
104	¹⁹⁹ Hg(n,n') ^{199m} Hg	Hg199nm	8034	E_{th} -20 MeV 20–60 MeV	INDC(NDS)-0526 TENDL-2010(r)	2008	new new	old old
105	²⁰⁴ Pb(n,n') ^{204m} Pb	Pb204nm	8225	E_{th} -20 MeV 20–60 MeV	INDC(CCP)-0431 TENDL-2011(r)	2002	new new	old old
106	²⁰⁹ Bi(n,2n) ²⁰⁸ Bi	Bi2092	8325	E_{th} -400MeV	V.G.Pronyaev	2019	–	new
107	²⁰⁹ Bi(n,3n) ²⁰⁷ Bi	Bi2093	8325	E_{th} -400MeV	V.G.Pronyaev	2019	–	new
108	²⁰⁹ Bi(n,4n) ²⁰⁶ Bi	Bi2094	8325	E_{th} -400MeV	V.G.Pronyaev	2019	–	new
109	²⁰⁹ Bi(n,5n) ²⁰⁵ Bi	Bi2095	8325	E_{th} -400MeV	V.G.Pronyaev	2019	–	new
110	²⁰⁹ Bi(n,6n) ²⁰⁴ Bi	Bi2096	8325	E_{th} -400MeV	V.G.Pronyaev	2019	–	new
111	²³² Th(n,f)	Th232f	9040	0–20 MeV 20–60 MeV	JENDL-4.0 IAEA CRP [34, 37]	2010 2009	new new	new old
112	²³² Th(n, γ) ²³³ Th	Th232g	9040	0–60 MeV	IAEA CRP [34, 37, 38]	2009	new	old
113	²³⁵ U(n,f)	U235f	9228	0–60 MeV	ENDF/B-VIII.0,CIELO [39–41]	2018	new	new
114	²³⁸ U(n,2n) ²³⁷ U	U2382	9237	E_{th} -30 MeV 30–60 MeV	ENDF/B-VIII.0,CIELO [39–41] TENDL-2015(r)	2018 2019	new new	new old
115	²³⁸ U(n,f)	U238f	9237	0–60 MeV	ENDF/B-VIII.0,CIELO [39–41]	2018	new	new
116	²³⁸ U(n, γ) ²³⁹ U	U238g	9237	0–60 MeV	ENDF/B-VIII.0,CIELO [39–41]	2018	new	new
117	²³⁷ Np(n,f)	Np237f	9346	0–0.5 MeV 0.5–20 MeV 20–60 MeV	JEFF-3.3 IRDF-II-v1.05 JENDL-4/HE TENDL-2015(r) covariance	2017 2012 2011 2019	new old new new	new new new new
118	²³⁹ Pu(n,f)	Pu239f	9437	0–60 MeV	ENDF/B-VIII.0,CIELO [39, 40]	2018	new	new
119	²⁴¹ Am(n,f)	Am241f	9543	0–20 MeV 20–60 MeV	JENDL-3.2 JENDL-4/HE TENDL-2015(r) covariance	1989 2011 2019	old old new	old new new

¹ Column (v2002 to v-1.05) marks any change in IRDF-II-v1.05 compared to IRDF-2002 as a reference up to 20 MeV.² Column (v-1.05 to IRDF-II) marks any change in IRDF-II compared to IRDF-II-v1.05 as a reference.

Consistency legend: new = new evaluation; old = unchanged; – = not present in reference.

TABLE 1: (continued). IRDF-II nuclear data contents and evaluation sources. (r) denotes renormalization.

ID No.	Reaction	Label	MAT	Energy Interval	Evaluation source	Eval. Date	Consistency	
							v2002 to v-1.05 ¹	v-1.05 to IRDF-II ²
Cross sections of cover materials: Both σ_{tot} , σ_{el} and σ_{abs} are tabulated								
1	^{nat} B(n,tot)	Btot	500	0–20 MeV 20–60 MeV	ENDF/B-VIII.0 TENDL-2010(r)	2018	old new	new
2	¹⁰ B(n,tot)	B10tot	525	0–20 MeV 20–60 MeV	ENDF/B-VIII.0 TENDL-2010(r)	2018	old new	new
3	^{nat} Cd(n,tot)	Cdtot	4800	0–20 MeV 20–60 MeV	JEFF-3.3 TENDL-2010(r)	2018	old new	new
4	^{nat} Gd(n,tot)	Gdtot	6400	0–20 MeV 20–60 MeV	ENDF/B-VII.1 [33] TENDL-2010(r)	2011	old new	new
Damage cross sections								
1	^{nat} Si(n,1-MeV)	Si1MeV	1400	0–20 MeV	ASTM E722-14 [42]	2015	old	new
2	^{nat} Fe(n,X)dpa	FeNRTdpa-A	2600	0–20 MeV	ASTM E693-17 [43]	2017	old	new
3	^{nat} Fe(n,X)dpa	FeNRTdpa-E	2600	0–60 MeV	JEFF-3.3	2017	old	new
4	^{nat} Fe(n,X)dpa	FeNRTdpa-I	2600	0–20 MeV	IRDF-2002	2005	old	old
5	GaAs(n,1-MeV)	GaAs1MeV-A	—	0–20 MeV	ASTM E722-14 [42]	2015	old	new
Decay data fission products (FP) from neutron-induced fission on ²³² Th, ^{235,238} U, ²³⁷ Np, and ²³⁹ Pu targets								
1	(n,f)→ ⁹⁵ Zr	FPZr95	—		JEFF-3.3, ENDF/B-VIII.0	2018	old	new
2	(n,f)→ ⁹⁹ Mo	FPMo99	—		JEFF-3.3, ENDF/B-VIII.0	2018	old	new
3	(n,f)→ ¹⁰³ Ru	FPRu103	—		JEFF-3.3, ENDF/B-VIII.0	2018	old	new
4	(n,f)→ ¹⁰⁶ Ru→ ¹⁰⁶ Rh	FPRu106	—		JEFF-3.3, ENDF/B-VIII.0	2018	old	new
5	(n,f)→ ¹³⁷ Cs→ ^{137m} Ba	FPCs137	—		JEFF-3.3, ENDF/B-VIII.0	2018	old	new
6	(n,f)→ ¹⁴⁰ Ba→ ¹⁴⁰ La	FPBa140	—		JEFF-3.3, ENDF/B-VIII.0	2018	old	new
7	(n,f)→ ¹⁴⁴ Ce	FPCe144	—		JEFF-3.3, ENDF/B-VIII.0	2018	old	new

¹ Column (v2002 to v-1.05) marks any change in IRDF-*v*1.05 compared to IRDF-2002 as a reference up to 20 MeV.

² Column (v-1.05 to IRDF-*II*) marks any change in IRDF-*II* compared to IRDF-*v*1.05 as a reference.

Consistency legend: new = new evaluation; old = unchanged; – = not present in reference.

These composite reactions include: ^{nat}Mg(n,X)²⁷Na, ^{nat}Si(n,X)²⁸Al, ^{nat}S(n,X)³²P, ^{nat}Ti(n,X)⁴⁵Ti, ^{nat}Ti(n,X)⁴⁶Sc, ^{nat}Ti(n,X)⁴⁷Sc, ^{nat}Ti(n,X)⁴⁸Sc, ^{nat}Cr(n,X)⁵¹Cr, ^{nat}Fe(n,X)⁵³Fe, ^{nat}Fe(n,X)⁵¹Cr, ^{nat}Fe(n,X)⁵⁴Mn, ^{nat}Fe(n,X)⁵⁶Mn, ^{nat}Ni(n,X)⁵⁷Ni, ^{nat}Ni(n,X)⁵⁸Co, ^{nat}Ni(n,X)⁶⁰Co, ^{nat}Cu(n,X)⁶⁰Co, ^{nat}Cu(n,X)⁶²Cu, ^{nat}Cu(n,X)⁶⁴Cu, ^{nat}Zn(n,X)⁶⁴Cu, ^{nat}Zn(n,X)⁶⁷Cu, ^{nat}Zr(n,X)⁸⁹Zr, ^{nat}Mo(n,X)^{92m}Nb, and ^{nat}In(n,X)^{114m}In.

In the new IRDF-*II* library the energy range was extended to consistently address incident neutron energies up to 60 MeV, so most of the library cross sections are a composite of separate existing nuclear data evaluations. Columns 5, 6, and 7 of Table 1 provide information on the source of the recommended nuclear data evaluations in the applicable incident neutron energy ranges.

Since clear documentation of the consistency of nuclear data used in an evaluation is important in the use of these data in metrology applications, Column 6 of Table 1 points to supplemental documentation for many of the entries in the IRDF-*II* library.

B. Reactions on Cover Materials

Activation dosimeters can be used to characterize the neutron spectra through spectrum unfolding [44] or the least-squares-based spectrum adjustment approaches [45]. Frequently they are shielded with cover materials to change the effective threshold energy and to move a dosimeters sensitivity from the thermal into the epithermal neutron energy region. Some commonly used approaches to model the effective cover material flux attenuation is to apply exponential attenuation, based on the energy-dependence of the total cross section of the cover material and the thickness of cover used [44]. Other approaches [46] to the cover corrections use a more rigorous treatment based on radiation transport models of the neutron scattering within the cover. In order to support this variety of approaches to the treatment of the cover material, the IRDF-*II* library provides the total, elastic and absorption cross sections for commonly used cover materials. Total, elastic, capture (and fission) cross sections are also provided for all monitor materials that exhibit resonance structure so that self-shielding corrections can be made, if needed. Table 1 lists the four cover materials that are addressed within the library.

C. Dosimetry Metrics

Various radiation dosimetry metrics are employed in the interpretation of radiation test results that supports various application areas. A dosimetry metric is the result of folding a calculated dosimetry-related energy-dependent response function with the incident neutron energy-dependent fluence. These energy-dependent response functions are treated within the IRDF-II library in a manner similar to reaction-dependent cross sections. The dosimetry metrics used by the radiation effects community have been demonstrated to provide a correlation between the calculated quantity and a specific damage/failure mode of interest to an application area, *i.e.*, iron embrittlement, that can result in failure of a critical weld in a reactor pressure vessel or gain degradation in a silicon bipolar junction transistor (BJT). Monitor dosimetry measurements used during the radiation testing are frequently combined with knowledge of the neutron spectrum during the irradiation to directly report the radiation dosimetry metrics of interest in the application. Since the IRDF-II library is intended to support dosimetry applications for the fission and fusion communities, we have elected to include, for reference purposes, the primary radiation dosimetry metrics of interest to this community – but to restrict this inclusion to just those dosimetry metrics that have been endorsed by a national nuclear regulator and/or by an international standards organization. Table 1 lists the five dosimetry metrics, called Damage cross sections, that are included in the IRDF-II library.

The primary dosimetry metric of interest to the commercial light water reactor (LWR) community is the displacement-per-atom (dpa) in iron. This metric is used to assess the embrittlement of structural materials near the critical structural areas, such as the beltline weld for the pressure vessel wall, where any loss of mechanical strength could result in safety concerns. The neutron energy-dependence of the iron dpa metric is set by the appropriate regulatory body for the nuclear reactor, thus, there are different iron dpa dosimetry metrics used by the community. The IRDF-II contains the iron dpa damage metric from ASTM E693-17 Standard Practice for Characterizing Neutron Exposures in Iron and Low Alloy Steels in Terms of Displacement Per Atom (DPA) [43]. This damage metric was calculated by looking at the primary Frenkel pair production from each of the four naturally occurring iron isotopes and applying the Norgett-Robinson-Torrens (NRT) damage partition function [47]. The IRDF-II library also contains the JEFF-3.3 library [31] NRT-based iron damage metric and the legacy iron damage metric from the IRDF-2002 library.

In addition to the iron dpa metric, the radiation damage to electronics community has silicon and GaAs displacement damage standards defined in ASTM E722-14 Standard Practice for Characterizing Neutron Fluence Spectra in Terms of an Equivalent Monoenergetic Neutron Fluence for Radiation-Hardness Testing of Electronics [42]. The same E722 silicon response functions are also supported in ASTM E1855-15 Standard Test Method

for use of 2N2222A Silicon Bipolar Transistors and Neutron Spectrum Sensors and Displacement Damage Monitors [48].

D. Nuclear Decay Data

The recommended source of nuclear decay data for the dosimetry community is the series of monographs by the Bureau International des Poids et Mesures (BIPM). This eight volume BIPM-5 Table of Radionuclides represents an international collaboration that captures primary recommended nuclear data for most reactions of interest to the dosimetry community. Updates to the BIPM-5 are coordinated through the Decay Data Evaluation Project (DDEP) [49] working group led by the Laboratoire National de Metrologie et d'Essais Laboratoire National Henri Becquerel (LNE-LNHB). The recommended nuclear data are periodically updated in separate volumes of the BIPM-5 and the ongoing recommended values are kept up to date at the [DDEP website](#). Not all nuclides are addressed within the DDEP library. For nuclides that are not present, the recommended values are taken from the ENSDF “ADOPTED” dataset [50]². While the DDEP/BIPM-5 represents an international consensus on the current recommended decay data to be used for metrology applications, it is critical in dosimetry applications that all nuclear data used in a given application be consistent. This means that it is critical that the nuclear decay data be consistent with the associated cross section data. An evaluator of nuclear reaction cross section data would normally use values that correspond to the BIPM-5 data at the time of the evaluation but these values may not necessarily correspond to the BIPM-5 recommended values seen by a reader at a later time when the cross sections are being used in a metrology application. There are also cases where the evaluator, after consulting with members of the DDEP community, elected to weigh underlying measurements for the DDEP evaluation in a different way and ended up with slightly different decay data. Thus, in order to assist the users of the IRDF-II cross sections, the corresponding nuclear decay data that are consistent with the recommended cross sections are also documented within the IRDF-II library files. Details are given in Sec. IV C.

E. Atomic Mass Data

Metrology applications also require data on the elemental and isotopic mass used for the target materials. The most common metrological application for these data is the determination of the number of target atoms in an activation foil of a given mass.

² ENSDF nuclear structure and decay data can be easily extracted, understood and studied in an attractive user-friendly manner by means of LiveChart of Nuclides [51] and NuDat [52].

1. *Elemental*

The recommended elemental atomic weights are taken from the International Union of Pure and Applied Chemistry (IUPAC) Commission on Isotopic Abundances and Atomic Weights (CIAAW) [53]. These data represents revisions made by the CIAAW in 2017 and published in the Pure and Applied Chemistry.

2. *Isotopic*

The recommended atomic weights for individual isotopes are taken from the recommendations of the Atomic Mass Data Center and are represented by the Atomic Mass evaluation 2016 (AME2016) [54]. It is noted that geological materials are known for which the element has an isotopic composition that is outside the uncertainties cited here for normal materials. We should also note that many dosimetry applications may use enriched or depleted materials that are the result of isotopic separation techniques, in which case the user has to resort to the isotopic data that are provided.

F. Natural Abundance Data

The natural isotopic abundance data are also required in order to convert the densities of target materials into the number densities of the constituting isotopes. The recommended natural abundance data come from the CIAAW recommendations that refer to the 2016 IUPAC Technical Report [55]. These values represent the best measurement of isotopic abundances from a single terrestrial source.

Some of the dosimetry reactions addressed in the IRDF-II are for natural materials and include the production of a residual isotope from all contributing reactions. The abundance data used in the compilation of these multi-reaction combinations are given in Table 6 of Sec. IV A.

G. Fission Product Yields

Since many important dosimetry metrics are for fission reactions, it is also important the fission yields used to support the dosimetry measurements be clearly documented. Thus, the recommended nuclear data contained in the IRDF-II library include the cumulative fission yields for all the relevant fission products used for metrology applications which are listed in Tables 2, 3, and 4 in the three generic incident energy regions.

For the measurements of time-integrated fluence rates with fission products up to periods of about six weeks, the most common measured fission product used for fission research reactor dosimetry applications is ^{140}Ba (half-life of 12.7527 days). In order to avoid isotopic separation techniques, the ^{140}Ba activity is often determined by directly counting its daughter product, ^{140}La (half-life of 1.6781 days). The independent fission yields are required

TABLE 2. IRDF-II thermal-neutron fission yields ($E = 0.0253$ eV).

Target	Reaction Product	Cumulative Yield
^{235}U	^{95}Zr	$6.5042\text{E-}02 \pm 1.00 \%$
	^{99}Mo	$6.1399\text{E-}02 \pm 1.30 \%$
	^{103}Ru	$3.1118\text{E-}02 \pm 2.10 \%$
	^{106}Ru	$4.0958\text{E-}03 \pm 2.30 \%$
	^{137}Cs	$6.0897\text{E-}02 \pm 1.04 \%$
	^{140}Ba	$6.3444\text{E-}02 \pm 1.00 \%$
	^{144}Ce	$5.4781\text{E-}02 \pm 0.90 \%$
^{237}Np	^{95}Zr	$5.9364\text{E-}02 \pm 11.80 \%$
	^{99}Mo	$6.6813\text{E-}02 \pm 6.80 \%$
	^{103}Ru	$5.8640\text{E-}02 \pm 7.30 \%$
	^{106}Ru	$1.5421\text{E-}02 \pm 19.40 \%$
	^{137}Cs	$7.0334\text{E-}02 \pm 11.71 \%$
	^{140}Ba	$5.8591\text{E-}02 \pm 9.90 \%$
	^{144}Ce	$4.2716\text{E-}02 \pm 12.76 \%$
^{239}Pu	^{95}Zr	$4.8805\text{E-}02 \pm 1.10 \%$
	^{99}Mo	$6.1410\text{E-}02 \pm 0.70 \%$
	^{103}Ru	$6.8709\text{E-}02 \pm 1.20 \%$
	^{106}Ru	$4.3404\text{E-}02 \pm 2.30 \%$
	^{137}Cs	$6.5796\text{E-}02 \pm 1.22 \%$
	^{140}Ba	$5.2880\text{E-}02 \pm 1.10 \%$
	^{144}Ce	$3.7623\text{E-}02 \pm 0.90 \%$
^{241}Am	^{95}Zr	$3.9643\text{E-}02 \pm 3.00 \%$
	^{99}Mo	$6.5724\text{E-}02 \pm 3.20 \%$
	^{103}Ru	$7.1423\text{E-}02 \pm 6.10 \%$
	^{106}Ru	$5.0867\text{E-}02 \pm 16.60 \%$
	^{137}Cs	$7.2069\text{E-}02 \pm 6.60 \%$
	^{140}Ba	$5.8045\text{E-}02 \pm 2.50 \%$
	^{144}Ce	$3.3905\text{E-}02 \pm 3.70 \%$

to treat the transient equilibrium between the ^{140}La and ^{140}Ba .

Other fission products are commonly used to support reactor irradiations of different periods [56]. The isotope ^{95}Zr (half-life 64.032 days) is commonly used for shorter irradiation durations, up to 6 months while ^{103}Ru (half-life of 39.247 days) is used for irradiations up to 4 months. The isotope ^{144}Ce (half-life 284.91 days) is also a commonly measured fission product in reactor applications and can be used for irradiations periods of 2 to 3 years. The isotope ^{137}Cs (half-life of 30.08 years) is used to support reactor irradiations of 30 to 40 years. The ^{137}Cs activity is often measured by counting the activity in its ^{137m}Ba decay product [57] (half-life of 2.552 minutes). The ^{106}Ru fission product (half-life 371.8 days) is also often counted [57]. The transient equilibrium of its ^{106}Rh decay product (half-life of 30.07 seconds) with the ^{106}Rh independent yield can be ignored for most applications.

In summary, the relevant fission products currently addressed in dosimetry-related standards applications include the nine isotopes: ^{95}Zr , ^{99}Mo , ^{103}Ru , ^{106}Ru , ^{137}Cs , ^{137m}Ba , ^{140}Ba , ^{140}La , and ^{144}Ce , but cumulative fission yields are needed only for seven of them.

TABLE 3. IRDF-II fast-neutron fission yields ($E \approx 400 - 500$ keV).

Target	Reaction Product	Cumulative Yield
^{232}Th	^{95}Zr	$5.4494\text{E-}02 \pm 2.90 \%$
	^{99}Mo	$2.8740\text{E-}02 \pm 2.80 \%$
	^{103}Ru	$1.5179\text{E-}03 \pm 6.30 \%$
	^{106}Ru	$5.3236\text{E-}04 \pm 5.70 \%$
	^{137}Cs	$6.1790\text{E-}02 \pm 5.12 \%$
	^{140}Ba	$7.6222\text{E-}02 \pm 3.19 \%$
	^{144}Ce	$7.6334\text{E-}02 \pm 6.12 \%$
^{235}U	^{95}Zr	$6.4589\text{E-}02 \pm 1.30 \%$
	^{99}Mo	$5.8957\text{E-}02 \pm 1.90 \%$
	^{103}Ru	$3.2809\text{E-}02 \pm 1.40 \%$
	^{106}Ru	$4.6597\text{E-}03 \pm 7.60 \%$
	^{137}Cs	$5.8572\text{E-}02 \pm 1.92 \%$
	^{140}Ba	$6.0586\text{E-}02 \pm 1.10 \%$
	^{144}Ce	$5.1578\text{E-}02 \pm 1.80 \%$
^{238}U	^{95}Zr	$5.2506\text{E-}02 \pm 1.60 \%$
	^{99}Mo	$6.2147\text{E-}02 \pm 1.60 \%$
	^{103}Ru	$6.0331\text{E-}02 \pm 1.80 \%$
	^{106}Ru	$2.5063\text{E-}02 \pm 5.30 \%$
	^{137}Cs	$6.0045\text{E-}02 \pm 2.33 \%$
	^{140}Ba	$6.0457\text{E-}02 \pm 1.29 \%$
	^{144}Ce	$4.6916\text{E-}02 \pm 2.25 \%$
^{237}Np	^{95}Zr	$5.6715\text{E-}02 \pm 2.70 \%$
	^{99}Mo	$7.7238\text{E-}02 \pm 16.90 \%$
	^{103}Ru	$5.3778\text{E-}02 \pm 11.20 \%$
	^{106}Ru	$2.2333\text{E-}02 \pm 10.80 \%$
	^{137}Cs	$6.2129\text{E-}02 \pm 3.62 \%$
	^{140}Ba	$5.7593\text{E-}02 \pm 2.00 \%$
	^{144}Ce	$4.1743\text{E-}02 \pm 4.79 \%$
^{239}Pu	^{95}Zr	$4.6909\text{E-}02 \pm 2.50 \%$
	^{99}Mo	$5.8366\text{E-}02 \pm 2.30 \%$
	^{103}Ru	$6.5709\text{E-}02 \pm 2.50 \%$
	^{106}Ru	$4.0375\text{E-}02 \pm 7.80 \%$
	^{137}Cs	$6.3098\text{E-}02 \pm 2.21 \%$
	^{140}Ba	$5.2916\text{E-}02 \pm 1.50 \%$
	^{144}Ce	$3.4994\text{E-}02 \pm 1.70 \%$
^{241}Am	^{95}Zr	$4.1632\text{E-}02 \pm 8.20 \%$
	^{99}Mo	$5.3596\text{E-}02 \pm 5.50 \%$
	^{103}Ru	$6.5234\text{E-}02 \pm 6.70 \%$
	^{106}Ru	$4.8254\text{E-}02 \pm 6.90 \%$
	^{137}Cs	$6.2281\text{E-}02 \pm 18.80 \%$
	^{140}Ba	$4.8853\text{E-}02 \pm 4.50 \%$
	^{144}Ce	$3.4353\text{E-}02 \pm 6.80 \%$

TABLE 4. IRDF-II fusion-peak fission yields ($E \approx 14$ MeV).

Reaction Label	Reaction Product	Cumulative Yield
^{232}Th	^{95}Zr	$5.0470\text{E-}02 \pm 13.60 \%$
	^{99}Mo	$1.9747\text{E-}02 \pm 5.30 \%$
	^{103}Ru	$8.9252\text{E-}03 \pm 7.20 \%$
	^{106}Ru	$1.0873\text{E-}02 \pm 10.40 \%$
	^{137}Cs	$6.0482\text{E-}02 \pm 8.81 \%$
	^{140}Ba	$5.7087\text{E-}02 \pm 4.00 \%$
	^{144}Ce	$4.2355\text{E-}02 \pm 8.66 \%$
^{235}U	^{95}Zr	$5.0620\text{E-}02 \pm 4.30 \%$
	^{99}Mo	$5.0301\text{E-}02 \pm 2.80 \%$
	^{103}Ru	$3.1210\text{E-}02 \pm 4.40 \%$
	^{106}Ru	$1.8000\text{E-}02 \pm 16.80 \%$
	^{137}Cs	$5.7013\text{E-}02 \pm 24.00 \%$
	^{140}Ba	$4.4929\text{E-}02 \pm 1.90 \%$
	^{144}Ce	$3.1575\text{E-}02 \pm 1.30 \%$
^{238}U	^{95}Zr	$4.6389\text{E-}02 \pm 1.22 \%$
	^{99}Mo	$5.7713\text{E-}02 \pm 0.80 \%$
	^{103}Ru	$4.4881\text{E-}02 \pm 2.30 \%$
	^{106}Ru	$2.4945\text{E-}02 \pm 5.30 \%$
	^{137}Cs	$5.4359\text{E-}02 \pm 8.21 \%$
	^{140}Ba	$4.6437\text{E-}02 \pm 0.70 \%$
	^{144}Ce	$3.6147\text{E-}02 \pm 3.57 \%$
^{237}Np	^{95}Zr	$5.5453\text{E-}02 \pm 16.00 \%$
	^{99}Mo	$4.8632\text{E-}02 \pm 11.00 \%$
	^{103}Ru	$4.3312\text{E-}02 \pm 16.00 \%$
	^{106}Ru	$2.9975\text{E-}02 \pm 23.00 \%$
	^{137}Cs	$5.1076\text{E-}02 \pm 23.00 \%$
	^{140}Ba	$4.8131\text{E-}02 \pm 11.00 \%$
	^{144}Ce	$2.9997\text{E-}02 \pm 16.00 \%$
^{239}Pu	^{95}Zr	$4.3837\text{E-}02 \pm 6.00 \%$
	^{99}Mo	$5.6556\text{E-}02 \pm 5.00 \%$
	^{103}Ru	$5.8641\text{E-}02 \pm 6.00 \%$
	^{106}Ru	$3.5345\text{E-}02 \pm 8.00 \%$
	^{137}Cs	$5.0107\text{E-}02 \pm 8.00 \%$
	^{140}Ba	$4.0556\text{E-}02 \pm 5.00 \%$
	^{144}Ce	$2.9598\text{E-}02 \pm 5.00 \%$
^{241}Am	^{95}Zr	$3.1194\text{E-}02 \pm 6.00 \%$
	^{99}Mo	$4.3358\text{E-}02 \pm 6.00 \%$
	^{103}Ru	$5.0404\text{E-}02 \pm 6.00 \%$
	^{106}Ru	$4.0805\text{E-}02 \pm 6.00 \%$
	^{137}Cs	$4.2719\text{E-}02 \pm 6.00 \%$
	^{140}Ba	$3.3620\text{E-}02 \pm 6.00 \%$
	^{144}Ce	$2.4968\text{E-}02 \pm 6.00 \%$

The cumulative yields are used because they are generally known with a higher precision than the independent yields. However, care must be taken to ensure that the cumulative yields reach equilibrium. The typical cooling times that are necessary are listed below:

^{95}Zr	The longest lived member of the chain is ^{95}Y at 10.3 minutes. All the others are less than 1 minute. If we wait for 6 half-lives that would be 1 hour.
^{99}Mo	Longest half-life has $^{99\text{m}}\text{Nb}$ at 2.6 m, requiring cooling time of about 15 minutes.
$^{103}\text{Ru} - ^{103}\text{Tc}$	Half-life 54 s, ^{103}Mo half life 1.13 m, requiring cooling time of about 13 minutes.
$^{106}\text{Ru} - ^{106}\text{Tc}$	Half-life 36 s, requiring cooling time of about 3.6 minutes.
$^{137}\text{Cs} - ^{137}\text{Xe}$	Half-life 3.82 m, requiring cooling time of about 23 minutes.
$^{140}\text{Ba} - ^{140}\text{Cs}$	Half-life 1.06 m, requiring cooling time of about 6 minutes.
$^{144}\text{Ce} - ^{144}\text{La}$	Half-life 40.7 s, requiring cooling time of about 4.1 minutes.

It should also be noted that several radionuclides of these chains have Kr or Xe precursors. This means that part or all of the chain yield can be lost as Kr or Xe gas unless the fissionable material is irradiated in a gas tight container. This is a very well-known effect and actinides are commonly sealed in quartz to avoid losing any Kr or Xe, which would lead to a low reading of the cumulative fission product.

Some activities, such as nonproliferation monitoring [58, 59], have a wider range of application-specific fission yield products that are monitored. This application area has independently standardized the underlying fission products yields to be used, hence this application area is not addressed by these IRDF-II fission-yield recommended values.

Work continues in refining the recommended yields and the treatment of their dependence on the energy of the incident neutron. There is an ongoing international collaboration, sponsored through an IAEA coordinated research project (CRP) (project number F42007 approved 19 December 2018), to provide recommended fission yields and to address recommended approaches for handling the energy-dependence of the fission yields. The results of this CRP will not be ready in time for inclusion in the present release of IRDF-II.

For present dosimetry applications, pending more investigation and validation of the energy-dependent modeling, the recommendation is to map the application need into one of the three generic energy regions supported by the commonly available nuclear data libraries: thermal, fast, and 14 MeV. The thermal energy generally equates to evaluated data for 0.0253 eV incident neutrons. The fast energy equates to evaluated data gathered using a range of broad-spectrum fast fission spectra. Within the commonly available tabulated nuclear data files, the fast region is characterized with a reference energy of 500 keV in the ENDF/B-VIII.0 library and with an energy of 400 keV in the JEFF-3.3 library. The 14 MeV energy equates to

evaluated data validated using data gathered from DT fusion neutron sources.

Pending results from ongoing international collaboration on updating the fission product yields, the IRDF-II library has adopted a set of recommended fission yields. These recommended fission yields are taken from the JEFF-3.3 library, if available. Alternatively, *e.g.*, in the case of the 14 MeV fission yields for ^{237}Np , ^{239}Pu and ^{241}Am , the recommended yields are taken from the ENDF/B-VIII.0 library.

H. Neutron Benchmark Fields

As part of the library evaluation and acceptance process for reactions accepted for inclusion within the IRDF-II library, data validation was addressed using experimental measurements in integral neutron benchmark fields. Such measurements are independent of the differential data (measured as a function of incident energy), used by the evaluators in conjunction with nuclear data models to generate the recommended nuclear reaction cross section data files. Since these neutron benchmark fields are tied to the validation evidence for the IRDF-II library, the decision was made to include the neutron spectral characterizations for these integral benchmark fields as part of the IRDF-II package. Table 5 summarizes the 45 neutron fields that were considered in the IRDF-II validation and provides brief descriptions of the selected 29 neutron benchmark fields. These benchmark fields are addressed in further detail in Secs. VII and VIII on sensitivity, data validation and data consistency in this paper. The IRDF-II library provides recommended spectra for these neutron fields as well as for several other spectra not selected as benchmarks, but useful in making data comparisons, *e.g.*, a constant spectrum, a linear spectrum, and a $1/E$ spectrum in the region $0.5 \text{ eV} < E < 20 \text{ MeV}$. Maxwellian spectra corresponding to different temperatures are also included that cover a range of temperatures (in the laboratory coordinate system) from 25 keV to 60 keV, which are of interest for astrophysics. The spectra are part of the IRDF-II distribution. Note that the spectra calculated with detailed computational models of the experiments by the Monte Carlo technique (*e.g.*, reactor spectra) are used as *prior* spectra in the least-squares fit. The uncertainty in the *prior* spectrum, in addition to being a subjective evaluation at this time, includes large uncertainty contributors which make direct comparison with measured data very difficult due to the large associated uncertainties. The total uncertainty of the calculated *prior* obviously includes the statistical uncertainty of the calculations, which usually is less than 10 %, except in the tails, where the spectrum approaches zero. The full spectrum covariance matrices from computational models being subjective and not available for all cases, the covariances in IRDF-II only include the statistical component for cross-checking the uncertainties in the calculated reaction rates of monitor reactions with median energies close to the tails of the spectrum.

TABLE 5. List of IRDF-II benchmark neutron fields

No.	Name	MAT	$E_{aver}[MeV]$	Description
Measured by Time-of-Flight neutron fields including the $^{252}Cf(sf)$ standard				
1	$^{252}Cf(sf)$	9861	2.121	Spontaneous fission neutron spectrum from ^{252}Cf
2	^{235}U PFNS	9228	2.000	Thermal-neutron induced prompt fission spectrum from ^{235}U
3	$^9Be(d,n)$ 16MeV	9408	5.608	Spectrum of neutrons from 16 MeV deuterons incident on a beryllium target
4	$^9Be(d,n)$ 40MeV	9409	15.58	Spectrum of neutrons from 40 MeV deuterons incident on a beryllium target
Measured by Time-of-Flight neutron fields not accepted as benchmark fields				
1	^{233}U PFNS	9222	2.030	Thermal-neutron induced prompt fission spectrum from ^{233}U
2	^{239}Pu PFNS	9437	2.073	Thermal-neutron induced prompt fission spectrum from ^{239}Pu
Neutron benchmark fields from detailed computational models				
1	ACRR-FF-32	9010	0.575	ACRR-FF-32 Reactor Extended Cavity Spectrum 640-group
2	ACRR-CdPoly	9011	0.657	ACRR-CdPoly Reactor Bucket Spectrum 640-group
3	ACRR-PLG	9012	0.439	ACRR-PLG Reactor Bucket Spectrum 640-group
4	ACRR-LB44	9013	0.715	ACRR-LB44 Reactor Bucket Spectrum 640-group
5	FREC-II	9015	0.545	FREC-II Spectrum (external cavity attached to ACRR) 640-group
6	SPR-III	9014	1.251	SPR-III Reactor Central Cavity Spectrum 640-group
7	Mol BR1 Mark-III	9020	1.864	Mol BR1 Mark-III, ^{235}U converter in Cd and Graphite cavity, 640-groups
8	LR0-Rez	9032	0.646	Rez-LR0 Reactor spectrum, 640-group
9	TRIGA-JSI	9041	0.389	TRIGA Mark-II Pneumatic tube (bare), 640-group
10	TRIGA-JSI/BN	9042	0.848	TRIGA Mark-II boron nitride cover, 640-group
11	TRIGA-JSI/B4C	9043	0.923	TRIGA Mark-II boron carbide cover, 640-group
12	TRIGA-JSI/10B4C	9044	1.090	TRIGA Mark-II enriched boron carbide cover, 640-group
13	ISNF	9004	1.058	ISNF Reactor Spectrum 725-group
14	CFRMF	9005	0.741	CFRMF Reactor Spectrum from IRDF-2002
15	Sigma-Sigma	9007	0.763	Sigma-Sigma facility in ^{nat}U and BC spheres inside Graphite column, 725g
16	HMF001	9101	1.433	Godiva, central region, 725-group
17	HMF028	9102	1.343	Flattop-25, central region, 725-group
18	IMF007	9103	0.570	Big-Ten 725-group
19	FMR001	9110	1.483	IPPE-BR1, central region, 725-group
20	FNS-Grph-096mm	9201	5.267	FNS-Graphite block with a D-T source and monitors at 96 mm, 725 group
21	FNS-Grph-293mm	9202	1.957	FNS-Graphite block with a D-T source and monitors at 293 mm, 725 group
ICSBEP spectra not accepted as benchmark fields				
1	PMF001	9104	1.797	Jezebel, central region, 725-group
2	PMF002	9105	1.747	Jezebel-240, central region, 725-group
3	PMF006	9106	1.589	Flattop-Pu, central region, 725-group
4	PMF008	9107	1.681	Thor, central region, 725-group
Analytical spectrum functions accepted as benchmark fields				
1	Thermal Maxw.	9901		Thermal Maxwellian at 293.6 K
2	1/E [0.55eV–2MeV]	9902		Pure 1/E between E_{cd} and E_2 ($0.55 \text{ eV} < E < 2 \text{ MeV}$)
3	Maxw.(25 keV)	9925		Maxwellian at 25 keV
4	Maxw.(30 keV)	9930		Maxwellian at 30 keV
Analytical spectrum functions not used as benchmark fields				
1	Const.	9900		Constant spectrum $\Phi=1$
2	1/E [0.5eV–20MeV]	9904		Pure 1/E between E_{cd} and E_2 ($0.5 \text{ eV} < E < 20 \text{ MeV}$)
3	Maxw.Fiss.	9905		Pure Maxwellian fission spectrum at temperature 2.03 MeV
4	Linear	9910		Linear spectrum $\Phi=E$ ($1.E-5 \text{ eV} < E < 20 \text{ MeV}$)
5	Maxw.(32 keV)	9932		Maxwellian at 32 keV
6	Maxw.(35 keV)	9935		Maxwellian at 35 keV
7	Maxw.(40 keV)	9940		Maxwellian at 40 keV
8	Maxw.(45 keV)	9945		Maxwellian at 45 keV
9	Maxw.(50 keV)	9950		Maxwellian at 50 keV
10	Maxw.(60 keV)	9960		Maxwellian at 60 keV

III. IRDF-II FORMAT AND DATA VERIFICATION

In order to support the user community, the cross sections in the IRDF-II library are distributed in several different formats, described in the sections below.

The baseline library is in the ENDF-6 format, where the full library data file contains several logical files for different nuclides, identified by the so-called MAT numbers. Each nuclide file is further subdivided into subfiles identified by the MF numbers that designate the type of data and the MT numbers that designate the reaction

type. A brief extract is given below for convenience. Full details are given in the ENDF-6 manual [60].

MAT Material/nuclide identifier is a four-digit number that uniquely identifies the target nuclide.

MF Data type identifier:

- 1 General information. The first section in this subfile is identified by MT=451 and contains evaluation description in plain text, including eigenvalues of the covariance matrices.
- 3 Cross sections and other parameters as a function of incident particle energy.
- 10 Cross sections for the excitation of discrete states, or radionuclide production cross sections.
- 33 Covariance data of the quantities in MF=3.
- 40 Covariance data for the quantities in MF=10.

MT Reaction type number (*e.g.*, $16 \equiv (n,2n)$, $102 \equiv (n,\gamma)$, $103 \equiv (n,p)$, *etc.*). For full details see Appendix B of the ENDF-6 Manual [60].

The derived libraries in other formats are generated by processing baseline libraries in point-wise cross-section representation with codes such as NJOY-2016 [61] or PREPRO [62]. At the request of the users, the recommended cross sections are also available in three column format (energy, cross section and the total uncertainty) to be used as reference activation cross sections for relative activation measurements³. If a user requires the cross sections at a different temperature, the threshold reactions would be practically unaffected. Reaction cross sections with resonance structure might need additional processing with codes like NJOY or PREPRO.

Note that residual-production covariance matrices in ENDF subfile MF=40 that contain more than one product cannot be processed with the current version of NJOY [61]. This is applicable to many of the natural element evaluations in IRDFF-II. A way around this problem is to split the evaluated file in such a way that only one product is present at a time, and process the partial files separately. Hopefully, an update to the processing code will be available soon that will support legitimate ENDF-6 format covariance representation.

A. ENDF-6 Format Point Cross Sections

The baseline library contains point cross sections at 293.6 K in ENDF-6 format representation that is well-documented and has the flexibility to capture the highest

fidelity characterization of neutron cross sections, including the resonance region, as well as the associated covariance files. This format can also be used to capture the nuclear decay data and the neutron spectral characterization data. The ENDF-6 file structure places reaction cross sections in MF=3 and production cross sections in MF=10. Both of these file types are used to capture the cross sections within the IRDFF-II library. The covariance data that correspond to data found in these sections are stored in MF=33 and MF=40, respectively. Radioactive decay data are given in MF=8. The file structure permits the inclusion of in-line documentation using MF=1, MT=451 entries.

B. ENDF-6 Format Neutron Spectra

Whereas the ENDF-6 file structure permits the energy distributions of secondary particles to be presented in MF=5 and MF=6, this field does not fit the IRDFF-II characterization for the benchmark neutron spectra. These data appear in a separate IRDFF-II library file in pseudo-ENDF format as MF=3, MT=261 sub-files. The spectra are differentiated by the MAT number. The covariance information is stored in the corresponding ENDF subfiles MF=33, MT=261. This is a non-standard use of the ENDF format, which uses MT=261 that is formally undefined in the ENDF-6 manual []. Storing the spectra in this format is convenient because for many operations on the data and for generating the comparison plots the same software can be used as for the cross sections.

C. ENDF-6 Format 640 and 725 Group Representation

Whereas a point cross section format is best for capturing the physics-based structure behind the cross section representation, this format does not readily support post-processing of the cross sections and the extraction of spectrum-averaged cross sections, reaction rates from integral neutron fields, or many other relevant material dosimetry metrics. For these applications, a multigroup representation is generally used. An important consideration with multigroup representations is the energy group structure. Whereas few-group representation may be sufficient for some applications, other applications require a fine-group representation.

A common fine-group structure that was found to be sufficient for capturing the energy-dependent shape of most responses relevant to the fission reactor community is the extended SAND-II group structure [44]. The extension here represented an increase from the original 18 MeV SAND-II maximum energy up to 20 MeV. Thus, this 640-group energy group structure starts at 10^{-4} eV and goes up to 20 MeV. Since the IRDFF-II library goes up to 60 MeV, in this representation the IRDFF-II data are essentially truncated at 20 MeV and collapsed into the 640-group structure.

The IRDFF-II application addresses the needs of the

³ Note that a full uncertainty propagation of these cross sections for a broad spectrum requires the use of the complete covariance matrix, which is only given in ENDF-6 format.

fusion and low temperature material science communities as well as the fission reactor community, hence a second 725-group representation that extends up to 60 MeV is also used to characterize the data. The energy group structure for this 725-group representation is formed by:

- starting at 10^{-5} eV and adding 45 groups to cover the energy range from 10^{-5} eV up to the 10^{-4} eV, where the 640-group SAND-II structure starts;
- using the same 640-groups between 10^{-4} eV and 20 MeV;
- adding 20 groups of 0.5 MeV width in energy to extend the range up to 30 MeV;
- adding 10 groups of 1 MeV width in energy to further extend the range up to 40 MeV;
- adding 10 groups of 2 MeV width in energy to further extend the range up to 60-MeV.

In this collapsing of the point cross sections to the 640 or 725 group structure, an energy-dependent weighting function can be used. The two most common weighting functions are:

1. A thermal Maxwellian shape in the thermal region, a $1/E$ shape in the reactor neutron slowing down region, a fission spectrum shape in the fast fission reactor region, and a fusion peak at high energies of the fusion blanket region. The GROUPE module of the NJOY-2012 code with the IWT=8 weighting function using the default weighting function shape parameters (thermal breakpoint of 0.1 eV, thermal temperature of 0.025 eV, fission breakpoint of 0.1 MeV, fission temperature of 1.4 MeV) can be used to collapse the data that appears in the file. The disadvantage of this weighting function is that it might not be representative of reactor spectra at epithermal energies and could even have a wrong gradient.
2. A flat weighting across the energy bin. Given the fine group nature of the two energy bin structures, this flat weighting is more general at high energies and mostly sufficient at lower energies as well.

In the preparation of the IRDFE-II multigroup data the second option, flat weighting, was used. This collapsing was performed with the GROUPE module of the PREPRO-2018 data processing package [62]. The same procedures are applicable also to the IRDFE-II file of neutron spectra in ENDF-6 format.

D. MCNP ACE Format

The ACE (A Compact ENDF) format is a common data format that is used in continuous-energy neutron-photon Monte Carlo codes like MCNP [63] and SERPENT [64]. The virtue of this format is that it can capture the continuous energy nature of the underlying cross section without resorting to a multi-group representation. Since many fission and fusion community applications

use the MCNP code to model the neutron spectrum at locations of interest for dosimetry measurements, the inclusion of this format increases the fidelity supported by the application and makes it easy for the user to interface with the recommended IRDFE-II library.

E. Data Verification

A critical step in the verification of the formal correctness of the IRDFE-II library files was to ensure that the ENDF-6 format rules were obeyed. For this purpose, the ENDF Utility Codes were employed [65], namely:

CHECKR checks the syntax;

FIZCON checks the internal consistency of the data;

PSYCHE does an in-depth checking, particularly regarding the covariance data.

In addition, the COVEIG code [66] was used to check for symmetry, large correlation coefficients and negative eigenvalues in the covariance matrices.

The ability to process the files in ENDF-6 format was checked by running the PREPRO codes [62]. The following modules are applicable:

LINEAR makes all cross sections linearly interpolable;

GROUPE generates group-averaged cross sections;

COMPLIT makes graphical comparison with other evaluated data files.

Finally, the NJOY code [61] was employed to prepare plots of the covariance matrices and to generate the IRDFE-II library in ACE format. Quality assurance procedure INDC(SEC)-0107 [67] was employed to minimize the possibility of data processing errors when generating the files in ACE format.

IV. RECOMMENDED NUCLEAR DATA FOR NEUTRON METROLOGY

This Sec. provides a more in-depth examination of the various nuclear data components within the IRDFE-II library and provides details on the process used to arrive at the recommendations.

A. Atomic Mass and Isotopic Natural Abundance Data

The atomic weights for the target elements are given in Table 6 in atomic mass units. The reported uncertainties correspond to those for isotopes in the normal terrestrial materials. Column 2 of Table 6 gives the recommended natural abundance value for all targets in the IRDFE-II library. Column 3 of Table 6 shows the isotopic weights represented as the mass excess in energy units and Column 4 gives the corresponding values in micro-atomic mass units, as quoted in the original publication [54].

TABLE 6. IRDF-II isotopic abundances, nuclear and elemental mass data. Numbers in parentheses indicate the absolute uncertainties, *e.g.*, 92.411(24) \equiv 92.411 \pm 0.024.

Target nucleus	Abundance [atom %]	Isotopic Atomic Mass Excess [keV]	Isotopic Atomic Mass [μ amu]	Elemental weight value or range [amu]
⁶ Li	7.589 (24)	+14086.8789 (14)	6015122.8874 (15)	[6.938, 6.997]
⁷ Li	92.411 (24)	+14907.105 (4)	7016003.437 (5)	[6.938, 6.997]
¹⁰ B	19.82 (2)	+12050.609 (15)	10012936.862 (16)	[10.806, 10.821]
¹¹ B	80.18 (2)	+8667.707 (12)	11009305.167 (13)	[10.806, 10.821]
¹⁹ F	100. (0)	-1487.4442 (09)	18998403.1629 (9)	18.998403 163(6)
²³ Na	100. (0)	-9529.8525 (18)	22989869.2820 (19)	22.98976928 (2)
²⁴ Mg	78.951 (12)	-13933.569 (13)	23985041.697 (14)	[24.304, 24.307]
²⁷ Al	100. (0)	-17196.86 (5)	26981538.41 (5)	26.9815385 (7)
²⁸ Si	92.22968 (44)	-21492.7943 (5)	27.9769265350 (5)	[28.084, 28.086]
²⁹ Si	4.68316 (32)	-21895.0784 (6)	28.9764946653 (6)	[28.084, 28.086]
³¹ P	100. (0)	-24440.5410 (7)	30973761.9986 (7)	30.973761 998 (5)
³² S	95.04074 (88)	-26015.5336 (13)	31972071.1744 (14)	[32.059, 32.076]
⁴⁵ Sc	100. (0)	-41071.9 (7)	44955907.5 (7)	44.955908 (5)
⁴⁶ Ti	8.249 (21)	-44127.80 (16)	45952626.86 (18)	47.867 (1)
⁴⁷ Ti	7.437 (14)	-44937.36 (12)	46951757.75 (12)	47.867 (1)
⁴⁸ Ti	73.720 (22)	-48492.71 (11)	47947940.93 (12)	47.867 (1)
⁵¹ V	99.7503 (6)	-52203.8 (4)	50 943956.9 (4)	50.9415 (1)
⁵⁰ Cr	0.2497 (6)	-50262.1 (4)	49 946041.4 (5)	51.9961 (6)
⁵² Cr	83.7895 (117)	-55419.2 (3)	51 940505.0 (4)	51.9961 (6)
⁵⁵ Mn	100. (0)	-57712.4 (3)	54938043.2 (3)	54.938 044 (3)
⁵⁴ Fe	5.8450 (230)	-56254.5 (4)	53939608.3 (4)	55.845 (2)
⁵⁴ Fe	5.8450 (230)	-56254.5 (4)	53939608.3 (4)	55.845 (2)
⁵⁶ Fe	91.7540 (240)	-60607.1 (3)	55934935.6 (3)	55.845 (2)
⁵⁸ Fe	0.2819 (27)	-62155.1 (3)	57933273.7 (4)	55.845 (2)
⁵⁹ Co	100. (0)	-62229.7 (4)	58933193.7 (4)	58.933194 (4)
⁵⁸ Ni	68.0769 (59)	-60228.7 (4)	57935341.8 (4)	58.6934 (4)
⁶³ Cu	69.174 (20)	-65579.8 (4)	62929597.2 (5)	63.546 (3)
⁶⁵ Cu	30.826 (20)	-67263.7 (6)	64927789.5 (7)	63.546 (3)
⁶⁴ Zn	49.1704 (83)	-66004.0 (6)	63929141.8 (7)	65.38 (2)
⁶⁷ Zn	4.0401 (18)	-67880.3 (8)	66 927127.5 (8)	65.38 (2)
⁷⁵ As	100. (0)	-73034.2 (9)	74 921594.6 (9)	74.921595 (6)
⁸⁹ Y	100. (0)	-87708.4 (16)	88 905841.2 (17)	88.90584 (2)
⁹⁰ Zr	51.452 (9)	-88772.54 (12)	89904698.76 (13)	91.224 (2)
⁹³ Nb	100. (0)	-87212.8 (15)	92906373.2 (16)	92.90637 (2)
⁹² Mo	14.649 (17)	-86808.58 (16)	91906807.16 (17)	95.95 (1)
¹⁰³ Rh	100. (0)	-88031.7 (23)	102905494.1 (25)	102.90550 (2)
¹⁰⁹ Ag	48.1608 (51)	-88719.4 (13)	108904755.8 (14)	107.8682 (2)
¹¹³ In	4.281 (17)	-89367.12 (19)	112904060.45 (20)	114.818(1)
¹¹⁵ In	95.719 (17)	-89536.346 (12)	114903878.774 (13)	114.818(1)
¹²⁷ I	100.	-88984 (4)	126904472 (4)	126.90447 (3)
¹³⁹ La	99.91119 (24)	-87226.2 (20)	138906358.8 (22)	138.90547 (7)
¹⁴¹ Pr	100.0 (0)	-86015.6 (17)	140907658.4 (18)	140.90766 (2)
¹⁶⁹ Tm	100.0 (0)	-61275.2 (08)	168934218.4 (09)	168.93422 (2)
¹⁸¹ Ta	99.98799 (8)	-48438.3 (14)	180947999.3 (15)	180.94788 (2)
¹⁸⁶ W	28.4259 (62)	-42508.5 (12)	185954365.2 (13)	183.84 (1)
¹⁹⁷ Au	100.0 (0)	-31139.7 (05)	196966570.1 (06)	196.966569 (5)
¹⁹⁹ Hg	16.938 (9)	-29546.1 (05)	198968281.0 (06)	200.592 (3)
²⁰⁴ Pb	1.4245 (12)	-25109.9 (11)	203973043.4 (12)	207.2 (1)
²⁰⁹ Bi	100.0 (0)	-18258.7 (14)	208980398.5 (15)	208.98040 (1)
²³² Th	99.998862 (2)	+35446.8 (14)	232038053.7 (15)	232.0377 (4)
²³⁵ U	0.72041 (36)	+40918.8 (11)	235043928.2 (12)	238.02891 (3)
²³⁸ U	99.27417 (36)	+47307.8 (15)	238050787.0 (16)	238.02891 (3)
²³⁷ Np	—	+44871.7 (11)	237048171.7 (12)	—
²³⁹ Pu	—	+48588.3 (11)	239052161.7 (12)	—
²⁴¹ Am	—	+52934.4 (11)	241056827.4 (12)	—

TABLE 7. IRDF-II recommended nuclear decay data from BIPM for pure beta emitters. Numbers in parentheses indicate the absolute uncertainties, *e.g.*, 12.312(25) \equiv 12.312 \pm 0.025.

Reaction Product	Beta Data				
	Half-life (recomm.)	Half-life (DD eval.)	Time unit	Aver. E_β [keV]	Emiss. Prob. [%]
^3H	12.32(2)	12.312(25)	a	5.68(1)	100.
^{32}P	14.263(36)	14.284(36)	d	695.5(3)	100.
^{106}Ru	371.8(18)	371.5(21)	d	10.03(6)	100.

B. Dosimetry Cross Sections

The IRDF-II recommended cross sections are chosen from available ENDF-6 [60] formatted point cross sections from different sources as discussed below. Whereas some applications, such as radiation transport calculations, require the use of a complete nuclear data evaluation that includes all potential reaction channels, the IRDF-II neutron metrology library only requires the specification of a cross section for the channels used within the metrology application. However, unlike many applications, it requires a consistent uncertainty characterization for the cross section in the form of an energy-dependent covariance matrix. Because of the demanding requirements on neutron metrology cross sections and corresponding uncertainties, the recommended evaluations, rather than being taken from existing nuclear data evaluations, are often the result of a specifically funded metrology-related activity. Table 1 provides a list of the cross sections addressed within the IRDF-II library and identifies the source evaluations. Columns 8 of Table 1 also shows how the IRDF-*v*1.05 library differs from the previous IRDF-2002.

TABLE 8: IRDF-II recommended nuclear decay data content for gamma emitters. (DD eval.) refers to the latest Chechev compilation at <https://www-nds.iaea.org/IRDFtest/RCM3/Chechev-RCM3.pdf>. (*) indicates data taken from ENSDF. New DDEP evaluations undertaken within this project are highlighted in blue. Numbers in parentheses indicate the absolute uncertainties, *e.g.*, 109.734(14) \equiv 109.734 \pm 0.014.

Reaction Product	Half-life (recomm.)	Half-life (DD eval.)	Time unit	Gamma/X-ray Energy [keV]	Gamma/X-ray		Source Document
					Emiss. Prob. [%] (recomm.)	Emiss. Prob. [%] (DD eval.)	
^{18}F	109.77(5)	109.734(14)	m	511.	193.72(38)		BIPM
				0.525 $XK_\alpha 1$	0.013(4)		
				0.525 $XK_\alpha 2$	0.007(2)		
^{22}Na	2.6018(22)	2.6020	a	1274.537(7)	99.94(13)	99.940(14)	BIPM
				511.	180.7(2)	180.76(4)	
^{24}Na	14.997(12)	14.958(2)	h	1368.630(5)	99.9934(5)	99.9936(15)	BIPM
				2754.049(13)	99.862(3)	99.855(5)	
^{27}Mg	9.458(12)		m	843.76(10)	71.800(20)	71.8(4)	ENSDF
				1014.52(10)	28.200(20)	28.0(4)	
^{26g}Al	717 000(24)		a	11129.67(10)	2.5(2)		ENSDF
				1808.65(7)	99.76(4)		
				511.0	163.5(4)		
^{28}Al	(*)2.245(2)	2.245(2)	m	1778.987(15)	100.0		ENSDF
^{31}Si	(*)157.36(26)		m	1266.2(14)	0.0554(7)		ENSDF
^{46}Sc	83.787(16)	83.787(16)	d	889.271(2)	99.98374(25)		BIPM
				1120.537(3)	99.97(2)		
^{47}Sc	3.3485(9)	3.3485(9)	d	159.373(12)	68.1(5)		BIPM
^{48}Sc	(*)43.67(9)	43.70	h	983.526(12)	100.1(6)		ENSDF
				1037.522(12)	97.6(7)		
				1312.120(12)	100.1(7)		
				511.0	169.6(3)		
^{45}Ti	(*)184.8(5)	184.8	m	511.0	169.6(3)		ENSDF
^{51}Cr	27.704	27.704	d	320.0835(4)	9.89(2)		BIPM

Note that “old” means no change in the data, symbol “new” means the data were newly evaluated. Otherwise, Column 9 indicates how the current IRDF-II compares with IRDF-*v*1.05. Sometimes the recommended evaluation represents a combination where different energy regions are derived from different evaluated data sources. Column 6 provides reference to auxiliary documentation that describes the derivation of some of the recommended evaluations.

C. Nuclear Decay Data

This Sec. addresses the IRDF-II recommended nuclear decay data for activation products and for fission products. Tables 7 (beta emitters) and 8 (gamma emitters) give the half-lives and emission probabilities associated with the residual nuclei (activation product) for all of the radionuclides according to the detected decay radiation. It also shows the most recent recommended gamma and beta decay data, namely the emission energies and emission probabilities for all of the residual nuclei associated with the reactions addressed within IRDF-II.

Note that recommended data are those used in the IRDF-II cross-section evaluation. These data agree within uncertainties with the latest decay data evaluations (labeled **DD eval.** in Table headers). The latest decay data evaluations have been compiled into associated IRDF-II decay data file that is released together with the IRDF-II neutron metrology library.

TABLE 8: (continued). IRDF-II recommended nuclear decay data content for gamma emitters. (DD eval.) refers to the latest Chechev compilation at <https://www-nds.iaea.org/IRDFtest/RCM3/Chechev-RCM3.pdf>. (*) indicates data taken from ENSDF. New DDEP evaluations undertaken within this project are highlighted in blue.

Reaction Product	Half-life (recomm.)	Half-life (DD eval.)	Time unit	Gamma/Xray		Source Document	
				Gamma/X-ray Energy[keV]	Emiss.Prob. [%] (recomm.)		Emiss.Prob. [%] (DD eval.)
⁵⁴ Mn	312.12(10)	312.19(3)	d	834.855(3)	99.9997(3)	99.976(1)	BIPM
⁵⁶ Mn	2.5785(2)	2.57878(46)	h	846.7638(19) 1810.726(4) 2113.092(6)	98.85(3) 26.9(4) 14.2(3)	98.87(30) 27.189(791) 14.336(395)	BIPM
⁵³ Fe	(*)8.51(2)		m				ENSDF
⁵⁹ Fe	44.495(9)	44.494(12)	d	1099.245(3) 1291.590(6)	56.51(31) 43.23(33)	56.6(18) 43.2(14)	BIPM
⁵⁷ Co	(*)271.74(6)	271.81	d	14.41295(31) 122.06065(12) 136.47356(29)	9.18(12) 85.49(14) 10.71(15)	85.60(17) 10.68(8)	BIPM
⁵⁸ Co	70.86(6)	70.85(3)	d	810.7602(20)	99.44(2)	99.450(10)	BIPM
⁶⁰ Co	5.2711(8)	5.2711(8)	a	1173.228(3) 1332.492(4)	99.85(3) 99.9826(6)		BIPM
⁵⁷ Ni	35.60(6)	35.60(6)	h	127.164(3) 1377.62(4) 1757.55(3) 1919.62(14)	16.0(5) 81.2(6) 6.1(4) 12.5(5)	81.7(24) 12.3(4)	BIPM
⁶² Cu	9.73(3)	9.67(3)	m	1172.97(10) 511.0	0.342 195.66(5)	2 x 94.86(5)	ENSDF
⁶⁴ Cu	12.701(2)	12.7004(20)	h	1345.77(6) 511.	0.4748(34) 35.2(4)	34.8(4)	BIPM
⁶⁷ Cu	(*)61.83(12)	61.86	h	184.577(10)	48.7(3)		ENSDF
⁷⁴ As	(*)17.77(2)		d	595.83(8)	59(3)		ENSDF
⁸⁸ Y	(*)106.626(21)	106.63	d	1836.070(8) 898.042(11)	99.346(25) 93.7(3)	99.2(3)	BIPM
⁸⁹ Zr	78.41(12)	78.42(13)	h	908.97(3) 511.	99.03(2) 45.6(6)	99.04(1) 45.5(5)	BIPM
⁹⁵ Zr	64.032(6)	64.032(6)	d	756.729(12) 724.193(3)	54.38(22) 44.27(22)		BIPM
⁹⁵ Nb	34.991(6)	34.991(6)	d	765.803(6)	99.808(7)		BIPM
⁹⁴ Nb	20 300(16)	20 200	a	871.091(18) 702.65(6)	99.892(0) 99.814(0)		ENSDF
^{94m} Nb	(*)6.263(4)		m	871	0.50(6)		ENSDF
^{93m} Nb	16.1(2)	16.12(15)	a	30.77(2) 16.5213 $XK_{\alpha 2}$ 16.6152 $XK_{\alpha 1}$ 18.67 $XK_{\beta 1}$ 18.967 $XK_{\beta 2}$	0.000591(9) 3.32(8) 6.34(15) 1.64(4) 0.246(11)		BIPM
^{92m} Nb	10.15(2)		d	934.44(10)	99.15(4)	99.07(4)	ENSDF
⁹⁹ Mo	(*)65.924(6)		h	739.500(17)	12.20		ENSDF
¹⁰³ Ru	(*)39.247(13)		d	610.333(10) 497.085(10)	5.76(6) 91.0(12)		ENSDF
^{103m} Rh	(*)56.114(9)		m	39.755(12) 20.073 $XK - L_2$ 20.215 $XK - L_3$ 22.699–22.912 $XK - M_{2,3,4}$ 22.699–23.215 $XK - MN$ 23.167–23.172 $XK - N_{2,3,4,5}O_{2,3}$	0.068(5) 2.00(18) 3.8(3) 1.03(9) 1.20(11) 0.171(16)		ENSDF
^{110m} Ag	249.78(2)		d	1505.028(2) 1384.2931(20) 937.485(3) 884.6781(13) 763.9424(17) 706.6760(15) 657.7600(11)	13.16(16) 24.7(5) 34.51(27) 74.0(12) 22.31(9) 16.48(8) 94.38(8)		BIPM

TABLE 8: (continued). IRDF-II recommended nuclear decay data content for gamma emitters. (DD eval.) refers to the latest Chechev compilation at <https://www-nds.iaea.org/IRDFtest/RCM3/Chechev-RCM3.pdf>. (*) indicates data taken from ENSDF. New DDEP evaluations undertaken within this project are highlighted in blue.

Reaction Product	Half-life (recomm.)	Half-life (DD eval.)	Time unit	Gamma/Xray		Source Document	
				Gamma/X-ray Energy[keV]	Emiss.Prob. [%] (recomm.)		Emiss.Prob. [%] (DD eval.)
^{113m}In	(*)99.476(23)	99.48	m	391.698(3)	64.94(17)	ENSDF	
^{114m}In	(*)49.51(1)	49.51	d	190.27(3) 24.002 24.21	15.56(15) 9.8(3) 18.2(6)	ENSDF	
^{115m}In	4.486(4)		h	336.244(17)	45.9(1)	ENSDF	
^{116m}In	54.29(17)		m	1293.56(2) 1097.28(2)	84.8(12) 58.5(8)	ENSDF	
^{126}I	(*)12.93(5)	12.93	d	666.331(12) 388.633(11)	32.9(7) 35.6(6)	ENSDF	
^{137m}Ba	(*)2.552(1)	2.552	m	661.657(3)	90.07(20)	BIPM	
^{137}Cs	(*)30.08(9)		a	661.657(3)	84.99(20)	BIPM	
^{140}La	1.6781(3)	1.67855(12)	d	1596.203(13) 815.784(6) 328.761(4)	95.40(5) 23.72(20) 20.8(3)	95.40(8) BIPM	
^{140}Ba	(*)12.7527(23)	12.7527	d	537.261(25)	24.6(5)	BIPM	
^{144}Ce	(*)284.91(5)		d	133.5152(20)	10.83(12)	BIPM	
^{140}Pr	(*)3.39(1)	3.39	m	1596.1(2) 511.0 39.258	0.49 102.0(6) 3.68(8)	ENSDF	
^{167}Tm	(*)9.25(2)	9.25	d	207.801(5)	42(8)	BIPM	
^{168}Tm	(*)93.1(2)		d	815.89(5) 447.515(3) 198.251(2)	50.95(16) 23.98(11) 54.49(16)	48.99(150) 23.06(71) 53.4(16)	ENSDF
^{182}Ta	114.43	114.74(12)	d	1221.395(3) 1189.040(3) 1121.290(3)	27.35(27) 16.66(16) 35.30(33)	BIPM	
^{187}W	23.72(6)	24.00	h	685.81(1) 551.55(1) 479.53(1)	33.2(5) 6.14(10) 26.6(4)	27.30(93) 21.84(71)	ENSDF
^{196}Au	6.183(10)	6.1669(6)	d	333.03(5) 355.73(5) 426.10(6)	22.9(9) 87. 6.6	ENSDF	
^{198}Au	2.6943(8)	2.6943(3)	d	411.80205(17) 675.8836(7) 1087.6842(7)	95.62(6) 0.804(7) 0.1591(21)	BIPM	
^{199m}Hg	42.6(2)	42.67(9)	m	158.3(1) 374.1(1)	52.3 13.8(11)	ENSDF	
^{204m}Pb	67.2(3)	66.93(10)	m	374.76(7) 911.74(15)	94.20(14) 91.5(13)	89.253(14876) 90.0691(2999)	ENSDF
^{204}Bi	11.22(10)	11.28	h	1755.28(6) 1703.27(5) 983.98(3) 899.15(3) 670.72(3)	1.23(16) 2.00(24) 59(6) 99(12) 11.4(13)	ENSDF	
^{205}Bi	15.31(4)	15.11	d	703.45(5) 987.66(5) 1764.30(10)	31.1 16.1(3) 32.5(7)	ENSDF	
^{206}Bi	6.243(3)		d	803.10(5) 537.45(4) 343.51(3)	99.0(14) 30.5(4) 23.5(4)	ENSDF	
^{207}Bi	32.9(14)	31.55(4)	a	1063.656(3) 1770.228(9) 569.698(2)	74.58(22) 6.871(26) 97.76(3)	74.6(5) 6.87(3)	BIPM
^{208}Bi	(*)368 000(4)		a	2614.5	99.785	ENSDF	
^{233}Th	(*)21.83(4)	22.15(8)	m	595.39(6) 170.60(6) 86.477(10)	0.1178(16) 0.507(9) 1.843(22)	BIPM	

TABLE 8: (continued). IRDF-II recommended nuclear decay data content for gamma emitters. (DD eval.) refers to the latest Chechev compilation at <https://www-nds.iaea.org/IRDFtest/RCM3/Chechev-RCM3.pdf>. (*) indicates data taken from ENSDF. New DDEP evaluations undertaken within this project are highlighted in blue.

Reaction Product	Half-life (recomm.)	Half-life (DD eval.)	Time unit	Gamma/X-ray Energy [keV]	Gamma/Xray		Source Document
					Emiss.Prob. [%] (recomm.)	Emiss.Prob. [%] (DD eval.)	
^{233}Pa	(*)26.975(13)	26.98(2)	d	415.764(5)	1.97(12)		BIPM
				311.904(5)	68.9(12)		
				300.129(5)	12.3(4)		
^{237}U	(*)6.752(2)	6.749(16)	d	332.376(16)	1.199(16)	21.2(3)	BIPM
				208.00(1)	21.3(3)		
				59.54091(10)	34.1(9)		
^{239}U	23.46		m	844.10(3)	0.139(3)		BIPM
				662.28(2)	0.170(5)		
				74.664(1)	51.6(13)		
^{239}Np	2.356(3)		d	106.125(2)	25.9(3)		BIPM
				228.183(1)	11.32(22)		
				334.310(3)	2.04(2)		

D. Fission Yield Data

The fission cross sections of the actinides included in the IRDF-II library differ significantly in the thermal and the epithermal neutron energy range, which determines the relative importance of the fission yield data. Comparison of the fission cross sections is given in Fig. 1 for ^{232}Th , $^{235,238}\text{U}$, ^{237}Np , ^{239}Pu , and ^{241}Am targets.

The baseline recommended fission yields for dosimetry applications are taken from the JEFF-3.3 library. These data are modifications to the JEFF-3.1.1 library and are also identified as UKFY3.7 [68]. When the fission yields do not appear in the JEFF-3.3 library, *e.g.*, the 14 MeV fission yields for ^{239}Pu and ^{237}Np , the recommended values are taken from the ENDF/B-VIII.0 library.

While the dosimetry community looks forward to the future use of a more refined energy-dependence to the recommended fission yields, the current IRDF-II library recommendation is based on a selection of the appropriate yield from one of the three energy-dependent groups: thermal, fast (average energy in the range 400 keV to 500 keV), and 14 MeV.

E. Thermal Neutron Cross Sections and Resonance Integrals

Neutron dosimetry is the inverse of neutron activation analysis (NAA). Independent data sets exist like the Kayzero library for NAA by the k_0 standardization method [69], for example. There is a direct relation between the constants in the Kayzero data-base and the differential cross sections [70]. Since the constants were measured independently of the differential cross sections, the k_0 and Q_0 constants contained in the Kayzero library can be used for the validation of the thermal capture cross sections and resonance integrals in the IRDF-II library, in addition to the comparison with other compilations like the Atlas of Resonance Parameters [71]. The comparison is discussed and presented in Sec. VI G.

V. CROSS-SECTION COMPARISON BETWEEN AVAILABLE EVALUATIONS FOR SELECTED REACTIONS

The following Secs. provide a commented review and details on some of the recommended cross sections that appear within the IRDF-II library. The reactions addressed below focus on the new and updated content of the library. Some of the reaction-specific verification evidence is summarized below and comparisons are made with other evaluations. Note that all reaction evaluations present in the IRDF-II library feature both the evaluated mean values and the corresponding covariance matrices. The definition of the covariance matrix requires that it be positive semi-definite. The eigenvalues of all covariance matrices have been checked to be non-negative in the evaluator's energy grids; all calculated eigenvalues are listed in the IRDF-II files in ENDF-6 format in the comment section (MT=451) for each dosimetry reaction. Users should be aware that numerical processing of covariance matrices may introduce very small negative eigenvalues due to round-off errors; processed covariance matrices require to be checked for negative eigenvalues, and regularized if needed by adding a very small number to the diagonal elements.

Whereas the ordering of the nuclear data reaction content in Table 1 is with increasing atomic number of the target nuclei, the ordering of Secs. here is different. This new library includes composite reactions and extends earlier evaluation up to 60 MeV. The ordering of Secs. below is designed to most easily illustrate the verification steps taken in the preparation of the library. It starts with a simple high energy reaction, the $^{58}\text{Ni}(n,p)^{58}\text{Co}$ reaction, then proceeds to address the related composite reaction, $^{\text{nat}}\text{Ni}(n,X)^{58}\text{Co}$, that is now added to the library. The following Secs. then address more complicated high energy composite dosimetry reactions such as $^{\text{nat}}\text{Fe}(n,X)^{54}\text{Mn}$, resonance reactions and new reactions that have a reaction threshold above 20 MeV. The survey of reactions then ends with a discussion of the very complex composite reactions, such as gas-production reactions $^{\text{nat}}\text{Li}(n,X)^4\text{He}$ and $^{\text{nat}}\text{B}(n,X)^4\text{He}$, where there can be multiple target

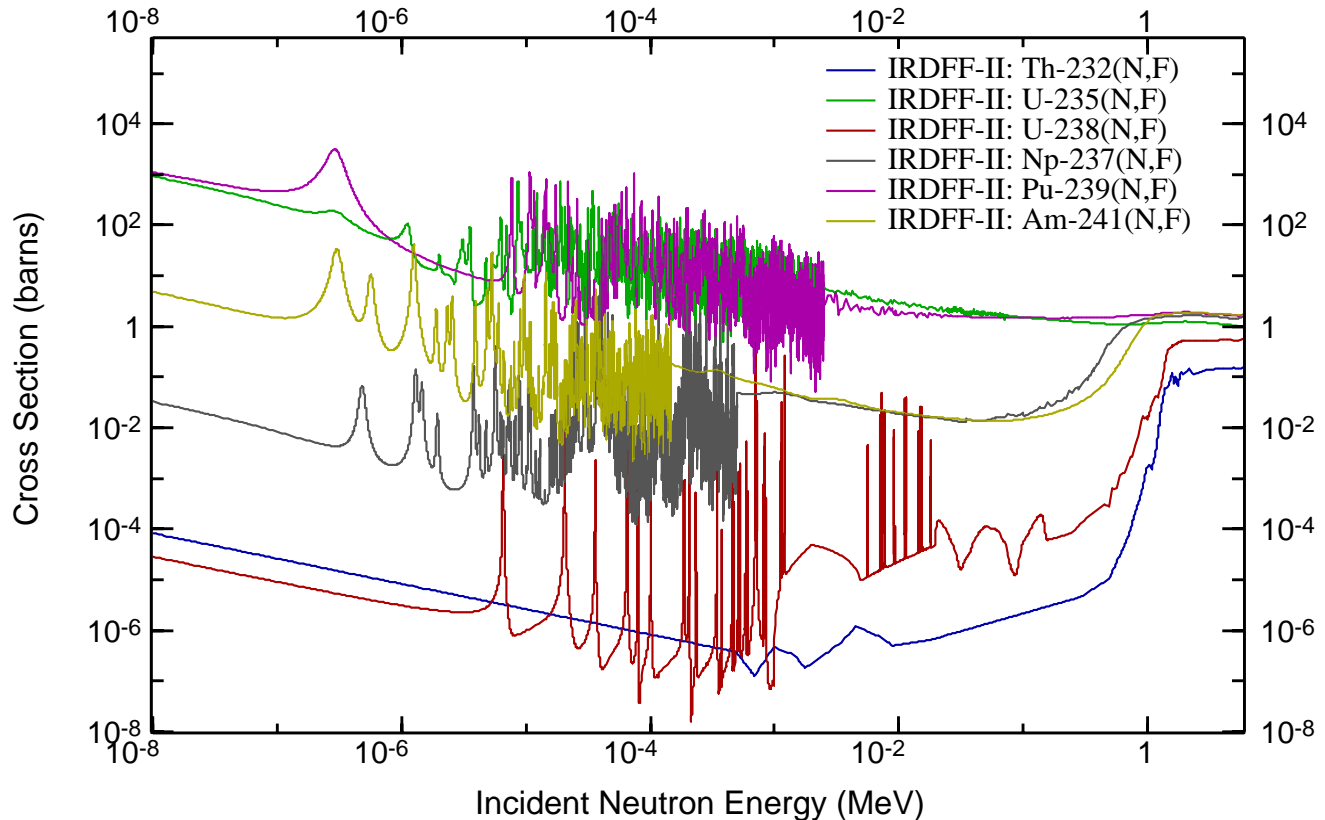


FIG. 1. (Color online) Comparison of the neutron-induced fission cross-section data of the actinides in the IRDF-II Library.

isotopes and multiple contributing reaction channels.

A. Reaction Q-values and Thresholds

Reaction threshold is directly related to its Q-value. As a rule, exothermic reactions do not have a threshold, but in some cases there is an effective threshold due to the Coulomb barrier in the outgoing charged-particle emission, and the probability of the reaction is very small. However, “small” is actually case-dependent, since in “well-thermalised systems” the neutron flux at thermal energies can be many orders of magnitude higher than the fast flux, where the cross section is significant. The special cases in the IRDF-II library are listed in Table 9, together with the Q-values and the adopted effective threshold energies.

TABLE 9. Reaction Q-values and thresholds

ID	Reaction	Label	MAT	Q-Value [keV]	Eff. Thr. [keV]
1	$^{10}\text{B}(n,\alpha)^7\text{Li}$	B10a	525	2789.323	0.0
2	$^{47}\text{Ti}(n,p)^{47}\text{Sc}$	Ti47p	2228	181.580	800.0
3	$^{54}\text{Fe}(n,p)^{54}\text{Mn}$	Fe54p	2625	85.2127	0.0
4	$^{54}\text{Fe}(n,\alpha)^{51}\text{Cr}$	Fe54a	2625	842.760	0.0
5	$^{58}\text{Ni}(n,p)^{58}\text{Co}$	Ni58p	2825	400.800	400.0
6	$^{59}\text{Co}(n,\alpha)^{56}\text{Mn}$	Co59a	2725	328.400	334.016
7	$^{63}\text{Cu}(n,\alpha)^{60}\text{Co}$	Cu63a	2925	1714.400	2250.0
8	$^{64}\text{Zn}(n,p)^{64}\text{Cu}$	Zn64p	3025	203.500	500.0
9	$^{67}\text{Zn}(n,p)^{67}\text{Cu}$	Zn67p	3034	220.690	0.0
10	$^{92}\text{Mo}(n,p)^{92m}\text{Nb}$	MoNb92pm	4225	425.680	1000.0

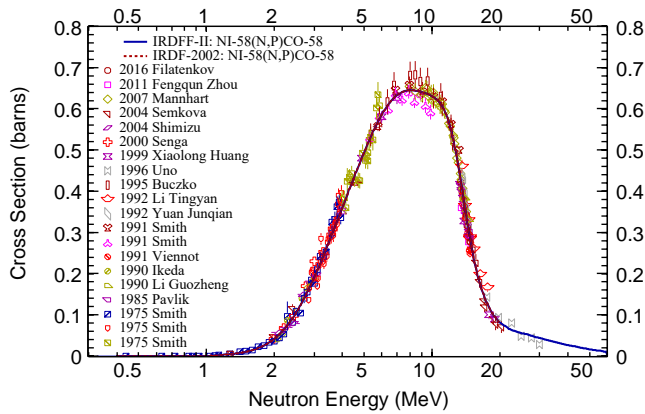
B. Description of Selected Cross-Section Evaluations

1. $^{58}\text{Ni}(n,p)^{58}\text{Co}$

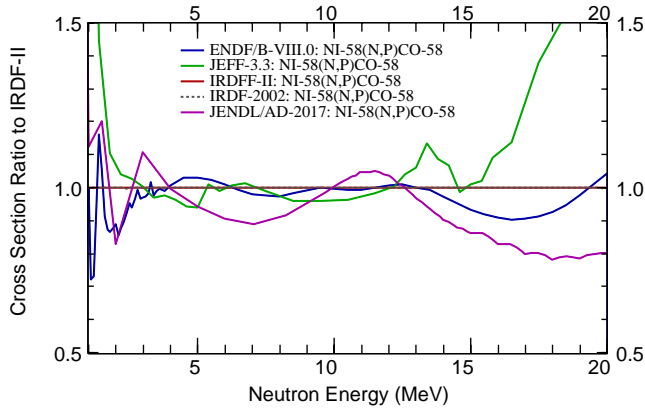
The zero to 20 MeV portion of the selected cross section comes from an IAEA-sponsored evaluation performed by K. Zolotarev in December 2002, documented as part of the RRDF-2002 library, and released within the IAEA-generated IRDF-2002 library. The IAEA sponsored the extension of this library from 20 MeV to 60 MeV, which was incorporated into the evaluation in February 2014. This extension used the model-based sub-library TENDL-2013(s60) of the TENDL-2013 library with explicit representation of reaction cross sections up to 60 MeV, while renormalizing (labelled in this paper as (r)) the calculations in the energy range 20 MeV to 60 MeV to ensure continuity of the data at 20 MeV. The associated IRDF-II covariance matrices correctly reflect the derivation of the evaluation. These development details can be seen in the entry in Table 1 for this reaction.

As noted in Sec. I, it is critical that recommended cross sections for dosimetry applications have evidence that the evaluation took into consideration the available body of experimental data. No experimental data exists for neutron energies greater than 30 MeV. Fig. 2(a) shows good agreement of the selected EXFOR data with the IRDF-II cross sections. This comparison provides strong verification evidence for the consistency of the evaluation with measured differential data.

Fig. 2(a) also shows that the cross sections for this reaction are practically equal to those in the older IRDF-2002.



(a) Comparison to selected experimental data from EXFOR [72].



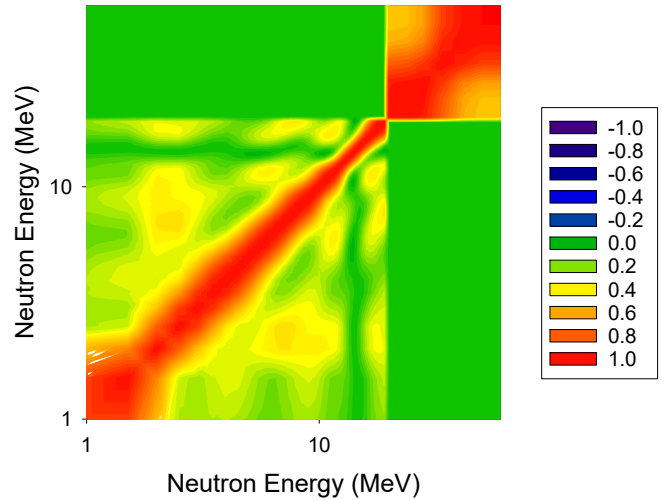
(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

 FIG. 2. (Color online) $^{58}\text{Ni}(n,p)^{58}\text{Co}$ cross-section evaluation in IRDFF-II library relative to other data.

For most fission reactor applications, the relevant 10 %-90 % sensitivity region for this reaction is between 2 MeV and 7 MeV. In order to also illustrate the variations between evaluations within this region, Fig. 2(b) shows the ratio of the various evaluations to the IRDFF-II recommended cross section, namely ENDF/B-VIII.0, JEFF-3.3, and JENDL/AD-2017. Variations on the order of 10 % are observed. The curve for the IRDFF-II ratio is unity by definition in this figure. The JEFF-3.3 evaluation shows significant differences near threshold and above 15 MeV.

The inclusion of uncertainty data, in the form of a covariance matrix, is a required feature of dosimetry cross sections. Fig. 3(b) shows the energy-dependent 1-sigma uncertainty in the cross section using the energy group structure provided by the evaluator. The energy structure used in the uncertainty characterization is seen to only have a single large-width bin in the 0.4 MeV to 1.5 MeV region, so, due to the correlation of the cross section within this region, the energy-grouped uncertainty in Fig. 3(b) does not exhibit the large uncertainty suggested by the variation between different evaluations in Fig. 2(b). Fig. 3(a) shows the correlation matrix for the evaluation. The covariance matrix elements in the >20 MeV region are seen to be uncorrelated with those at lower energies - as expected based on the evaluation methodology, since the two regions originate from different evaluations.

As discussed above, the definition of the covariance matrix requires that it be positive semi-definite. In order



(a) Cross-section correlation matrix.

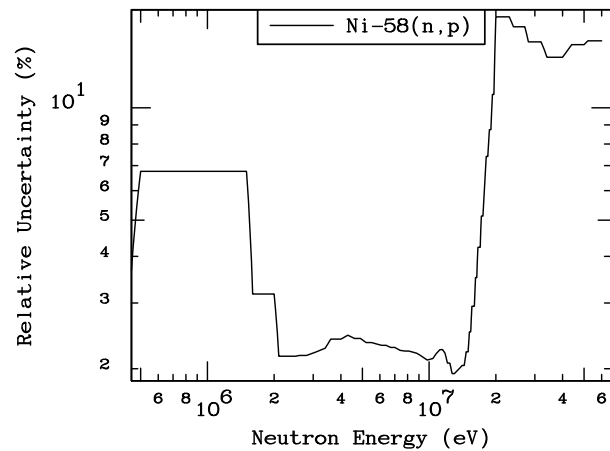

 (b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

 FIG. 3. (Color online) Uncertainties and correlations of the $^{58}\text{Ni}(n,p)^{58}\text{Co}$ cross-section evaluation in IRDFF-II library.

 TABLE 10. List of lowest eigenvalues in ascending order of the $^{58}\text{Ni}(n,p)^{58}\text{Co}$ covariance matrix.

No. Row	Eigenvalue
1	1 0.00000E+00
2	11 2.98127E-08
3	17 3.02008E-08
4	24 3.03488E-08
5	20 3.17274E-08
6	8 3.23051E-08
7	21 3.27524E-08
8	22 3.37126E-08
9	12 3.53861E-08
10	14 3.71190E-08
...	
44	39 1.16138E-01

to verify this mathematical property, the smallest and the largest eigenvalues of the covariance matrix are shown in Table 10 for this reaction; all eigenvalues are positive numbers.

Another verification step is the comparison of the calculated-to-experimental ratio of the spectrum-

TABLE 11. C/E of SACS for $^{58}\text{Ni}(n,p)^{58}\text{Co}$ reaction and data used by the nuclear data evaluator.

Spectrum	Experiment		IRDF-II Calculated x.s. (mb)
	Cross Section (mb)	Source	
$^{235}\text{U}(n_{th},f)$	108.9 ± 4.78 %	[73]	107.44
	106.0 ± 6.60 %	[74]	
	108.5 ± 1.29 %	[75]	
$^{252}\text{Cf}(s.f.)$	117.6 ± 1.28 %	[76]	117.36
	117.5 ± 1.30 %	[77]	

averaged cross section (SACS) in the ^{252}Cf spontaneous fission and $^{235}\text{U}(n_{th},f)$ prompt fission neutron benchmark fields. We emphasize that this, within the context of the evaluation, is treated purely as a verification step since the nuclear data evaluator also commonly examines this metric as part of their verification process for the evaluation. The treatment of validation in these benchmark fields is addressed later in Sec. VII. Table 11 compares the spectrum-averaged cross sections produced with the IRDF-II library to that reported by the nuclear data evaluator in the evaluation documentation.

The residual nucleus produced in this reaction is ^{58}Co which, in addition to its 70.86 day half-life for the ground state, also has a 9.04 hour half-life isomer. It is critical to note that, consistent with adopted notation, this evaluation captures the complete $^{58}\text{Ni}(n,p)^{58(m+g)}\text{Co}$ cross section, that is, it represents the combined (n,p) reaction that populates the ^{58}Co ground state and the meta-stable states. An important consideration in the use of this reaction for dosimetry in commercial fission reactors is the consideration of the thermal neutron burn-up correction that can be a factor in long irradiations in the presence of a thermal neutron environment. The effective burn-up cross section, via the $^{58}\text{Co}(n,\gamma)$ reaction, is 1650 barns for ^{58g}Co while that for the ^{58m}Co residual nucleus is 1.4×10^5 barns. The caveats on the need to make thermal neutron burn-up corrections for applications in light water reactors are found in ASTM E264 [78]. The cross sections and the decay data of the metastable isotope ^{58m}Co are not included in the present version of IRDF-II.

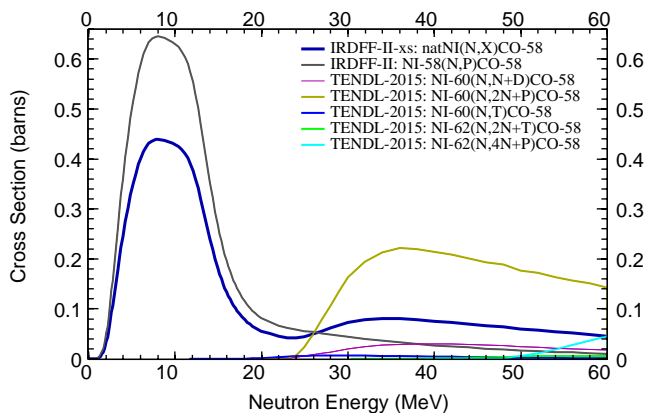


FIG. 4. (Color online) Contributing reaction cross sections to the ^{58}Co production from neutrons incident on a ^{nat}Ni target. Note that the cross sections are not scaled by the isotopic abundance.

2. $^{nat}\text{Ni}(n,X)^{58}\text{Co}$

Nickel is an element that has several naturally occurring isotopes and many reactions on different isotopes may lead to the same residual. Monitor samples are usually made of natural elements, so it is advantageous to provide the data for the residual production from the element, since production paths may include a multitude of minor channels which a user would need to consider. The cross sections and the covariance data for the $^{nat}\text{Ni}(n,X)^{58}\text{Co}$ reaction were assembled from the IRDF-II evaluation for the $^{58}\text{Ni}(n,p)$ reaction (described in the previous section), which is the dominant contribution at low energies. The contributing reactions at higher energies were taken from the TENDL-2015 library, as shown in Fig. 4.

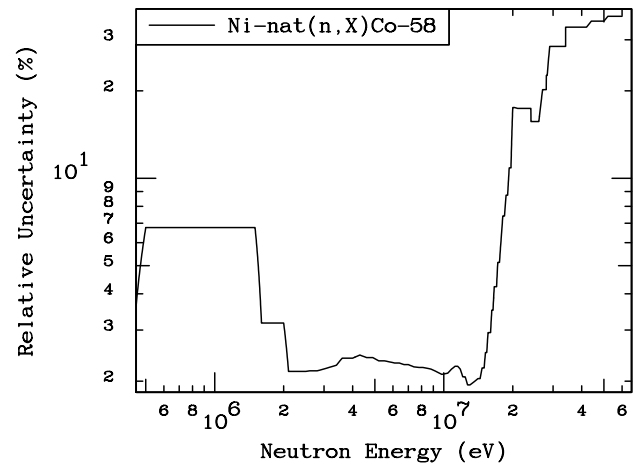


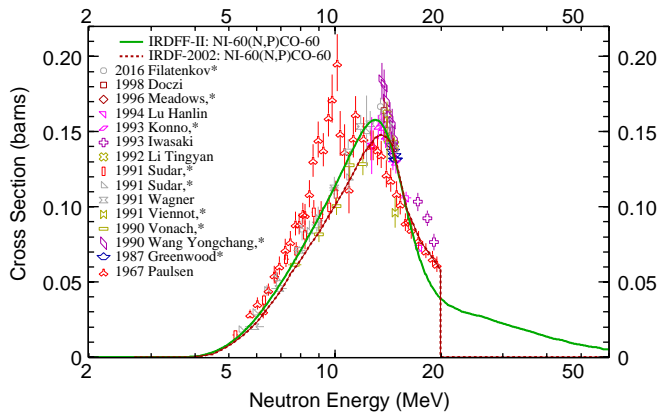
FIG. 5. (Color online) Uncertainties in the $^{nat}\text{Ni}(n,X)^{58}\text{Co}$ reaction cross sections.

The covariance plot and the uncertainty are practically the same as for the $^{58}\text{Ni}(n,p)^{58}\text{Co}$ reaction except for the high energy part. The 1-sigma standard deviation is shown in Fig. 5.

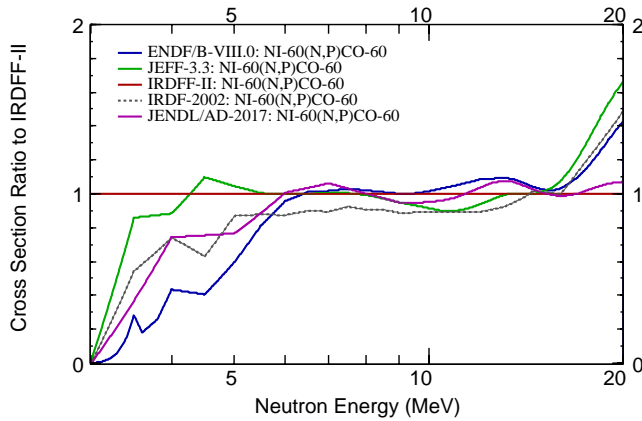
3. $^{60}\text{Ni}(n,p)^{60}\text{Co}$

The cross sections and the covariance data for the $^{60}\text{Ni}(n,p)^{60}\text{Co}$ reaction in the energy range from threshold to 21 MeV were evaluated by Zolotarev [79] in 2008. Extension to 60 MeV was done at the IAEA based on TENDL-2011, renormalizing the cross sections at 21 MeV for continuity. Comparison of the cross sections with experimental data is shown in Fig. 6(a). The figure also shows the comparison between IRDF-2002 and IRDF-II, which remained unchanged from version IRDF-II-v1.05.

Cross sections in different evaluated libraries differ significantly near threshold as shown in Fig. 6(b), where the ratios of the various evaluations to the IRDF-II recommended cross section are given. These differences may be important if this reaction monitor is used in reactor spectra. It is interesting that the cross section curves differ also at the plateau of the cross section near 13 MeV. The plot of the 1-sigma uncertainty is shown in Fig. 7(b), and the



(a) Comparison to selected experimental data from EXFOR [72].

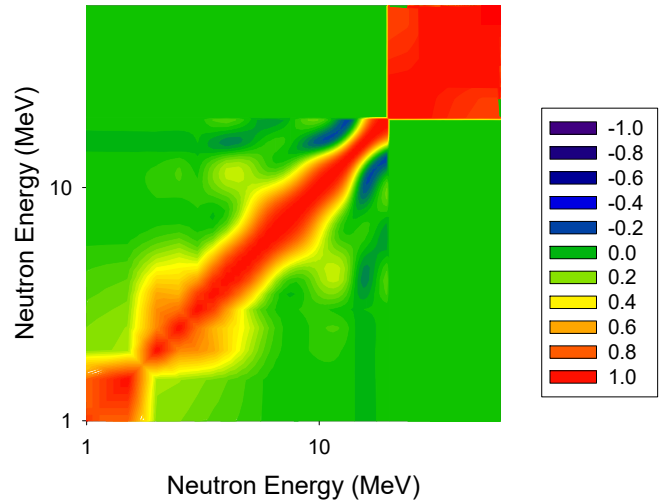


(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

 FIG. 6. (Color online) $^{60}\text{Ni}(n,p)^{60}\text{Co}$ cross-section evaluation in IRDFF-II library relative to other data.

corresponding correlation matrix for the $^{60}\text{Ni}(n,p)^{60}\text{Co}$ reaction is shown in Fig. 7(a).

The $^{60}\text{Ni}(n,p)^{60}\text{Co}$ cross sections are traditionally included in dosimetry libraries, but the users should note that several neutron induced reactions from the heavier nickel isotopes lead to the same product, as shown in Fig. 8. Although the abundance of ^{61}Ni is about twenty times smaller than the abundance of ^{60}Ni , the latter may start contributing to the ^{60}Co production at energies above 12 MeV through the $^{61}\text{Ni}(n,np)^{60}\text{Co}$ reaction. The user is advised to use the $^{\text{nat}}\text{Ni}(n,X)^{60}\text{Co}$ cross sections that are newly introduced into the IRDFF-II library, noting that the cross sections are not validated for dosimetry purposes in neutron fields with a significant contribution of the flux above 12 MeV.



(a) Cross-section correlation matrix.

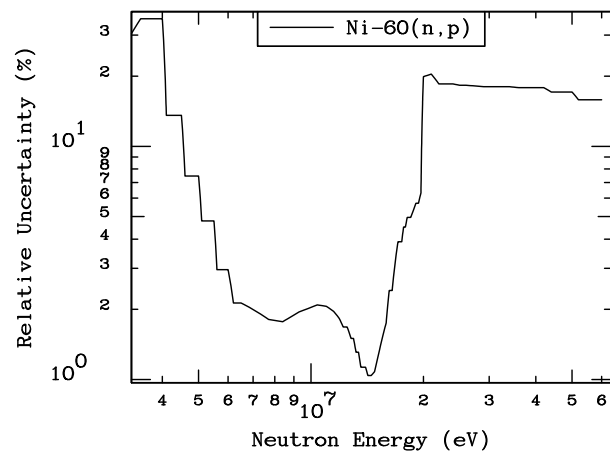
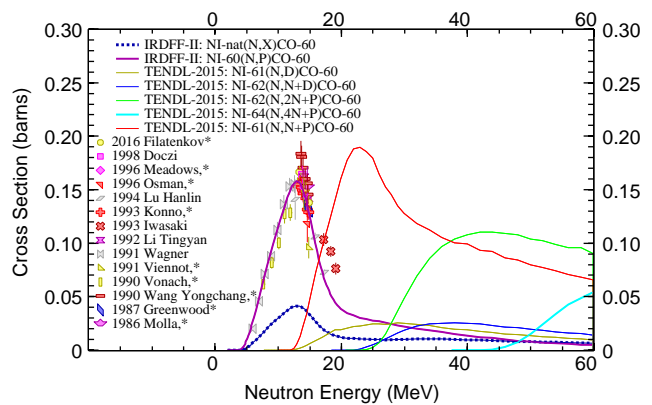
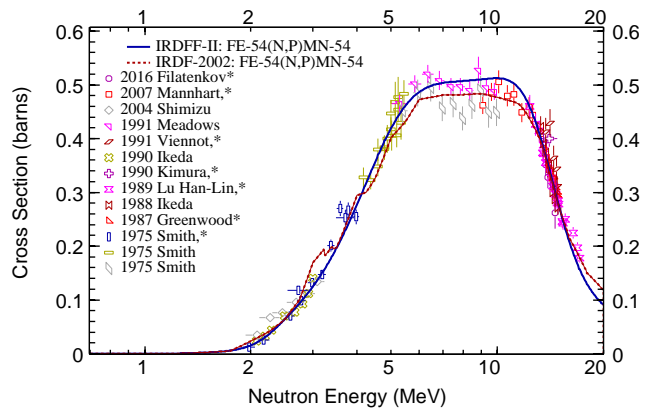

 (b) One-sigma uncertainties ($\equiv \sqrt{\text{cov}(i, i)}$).

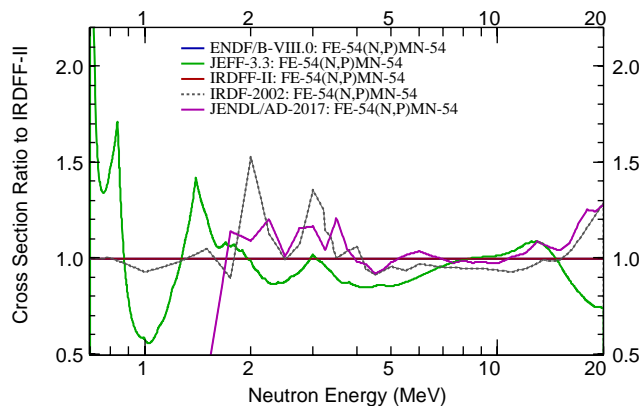
 FIG. 7. (Color online) Uncertainties and correlations of the $^{60}\text{Ni}(n,p)^{60}\text{Co}$ cross-section evaluation in IRDFF-II library.

 FIG. 8. (Color online) Contributing reactions to the ^{60}Co production from neutrons incident on a $^{\text{nat}}\text{Ni}$ target. Note that the cross sections are not scaled by the isotopic abundance.

4. $^{54}\text{Fe}(n,p)^{54}\text{Mn}$

The $^{54}\text{Fe}(n,p)^{54}\text{Mn}$ reaction cross sections and covariances below 20 MeV were evaluated by Zolotarev [81] in 2013. Extension to 60 MeV was done at the IAEA based on TENDL-2013, renormalizing the cross sections at 20 MeV for continuity. Comparison of the cross sections with experimental data is shown in Fig. 9(a). The figure also shows the comparison between IRDF-2002 and IRDF-2002 and IRDF-2002, which remained unchanged from version IRDF-2002-v1.05. Cross sections in different evaluated libraries differ significantly near threshold as shown in Fig. 9(b), where the ratios of the various evaluations to the IRDF-2002 recommended cross section are given. These differences may be important if this reaction monitor is used in reactor spectra.



(a) Comparison to selected experimental data from EXFOR [72].

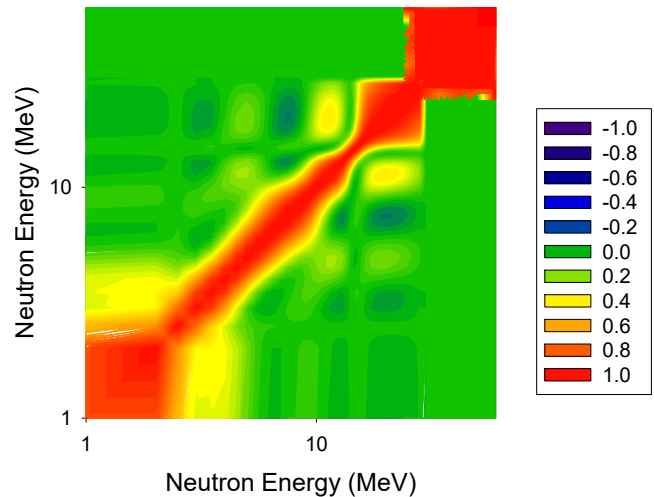


(b) Ratio of cross-section evaluations to IRDF-2002 evaluation.

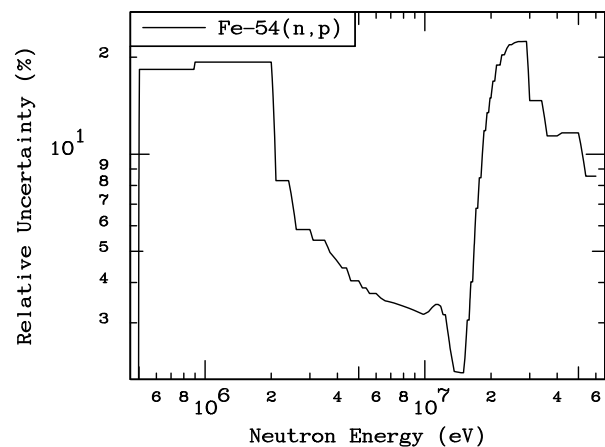
FIG. 9. (Color online) $^{54}\text{Fe}(n,p)^{54}\text{Mn}$ cross-section evaluation in IRDF-2002 library relative to other data.

The plot of the correlation matrix for the $^{54}\text{Fe}(n,X)^{54}\text{Mn}$ reaction is shown in Fig. 10(a). The plot of the 1-sigma uncertainty is shown in Fig. 10(b).

As in the case of nickel, the $^{54}\text{Fe}(n,p)^{54}\text{Mn}$ reaction cross sections are traditionally included in dosimetry libraries, but the users should note that several neutron induced reactions from the other isotopes lead to the same product. The thresholds for the competing reactions are fairly high and they become significant only well above 20 MeV. Nevertheless, the user is advised to use the $^{\text{nat}}\text{Fe}(n,X)^{54}\text{Mn}$ cross sections that are newly introduced into the IRDF-2002 library and are described



(a) Cross-section correlation matrix.



(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

FIG. 10. (Color online) Uncertainties and correlations of the $^{54}\text{Fe}(n,p)^{54}\text{Mn}$ cross-section evaluation in IRDF-2002 library.

below, noting that the cross sections are not validated for dosimetry purposes in neutron fields with a significant contribution of the flux above 20 MeV.

5. $^{\text{nat}}\text{Fe}(n,X)^{54}\text{Mn}$

Elemental iron consists mainly of isotope ^{56}Fe , but the minor isotopes are sometimes important as they may lead to reactions that produce the same residual. Monitor samples are usually made of natural elements, so it is advantageous to provide the data for the residual production from the element, since production paths may include a multitude of minor channels which a user would need to consider. The cross sections and the covariance data for the $^{\text{nat}}\text{Fe}(n,X)^{54}\text{Mn}$ reaction were assembled from the IRDF-2002 evaluation for the $^{54}\text{Fe}(n,p)$ reaction (described in the previous section), which is the dominant contribution at low energies. The contributing reactions at higher energies were taken from the TENDL-2015 library, as shown in Fig. 11. Note that above 20 MeV the ^{54}Mn production cross section is dominated by the con-

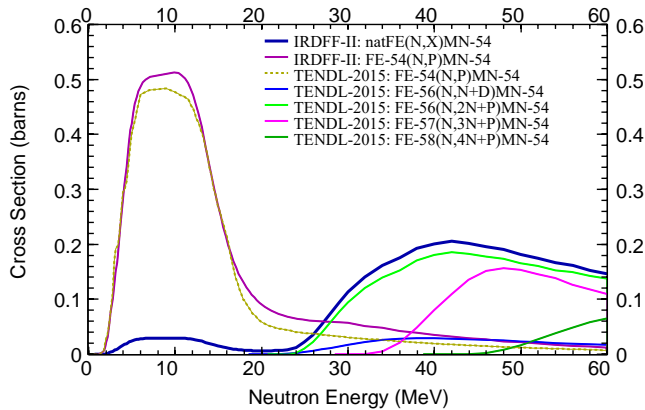


FIG. 11. Contributing reaction cross sections to the ^{54}Mn production from neutrons incident on a $^{\text{nat}}\text{Fe}$ target. Note that the cross sections are not scaled by the isotopic abundance.

tributions from reactions on ^{56}Fe . For neutron dosimetry at these energies the use of the elemental radionuclide production cross sections is essential.

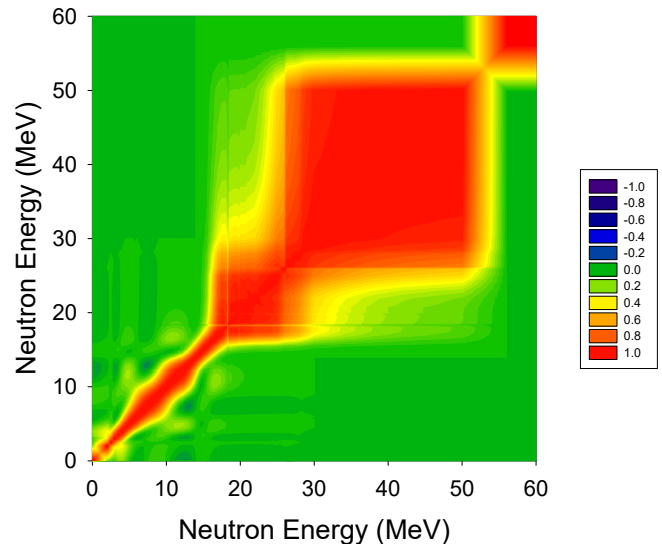
The plot of the correlation matrix for the $^{\text{nat}}\text{Fe}(n,X)^{54}\text{Mn}$ reaction is shown in Fig. 12(a). The plot of the 1-sigma uncertainty is shown in Fig. 12(b).

6. $^{58}\text{Fe}(n,\gamma)^{59}\text{Fe}$

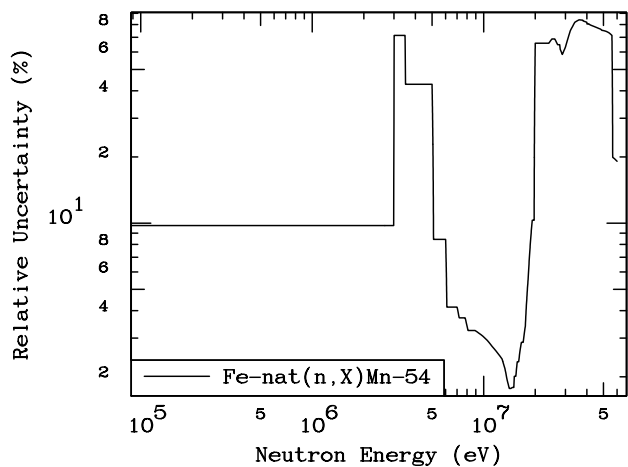
The $^{58}\text{Fe}(n,\gamma)^{59}\text{Fe}$ reaction belongs to a minor isotope of iron, with abundance of less than 0.3 % and pronounced resonance structure, hence the measured cross section data are less precise. Through an IAEA-sponsored activity M. Moxon re-evaluated the resonance parameters of ^{58}Fe in 2004 [82], which were adopted in the JEFF and ENDF/B libraries. The evaluation removed some of the discrepancies with the measured thermal capture cross section and the resonance integral. The cross sections of ^{58}Fe up to 60 MeV for IRDF-II were taken from the JEFF-3.1 library. Comparison of the cross sections with experimental data in the fast energy range is shown in Fig. 13; big discrepancies are shown among different evaluated libraries. The cross section comparison in the resonance range is not shown, since many libraries adopt the Moxon evaluation. The figure also shows the comparison between IRDF-2002 and IRDF-II, which remained unchanged from version IRDF-II-v1.05.

The plot of the correlation matrix for the $^{58}\text{Fe}(n,\gamma)^{59}\text{Fe}$ reaction is shown in Fig. 14(a). The plot of the 1-sigma uncertainty is shown in Fig. 14(b).

The $^{58}\text{Fe}(n,\gamma)^{59}\text{Fe}$ reaction is interesting for dosimetry applications because it is “clean” in the sense that it does not have any contributions from other isotopes of the elemental iron. The drawback is its low abundance, which is inherently known with a lower relative precision and this makes precise differential measurements more difficult, because they require enriched samples. The drawbacks are reflected in the lower accuracy of the evaluated cross sections.



(a) Cross-section correlation matrix.



(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

FIG. 12. (Color online) Uncertainties and correlations of the $^{\text{nat}}\text{Fe}(n,X)^{54}\text{Mn}$ cross-section evaluation in IRDF-II library.

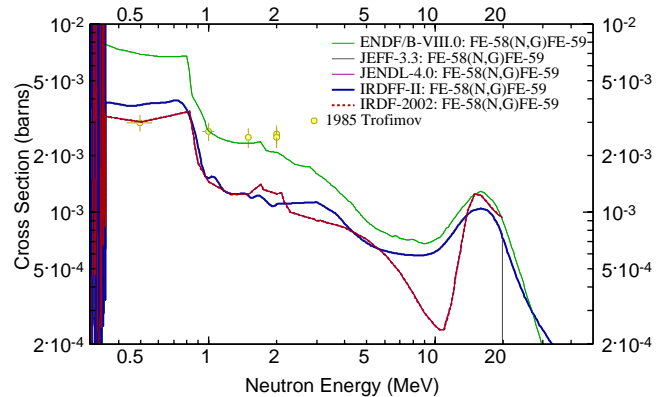
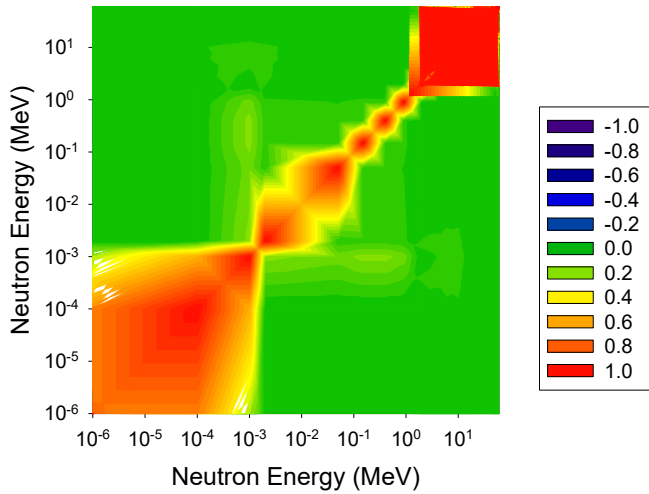
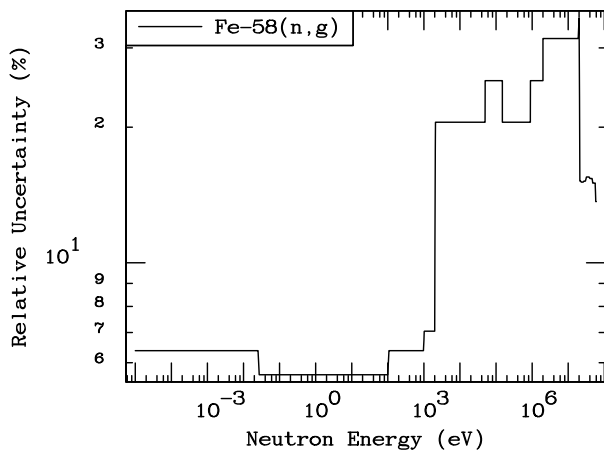


FIG. 13. (Color online) Comparison of $^{58}\text{Fe}(n,\gamma)^{59}\text{Fe}$ cross-section evaluations with experimental data from EXFOR.



(a) Cross-section correlation matrix.

(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.FIG. 14. (Color online) Uncertainties and correlations of the $^{58}\text{Fe}(n,\gamma)^{59}\text{Fe}$ cross-section evaluation in IRDFF-II library.

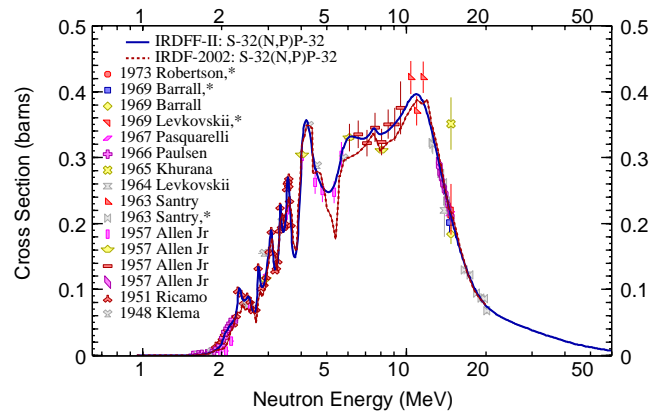
7. $^{32}\text{S}(n,p)^{32}\text{P}$

The residual nuclide from the $^{32}\text{S}(n,p)^{32}\text{P}$ reaction is ^{32}P , which, as seen from Table 7, is a pure beta emitter. As with all beta dosimetry, care must be taken in the dosimetry measurement process to ensure that only the ^{32}P , and no other activation products are being counted. Potential interfering reaction products may arise from activation of any of the naturally occurring isotopes in the target material, *e.g.*, ^{32}S , ^{33}S , or ^{35}S . There are many ways to addressing the potential interferences [80]. For beta measurements performed on samples irradiated in fission reactor environments, a common practice for eliminating interference from other beta emitters is to use a cadmium thermal neutron shield to reduce the generation of capture products and to wait 24 hours between irradiation and counting so that residual reaction products, such as ^{31}S , ^{34}P , ^{31}Si , and ^{37}S , can die away before the counting is started. To increase the efficiency of the counting, the irradiated sulfur can also be burned, leaving a residue of ^{32}P that can be counted without the need for corrections

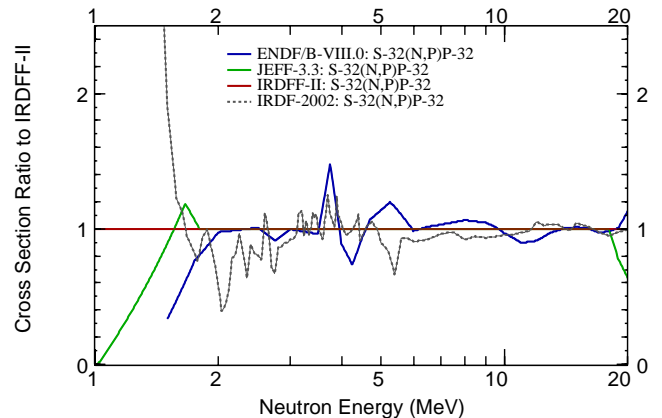
to account for absorption of the beta particles by the bulk sulfur in the irradiated material [80].

The $^{32}\text{S}(n,p)^{32}\text{P}$ reaction cross sections and covariances below 20 MeV were evaluated by Zolotarev [81] in 2013. Extension to 60 MeV was done at the IAEA based on TENDL-2013, renormalizing the cross sections at 20 MeV for continuity.

Comparison of the cross sections with experimental data is shown in Fig. 15(a). The figure also shows the comparison between IRDF-2002 and IRDFF-II, which remained unchanged from IRDFF-v1.05. Cross sections in different evaluated libraries differ significantly near threshold as shown in Fig. 15(b), where the ratios of the various evaluations to the IRDFF-II recommended cross section are given.



(a) Comparison to selected experimental data from EXFOR [72].

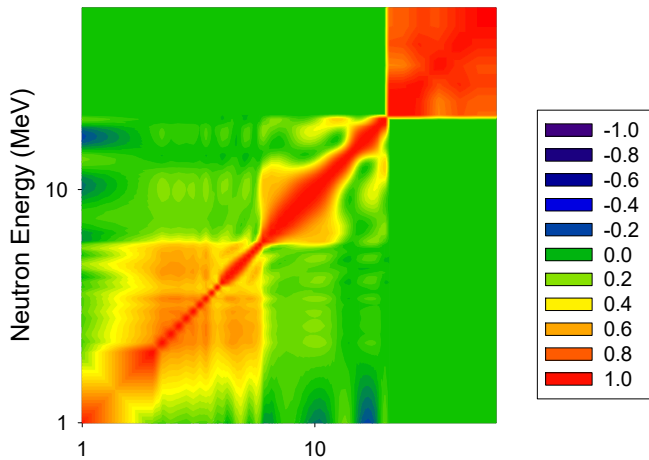


(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

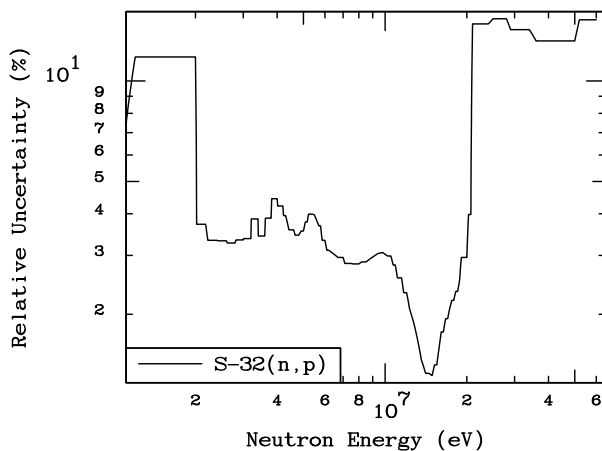
FIG. 15. (Color online) $^{32}\text{S}(n,p)^{32}\text{P}$ cross-section evaluation in IRDFF-II library relative to other data.

The plot of the correlation matrix for the $^{32}\text{S}(n,p)^{32}\text{P}$ reaction is shown in Fig. 16(a). The plot of the 1-sigma uncertainty is shown in Fig. 16(b).

Reactions on the heavier isotopes of sulphur may lead to the same reaction product ^{32}P . In particular, the cross sections for the $^{33}\text{S}(n,np)^{32}\text{P}$ reaction rises sharply above 12 MeV, but the natural abundance of ^{33}S is only 0.75%. However, at energies above 30 MeV the $^{34}\text{S}(n,2np)^{32}\text{P}$ may contribute to the ^{32}P production, so the $^{32}\text{S}(n,p)^{32}\text{P}$ reaction monitor should not be used for high energy applications. The reaction $^{\text{nat}}\text{S}(n,X)^{32}\text{P}$ should be used instead.



(a) Cross-section correlation matrix.



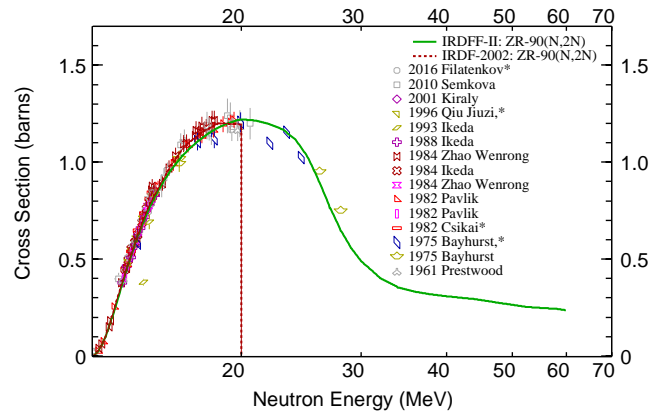
(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

FIG. 16. (Color online) Uncertainties and correlations of the $^{32}\text{S}(n,p)^{32}\text{P}$ cross-section evaluation in IRDF-II library.

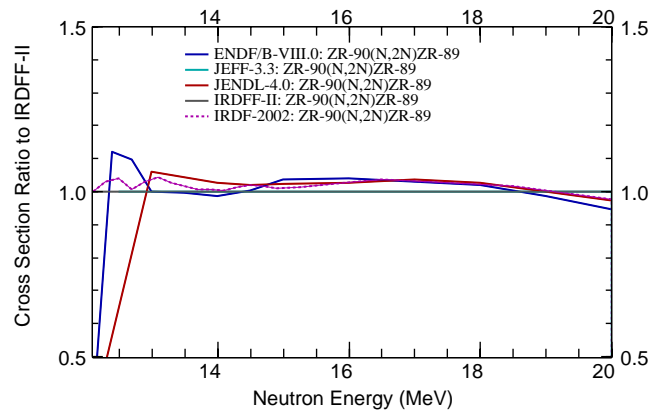
8. $^{90}\text{Zr}(n,2n)^{89}\text{Zr}$

The $^{90}\text{Zr}(n,2n)^{89}\text{Zr}$ reaction cross sections and covariances below 20 MeV were evaluated by Zolotarev [83] in 2009. Extension to 60 MeV was done at the IAEA based on TENDL-2013, renormalizing the cross sections at 20 MeV for continuity. Comparison of the cross sections with experimental data is shown in Fig. 17(a). The figure also shows the comparison between IRDF-2002 and IRDF-II, which remained unchanged from version IRDF-II-v1.05. Cross sections in different evaluated libraries differ significantly near threshold as shown in Fig. 17(b), where the ratios of the various evaluations to the IRDF-II recommended cross section are given. These differences may be important if this reaction monitor is used in reactor spectra.

The plot of the correlation matrix for the $^{90}\text{Zr}(n,2n)^{89}\text{Zr}$ reaction is shown in Fig. 18(a). The plot of the 1-sigma uncertainty is shown in Fig. 18(b). Reactions on the heavier isotopes of Zr may lead to the same reaction product ^{89}Zr . In particular, the cross



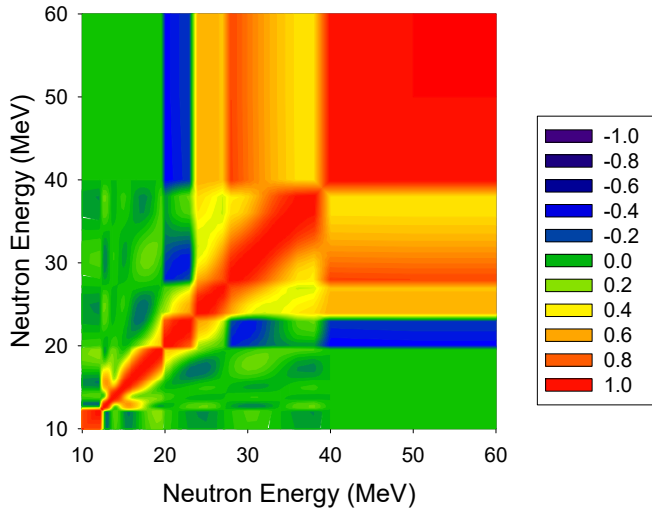
(a) Comparison to selected experimental data from EXFOR [72].



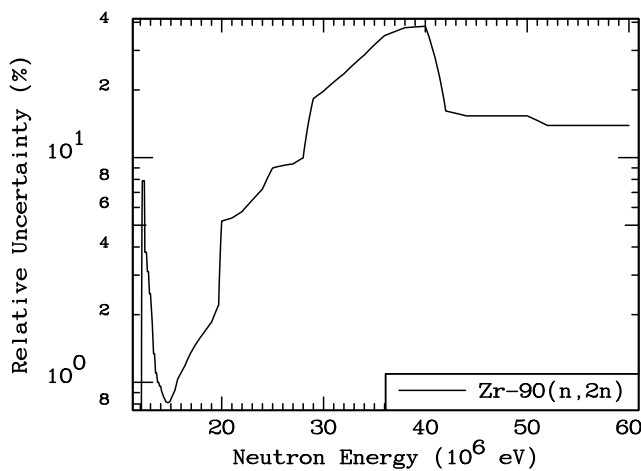
(b) Ratio of cross-section evaluations to IRDF-II evaluation.

FIG. 17. (Color online) $^{90}\text{Zr}(n,2n)^{89}\text{Zr}$ cross-section evaluation in IRDF-II library relative to other data.

sections for the $^{91}\text{Zr}(n,3n)^{89}\text{Zr}$ reaction rises sharply above 20 MeV. The use of the $^{\text{nat}}\text{Zr}(n,X)^{89}\text{Zr}$ reaction cross sections, which are present in the IRDF-II library, is strongly recommended, even though the user should be aware that the minor contributing channels represent a correction and do not meet the dosimetry quality standards, so even the reaction $^{\text{nat}}\text{Zr}(n,X)^{89}\text{Zr}$ should be used with caution in neutron fields with a significant contribution of neutrons with energies above 20 MeV.



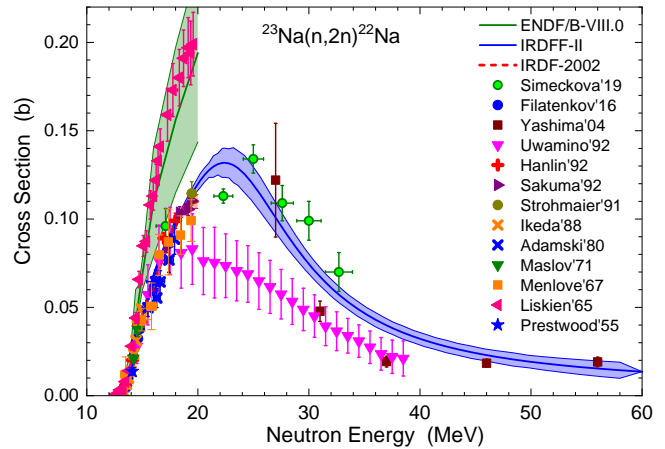
(a) Cross-section correlation matrix.

(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.FIG. 18. (Color online) Uncertainties and correlations of the $^{90}\text{Zr}(n,2n)^{89}\text{Zr}$ cross-section evaluation in IRDF-II library.

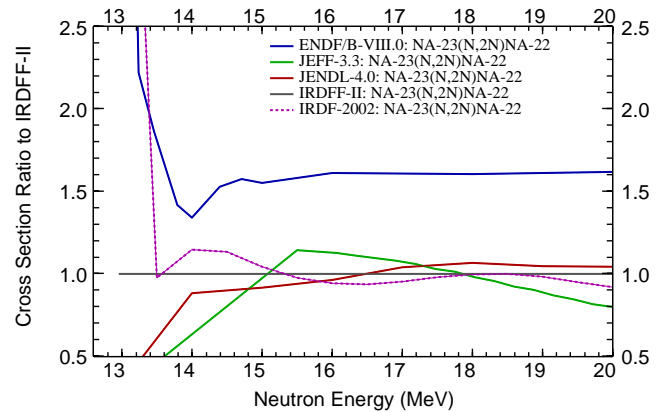
9. $^{23}\text{Na}(n,2n)^{22}\text{Na}$

Sodium is a mono-isotopic naturally occurring element and is convenient for neutron dosimetry because it does not have competing reactions from different isotopes producing the same residual. The $^{23}\text{Na}(n,2n)^{22}\text{Na}$ has a relatively high threshold and the plateau above 20 MeV; it can be used for probing the very high-energy tail of the fission spectrum.

The $^{23}\text{Na}(n,2n)^{22}\text{Na}$ reaction cross sections and covariances in the entire energy range up to 60 MeV were evaluated by Zolotarev [84] in 2017. In 2019 this reaction cross section was measured at Nuclear Physics Institute of the Czech Academy of Sciences employing the p-Li quasi-monoenergetic neutrons with energies between 17 and 33 MeV [85]. As seen in Fig. 19(a), these data confirm the latest Zolotarev evaluation. The figure also shows the comparison between IRDF-2002 and IRDF-II, which remained unchanged from version IRDFF-v1.05. Cross sections in different evaluated libraries differ significantly near threshold as shown in Fig. 19(b), where the ratios



(a) Comparison to selected experimental data from EXFOR [72] and new data from Ref. [85] (Simeckova'19).

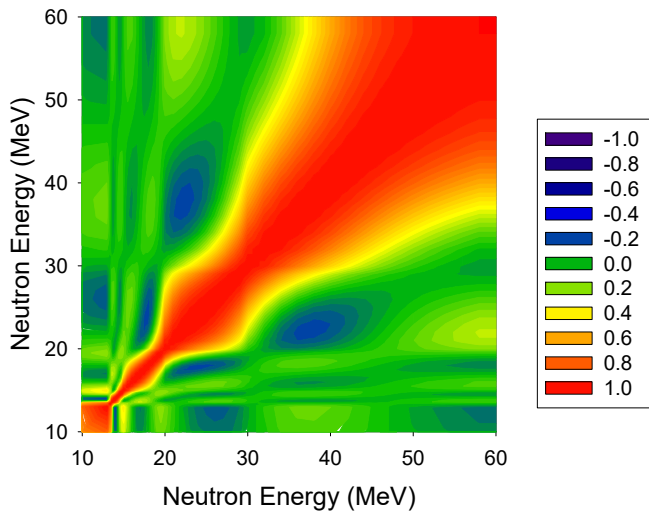


(b) Ratio of cross-section evaluations to IRDF-II evaluation.

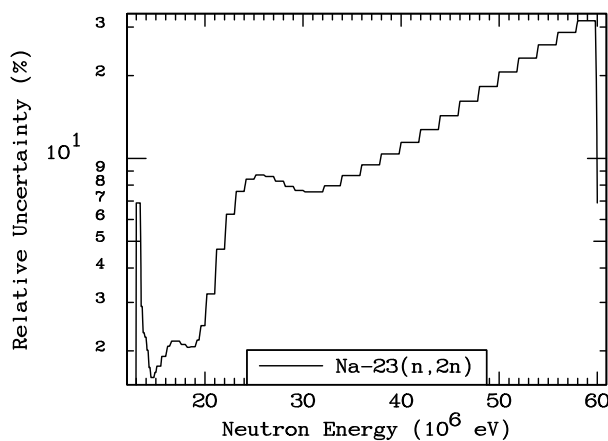
FIG. 19. (Color online) $^{23}\text{Na}(n,2n)^{22}\text{Na}$ cross-section evaluation in IRDF-II library relative to other data.

of the various evaluations to the IRDF-II recommended cross section are given. The ENDF/B-VIII.0 evaluation followed a different set of experimental data. Therefore, differences are observed between evaluations, but also between measurements.

The covariance matrix of the $^{23}\text{Na}(n,2n)^{22}\text{Na}$ reaction cross sections is shown in Fig. 20(a) and the corresponding 1-sigma uncertainty in Fig. 20(b).



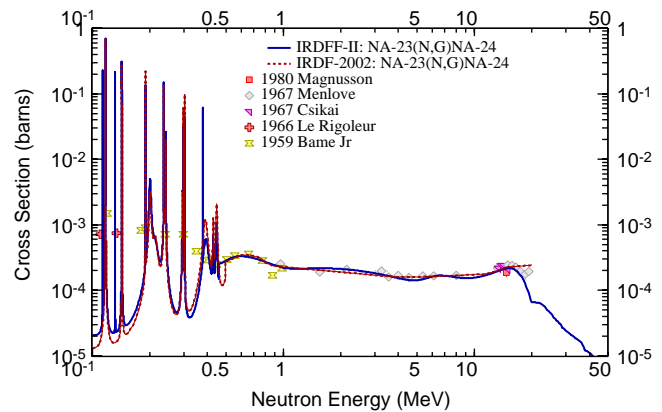
(a) Cross-section correlation matrix.

(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.FIG. 20. (Color online) Uncertainties and correlations of the $^{23}\text{Na}(n,\gamma)^{24}\text{Na}$ cross-section evaluation in IRDF-II library.

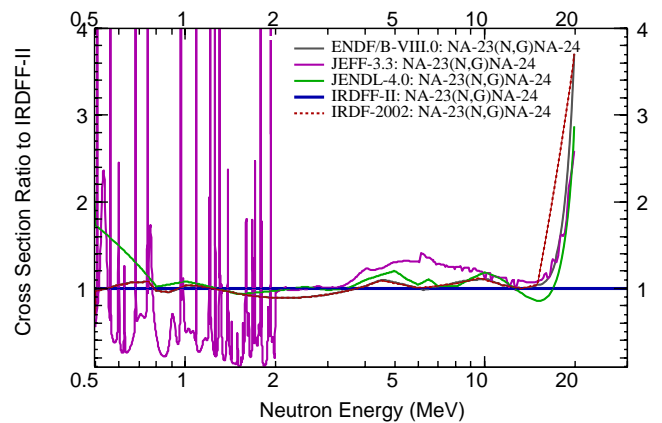
10. $^{23}\text{Na}(n,\gamma)^{24}\text{Na}$

The capture reaction has a strong resonance near 2.8 keV and is highly sensitive to the epithermal energies in reactor spectra. The $^{23}\text{Na}(n,\gamma)^{24}\text{Na}$ reaction resonance parameters, cross sections and covariances up to 20 MeV were also evaluated by Zolotarev [84]. Comparison of the cross sections with experimental data in the fast energy range is shown in Fig. 21(a). The figure also shows the comparison between IRDF-2002 and IRDF-II, which is different from IRDF-*v*1.05. Cross sections in different evaluated libraries differ significantly as shown in Fig. 21(b), where the ratios of the various evaluations to the IRDF-II recommended cross section are given.

The covariance matrix of the $^{23}\text{Na}(n,\gamma)^{24}\text{Na}$ reaction cross sections is shown in Fig. 22(a) and the corresponding 1-sigma uncertainty in Fig. 22(b).



(a) Comparison to selected experimental data from EXFOR [72].



(b) Ratio of cross-section evaluations to IRDF-II evaluation.

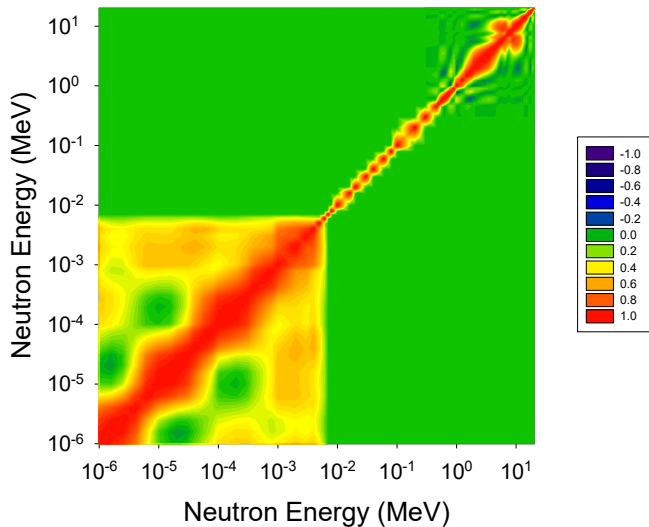
FIG. 21. (Color online) $^{23}\text{Na}(n,\gamma)^{24}\text{Na}$ cross-section evaluation in IRDF-II library relative to other data.

11. $^{24}\text{Mg}(n,p)^{24}\text{Na}$

The $^{24}\text{Mg}(n,p)^{24}\text{Na}$ reaction cross sections and covariances below 21 MeV were evaluated by Zolotarev [79] in 2008. Extension to 60 MeV was done at the IAEA based on TENDL-2010, renormalizing the cross sections at 21 MeV for continuity. Comparison of the cross sections with experimental data is shown in Fig. 23(a). The figure also shows the comparison between IRDF-2002 and IRDF-II, which remained unchanged from version IRDF-*v*1.05. Cross sections in different evaluated libraries differ significantly near threshold as shown in Fig. 23(b), where the ratios of the various evaluations to the IRDF-II recommended cross section are given. These differences may be important if this reaction monitor is used in reactor spectra.

The plot of the correlation matrix for the $^{24}\text{Mg}(n,p)^{24}\text{Na}$ reaction is shown in Fig. 24(a). The plot of the 1-sigma uncertainty is shown in Fig. 24(b).

Reactions on the heavier isotopes of Mg may lead to the same reaction product ^{24}Na . In particular, the cross sections for the $^{25}\text{Mg}(n,np)^{24}\text{Na}$ reaction rises sharply above 18 MeV. The use of the $^{\text{nat}}\text{Mg}(n,X)^{24}\text{Na}$ reaction cross sections, which are present in the IRDF-II library, is strongly recommended, especially for neutron fields that have a significant fraction of neutrons with energies above 18 MeV. However, the user should be aware that the minor



(a) Cross-section correlation matrix.

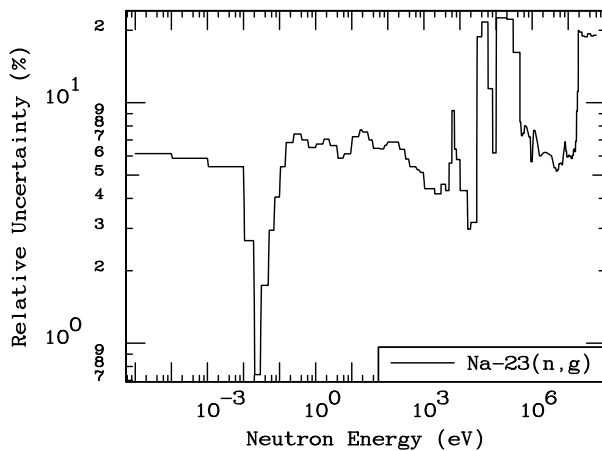
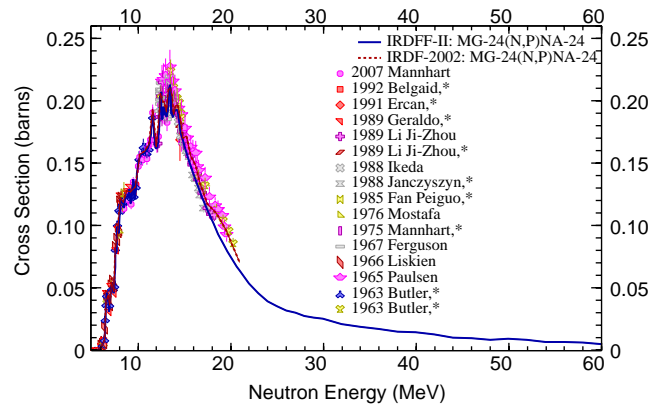
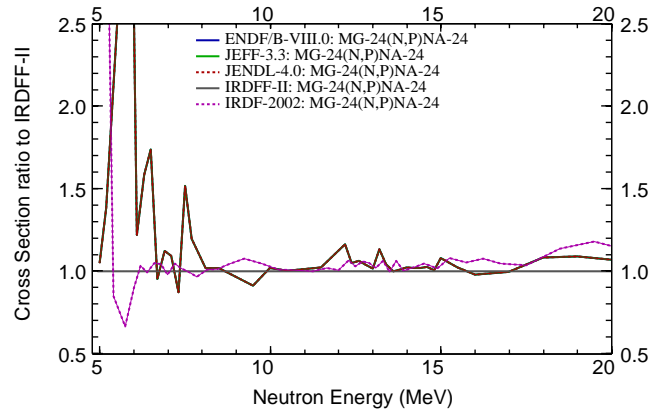

 (b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{cov(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

 FIG. 22. (Color online) Uncertainties and correlations of the $^{23}\text{Na}(n,\gamma)^{24}\text{Na}$ cross-section evaluation in IRDFF-II library.

contributing channels represent a correction and do not meet the same dosimetry quality standards.



(a) Comparison to selected experimental data from EXFOR [72].

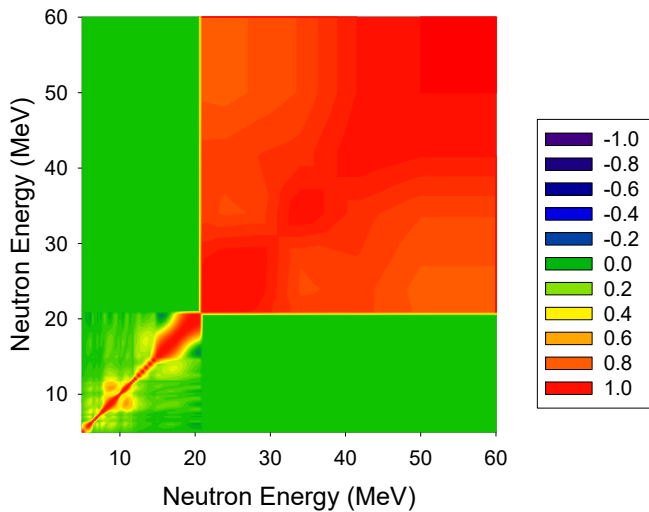


(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

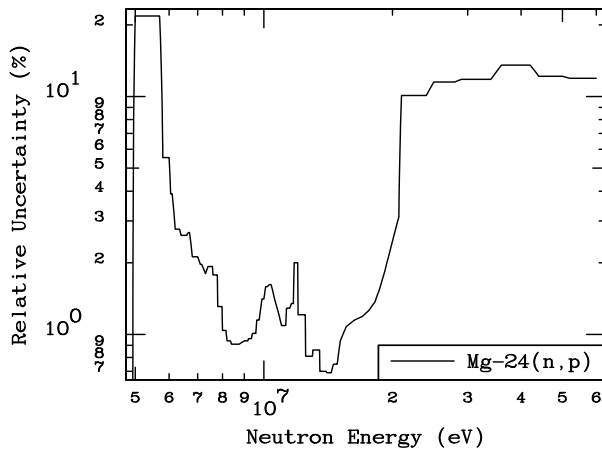
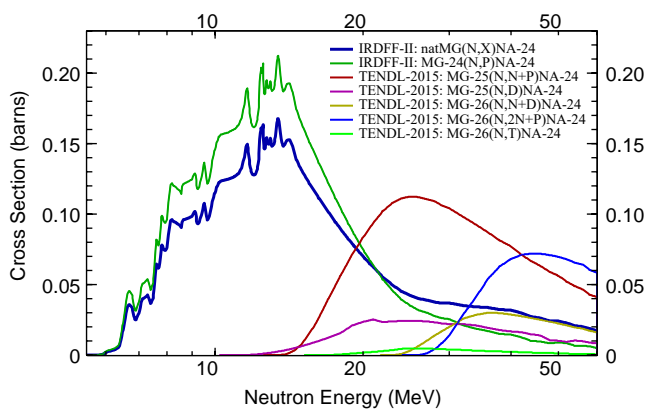
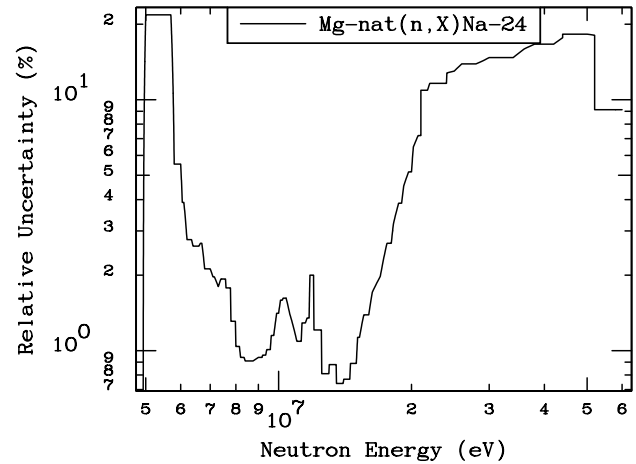
 FIG. 23. (Color online) $^{24}\text{Mg}(n,p)^{24}\text{Na}$ cross-section evaluation in IRDFF-II library relative to other data.

12. $^{nat}\text{Mg}(n,X)^{24}\text{Na}$

The major contribution to the ^{24}Na production comes from the $^{24}\text{Mg}(n,p)^{24}\text{Na}$ reaction, which is present in the IRDFF-II library, supplemented by the contributions of other channels, including $^{25}\text{Mg}(n,np)$, $^{25}\text{Mg}(n,d)$, $^{26}\text{Mg}(n,2np)$, $^{25}\text{Mg}(n,nd)$ and $^{25}\text{Mg}(n,t)$ that are taken from the TENDL-2015 library. The contribution from the $^{25}\text{Mg}(n,np)$ rises sharply above about 18 MeV. The uncertainty and the covariance matrix below this energy are practically the same as for $^{24}\text{Mg}(n,p)^{24}\text{Na}$; see the uncertainties plotted in Fig. 26. It is strongly recommended to use $^{nat}\text{Mg}(n,X)^{24}\text{Na}$ reaction cross sections, especially in neutron fields with a significant fraction of neutrons with energies above 15 MeV.



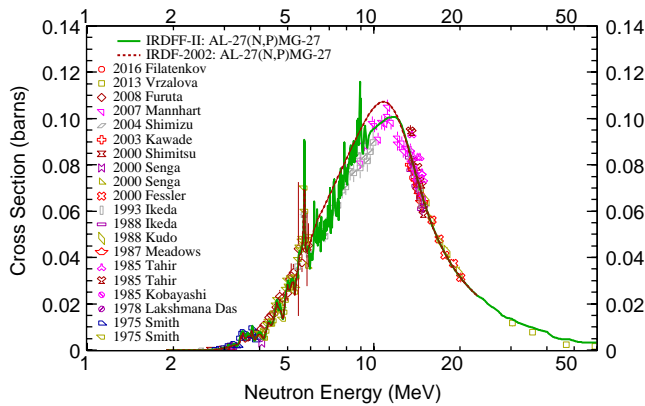
(a) Cross-section correlation matrix.

(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.FIG. 24. (Color online) Uncertainties and correlations of the $^{24}\text{Mg}(n,p)^{24}\text{Na}$ cross-section evaluation in IRDFF-II library.FIG. 25. (Color online) Contributing reactions to the ^{24}Na production from neutrons incident on $^{\text{nat}}\text{Mg}$ target. Note that the cross sections are not scaled by the isotopic abundance.FIG. 26. $^{\text{nat}}\text{Mg}(n,X)^{24}\text{Na}$ cross-section 1-sigma uncertainties.13. $^{27}\text{Al}(n,p)^{27}\text{Mg}$

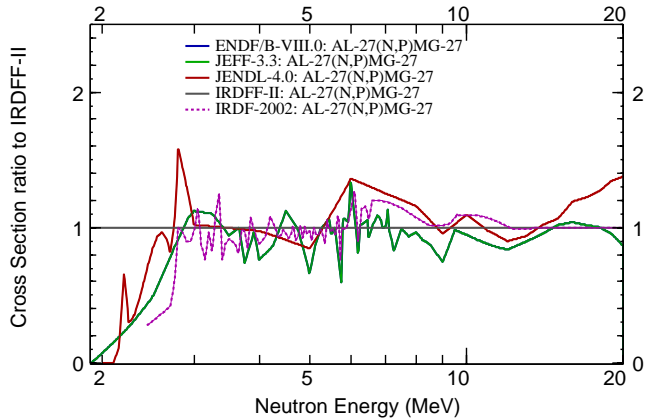
The $^{27}\text{Al}(n,p)^{27}\text{Mg}$ reaction cross sections were evaluated by Zolotarev up to 22 MeV in 2004 [86]. Extension to 60 MeV was done at the IAEA based on TENDL-2010, renormalizing the cross sections at 22 MeV for continuity. Comparison of the cross sections with experimental data is shown in Fig. 27(a). The figure also shows the comparison between IRDF-2002 and IRDFF-II, which remained unchanged from version IRDFF-v1.05.

Cross sections in different evaluated libraries are shown in Fig. 27(b) as ratios of the various evaluations to the IRDFF-II recommended cross section.

The plot of the correlation matrix for the $^{27}\text{Al}(n,p)^{27}\text{Mg}$ reaction is shown in Fig. 28(a). The plot of the 1-sigma uncertainty is shown in Fig. 28(b).



(a) Comparison to selected experimental data from EXFOR [72].



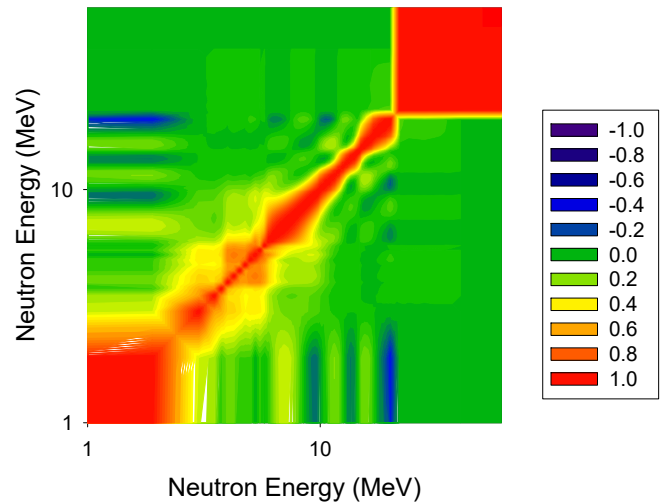
(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

 FIG. 27. (Color online) $^{27}\text{Al}(n,p)^{27}\text{Mg}$ cross-section evaluation in IRDFF-II library relative to other data.

14. $^{27}\text{Al}(n,\alpha)^{24}\text{Na}$

In 2004 the $^{27}\text{Al}(n,\alpha)^{24}\text{Na}$ reaction cross sections were evaluated by Zolotarev up to 40 MeV [83], but the cross sections had unphysical behaviour above 30 MeV. Extension between 30 MeV and 60 MeV was done at the IAEA based on TENDL-2010, renormalizing the cross sections at 30 MeV for continuity. Comparison of the cross sections with experimental data is shown in Fig. 29(a). The figure also shows the comparison between IRDF-2002 and IRDFF-II, which remained unchanged from version IRDFF-v1.05. Cross sections in different evaluated libraries are shown in Fig. 29(b) as ratios of the various evaluations to the IRDFF-II recommended cross section.

The plot of the correlation matrix for the $^{27}\text{Al}(n,\alpha)^{24}\text{Na}$ reaction is shown in Fig. 30(a). The plot of the 1-sigma uncertainty is shown in Fig. 30(b).



(a) Cross-section correlation matrix.

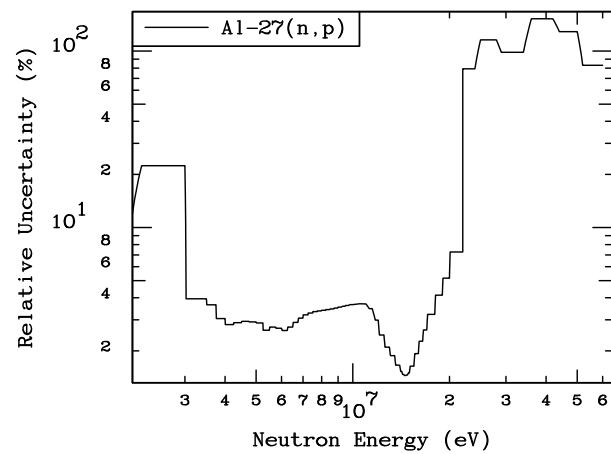
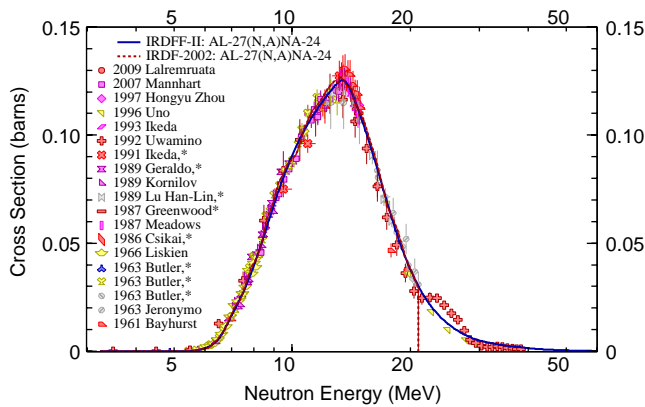
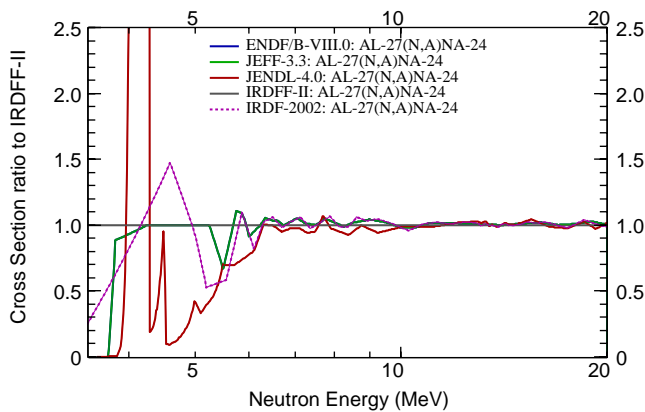

 (b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

 FIG. 28. (Color online) Uncertainties and correlations of the $^{27}\text{Al}(n,p)^{27}\text{Mg}$ cross-section evaluation in IRDFF-II library.

Aluminium is a convenient monitor for neutron dosimetry because it is a mono-isotopic element. However, at energies above 30 MeV there appear competing reactions that produce the same ^{24}Na residual. It is strongly recommended to use $^{27}\text{Al}(n,X)^{24}\text{Na}$ reaction cross sections, which are provided in IRDFF-II, especially in neutron fields with a significant fraction of neutrons with energies above 30 MeV.



(a) Comparison to selected experimental data from EXFOR [72].



(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

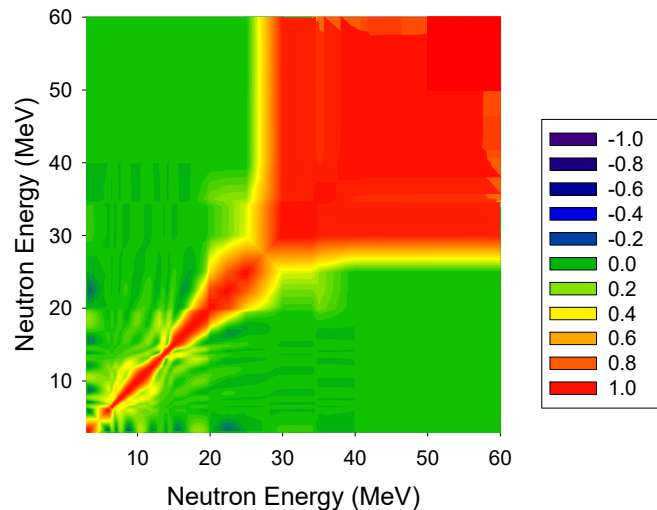
 FIG. 29. (Color online) $^{27}\text{Al}(n,\alpha)^{24}\text{Na}$ cross-section evaluation in IRDFF-II library relative to other data.

15. $^{47}\text{Ti}(n,p)^{47}\text{Sc}$

The $^{47}\text{Ti}(n,p)^{47}\text{Sc}$ reaction cross sections were evaluated by Zolotarev up to 20 MeV in 2002 for the Russian Dosimetry Library RRDF-2002 within the BROND-2 project. Extension to 60 MeV was done at the IAEA based on TENDL-2011, renormalizing the cross sections at 22 MeV for continuity. Comparison of the cross sections with experimental data is shown in Fig. 31(a). The figure also shows the comparison between IRDF-2002 and IRDFF-II, which remained unchanged from version IRDFF-v1.05. Cross sections in different evaluated libraries are shown in Fig. 31(b) as ratios of the various evaluations to the IRDFF-II recommended cross section.

The plot of the correlation matrix for the $^{47}\text{Ti}(n,p)^{47}\text{Sc}$ reaction is shown in Fig. 32(a). The plot of the 1-sigma uncertainty is shown in Fig. 32(b).

Titanium is a monitor elemental material that has a large number of naturally occurring isotopes that lead to the same residual. The $^{47}\text{Ti}(n,p)^{47}\text{Sc}$ reaction is just one example, where the residual ^{47}Sc can be produced from the heavier isotopes in reactions like $^{48}\text{Ti}(n,np)^{47}\text{Sc}$, *etc.* The heavier isotopes begin to contribute at energies around 14 MeV. For this reason, Reactions like $^{\text{nat}}\text{Ti}(n,X)^{47}\text{Sc}$ were introduced into the IRDFF-II library, as illustrated in the section below.



(a) Cross-section correlation matrix.

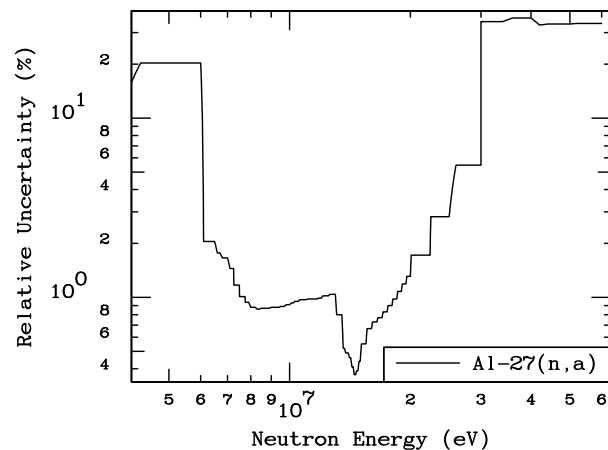
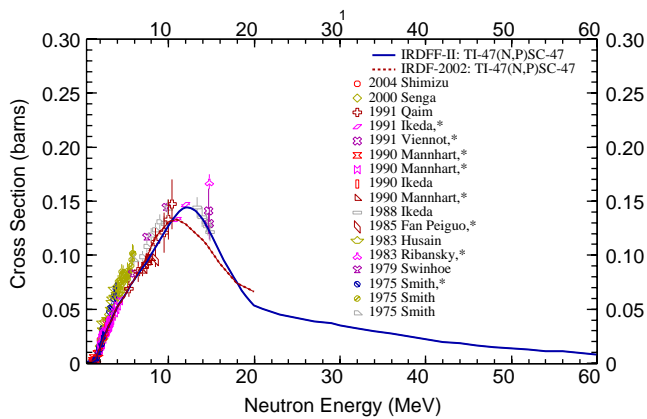

 (b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

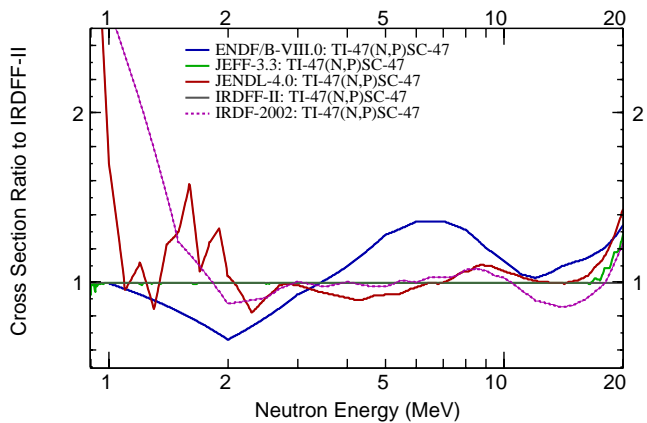
 FIG. 30. (Color online) Uncertainties and correlations of the $^{27}\text{Al}(n,\alpha)^{24}\text{Na}$ cross-section evaluation in IRDFF-II library.

16. $^{\text{nat}}\text{Ti}(n,X)^{47}\text{Sc}$

The major contribution to the ^{47}Sc production comes from the $^{47}\text{Ti}(n,p)^{47}\text{Sc}$ reaction, which is present in the IRDFF-II library, supplemented by the contributions of other channels, including $^{48}\text{Ti}(n,np)$, $^{48}\text{Ti}(n,d)$, $^{49}\text{Ti}(n,2np)$, $^{49}\text{Ti}(n,nd)$, $^{49}\text{Ti}(n,t)$, $^{50}\text{Ti}(n,2nd)$, $^{50}\text{Ti}(n,nt)$ and $^{50}\text{Ti}(n,3np)$ that are taken from the TENDL-2015 library. Relevant cross sections are compared to the available experimental data in Fig. 33. The contribution from the $^{48}\text{Ti}(n,np+pn)$ rises sharply above about 14 MeV. The uncertainty and the covariance matrix below this energy are practically the same as for the $^{47}\text{Ti}(n,p)^{47}\text{Sc}$ reaction. It is strongly recommended to use $^{\text{nat}}\text{Ti}(n,X)^{47}\text{Sc}$ reaction cross section for neutron dosimetry applications, especially in neutron fields with a significant fraction of neutrons with energies above 14 MeV.



(a) Comparison to selected experimental data from EXFOR [72].



(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

 FIG. 31. (Color online) $^{47}\text{Ti}(n,p)^{47}\text{Sc}$ cross-section evaluation in IRDFF-II library relative to other data.

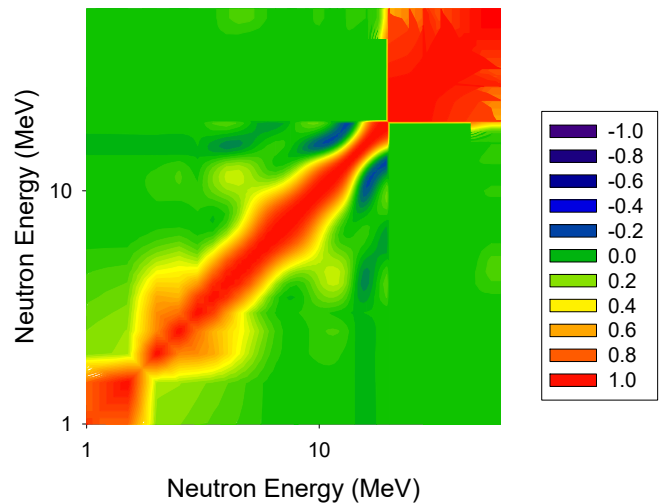
 17. $^{59}\text{Co}(n,2n)^{58}\text{Co}$ and $^{59}\text{Co}(n,3n)^{57}\text{Co}$

The $^{59}\text{Co}(n,2n)^{58}\text{Co}$ reaction cross sections and covariances in energy range up to 20 MeV were evaluated by Zolotarev [83] in 2009. Extension to 60 MeV was done at the IAEA based on TENDL-2010, renormalizing the cross sections at 20 MeV for continuity.

The $^{59}\text{Co}(n,3n)^{57}\text{Co}$ reaction cross sections and covariances in energy range up to 20 MeV were evaluated by Zolotarev [87] in 2010. Extension to 60 MeV was done at the IAEA based on TENDL-2010, renormalizing the cross sections at 20 MeV for continuity. Comparison of the cross sections with experimental data for both reactions is shown in Fig. 34(a). The reaction cross sections were not present in IRDF-2002. In IRDFF-II they remained unchanged from version IRDFF-v1.05. The $^{59}\text{Co}(n,2n)^{58}\text{Co}$ cross sections in different evaluated libraries differ significantly near threshold as shown in Fig. 34(b), where the ratios of the various evaluations to the IRDFF-II recommended cross section are given. The cross sections in JENDL/AD-2017 are significantly lower compared to other evaluations.

The plot of the 1-sigma uncertainty for the $^{59}\text{Co}(n,3n)^{57}\text{Co}$ reaction is shown in Fig. 35.

The plot of the correlation matrix for the $^{59}\text{Co}(n,2n)^{58}\text{Co}$ reaction is shown in Fig. 36(a). The plot



(a) Cross-section correlation matrix.

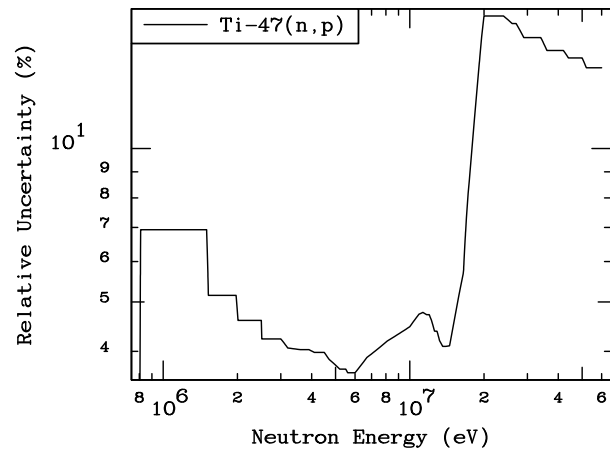
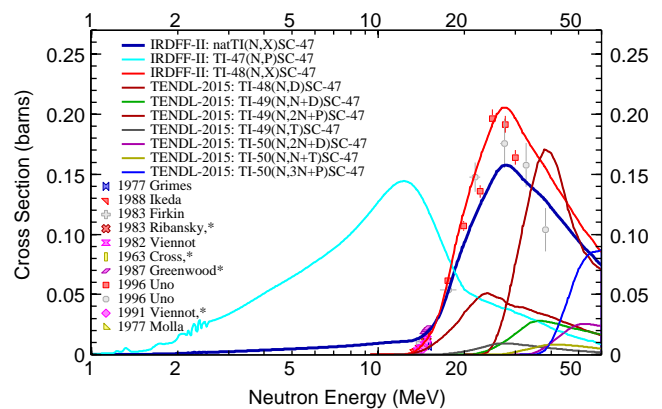
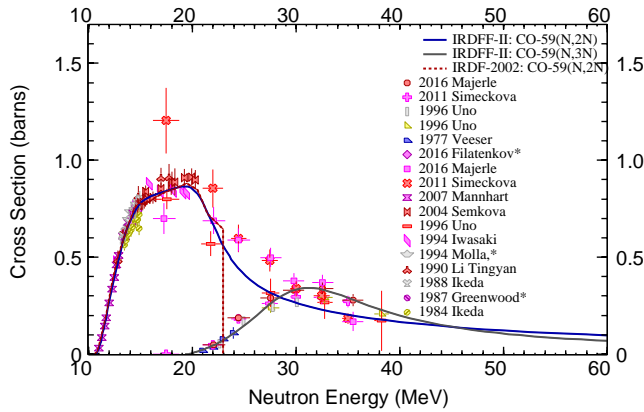
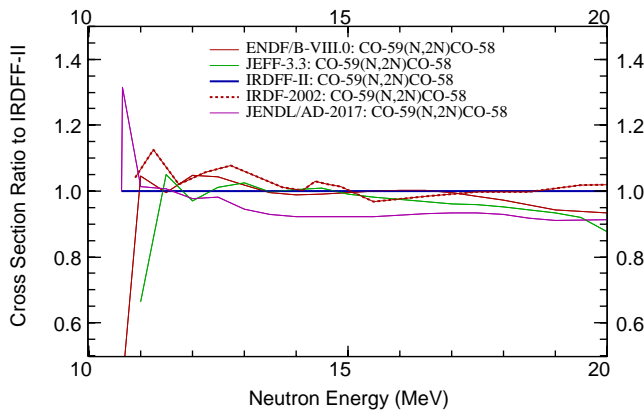

 (b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

 FIG. 32. (Color online) Uncertainties and correlations of the $^{47}\text{Ti}(n,p)^{47}\text{Sc}$ cross-section evaluation in IRDFF-II library.

 FIG. 33. (Color online) Contributing reactions to the ^{47}Sc production from neutrons incident on $^{\text{nat}}\text{Ti}$ target. Note that the cross sections are not scaled by the isotopic abundance.



(a) Comparison to selected experimental data from EXFOR [72].



(b) Ratio of cross-section evaluations to IRDFFF-II evaluation.

FIG. 34. (Color online) $^{59}\text{Co}(n,2n)^{58}\text{Co}$ and $^{59}\text{Co}(n,3n)^{57}\text{Co}$ cross-section evaluations in IRDFFF-II library relative to other data.

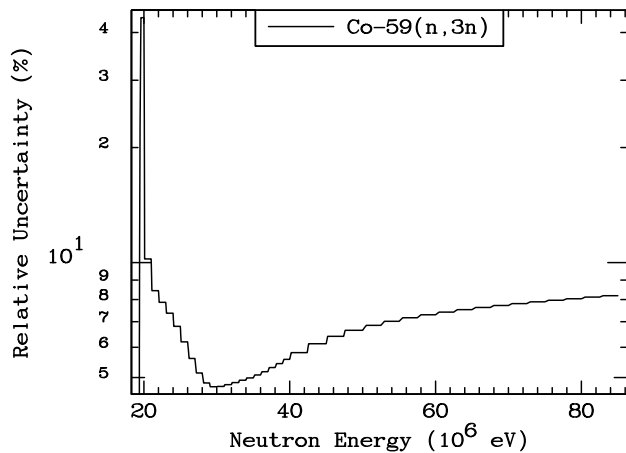
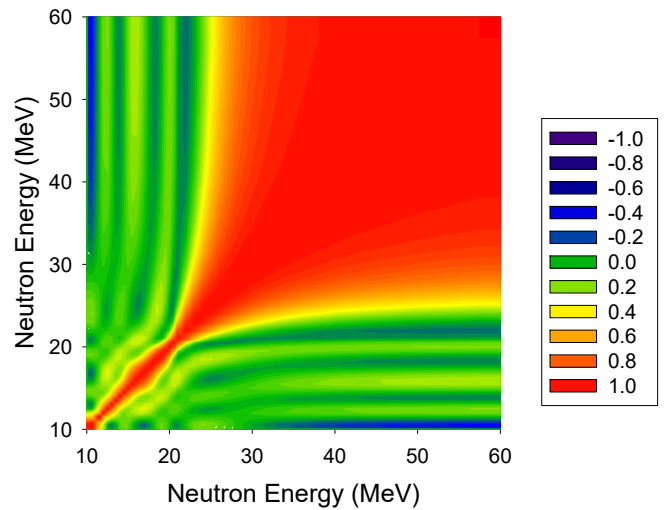


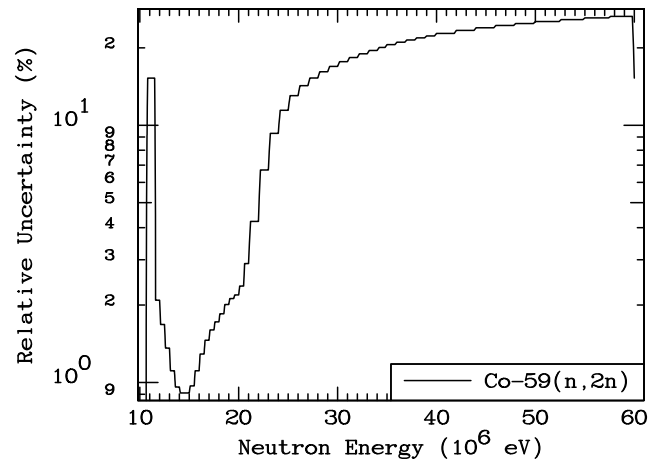
FIG. 35. $^{59}\text{Co}(n,3n)^{57}\text{Co}$ cross-section 1-sigma uncertainties.

of the 1-sigma uncertainty is shown in Fig. 36(b).

The $^{59}\text{Co}(n,2n)^{58}\text{Co}$ and $^{59}\text{Co}(n,3n)^{57}\text{Co}$ reactions are examples of monitor reactions that can be used to probe neutron fields at higher energies.



(a) Cross-section correlation matrix.

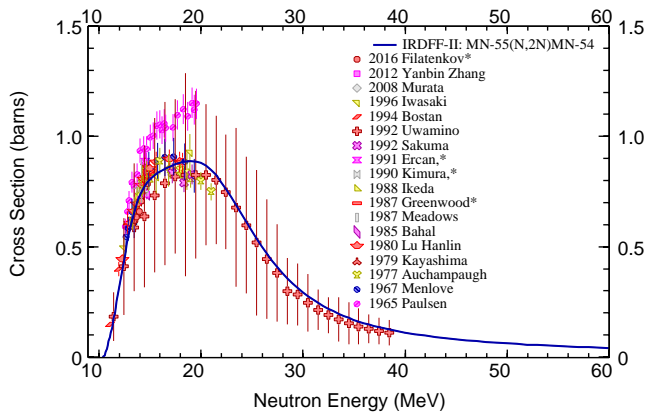


(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

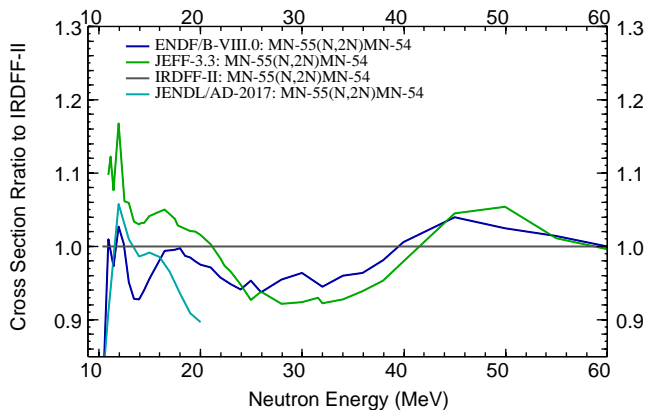
FIG. 36. (Color online) Uncertainties and correlations of the $^{59}\text{Co}(n,2n)^{58}\text{Co}$ cross-section evaluation in IRDFFF-II library.

18. $^{55}\text{Mn}(n,2n)^{54}\text{Mn}$

The $^{55}\text{Mn}(n,2n)^{54}\text{Mn}$ reaction cross sections and covariances in energy range up to 20 MeV were evaluated by Zolotarev [83] in 2009. Extension to 60 MeV was done at the IAEA based on TENDL-2010, renormalizing the cross sections at 20 MeV for continuity. Comparison of the cross sections with experimental data is shown in Fig. 37(a). The reaction cross sections were not present in IRDF-2002. In IRDFF-II they were updated in version IRDFF-v1.05. The $^{55}\text{Mn}(n,2n)^{54}\text{Mn}$ cross sections in different evaluated libraries differ significantly near threshold as shown in Fig. 37(b), where the ratios of the various evaluations to the IRDFF-II recommended cross section are given.



(a) Comparison to selected experimental data from EXFOR [72].

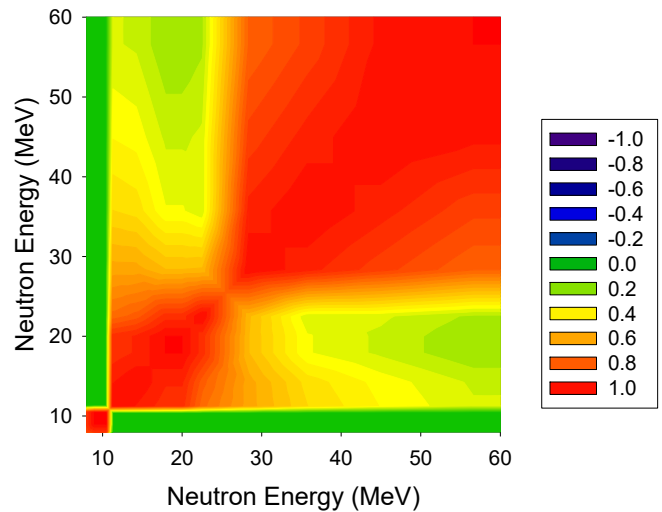


(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

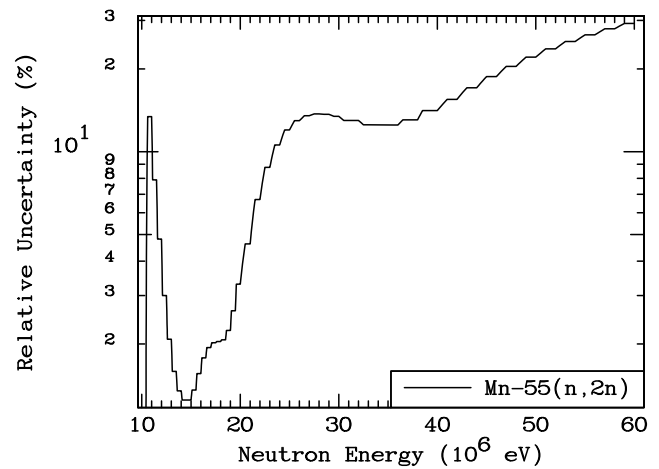
FIG. 37. (Color online) $^{55}\text{Mn}(n,2n)^{54}\text{Mn}$ cross-section evaluation in IRDFF-II library relative to other data.

The plot of the correlation matrix for the $^{55}\text{Mn}(n,2n)^{54}\text{Mn}$ reaction is shown in Fig. 38(a). The plot of the 1-sigma uncertainty is shown in Fig. 38(b).

The $^{55}\text{Mn}(n,2n)^{54}\text{Mn}$ reaction is also one that can be used to probe neutron fields at higher energies.



(a) Cross-section correlation matrix.

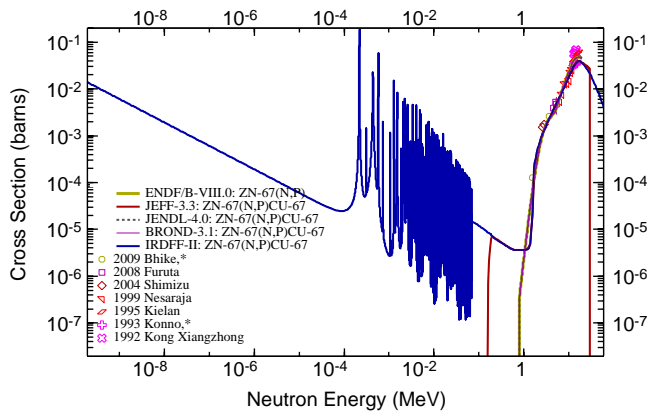


(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

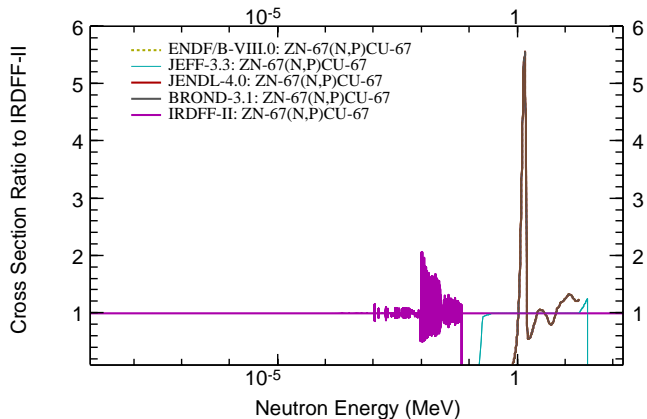
FIG. 38. (Color online) Uncertainties and correlations of the $^{55}\text{Mn}(n,2n)^{54}\text{Mn}$ cross-section evaluation in IRDFF-II library.

19. $^{67}\text{Zn}(n,p)^{67}\text{Cu}$

The $^{67}\text{Zn}(n,p)^{67}\text{Cu}$ reaction cross sections and covariances in energy range up to 20 MeV were evaluated by Zolotarev [79] in 2008. Extension to 60 MeV was done at the IAEA based on TENDL-2010, renormalizing the cross sections at 20 MeV for continuity. This reaction is exothermic and has a positive Q-value, therefore it has no real threshold. However, the difference is the relatively large thermal capture cross section. The capture cross sections from JENDL-4 were re-scaled to give a thermal value of 0.00123 barns and adjusted above 600 keV to join smoothly with the (n,p) cross section evaluation by Zolotarev. A fully correlated uncertainty of 25 % was assigned arbitrarily to the cross sections below the pseudo-threshold. The plot of the cross sections is shown in Fig. 39(a). The cross sections for this reaction were not present in IRDF-2002. In IRDF-II they remained unchanged from version IRDF-v1.05. The $^{67}\text{Zn}(n,p)^{67}\text{Cu}$ cross sections in different evaluated libraries differ significantly, especially near the apparent threshold as shown in Fig. 39(b), where the ratios of the various evaluations to the IRDF-II recommended cross section are given.



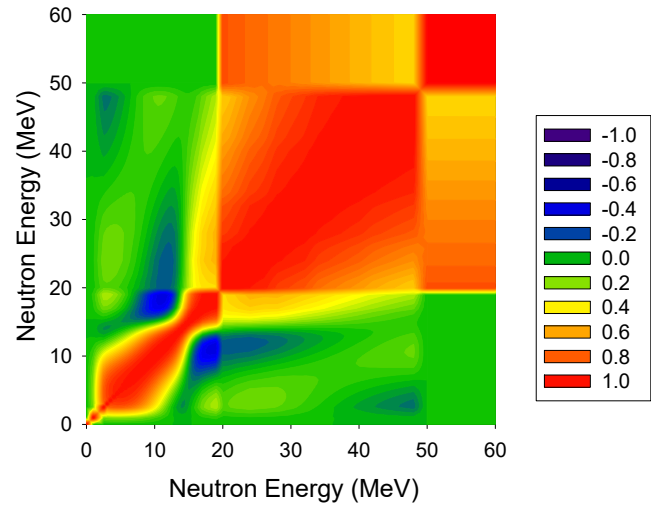
(a) Comparison to selected experimental data from EXFOR [72].



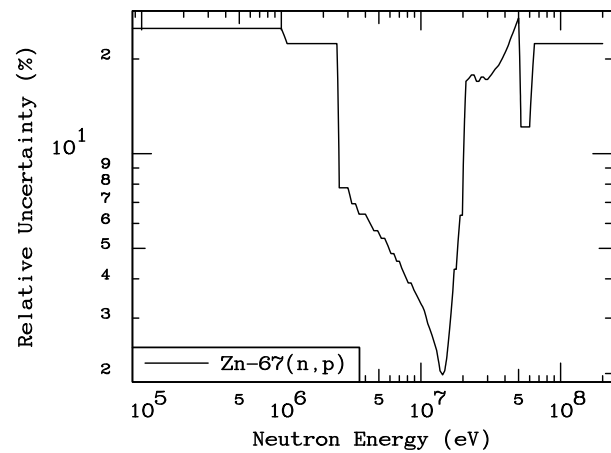
(b) Ratio of cross-section evaluations to IRDF-II evaluation.

FIG. 39. (Color online) $^{67}\text{Zn}(n,p)^{67}\text{Cu}$ cross-section evaluation in IRDF-II library relative to other data.

The plot of the correlation matrix for the $^{67}\text{Zn}(n,p)^{67}\text{Cu}$ reaction is shown in Fig. 40(a). The plot of the 1-sigma uncertainty is shown in Fig. 40(b).



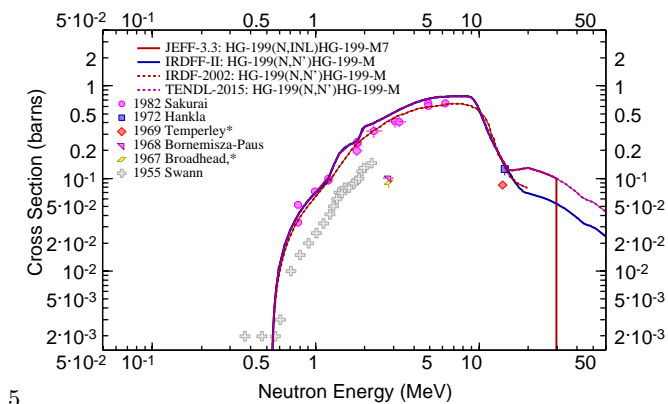
(a) Cross-section correlation matrix.

(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.FIG. 40. (Color online) Uncertainties and correlations of the $^{67}\text{Zn}(n,p)^{67}\text{Cu}$ cross-section evaluation in IRDF-II library.

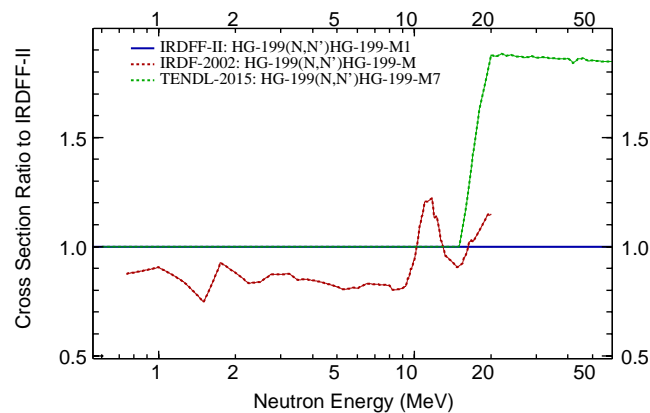
The $^{67}\text{Zn}(n,p)^{67}\text{Cu}$ reaction is an example of a reaction with a positive Q-value, where the cross sections were extended down to thermal energies and a resonance evaluation was included.

20. $^{199}\text{Hg}(n,n')^{199m}\text{Hg}$

The $^{199}\text{Hg}(n,n')^{199m}\text{Hg}$ reaction cross sections and covariances in energy range up to 20 MeV were evaluated by Zolotarev [79] in 2008. Extension to 60 MeV was done at the IAEA based on TENDL-2010, renormalizing the cross sections at 20 MeV for continuity. Comparison with experimental data is shown in Fig. 41(a). The figure also shows the comparison between IRDF-2002 and IRDFF-II, which remained unchanged from version IRDFF-v1.05. The $^{199}\text{Hg}(n,n')^{199m}\text{Hg}$ cross sections represent the excitation of a metastable state and are not present in other major libraries, hence only the comparison with IRDF-2002 is shown in Fig. 41(b), where the ratios to the IRDFF-II recommended cross section are given.



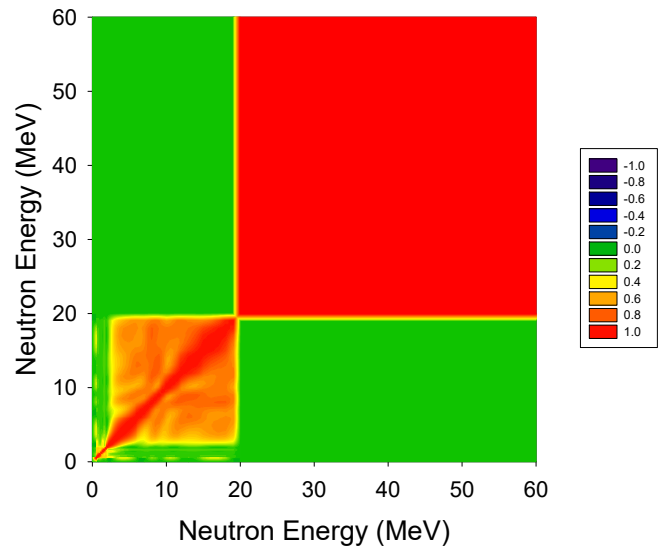
(a) Comparison to selected experimental data from EXFOR [72].



(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

 FIG. 41. (Color online) $^{199}\text{Hg}(n,n')^{199m}\text{Hg}$ cross-section evaluation in IRDFF-II library relative to other data.

The plot of the correlation matrix for the $^{199}\text{Hg}(n,n')^{199m}\text{Hg}$ reaction is shown in Fig. 42(a). The plot of the 1-sigma uncertainty is shown in Fig. 42(b).



(a) Cross-section correlation matrix.

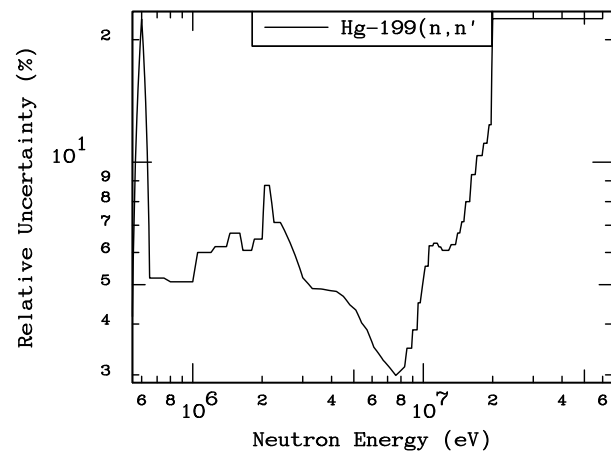
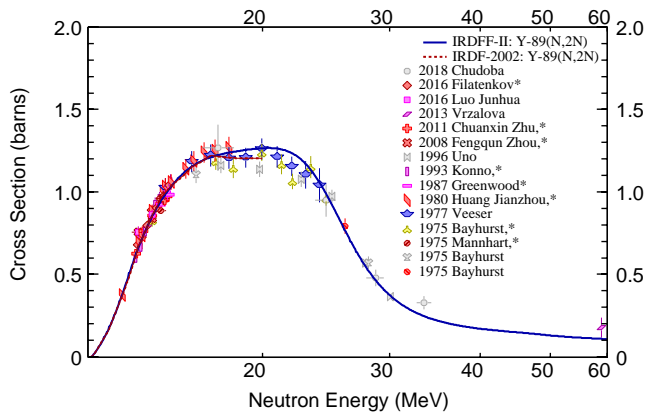

 (b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

 FIG. 42. (Color online) Uncertainties and correlations of the $^{199}\text{Hg}(n,n')^{199m}\text{Hg}$ cross-section evaluation in IRDFF-II library.

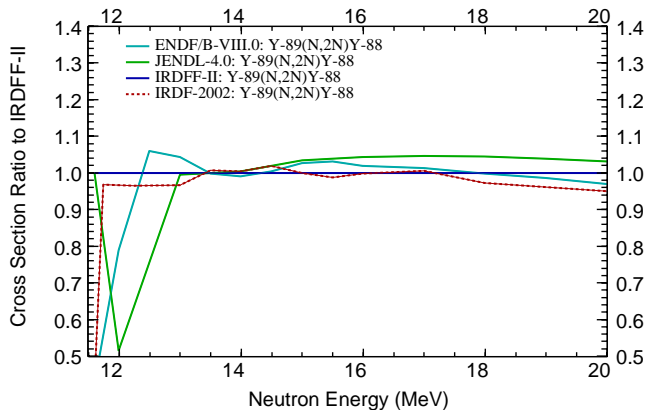
21. $^{89}\text{Y}(n,2n)^{88}\text{Y}$

The $^{89}\text{Y}(n,2n)^{88}\text{Y}$ reaction cross sections and covariances in energy range up to 20 MeV were evaluated by Zolotarev [83] in 2009. Extension to 60 MeV was done at the IAEA based on TENDL-2010, renormalizing the cross sections at 20 MeV for continuity. Comparison of the cross sections with experimental data is shown in Fig. 43(a). The figure also shows the comparison between IRDF-2002 and IRDFF-II, which remained unchanged from version IRDFF-v1.05.

The $^{89}\text{Y}(n,2n)^{88}\text{Y}$ cross sections in different evaluated libraries differ significantly near threshold as shown in Fig. 43(b), where the ratios of the various evaluations to the IRDFF-II recommended cross section are given.



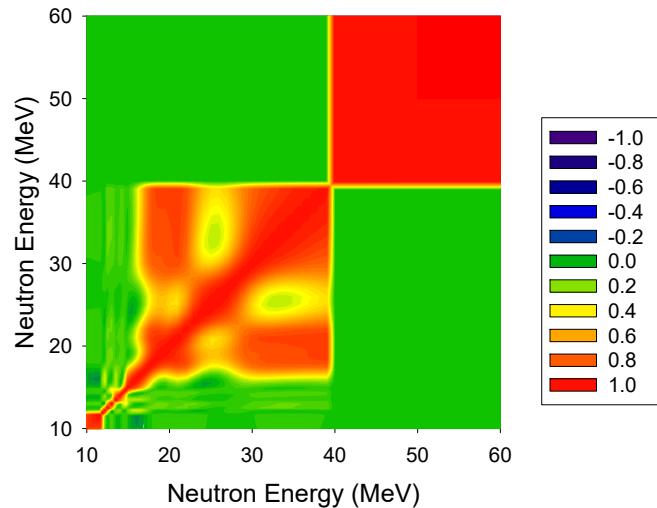
(a) Comparison to selected experimental data from EXFOR [72].



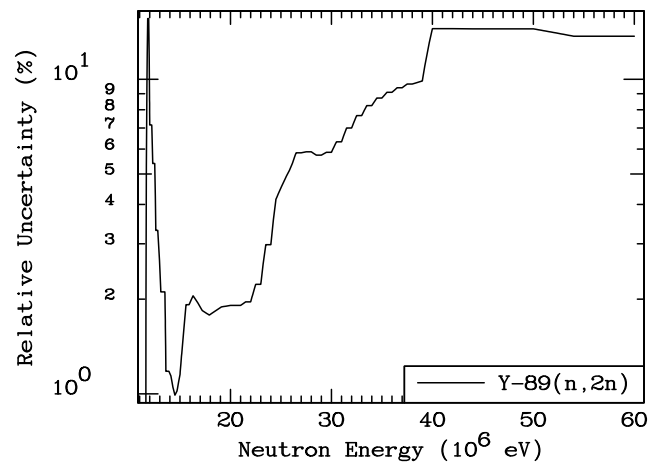
(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

FIG. 43. (Color online) $^{89}\text{Y}(n,2n)^{88}\text{Y}$ cross-section evaluation in IRDFF-II library relative to other data.

The plot of the correlation matrix for the $^{89}\text{Y}(n,2n)^{88}\text{Y}$ reaction is shown in Fig. 44(a). The plot of the 1-sigma uncertainty is shown in Fig. 44(b). The $^{89}\text{Y}(n,2n)^{88}\text{Y}$



(a) Cross-section correlation matrix.



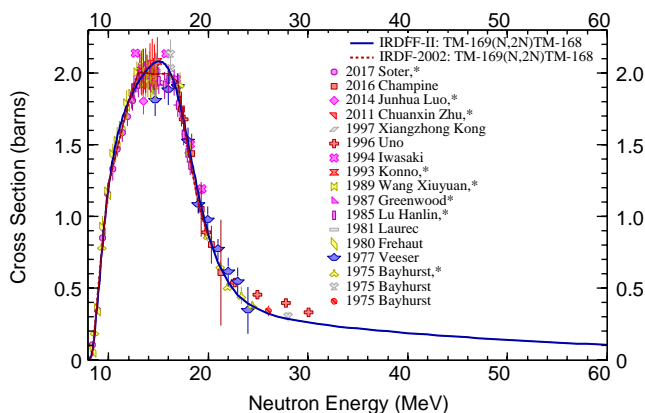
(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

FIG. 44. (Color online) Uncertainties and correlations of the $^{89}\text{Y}(n,2n)^{88}\text{Y}$ cross-section evaluation in IRDFF-II library.

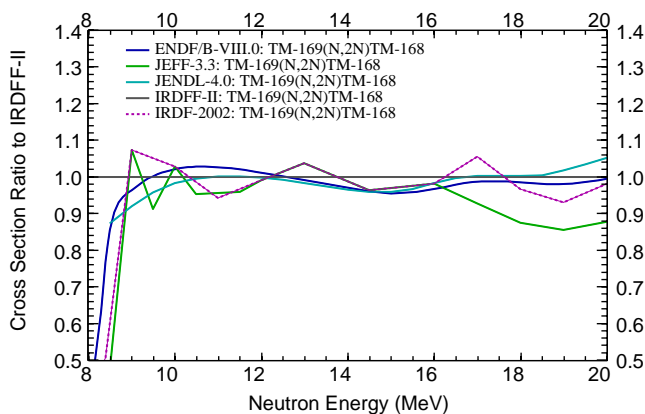
reaction is another one that can be used to probe neutron fields at higher energies.

22. $^{169}\text{Tm}(n,2n)^{168}\text{Tm}$ and $^{169}\text{Tm}(n,3n)^{167}\text{Tm}$

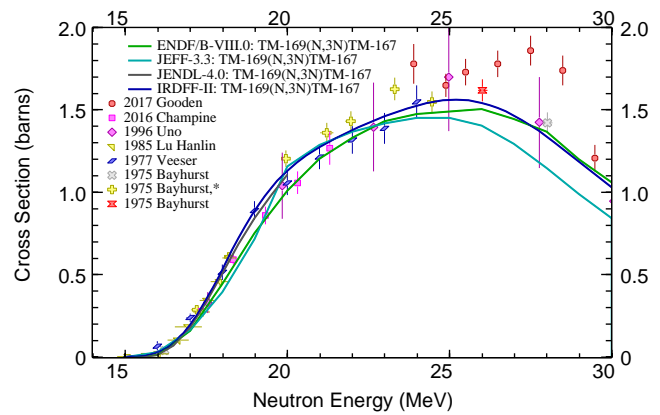
The $^{169}\text{Tm}(n,2n)^{168}\text{Tm}$ reaction cross sections and covariances in energy range up to 40 MeV were evaluated by Zolotarev [87]. Extension to 60 MeV was done at the IAEA based on TENDL-2010, renormalizing the cross sections at 40 MeV for continuity. The $^{169}\text{Tm}(n,3n)^{167}\text{Tm}$ reaction cross sections and covariances in energy range up to 60 MeV were evaluated by Zolotarev [81] in 2003; no extension in energy was necessary. Comparison of the cross sections with experimental data for $^{169}\text{Tm}(n,2n)^{168}\text{Tm}$ and $^{169}\text{Tm}(n,3n)^{167}\text{Tm}$ reactions is shown in Figs. 45(a) and 46(a), respectively. The differences in reaction cross sections for $^{169}\text{Tm}(n,2n)^{168}\text{Tm}$ from IRDF-2002 are shown, which were unchanged in IRDF-2002. The cross sections for $^{169}\text{Tm}(n,3n)^{167}\text{Tm}$ were not present in IRDF-2002; in IRDF-2002 they remained unchanged from version IRDF-v1.05. The $^{169}\text{Tm}(n,2n)^{168}\text{Tm}$ cross sections in different evaluated libraries differ near threshold are shown in Fig. 45(b), where the ratios of the various evaluations to the IRDF-2002 recommended cross section are given. Differences in the $^{169}\text{Tm}(n,3n)^{167}\text{Tm}$ cross sections are even larger. Fig. 46(b) includes experimental data, which tend to support the IRDF-2002 evaluation.



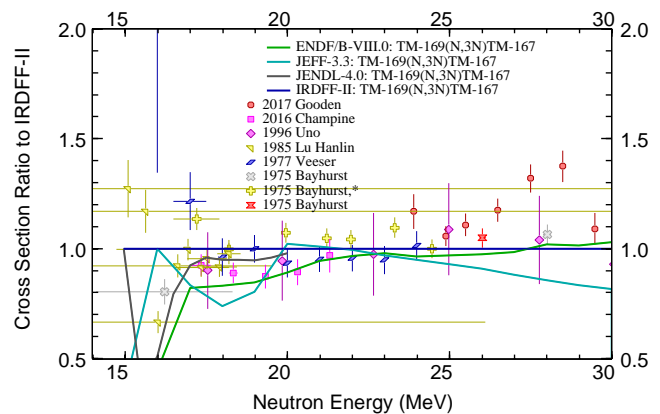
(a) Comparison to selected experimental data from EXFOR [72].



(b) Ratio of cross-section evaluations to IRDF-2002 evaluation.

FIG. 45. (Color online) $^{169}\text{Tm}(n,2n)^{168}\text{Tm}$ cross-section evaluation in IRDF-2002 library relative to other data.

(a) Comparison to selected experimental data from EXFOR [72].



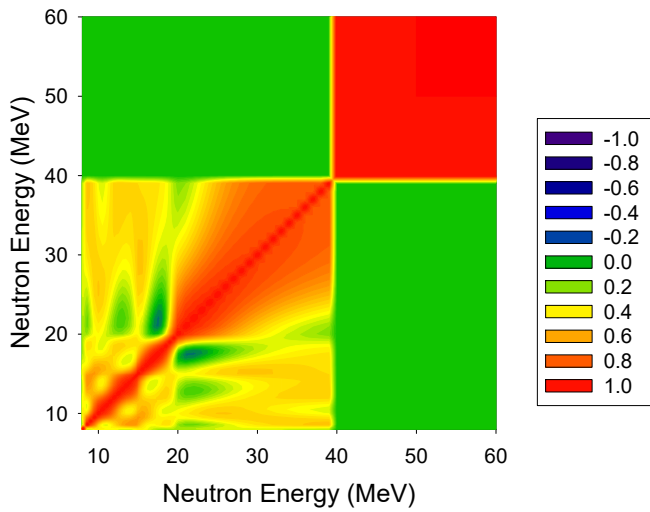
(b) Ratio of cross-section evaluations to IRDF-2002 evaluation.

FIG. 46. (Color online) $^{169}\text{Tm}(n,3n)^{167}\text{Tm}$ cross-section evaluation in IRDF-2002 library relative to other data.

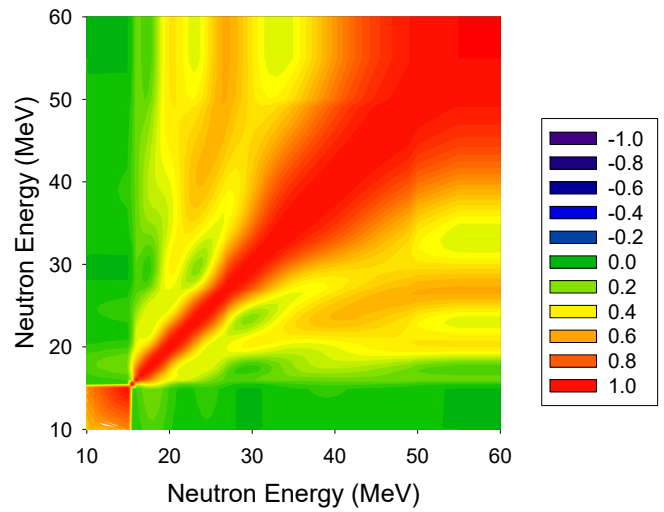
The plot of the correlation matrix for the $^{169}\text{Tm}(n,2n)^{168}\text{Tm}$ reaction is shown in Fig. 47(a). The plot of the 1-sigma uncertainty is shown in Fig. 47(b).

The plot of the correlation matrix for the $^{169}\text{Tm}(n,3n)^{167}\text{Tm}$ reaction is shown in Fig. 48(a). The plot of the 1-sigma uncertainty is shown in Fig. 48(b).

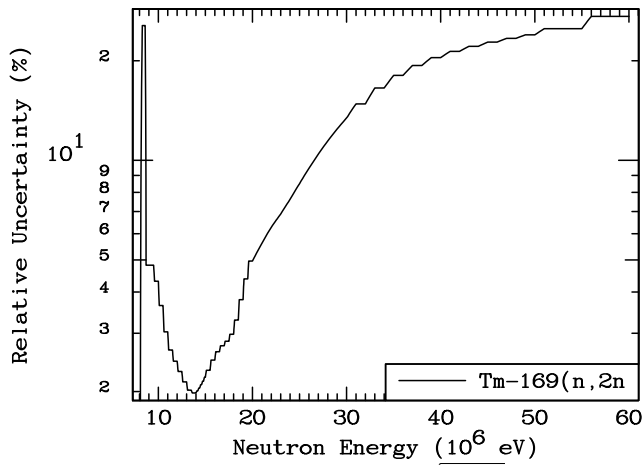
The $^{169}\text{Tm}(n,2n)^{168}\text{Tm}$ and $^{169}\text{Tm}(n,3n)^{167}\text{Tm}$ reactions are examples of monitor reactions that can be used to probe neutron fields at higher energies.



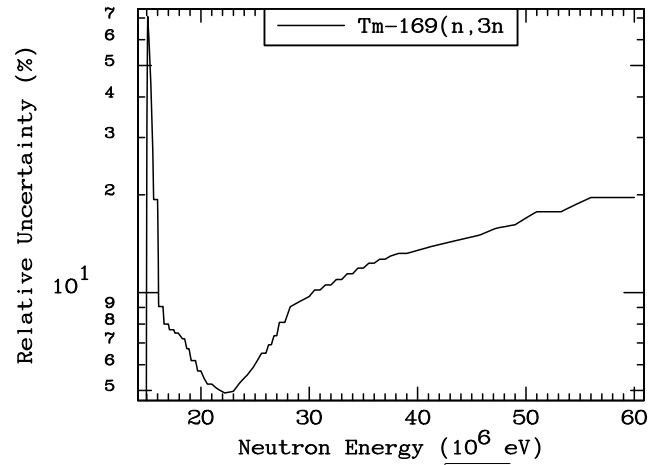
(a) Cross-section correlation matrix.



(a) Cross-section correlation matrix.



(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{cov(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.



(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{cov(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

FIG. 47. (Color online) Uncertainties and correlations of the $^{169}\text{Tm}(n,2n)^{168}\text{Tm}$ cross-section evaluation in IRDFF-II library.

FIG. 48. (Color online) Uncertainties and correlations of the $^{169}\text{Tm}(n,3n)^{167}\text{Tm}$ cross-section evaluation in IRDFF-II library.

23. $^{209}\text{Bi}(n,3n)^{207}\text{Bi}$, $^{209}\text{Bi}(n,4n)^{206}\text{Bi}$, and $^{209}\text{Bi}(n,5n)^{205}\text{Bi}$

The ^{209}Bi reaction cross sections for multiple neutron emission with $n=2-6$ were evaluated up to 100 MeV by Pronyaev in 2013 using a GMA code and smoothed and extended up to 400 MeV by Pronyaev in 2019. Comparison of the cross sections for $^{209}\text{Bi}(n,3n)$, $^{209}\text{Bi}(n,4n)$ and $^{209}\text{Bi}(n,5n)$ reactions with experimental data is shown in Fig. 49. No comparison with other libraries is given, since the evaluated cross sections from other libraries do not extend to energies above 20 MeV which are supported by the new IRDF-II library. The cross sections for these reactions were not present in IRDF-2002.

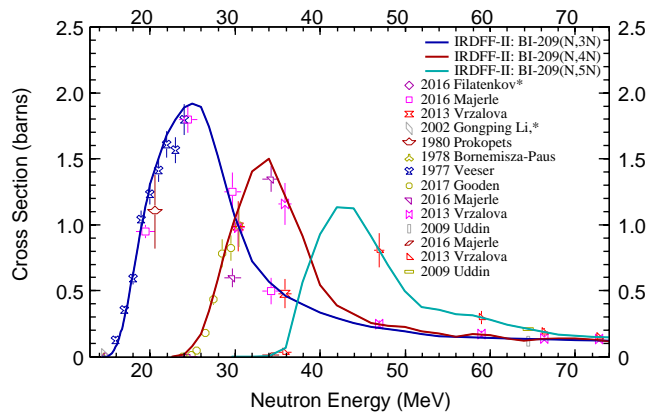
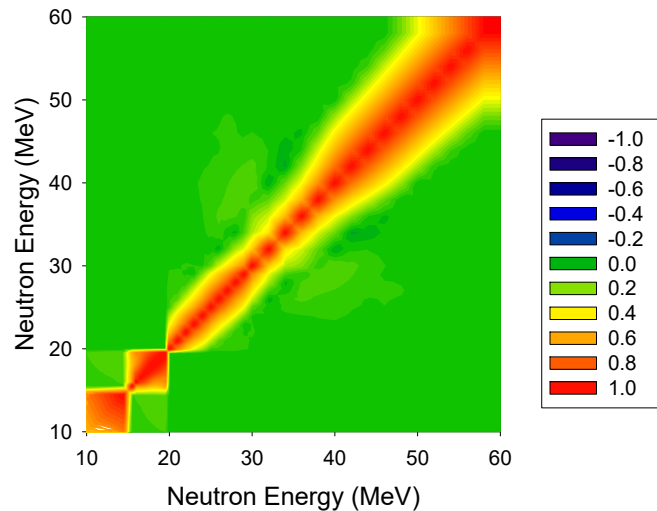


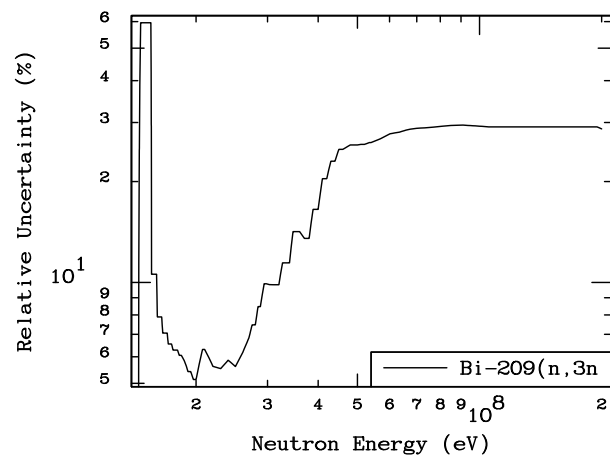
FIG. 49. (Color online) Comparison of $^{209}\text{Bi}(n,xn)$ cross-section evaluations with experimental data from EXFOR.

As an example for these reactions, the plot of the correlation matrix for the $^{209}\text{Bi}(n,3n)^{207}\text{Bi}$ reaction is shown in Fig. 50(a). The plot of the 1-sigma uncertainty is shown in Fig. 50(b).

The $^{209}\text{Bi}(n,xn)$ reactions in the IRDF-II library are examples of monitor reactions that are intended for probing high-energy neutron fields that extend well above 10 MeV up to the maximum energy of 60 MeV.



(a) Cross-section correlation matrix.

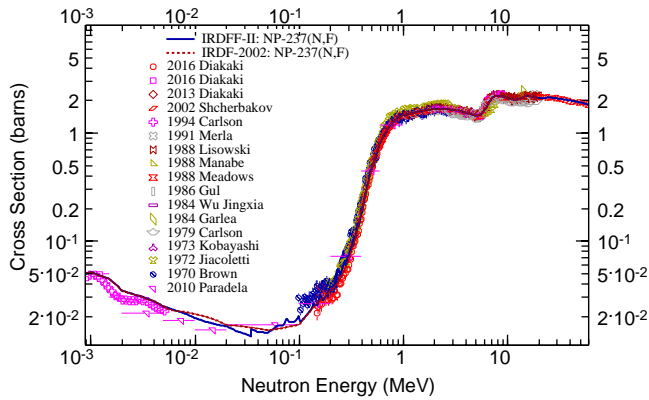


(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

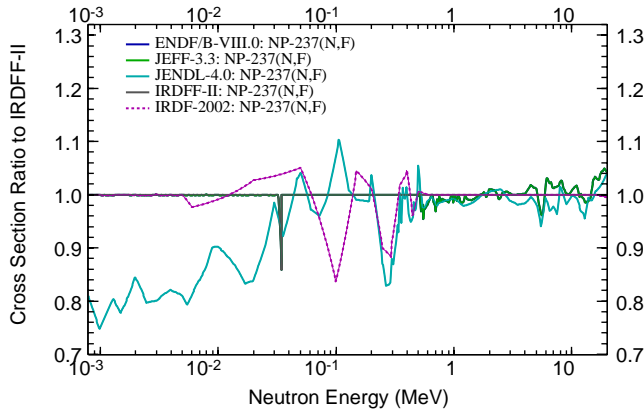
FIG. 50. (Color online) Uncertainties and correlations of the $^{209}\text{Bi}(n,3n)^{207}\text{Bi}$ cross-section evaluation in IRDF-II library.

24. $^{237}\text{Np}(n,f)$

The cross sections for the $^{237}\text{Np}(n,f)$ reaction below 0.5 MeV were taken from the JEFF-3.3 evaluation. In the energy interval 0.5–20 MeV the data were taken from RRDF-2002. Above this energy the JENDL-4/HE evaluation was used, but it does not include the covariances. The difference in the cross sections from TENDL-2015 in 11 subintervals was calculated and the relative difference was added to the diagonal term of the TENDL-2015 covariance matrix in the 20 MeV to 60 MeV energy interval to account for the fact that the original covariances do not correspond to the same cross section evaluation. Comparison of the cross sections with experimental data is shown in Fig. 51(a). The $^{237}\text{Np}(n,f)$ cross sections in different evaluated libraries are shown in Fig. 51(b), where the ratios of the various evaluations to the IRDF-II recommended cross section are given. Eigenvalues of covariances were checked to be positive, but it is worth noting that the quality of the evaluation above 20 MeV should be improved if this reaction will be used at those energies.



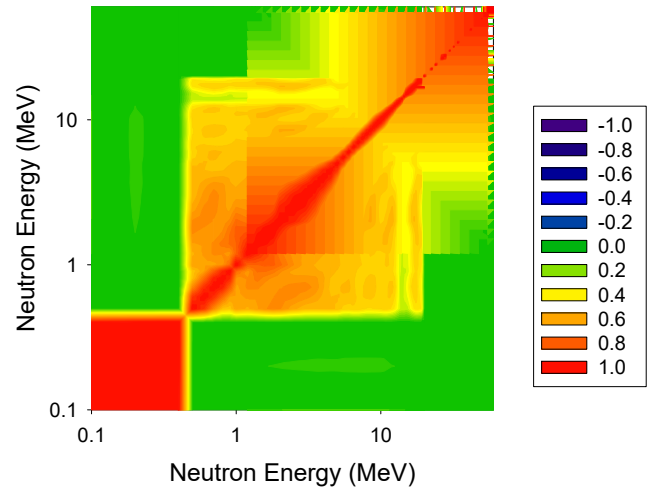
(a) Comparison to selected experimental data from EXFOR [72].



(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

 FIG. 51. (Color online) $^{237}\text{Np}(n,f)$ cross-section evaluation in IRDFF-II library relative to other data.

The plot of the correlation matrix for the $^{237}\text{Np}(n,f)$ reaction is shown in Fig. 52(a). The plot of the 1-sigma uncertainty is shown in Fig. 52(b).



(a) Cross-section correlation matrix.

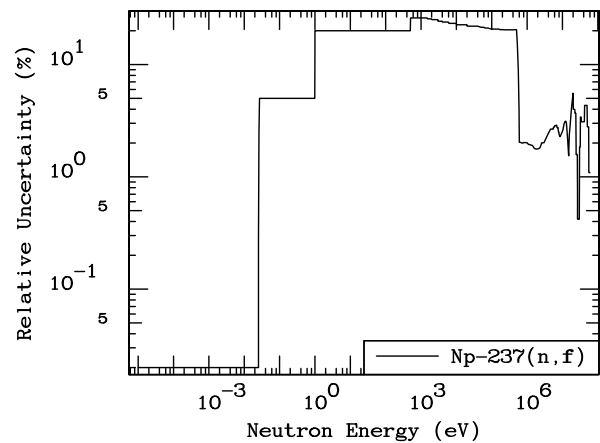
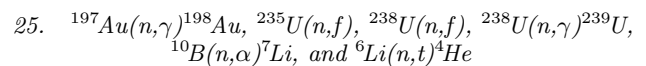
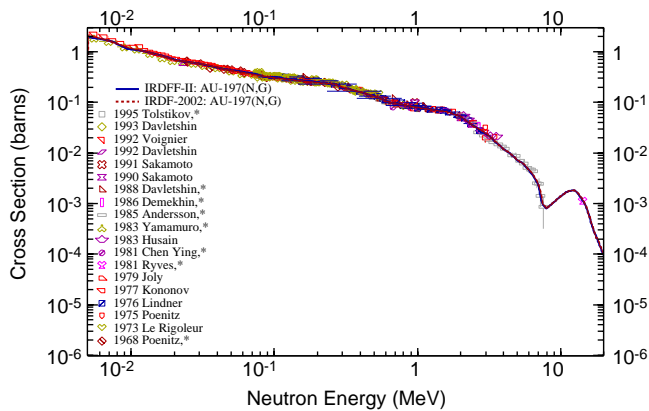

 (b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

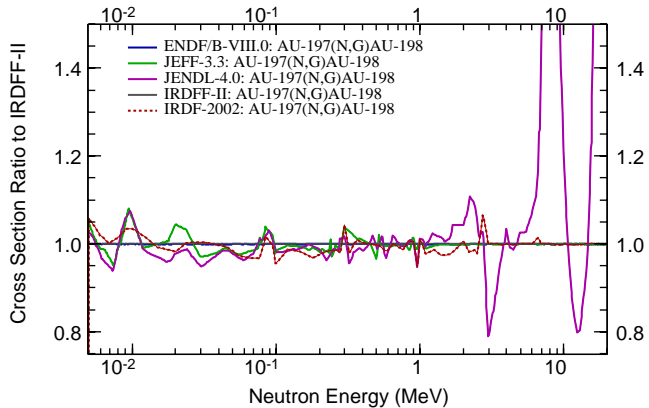
 FIG. 52. (Color online) Uncertainties and correlations of the $^{237}\text{Np}(n,f)$ cross-section evaluation in IRDFF-II library.


The reactions in this section are Neutron Standards (or reaction cross section evaluated together with the Standards) in certain limited energy ranges and are described in detail in a previous publication [32]. What is shown here are examples, where the content of the IRDFF-II library significantly exceeds the energy range of the IAEA Standards 2017.

The $^{197}\text{Au}(n,\gamma)^{198}\text{Au}$ reaction cross sections are compared to experimental data in Fig. 53(a). The capture cross sections of gold are probably the best known capture cross sections over a broad energy range; this is why they are declared a standard at thermal and in the energy range 0.2 MeV to 2.5 MeV [88]. The ENDF/B-VIII.0 evaluation is consistent with the IAEA Standards 2017 for this reaction, hence it was adopted for IRDFF-II. Comparison with other evaluations is shown as ratio to IRDFF-II in Fig. 53(b) for the fast energy range, since the comparison in the resonance range is not meaningful due to the resonances.



(a) Comparison to selected experimental data from EXFOR [72].



(b) Ratio of cross-section evaluations to IRDFF-II evaluation.

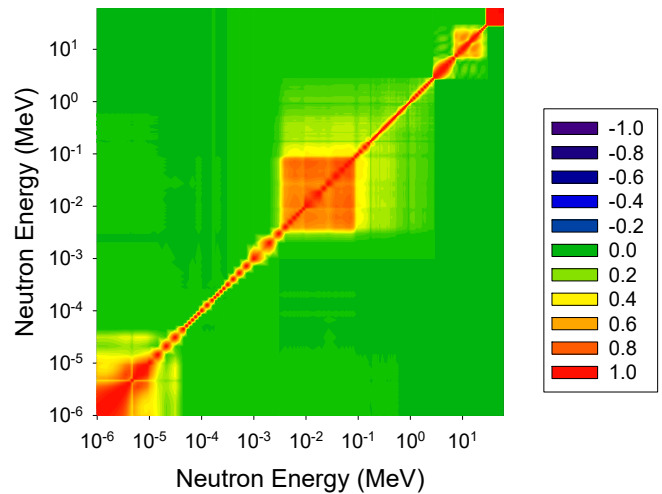
 FIG. 53. (Color online) $^{197}\text{Au}(n,\gamma)^{198}\text{Au}$ cross-section evaluation in IRDFF-II library relative to other data.

The correlation matrix is given in Fig. 54(a) and the corresponding uncertainty is shown in Fig. 54(b).

The tritium and ^4He production reaction $^6\text{Li}(n,t)^4\text{He}$ has also been evaluated for the IAEA Neutron Standards 2017 in the energy range from 0.0253 eV to 1.4 MeV, being the standard energy region from the thermal point up to 1 MeV. The actual evaluation was performed over a broader energy range and is consistent with the contents of ENDF/B-VIII.0 up to 20 MeV. However, by a careful consideration of available experimental evidence it was concluded that the structure in the cross section near 5 MeV that appears in the ENDF/B-VIII.0 is questionable. Note that this region is well outside the Neutron Standards range. At 2.75 MeV and between 8 MeV and 20 MeV the ENDF/B-VIII.0 cross sections for this reaction are equal to those in EAF-2010. To smooth out the structure in the ENDF/B-VIII.0 cross sections the data from EAF-2010 were adopted above 2.75 MeV up to 60 MeV. In this respect IRDFF-II differs from ENDF/B-VIII.0. Comparison of the cross sections for the $^6\text{Li}(n,X)^3\text{H}$ reaction with experimental data and with ENDF/B-VIII.0 and IRDFF-II is shown in Fig. 55, from which the anomalous behaviour of the ENDF/B-VIII.0 cross sections can be seen.

The correlation matrix is given in Fig. 56(a) and the corresponding uncertainty is shown in Fig. 56(b).

The $^{10}\text{B}(n,\alpha)$ reaction cross section was also evaluated as part of the IAEA Neutron Standards 2017 from ther-



(a) Cross-section correlation matrix.

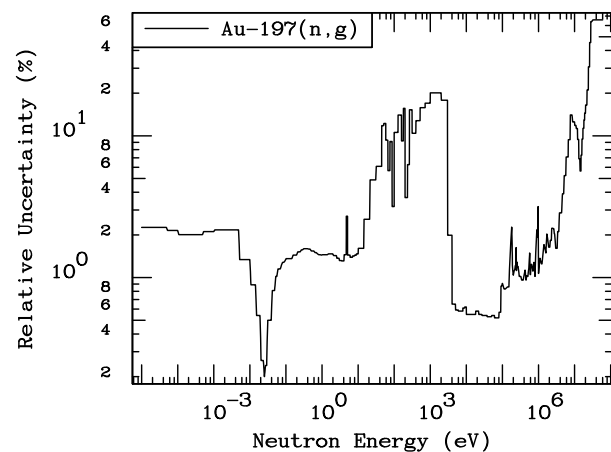
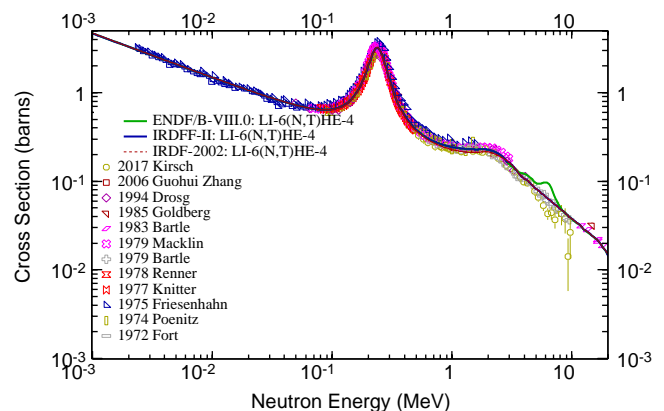
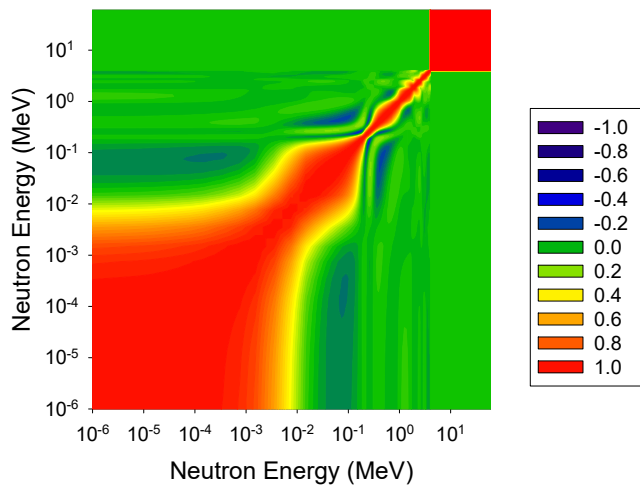
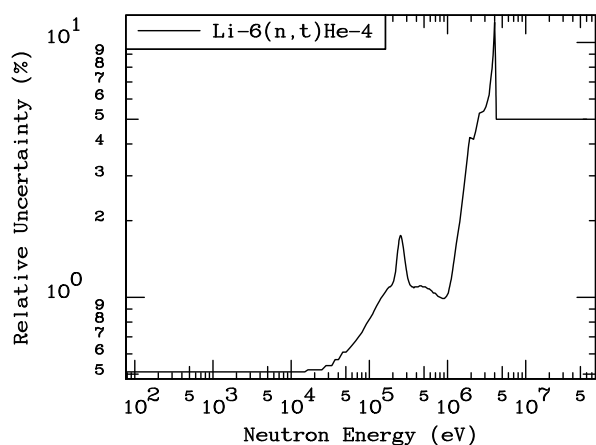

 (b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.

 FIG. 54. (Color online) Uncertainties and correlations of the $^{197}\text{Au}(n,\gamma)^{198}\text{Au}$ cross-section evaluation in IRDFF-II library.

 FIG. 55. (Color online) Comparison of $^6\text{Li}(n,t)^4\text{He}$ cross sections with experimental data from EXFOR.

mal to 1 MeV [32]. Both (n,α) reactions on the ground and the first excited state of the ^{10}B target were evaluated and coded as MT=800 and MT=801 in ENDF-6 format. The ENDF/B-VIII.0 evaluation is consistent with these Standards.



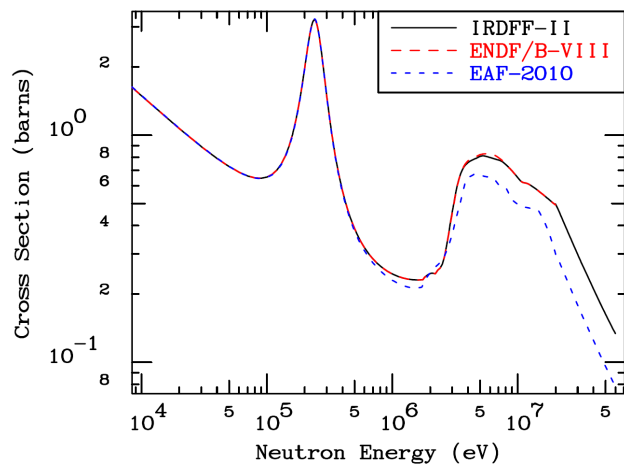
(a) Cross-section correlation matrix.

(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding cross-section mean value.FIG. 56. (Color online) Uncertainties and correlations of the ${}^6\text{Li}(n,t){}^4\text{He}$ cross-section evaluation in IRDF-II library.

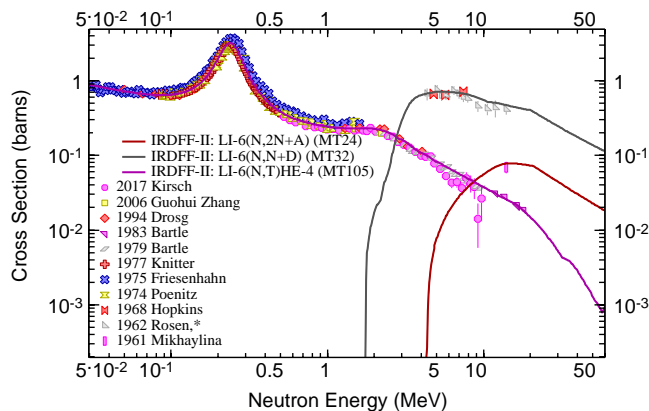
26. ${}^6\text{Li}(n,X){}^3\text{H}$ and ${}^6\text{Li}(n,X){}^4\text{He}$

The gas-production reactions from neutrons incident on light targets were not evaluated separately. Direct tritium production in ${}^6\text{Li}$, with ${}^4\text{He}$ as the residual was evaluated within IAEA Standards 2017 [32] and is described in Sec. VB 25. Other gas-production cross sections were assembled by reviewing existing evaluated data libraries and choosing the most appropriate combination of data, as described below. In particular, the libraries that were considered include ENDF/B-VIII.0, EAF-2010 European Activation File and TENDL-2015 sublibrary "s60", which contains explicit cross section representation with covariances up to 60 MeV. The TENDL-2015.s60 library comes from TALYS nuclear model calculations, which are not best suited for light nuclei, but the library was considered because of its completeness in terms of reactions. However, TENDL-2015 library was found inappropriate for neutron induced reactions on boron and lithium isotopes by extensive comparison with available experimental data and other available evaluations, and will not be discussed in this subsection.

Tritium production from ${}^6\text{Li}$ in IRDF-II corresponds



(a) Comparison of data from selected evaluated libraries relative to the IRDF-II evaluation.

(b) Contribution of different reaction channels to the total ${}^4\text{He}$ gas-production cross-sections in the IRDF-II evaluation.FIG. 57. (Color online) Comparison of the ${}^4\text{He}$ gas-production cross sections for the $n+{}^6\text{Li}$ reaction in the IRDF-II library.

uniquely to the ${}^6\text{Li}(n,t){}^4\text{He}$ reaction (ENDF-6 MT 105), which is a Neutron Standard up to 1 MeV and was discussed in the previous section; a comparison of the cross section with experimental data and with ENDF/B-VIII.0 is shown in Fig. 55.

Cross sections for the ${}^3\text{He}$ production are only available in the TENDL-2015 library, and were neglected for the assembly of the IRDF-II library.

Since the neutron metrology community also uses mass spectrometry methods to measure ${}^4\text{He}$ production in materials such as ${}^6\text{Li}$, the new IRDF-II library also includes this reaction. The ${}^4\text{He}$ nucleus is the residual of the ${}^6\text{Li}(n,t)$ reaction that is discussed in Sec. VB 26. There are other reaction channels that can produce ${}^4\text{He}$, as can be seen in Table 12. As a contributor to the ${}^4\text{He}$ production, all libraries contain the reaction ${}^6\text{Li}(n,2n\alpha)$ (MT 24). The ENDF/B-VIII.0 evaluation was adopted up to 20 MeV, and the EAF-2010 library from 20 up to 60 MeV, renormalized for continuity at 20 MeV. The EAF-2010 library contains reaction ${}^6\text{Li}(n,nd)$ MT 32 explicitly, while in the ENDF/B-VIII.0 library this reaction is represented by discrete level inelastic cross sections ${}^6\text{Li}(n,n)$ of the MT 50 series, with a consecutive breakup of the residual. Note that not all discrete levels contribute to the ${}^4\text{He}$ production, governed by the LR flag

TABLE 12. Gas production reactions in IRDF-2 library from neutrons interacting with ${}^6\text{Li}$ target. The MT numbers (reactions) used in IRDF-2 assembly following the ENDF-6 format, and the relevant reaction products are highlighted in bold.

MT	LR	Gas	Reaction	Products	Source evaluations of IRDF-2	
105		${}^3\text{H}$	${}^6\text{Li}(n,t){}^4\text{He} \rightarrow {}^3\text{H} + {}^4\text{He}$		EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0 ($E < 20$)
24		${}^4\text{He}$	${}^6\text{Li}(n,2n\alpha) \rightarrow 2n + {}^4\text{He} + {}^1\text{H}$		EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0 ($E < 20$)
32			${}^6\text{Li}(n,nd) \rightarrow n + {}^2\text{H} + {}^4\text{He}$		EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0 MT51,52,...
51,52,...	32		${}^6\text{Li}(n,n') \rightarrow n + ({}^6\text{Li} \rightarrow {}^2\text{H} + {}^4\text{He})$			ENDF/B-VIII.0 ($E < 20$)
105			${}^6\text{Li}(n,t){}^4\text{He} \rightarrow {}^3\text{H} + {}^4\text{He}$		EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0 ($E < 20$)

in the ENDF file. The MT 32 contribution reconstructed from the discrete inelastic levels in the ENDF/B VIII.0 library was adopted for IRDF-2, while renormalized EAF-2010 cross sections were adopted from 20 up to 60 MeV. The total ${}^4\text{He}$ production in IRDF-2 is higher than in EAF-2010 library due to the larger cross section from the ${}^6\text{Li}(n,n)$ channels. Fig. 57(a) compares the ${}^4\text{He}$ gas production cross sections in IRDF-2, ENDF/B-VIII.0 and EAF-2010 libraries. Fig. 57(b) shows the relative contribution to the ${}^4\text{He}$ production cross section from the various reaction channels in IRDF-2.

27. ${}^7\text{Li}(n,X){}^3\text{H}$ and ${}^7\text{Li}(n,X){}^4\text{He}$

Since elemental Li is composed of both ${}^6\text{Li}$ and ${}^7\text{Li}$, different enrichments of Li are often used in neutron metrology measurements. Thus, the IRDF-2 library also provides the tritium and ${}^4\text{He}$ gas-production cross sections in ${}^7\text{Li}$. This enables users to combine the cross sections in a manner that represents the composition of their dosimeters.

The list of reactions contributing to tritium production is shown in Table 13. In the EAF-2010 library the tritium production is represented by reaction ${}^7\text{Li}(n,n\alpha)$ MT 22, while in the ENDF/B VIII.0 library this reaction is represented by discrete level inelastic cross sections ${}^7\text{Li}(n,n)$ of the MT 50 series, with a consecutive break-up of the residual from most of the discrete levels into ${}^3\text{H}$ and ${}^4\text{He}$. Reaction MT 33 reconstructed from the discrete inelastic levels from ENDF/B VIII.0 was adopted for IRDF-2, extrapolated above 20 MeV with MT 22 from EAF-2010 since the two reactions produce the same set of emitted particles. The tritium production cross sections are plotted in Fig. 58. The bumps in the cross section near threshold come from the contributions of different discrete levels. The ENDF/B-VIII.0 cross sections are equal to IRDF-2 and are not included in the plot.

Cross sections for the ${}^3\text{He}$ production are only available in the TENDL-2015 library. The reaction threshold is near 30 MeV. The absolute magnitude of the cross sections is about 2 mb at the peak, according to TENDL-2015. Considering that the data from TALYS calculations for the gas production data in light elements in TENDL-2015.s60 are not reliable and there is no alternative source for the ${}^3\text{He}$ production reaction cross sections, these cross sections were not included in the IRDF-2 library.

The list of reactions contributing to ${}^4\text{He}$ production is also shown in Table 13. The libraries EAF-2010 and ENDF/B-VIII.0 contain reactions ${}^7\text{Li}(n,2n\alpha)$ MT 24 and ${}^7\text{Li}(n,\gamma)$ MT 102; the latter is adopted for IRDF-2. The

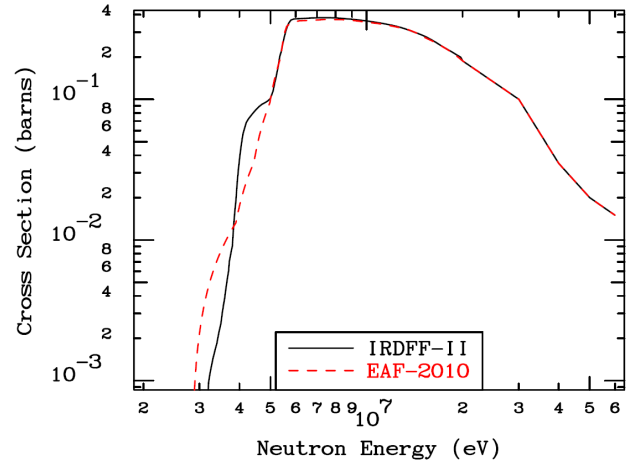


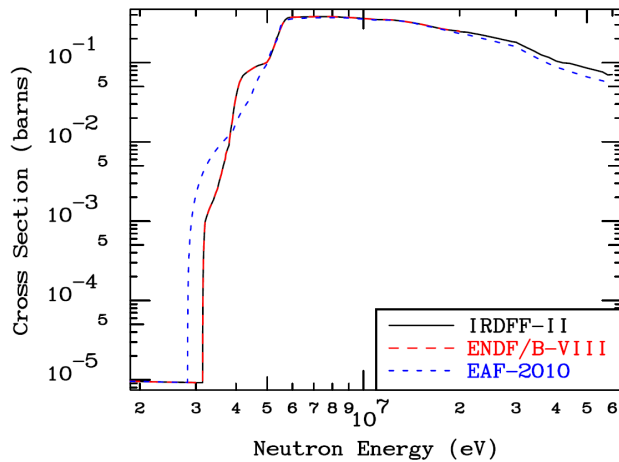
FIG. 58. (Color online) Comparison of the tritium production cross sections for the $n+{}^7\text{Li}$ reaction for different data libraries.

ENDF/B-VIII.0 library contains discrete inelastic levels where the residual is flagged to break up as ${}^7\text{Li}(n,nt)$ MT 33. The summed contributions from the respective levels are stored in IRDF-2 as MT 33. The EAF-2010 library takes a different approach, it contains the ${}^7\text{Li}(n,n\alpha)$ MT 22 reaction, which involves exactly the same outgoing particles. The ENDF/B-VIII.0 library also contains the MT 25 reaction, which has a higher threshold and is not present in EAF-2010. This reaction was adopted from ENDF/B-VIII.0 and extrapolated to 60 MeV with the shape of the ${}^7\text{Li}(n,np)$ reaction, which is mostly equivalent. The contribution of this reaction is mainly responsible for the higher total alpha-production cross section in IRDF-2 compared to EAF-2010. The ${}^4\text{He}$ production cross sections from different libraries are plotted in Fig. 59(a) and the contribution of different reaction channels in IRDF-2 are shown in Fig. 59(b). Note the capture cross section as a contributor to the ${}^4\text{He}$ production at low energies. Two alpha particles are produced after the disintegration of the capture product, hence the ${}^4\text{He}$ production cross section is by a factor of two higher than the capture cross section itself.

There is an issue regarding the ${}^7\text{Li}(n,\gamma)$ MT 102 reaction, which is found in all libraries and is also included in IRDF-2. The residual is ${}^8\text{Li}$, and this nuclide decays into ${}^8\text{Be}$ with a half-life of 0.5 s, which, in turn, breaks up almost instantaneously into two alpha particles. In general, the contribution of this reaction is small, but it is the only contribution to alpha production at thermal energies - a region where the cross section is about 0.1 b - and its multiplicity is two. Beware that the contribution of this reaction is not included in the gas production cross

TABLE 13. Gas production reactions in IRDFF-II library from neutrons interacting with ${}^7\text{Li}$ target. The MT numbers (reactions) used in IRDFF-II assembly following the ENDF-6 format, and the relevant reaction products are highlighted in bold.

MT	LR	Gas	Reaction	Products	Source evaluations of IRDFF-II	
22	33	${}^3\text{H}$	${}^7\text{Li}(n,n\alpha) \rightarrow n + {}^4\text{He} + {}^3\text{H}$	${}^3\text{H}$	EAF-2010(r) ($E > 20$)	
52,53,...			${}^7\text{Li}(n,n') \rightarrow n + ({}^7\text{Li}^* \rightarrow {}^3\text{H} + {}^4\text{He})$	${}^3\text{H}$		ENDF/B-VIII.0 ($E < 20$)
33			${}^7\text{Li}(n,nt) \rightarrow n + {}^3\text{H} + {}^4\text{He}$	${}^3\text{H}$	EAF-2010(r) MT22	ENDF/B-VIII.0 MT 52,53,...
24	33	${}^4\text{He}$	${}^7\text{Li}(n,2n\alpha) \rightarrow 2n + {}^4\text{He} + {}^2\text{H}$	${}^4\text{He}$	EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0 ($E < 20$)
25			${}^7\text{Li}(n,3n\alpha) \rightarrow 3n + {}^4\text{He} + {}^1\text{H}$	${}^4\text{He}$	EAF-2010(r) MT28	ENDF/B-VIII.0 ($E < 20$)
28			${}^7\text{Li}(n,np) \rightarrow 3n + {}^4\text{He} + {}^1\text{H}$	${}^4\text{He}$	EAF-2010(r) ($E > 20$)	
22			${}^7\text{Li}(n,n\alpha) \rightarrow n + {}^4\text{He} + {}^3\text{H}$	${}^4\text{He}$	EAF-2010(r) ($E > 20$)	
52,53,...			${}^7\text{Li}(n,n') \rightarrow n + ({}^7\text{Li}^* \rightarrow {}^3\text{H} + {}^4\text{He})$	${}^4\text{He}$		ENDF/B-VIII.0 ($E < 20$)
33			${}^7\text{Li}(n,nt) \rightarrow n + {}^3\text{H} + {}^4\text{He}$	${}^4\text{He}$	EAF-2010(r) MT22	ENDF/B-VIII.0 MT 52,53,...
102			${}^7\text{Li}(n,\gamma) \rightarrow {}^8\text{Li} \rightarrow (0.8\text{ s}) ({}^8\text{Be} \rightarrow 2\text{ }{}^4\text{He})$	${}^4\text{He}$	EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0 ($E < 20$)



(a) Comparison of data from selected evaluated libraries relative to the IRDFF-II evaluation

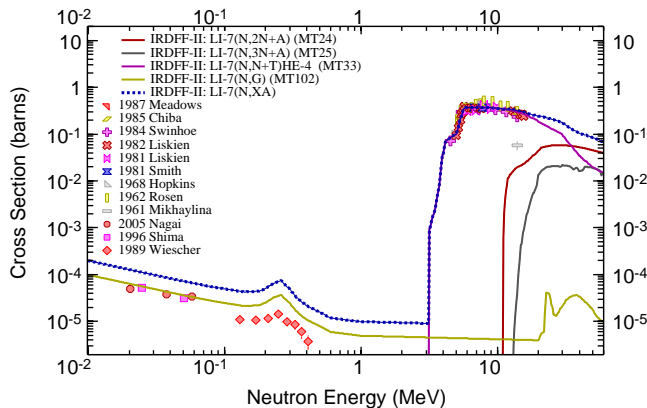
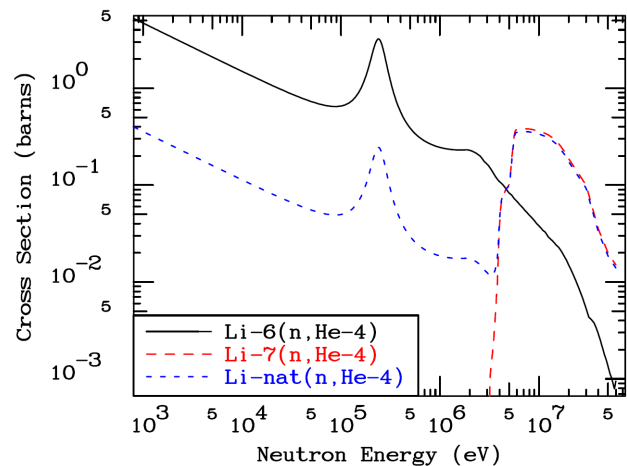

 (b) Contribution of different reaction channels to the total ${}^4\text{He}$ gas-production cross-sections in the IRDFF-II evaluation.

 FIG. 59. (Color online) Comparison of the ${}^4\text{He}$ gas-production cross sections for the $n+{}^7\text{Li}$ reaction in the IRDFF-II library.

sections generated from the ENDF/B-VIII.0 data with the current version of the NJOY-2016 code because the reaction is not flagged for break-up. Due to the very short half-life involved, the radiative capture reaction was included as a contributor to alpha production in the present work for IRDFF-II.

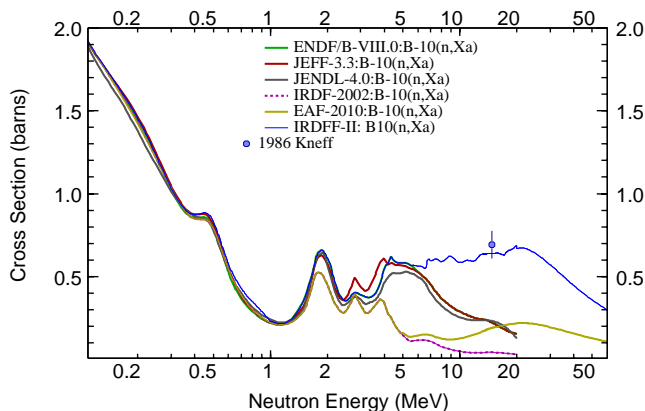
28. ${}^{\text{nat}}\text{Li}(n,X){}^4\text{He}$

In order to support application of the IRDFF-II library to irradiations that use a lithium dosimeter having naturally occurring lithium with nominal abundances for ${}^6\text{Li}$ and ${}^7\text{Li}$, the IRDFF-II library also provides an entry for the ${}^4\text{He}$ gas production on elemental lithium. The cross sections for the gas production in natural element lithium were constructed from the isotopic data in IRDFF-II using the recommended natural abundance data. Fig. 60 compares the ${}^4\text{He}$ gas production on ${}^6\text{Li}$, ${}^7\text{Li}$, and ${}^{\text{nat}}\text{Li}$ targets. The ${}^4\text{He}$ gas production cross sections of ${}^{\text{nat}}\text{Li}$ are normalized per Li atom of the element with an average mass.

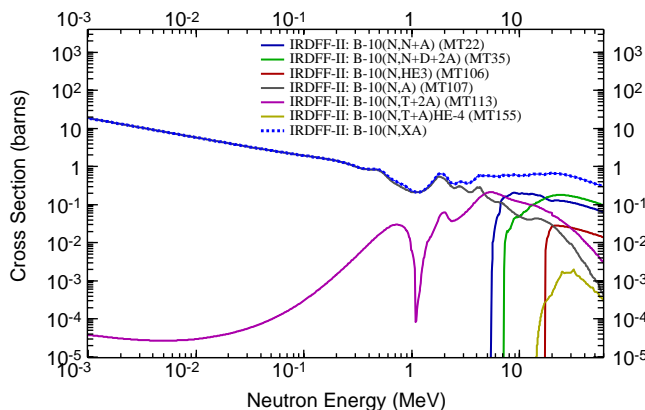

 FIG. 60. (Color online) Comparison of the IRDFF-II ${}^4\text{He}$ gas-production cross sections in neutron induced reactions on ${}^6\text{Li}$, ${}^7\text{Li}$, and ${}^{\text{nat}}\text{Li}$ targets.

29. $^{10}\text{B}(n,X)^4\text{He}$

Gas production in the boron isotopes is very complex because many reactions contribute and very often the residual nuclei undergo further break-up. The complexity of the gas production cross sections in terms of contributing reactions is captured in Table 14. As in the case of Li, the cross sections from the TENDL-2015 library are not considered in the analysis because boron is a light element and TALYS calculations underlying TENDL-2015.s60 are not appropriate in such cases. The discussion in this section is limited to the ^4He production from neutrons incident on ^{10}B , for tritium production relevant contributing reactions are listed in Table 14.



(a) Comparison of data from evaluated libraries relative to Kneff *et al.* measurement [89].



(b) Contribution of different reaction channels to the total ^4He gas-production cross-sections in the IRDFF-II evaluation.

FIG. 61. (Color online) Comparison of the ^4He gas-production cross sections for the $n+^{10}\text{B}$ reaction in the IRDFF-II library.

The $^{10}\text{B}(n,\alpha)$ reaction cross sections are part of the IAEA Neutron Standards 2017 from thermal to 1 MeV. The ENDF/B-VIII.0 evaluation is consistent with these Standards. The library also contains discrete level inelastic cross sections of the MT 50 series, which are flagged for break-up of the residual, equivalent to the reactions under MT 22 and 35. In addition there are also discrete level cross sections for the $^{10}\text{B}(n,t)$ MT 700 and $^{10}\text{B}(n,\alpha)$ MT 800 series, that can be summed into MT 105 and MT 107, respectively. For IRDFF-II the $^{10}\text{B}(n,\alpha)$ MT 107 cross sections are taken from the ENDF/B-VIII.0 library up to 1 MeV because these data agree with the IAEA Neu-

tron Standards 2017. Extrapolation to 60 MeV is done using the shape of the $^{10}\text{B}(n,\alpha)$ MT 107 cross sections from the EAF-2010 library.

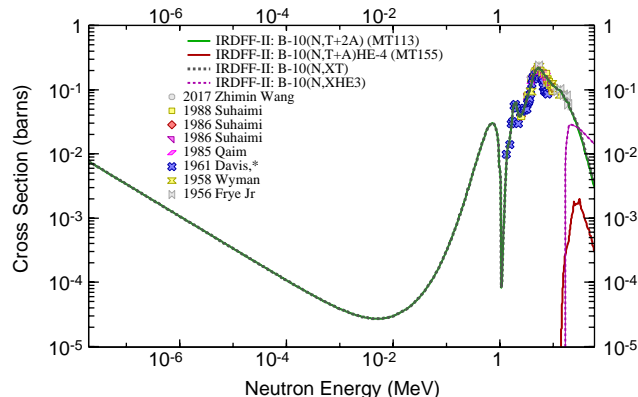


FIG. 62. (Color online) Comparison of tritium-gas production cross sections for the $n+^{10}\text{B}$ reaction for different reaction channels in the IRDFF-II library.

In the IRDFF-II library the $^{10}\text{B}(n,\alpha)$ MT 22 cross sections are taken from the summed contributions of the discrete level inelastic cross sections in ENDF/B-VIII.0 up to 20 MeV. Extrapolation to 60 MeV is done using the shape of the $^{10}\text{B}(n,\alpha)$ MT 22 cross sections from the EAF-2010 library, normalized for continuity at 20 MeV. The data from the ENDF/B-VIII.0 library do not have a dip near 10 MeV, which is observed in the EAF-2010 data.

The cross section curves from the various libraries are shown in Fig. 61(a); plots of selected contributing reactions to ^4He gas production are shown in Fig. 61(b). The plots of contributing reactions to tritium gas production are shown in Fig. 62.

 30. $^{11}\text{B}(n,X)^4\text{He}$

Many reaction channels from neutrons incident on ^{11}B result in ^4He production and/or tritium production, as seen from Table 15. There are 7 such reactions in the EAF-2010 library.

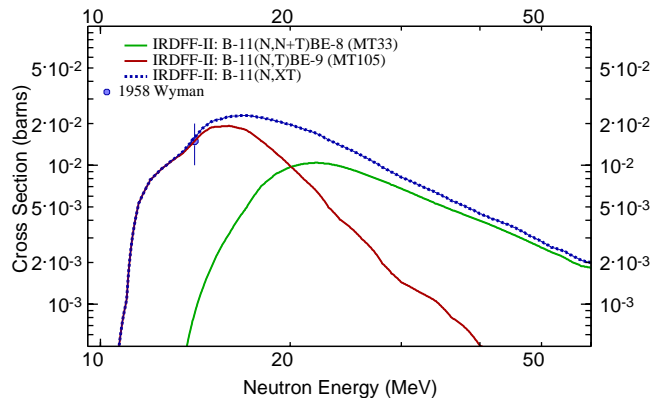


FIG. 63. (Color online) Comparison of tritium gas-production cross sections for the $n+^{11}\text{B}$ reaction for different reaction channels in the IRDFF-II library.

TABLE 14. Gas production reactions from neutrons interacting with ^{10}B target. The MT numbers (reactions) used in IRDF-II assembly following the ENDF-6 format, and the relevant reaction products are highlighted in bold.

MT	LR	Gas	Reaction	Products	Source evaluations of IRDF-II	
105		^3H	$^{10}\text{B}(n,t)$	$\rightarrow ^3\text{H} + (^8\text{Be}^* \rightarrow 2\ ^4\text{He})$	EAF-2010(r) ($E > 20$)	
700			$^{10}\text{B}(n,t)$	$\rightarrow ^3\text{H} + (^8\text{Be}^* \rightarrow 2\ ^4\text{He})$		ENDF/B-VIII.0 ($E < 20$)
113			$^{10}\text{B}(n,t2\alpha)$	$\rightarrow ^3\text{H} + 2\ ^4\text{He}$	EAF-2010(r) MT105	ENDF/B-VIII.0 MT700
155			$^{10}\text{B}(n,t\alpha)$	$\rightarrow ^3\text{H} + 2\ ^4\text{He}$	EAF-2010	
106		^3He	$^{10}\text{B}(n,^3\text{He})$	$\rightarrow ^3\text{He} + (^8\text{Li} \rightarrow ^8\text{Be}^* \rightarrow 2\ ^4\text{He})$	EAF-2010	
22		^4He	$^{10}\text{B}(n,n\alpha)$	$\rightarrow n + ^4\text{He} + ^6\text{Li}$	EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0 MT55,56,...
32			$^{10}\text{B}(n,nd)$	$\rightarrow n + ^2\text{H} + (^8\text{Be}^* \rightarrow 2\ ^4\text{He})$	EAF-2010(r) ($E > 20$)	
35			$^{10}\text{B}(n,nd2\alpha)$	$\rightarrow n + ^2\text{H} + 2\ ^4\text{He}$	EAF-2010(r) MT32	ENDF/B-VIII.0 MT55,56,...
55,56,...	22		$^{10}\text{B}(n,n)$	$\rightarrow n + ^4\text{He} + ^6\text{Li}$		ENDF/B-VIII.0 ($E < 20$)
55,56,...	35		$^{10}\text{B}(n,n)$	$\rightarrow n + ^2\text{H} + 2\ ^4\text{He}$		ENDF/B-VIII.0 ($E < 20$)
105			$^{10}\text{B}(n,t)$	$\rightarrow ^3\text{H} + (^8\text{Be}^* \rightarrow 2\ ^4\text{He})$	EAF-2010(r) ($E > 20$)	
700			$^{10}\text{B}(n,t)$	$\rightarrow ^3\text{H} + (^8\text{Be} \rightarrow 2\ ^4\text{He})$		ENDF/B-VIII.0 ($E < 20$)
106			$^{10}\text{B}(n,^3\text{He})$	$\rightarrow ^3\text{He} + (^8\text{Li}^* \rightarrow ^8\text{Be}^* \rightarrow 2\ ^4\text{He})$	EAF-2010	
107			$^{10}\text{B}(n,\alpha)$	$\rightarrow ^4\text{He} + ^7\text{Li}$	EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0 MT800,801
800			$^{10}\text{B}(n,\alpha)$	$\rightarrow ^4\text{He} + ^7\text{Li}$		ENDF/B-VIII.0 ($E < 20$)
801			$^{10}\text{B}(n,\alpha)$	$\rightarrow ^4\text{He} + ^7\text{Li}^*$		ENDF/B-VIII.0 ($E < 20$)
113			$^{10}\text{B}(n,t2\alpha)$	$\rightarrow ^3\text{H} + 2\ ^4\text{He}$	EAF-2010(r) MT105	ENDF/B-VIII.0 MT700
155			$^{10}\text{B}(n,t\alpha)$	$\rightarrow ^3\text{H} + 2\ ^4\text{He}$	EAF-2010	

TABLE 15. Gas production reactions from neutrons interacting with ^{11}B target. The MT numbers (reactions) used in IRDF-II assembly following the ENDF-6 format, and the relevant reaction products are highlighted in bold.

MT	LR	Gas	Reaction	Products	Source evaluations of IRDF-II	
33		^3H	$^{11}\text{B}(n,nt)$	$\rightarrow n + ^3\text{H} + (^8\text{Be} \rightarrow 2\ ^4\text{He})$	EAF-2010	
105			$^{11}\text{B}(n,t)$	$\rightarrow ^3\text{H} + ^9\text{Be}$	EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0 ($E < 20$)
11		^4He	$^{11}\text{B}(n,2nd)$	$\rightarrow 2\ n + ^2\text{H} + (^8\text{Be} \rightarrow 2\ ^4\text{He})$	EAF-2010	
22			$^{11}\text{B}(n,n\alpha)$	$\rightarrow n + ^4\text{He} + ^7\text{Li}$	EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0(r) ($E < 20$)
24			$^{11}\text{B}(n,2n\alpha)$	$\rightarrow 2\ n + ^4\text{He} + ^6\text{Li}$	EAF-2010	
33			$^{11}\text{B}(n,nt)$	$\rightarrow n + ^3\text{H} + (^8\text{Be} \rightarrow 2\ ^4\text{He})$	EAF-2010	
107			$^{11}\text{B}(n,\alpha)$	$\rightarrow ^4\text{He} + (^8\text{Li} \rightarrow ^8\text{Be} \rightarrow 2\ ^4\text{He})$	EAF-2010(r) ($E > 20$)	ENDF/B-VIII.0 ($E < 20$)
117			$^{11}\text{B}(n,d\alpha)$	$\rightarrow ^2\text{H} + ^4\text{He} + (^6\text{He} \rightarrow ^6\text{Li})$	EAF-2010	

The ENDF/B-VIII.0 library contains only $^{11}\text{B}(n,n\alpha)$ MT 22 and $^{11}\text{B}(n,\alpha)$ MT 107 reactions; the first one is much higher than the corresponding one in the EAF-2010 library because it was normalized to the measurement by Kneff (EXFOR #10933019); It can be understood from the original article that the measured quantity was actually the total helium production, therefore the normalisation in ENDF/B-VIII.0 is not correct.

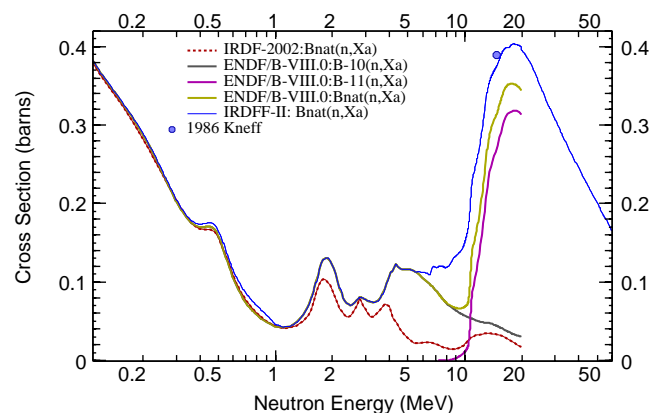


FIG. 64. (Color online) Comparison of the IRDF-II ^4He gas-production cross sections in neutron induced reactions on ^{nat}B target relative to Kneff *et al.* measurement [89].

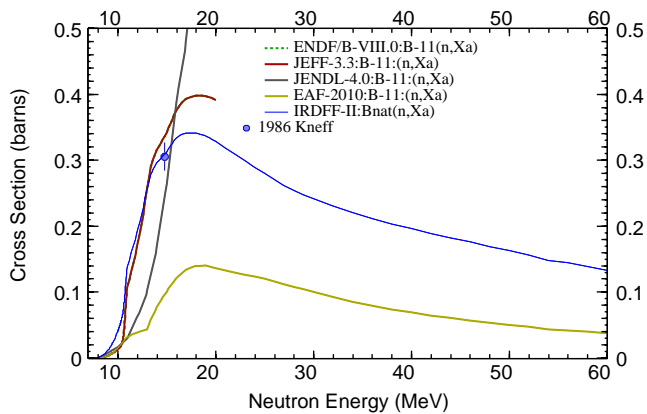
Reactions in the IRDF-II library are adopted from

the EAF-2010 library, except for those that are also present in ENDF/B-VIII.0. The $^{11}\text{B}(n,n\alpha)$ MT 22 from ENDF/B-VIII.0 was scaled down by 30 % to agree with the total helium-production measurement by Kneff. The cross sections above 20 MeV are taken from the EAF-2010 library, scaled for continuity at 20 MeV.

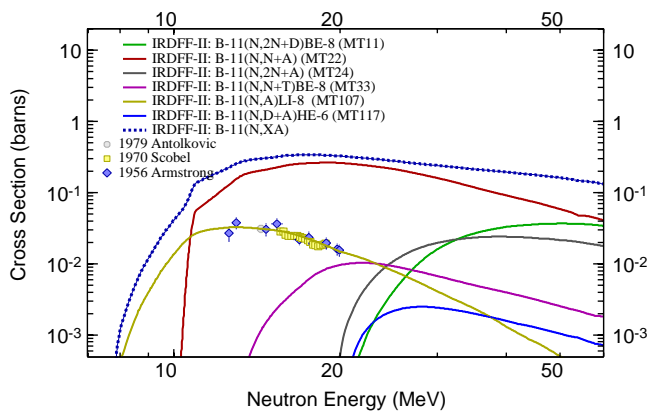
The cross-section curves are shown in Fig. 65(a). Examples of contributing reactions to ^4He and tritium gas production from neutrons interacting with ^{11}B are shown in Figs. 65(b) and 63.

31. $^{nat}\text{B}(n,X)^4\text{He}$

Since various ^{10}B enrichments can be used for the dosimeter, users of the library can combine ^4He gas generation contributions from ^{10}B and ^{11}B to get the dosimeter response. Some dosimeters use the boron in its natural abundance. To serve as a check for how the individual isotopic contributions should be combined, the IRDF-II library also includes the ^4He gas generation cross section from elemental boron with the naturally occurring isotopic abundances. Fig. 64 compares the ^4He gas production from the separate isotopes and from the elemental boron file and experimental data at 14 MeV measured by Kneff in 1986 [89]. The IRDF-II is in excellent agreement with measured data.



(a) Comparison of data from evaluated libraries relative to Kneff *et al.* measurement [89].



(b) Contribution of different reaction channels to the total ^4He gas-production cross-sections in the IRDF-II evaluation.

FIG. 65. (Color online) Comparison of the ^4He gas-production cross sections for the $n+^{11}\text{B}$ reaction in the IRDF-II library.

VI. NEUTRON BENCHMARK FIELDS

The following subsections describe the source for the neutron spectral data used to characterize the benchmark neutron fields.

A. Standard Field

A standard benchmark neutron field is defined [90, 92] as a permanent and reproducible neutron field that is characterized to state-of-the-art accuracy in terms of neutron fluence rate and energy spectra, and one where the associated spatial and angular distributions, and other important field quantities, have been verified by interlaboratory measurements. A given neutron field is considered to be a “standard” only over a specified neutron energy range and there is only one type of “standard neutron field” for a given energy range. The recognized “standard neutron field” is the ^{252}Cf spontaneous fission field. Secs. VII A and VIII A address how this field is used to provide supporting IRDF-II validation evidence and how consistent are those evidences.

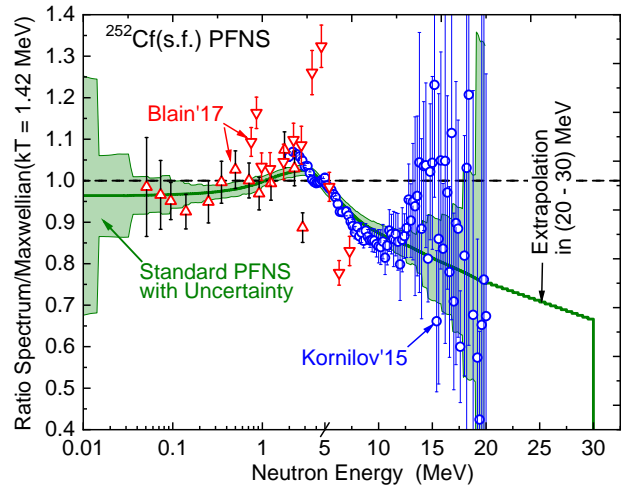


FIG. 66. (Color online) Standard $^{252}\text{Cf}(s.f.)$ prompt fission neutron spectrum (PFNS) and its uncertainties [32] as a ratio to the Maxwellian distribution with $kT = 1.42$ MeV (both spectra are normalised to unity). Spectrum extrapolation from 20 MeV to 30 MeV is shown by dash line. The recent new measurements [94] and [95] are plotted as symbols. Note that the scale changes from log to linear above 5 MeV.

1. $^{252}\text{Cf}(s.f.)$

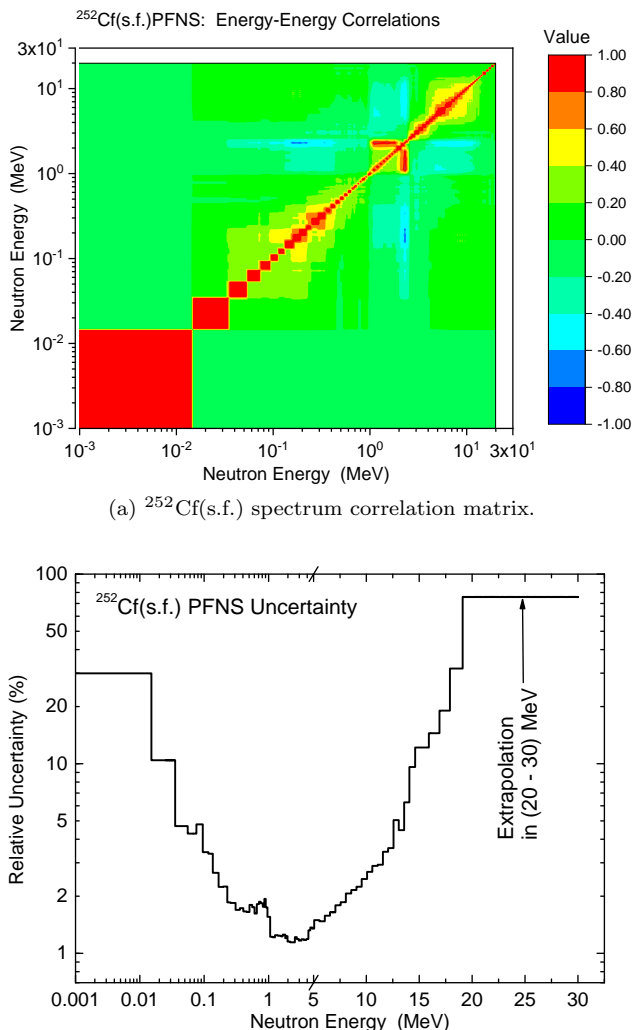
The ^{252}Cf spontaneous fission neutron field is defined to be a standard neutron field over the energy range from 0.1 MeV to 20 MeV.

The standard reference prompt fission neutron spectrum from spontaneous fission of $^{252}\text{Cf}(s.f.)$ [91] is characterized based on neutron time-of-flight measurements and subsequent model-independent evaluation. This work provided the spectrum definition incorporated into the IRDF-II library up to 20 MeV. The spectrum was then extrapolated up to 30 MeV of neutron outgoing energies by adding a point at 30 MeV, preserving the energy slope as below 20 MeV. All $^{252}\text{Cf}(s.f.)$ spectrum evaluations are based on original Mannhart work [93]. There are some minor differences above 10 MeV outgoing neutron energy between IRDF-II spectrum and the one included in the ENDF/B-VIII.0 library.

Figs. 67(a) 67(b) shows the correlation matrix and the energy-dependent standard deviation for the $^{252}\text{Cf}(s.f.)$ spectrum, correspondingly.

The spectrum is shown in Fig. 66. In the figure the y-axis is shown as a ratio to the Maxwellian distribution with $kT = 1.42$ MeV to better capture the large variation in neutron population with increasing energy. Since the time-of-flight (TOF) spectral characterization data only addressed the region up to 20 MeV, the higher energy portion of the spectrum is obtained from the extrapolation process described above. Significant increases in the spectral uncertainties are seen above 10 MeV and continuing up to the TOF cut-off energy at 20 MeV. Above this energy, the uncertainty in the extrapolated spectrum is assumed to be the same as that at the 20-MeV last measured value, which is 75 %.

Since the evaluation of the $^{252}\text{Cf}(s.f.)$ prompt fission spectrum performed by W. Mannhart in 1987 [93], several new measurements have been carried out by the time-



(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding spectrum mean value.

FIG. 67. (Color online) Uncertainties and correlations of the $^{252}\text{Cf}(\text{s.f.})$ standard neutron benchmark spectrum in IRDF-II library.

of-flight technique. The two newest measurements are discussed below. In 2015, N. Kornilov has measured the PFNS in the neutron range 2 MeV to 20 MeV [94]. In 2017, E. Blain and co-workers have reported two data sets measured by different detectors: from 0.05 to 2.8 MeV and from 0.75 to 7.3 MeV, however both - in arbitrary units [95]. Regrettably these new experiments, as seen in Fig. 66, have rather large uncertainties or fluctuations, thus contributing little to the improvement of the existing $^{252}\text{Cf}(\text{s.f.})$ standard.

B. Experimentally Measured Potential Reference Neutron Fields

Reference neutron benchmark fields are defined [90, 92] as permanent and reproducible neutron fields, less-well-characterized than a standard field, but acceptable as a measurement reference by a community of users. As part of the validation effort, an IAEA-sponsored Coordinated Research Project has focused on addressing improving the fidelity of the spectral characterization at candidate ref-

erence benchmark fields and using data gathered in these fields as part of the validation evidence for the IRDF-II library. The following subsections address the spectral characterization data that have been used in the supporting analysis reported here.

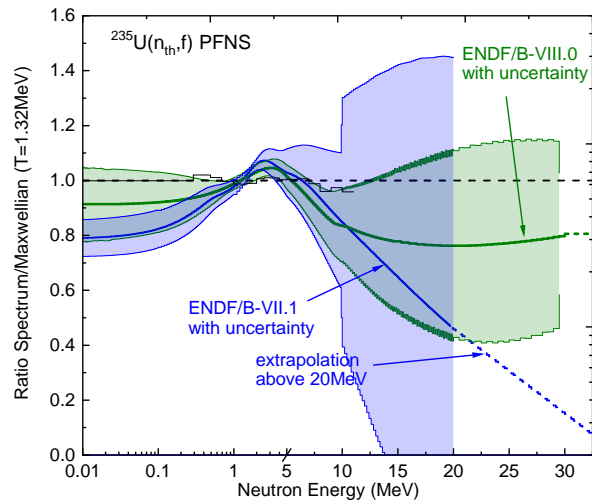


FIG. 68. (Color online) Reference $^{235}\text{U}(n_{th},f)$ prompt fission neutron spectrum (PFNS) [98–100] and its uncertainties as a ratio to the Maxwellian distribution with $kT = 1.32$ MeV (the spectra are normalised to unity). Red curve - from ENDF/B-VIII.0 [30], black curve - from ENDF/B-VII.1 [33]. Extrapolations beyond 20 MeV or 30 MeV are shown by dashed lines. Note that the scale changes from log to linear above 5 MeV.

1. $^{235}\text{U}(n_{th},f)$

A new ^{235}U thermal fission spectrum evaluation was developed within an IAEA-sponsored project using a least-square code GMAP in conjunction with shape data relative to the very-well-characterized $^{252}\text{Cf}(\text{s.f.})$ prompt fission neutron spectrum (PFNS). This new spectrum was the result of contributions from the IAEA CIELO collaboration [39, 40, 96, 97], the IAEA Neutron Standard committee [32], and the IAEA Prompt Fission Neutron Spectra project [98–100]. The spectrum incorporated into the IRDF-II library was converted from the ENDF/B-VIII.0 MF5/MT18 entry [30]. The associated covariance matrix was taken from the MF35/MT18 entry. Fig. 68 shows this ^{235}U thermal neutron fission spectrum - represented as a ratio to a Maxwellian distribution at a temperature of $kT=1.32$ MeV.

Figs. 69(a) 69(b) shows the correlation matrix and the energy-dependent standard deviation for the $^{235}\text{U}(n_{th},f)$ reference spectrum, correspondingly.

2. $^{233}\text{U}(n_{th},f)$ and $^{239}\text{Pu}(n_{th},f)$

The PFNS of thermal-neutron induced fission on ^{233}U and ^{239}Pu targets were evaluated together with the PFNS of thermal-neutron induced fission on ^{235}U target within the IAEA CRP on the Prompt Fission Neutron Spectra of Major Actinides [100]. A very few measured reaction rates are available, one set for the $^{233}\text{U}(n,f)$ PFNS, none

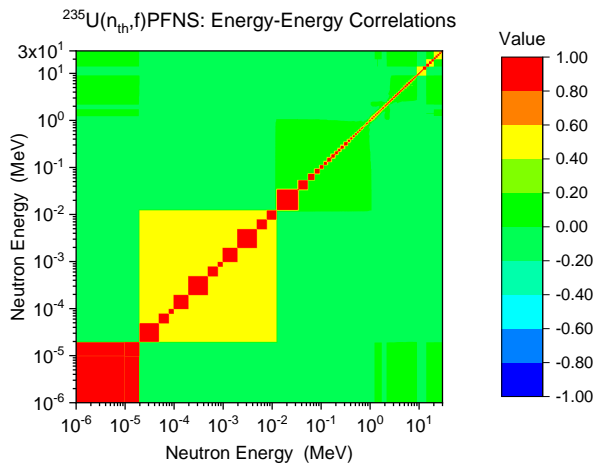
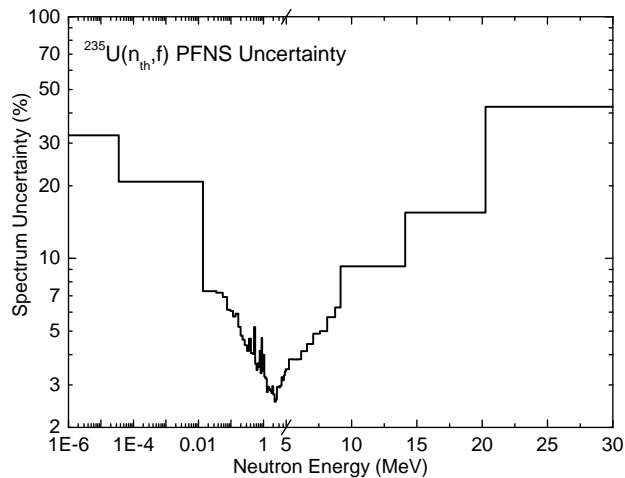
(a) $^{235}\text{U}(n_{th},f)$ spectrum correlation matrix.(b) One-sigma uncertainties in % ($\equiv 100 \times \frac{\sqrt{\text{cov}(i,i)}}{\mu_i}$), being μ_i the corresponding spectrum mean value.

FIG. 69. (Color online) Uncertainties and correlations of the $^{235}\text{U}(n_{th},f)$ standard neutron benchmark spectrum in IRDF-II library.

for $^{239}\text{Pu}(n,f)$, but some measurements were reported as double ratios of spectral indices to the ratio in PFNS of Pu. Since the spectra are not readily available elsewhere, they are included in the IRDF-II spectra file. The plot of the ^{233}U spectrum is shown in Fig. 70 as a ratio to a Maxwellian spectrum with temperature $kT=1.30$ MeV. The plot of the ^{239}Pu spectrum is shown in Fig. 71 as a ratio to a Maxwellian spectrum with temperature $kT=1.35$ MeV. These spectra were not modified above 10 MeV, but the proposed modification of the high-energy tail was described in detail in Ref. [100].

3. $^9\text{Be}(d,n)$ Reaction Neutron Field

The development and qualification of fusion reactor materials requires irradiation in a neutron spectrum that includes very high flux at 14 MeV along with lower energy neutrons to simulate the first wall spectrum at proposed fusion power reactors. Since fusion reactors do not exist for the testing and fission reactors lack any significant flux at 14 MeV, it has been proposed to produce suitable neu-

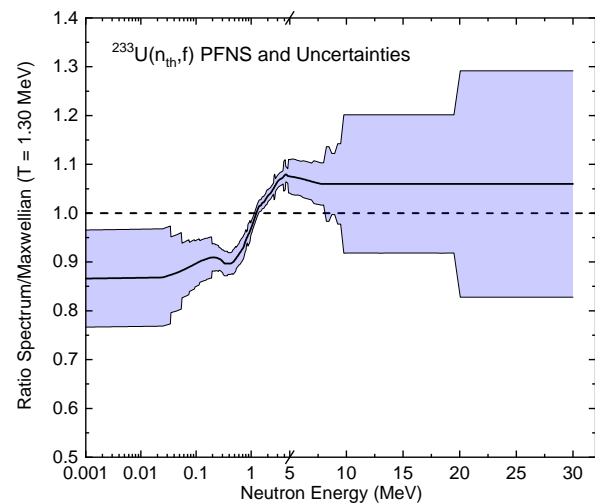


FIG. 70. (Color online) The $^{233}\text{U}(n_{th},f)$ prompt fission neutron spectrum (PFNS) ratios to the Maxwellian distribution with $kT=1.30$ MeV. Note that the scale changes from log to linear above 5 MeV.

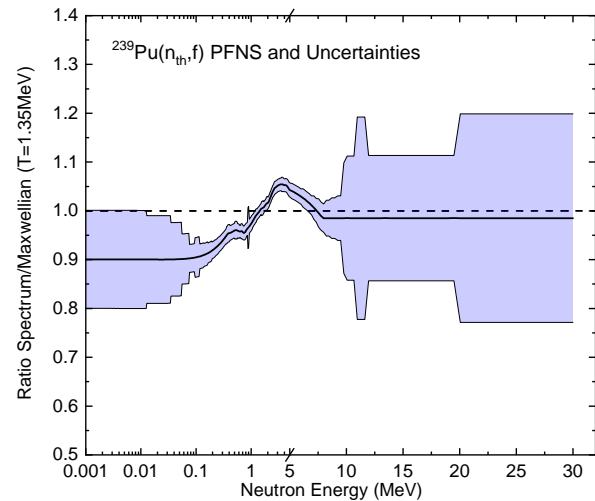


FIG. 71. (Color online) The ^{239}Pu prompt fission neutron spectra (PFNS) ratios to the Maxwellian distribution with $kT=1.35$ MeV. Note that the scale changes from log to linear above 5 MeV.

tron fields by stopping intense deuteron beams produced by particle accelerators in thick Be or Li targets. This produces a very high flux of neutrons over a continuous energy spectrum that can be designed to peak around 14 MeV. The IFMIF (International Fusion Materials Irradiation Facility) [26] is being proposed to study radiation damage in candidate fusion reactor materials.

Experiments were performed at deuteron energies of 16 MeV and 40 MeV producing neutron spectra as shown in Fig. 72. The neutron spectrum produced with deuteron energy of 16 MeV was measured at the Argonne National Laboratory tandem accelerator [101]. The neutron spectrum produced with deuteron energy of 40 MeV was measured at the Oak Ridge Isochronous Cyclotron (ORIC) [102]. The data from this facility can be found in the EXFOR entry No. C1832.002. The neutron spectra were very well characterized by using neutron time-of-flight technique. This method accurately determines the

energy of neutrons interacting with a neutron detector at a fixed distance from the beryllium target by measuring the time required for a neutron to be created in the target and then travel to the neutron detector. The deuteron beam is pulsed to provide the start time and the stop time is determined when a neutron is detected. The absolute accuracy of the neutron flux is determined by the efficiency of the neutron detector, which is about 10 %. Dosimetry materials were then irradiated in prepared geometries resulting in accurately measured reaction rates in well-characterized neutron spectra.

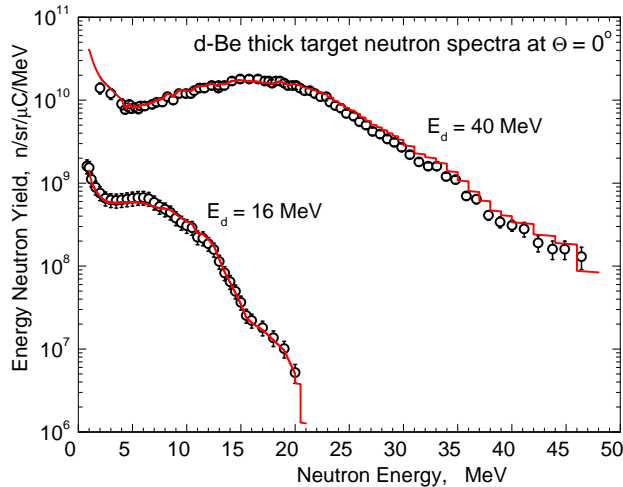


FIG. 72. (Color online) Neutron time-of-flight spectra are shown for thick Be targets at deuteron energies of 16 and 40 MeV at zero degrees. The red lines show neutron spectral adjustment in 725 groups using the new IRDFF-II natural element cross sections with STAYSL-PNNL.

C. Reference Neutron Fields from Detailed Computational Models

1. SPR-III Fast Burst Reactor Central Cavity

SPR-III is an advanced fast-burst Godiva-type reactor with a large 16.5-cm central cavity, commissioned in 1975 at Sandia National Laboratory, NM, U.S.A. [103]. This is an unmoderated cylindrical assembly of uranium enriched to 93 % ^{235}U and alloyed with 10-wgt-% molybdenum to ensure phase stability. The core consists of eighteen fuel plates mechanically fastened into two halves of nine plates each. The mass of the individual plates varies between 6.8 and 15.4 kg and the total mass of the core is about 258 kg. The nine upper plates are held stationary by the core support structure; the nine lower plates are attached to an electromechanical drive mechanism. Four reflector-type control devices are used; three for control and the fourth is the burst element. The control elements are used to establish a critical configuration of the core and to adjust the pulse yields. Although it was developed primarily for the radiation testing of electronic parts and systems, it has been used in a wide variety of research activities. It is positioned in the centre of an air-filled shield building called a Kiva. SPR-III has a ^{10}B shroud

that serves to decouple it from room return. The reactor can be operated in steady-state (up to 10 kW power) or pulsed mode (10 MJ in an 76 μs FWHM pulse that yields approximately 5×10^{14} n/cm 2 in the reactor central cavity).

The reference SPR-III spectrum adopted within the IRDFF-II library characterizes the fuel centreline of the central cavity. High fidelity calculations using the MCNP, version 6.2.0, point cross section Monte Carlo code with the ENDF/B-VIII.0 cross sections and a detailed 3D geometry were performed to provide the best possible calculated neutron spectrum. The reactor, support structures, Kiva, and dosimetry test fixture were modelled in detail. The coupled neutron/photon calculation was performed using the MPI version of the code with 40 processors and using the L'Ecuyer 63-bit random number generator. The calculation used 5015 cycles with 2000000 source neutrons per cycle, of which 15 cycles were used to establish the normal fission distribution before beginning tally accumulation, for a total neutron source tally sample size of 100 billion neutrons. The neutron spectrum was scored in the 640-group SAND-II energy structure.

For this benchmark neutron field to be used to support the validation of the IRDFF-II cross sections, it is important that there be an uncertainty estimate of the calculated “*a priori*” spectrum, in the form of an energy-dependent covariance matrix, and that this uncertainty estimate be derived independent of the activation measurements that are being used in the intended validation activity. This derivation of an energy-dependent uncertainty for a fine-group calculated spectrum is a very challenging task and a task where existing uncertainty quantification (UQ) methodologies are deficient and active work is still needed by the community. In addition to consideration of the numerical uncertainty, great care must be taken to not only address the uncertainty in the underlying transport cross sections and material geometry used in the radiation transport calculation, but to also investigate existing sources of “model defect”, *i.e.*, potentially correlated sources of uncertainty in the calculational models not directly captured in the user input parameters. A covariance matrix for this calculated spectrum was derived using the methods detailed in Reference [104] and subject-matter expertise based on previous modelling activities and subsequent comparison with the results obtained through spectrum adjustment methods. This “*a priori*” covariance matrix for the reference neutron, derived independent of the reaction activities addressed later in this paper as part of the IRDFF-II validation evidence, is included as part of the IRDFF-II file that was outlined in Sec. II.H. This content from the covariance matrix is seen in Figs. 73 and 74. Fig. 73 shows the energy-dependent uncertainty in the neutron spectrum. Fig. 74 shows the correlation matrix.

2. ACRR Pool-type Reactor Central Cavity

ACRR is a pulse and steady-state, pool-type research reactor located at Sandia National Laboratory, Albuquerque, U.S.A. The ACRR maintains a large, dry irradi-

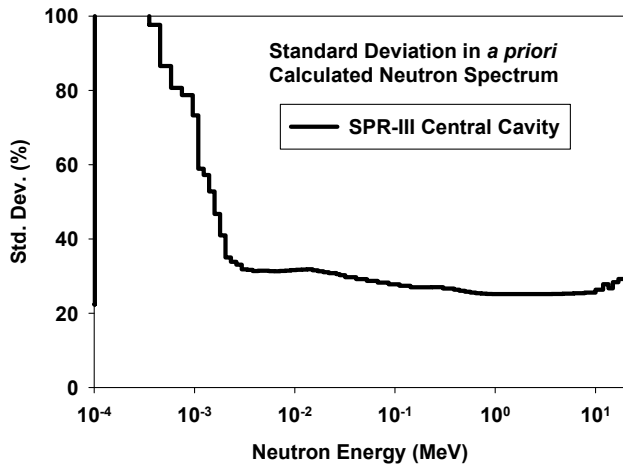


FIG. 73. Standard deviation for the calculated SPR-III fast burst reactor central cavity neutron spectrum.

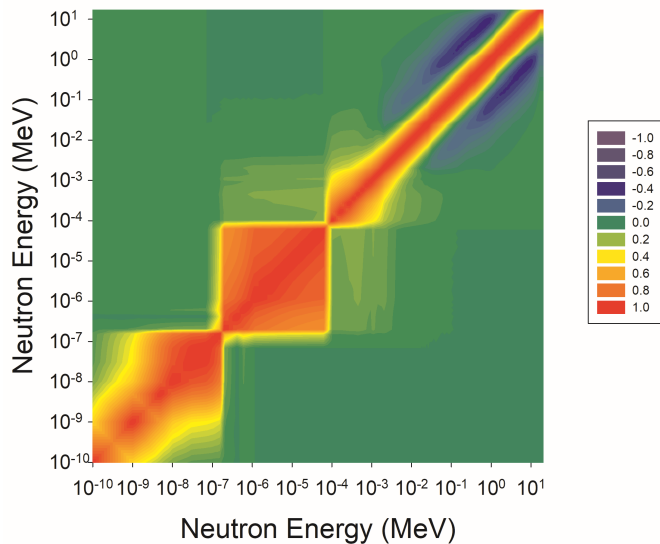


FIG. 74. (Color online) Correlation matrix for the Calculated SPR-III fast burst reactor central cavity neutron spectrum.

ation cavity at the centre of its core. ACRR is typically used to perform irradiation testing where a high neutron fluence is required for a short period of time. Historically, ACRR has been used for a wide variety of experiment campaigns including electronics damage testing, nuclear fuels testing, space nuclear thermal propulsion testing, and medical isotope production. ACRR is currently fully operational and is used to add validation evidence for new dosimetry-quality reactions with a response in the epithermal and fast fission energy regions. ACRR's main attributes include a large, dry central irradiation cavity, epithermal neutron fluence, and large pulsing capabilities that permit a high irradiation and rapid recovery of the dosimeter.

ACRR can operate in a steady-state, transient, or pulse mode. In the steady-state mode, the operating power level is limited to 4 MW. In the pulse mode, a maximum pulse size of 250 MJ with a full-width half-maximum (FWHM)

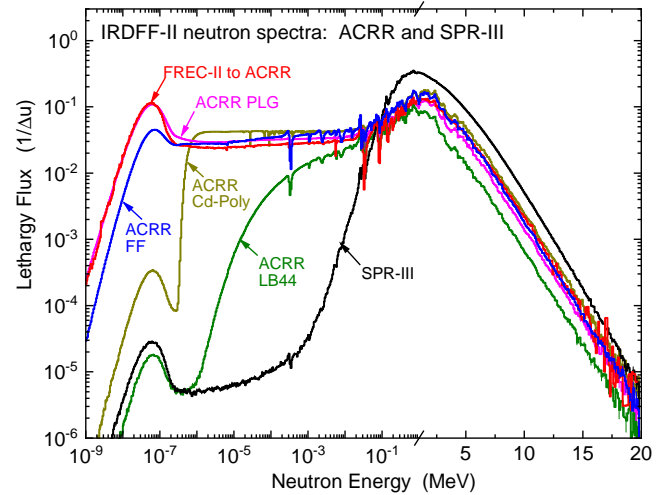


FIG. 75. (Color online) Neutron reference reactor spectra for ACRR fission fields with various “Bucket” modifying environments. Note that the scale changes from log to linear above 1 MeV.

of 6 ms. In the transient mode, the reactor power shape can be tailored to the desired requirements for a total reactor energy deposition of 300 MJ. The transient mode can be used to increase the reactor power quickly; for example, a ramp increase in power level linear with time from low power to high power. ACRR is fueled by a 236 element array of $\text{UO}_2\text{-BeO}$ fuel elements. The fuel is uranium enriched to 35 weight percent ^{235}U , with 21.5 weight percent UO_2 and 78.5 weight percent BeO . The ACRR fuel elements are stainless steel clad, 3.747 cm in outer diameter and 52.25 cm in fuel length. Within the elements are niobium cups that hold the UO_2BeO fuel pieces. ACRR is controlled by two fuel-followed safety rods, three poison (void-followed) transient rods, and six fuel-followed control rods. The control rods make up part of the 236 elements for the normal core configuration. Further information regarding the ACRR irradiation environment can be found in internal laboratory documents [105].

The ACRR reference spectrum [106] adopted within the IRDF-II library characterizes the core centreline of the central cavity with the 32-inch pedestal. High fidelity calculations were performed to provide the best possible calculated neutron spectrum. This is the spectrum that should be used as input to various spectrum adjustment codes, or as the trial spectrum in iterative unfolding codes. The radiation transport was performed with the continuous energy, three-dimensional Monte Carlo particle transport code MCNP [63]. The MCNP model of the ACRR central cavity with the 32-inch pedestal is shown in Fig. 76. The coupled neutron/photon calculation was performed using the MPI version of the code with 40 processors and using the L'Ecuyer 63-bit random number generator. The calculation used 5015 cycles with 2000000 source neutrons per cycle, of which 15 cycles were used to establish the normal fission distribution before beginning tally accumulation, for a total neutron source tally sample size of 100 billion neutrons. The neutron spectrum was scored in the 640-bin SAND-II energy group structure.

For this benchmark neutron field to be used in support of the validation of the IRDF-II cross sections, a

calculated "a priori" spectrum, in the form of an energy-dependent covariance matrix, and one that has been developed independent of the activation measurements that are being used in the intended validation activity, is required. The covariance matrix for this calculated spectrum was derived using the methods detailed in Reference [104] and subject-matter expertise based on previous modelling activities and subsequent comparison with the results obtained through spectrum adjustment methods. This "a priori" covariance matrix for the reference neutron, derived independent of the reaction activities is included as part of the IRDF-II file that was outlined in Sec. II.H. This content from the covariance matrix is seen in Figs. 77 and 78. Fig. 77 shows the energy-dependent uncertainty in the neutron spectrum. Fig. 78 shows the correlation matrix.

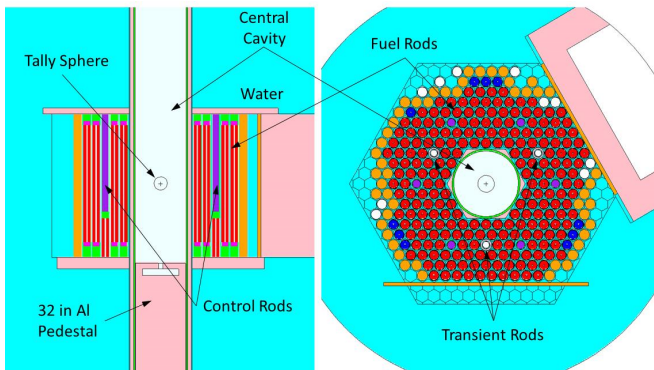


FIG. 76. (Color online) Computational model for the ACRR pool-type reactor central neutron spectrum.

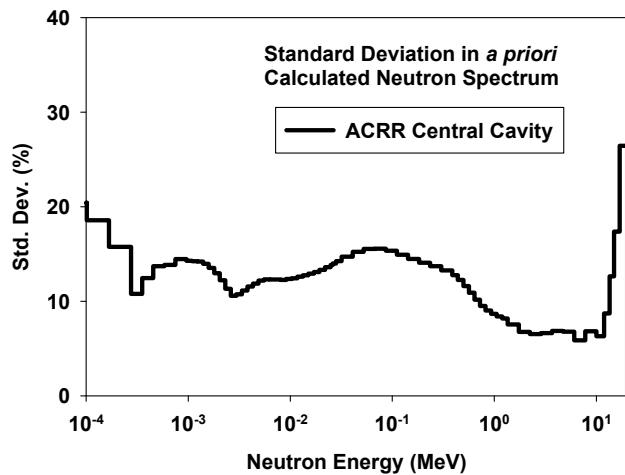


FIG. 77. Standard deviation for the calculated ACRR pool-type reactor central neutron spectrum.

3. ACRR Pool-type Reactor Pb-B₄C (LB₄₄) Bucket

The ACRR maintains an epithermal neutron fluence spectrum in the core and central cavity. This allows for the neutron energy fluence spectrum to be tailored to the desired specifications of an experiment by introducing various spectrum modifying buckets. Moderators can

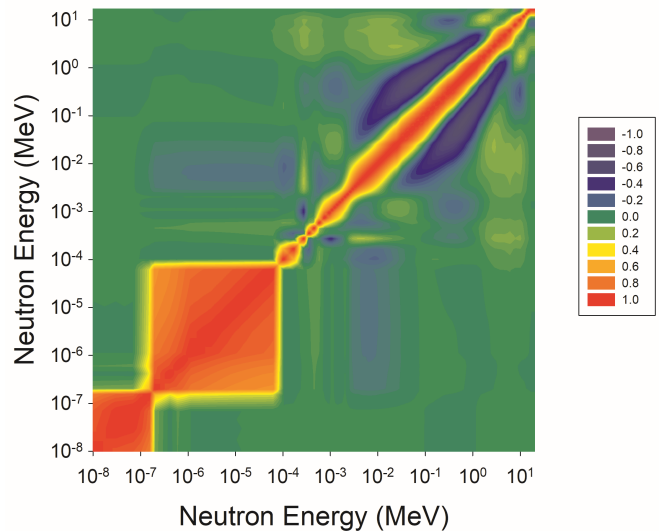


FIG. 78. (Color online) Correlation matrix for the calculated ACRR pool-type reactor central neutron spectrum.

be used within the cavity to thermalize the neutron spectrum. Boron and lead can be used to increase the fast neutron fluence ratio and decrease the gamma-ray fluence, respectively. The ACRR-LB44 neutron field uses a 44-inch tall lead and B₄C "bucket" to moderate neutron field seen at a 5-inch inner diameter experiment location. The primary purpose of this "bucket", depicted in Fig. 79, is to reduce the thermal component of the neutron spectrum as well as the associated gamma exposure. This neutron spectrum is calculated using the MCNP code in a manner similar to that described in Sec. VI.B.3. The calculated neutron spectrum is shown in Fig. 75.

4. ACRR Pool-type Reactor Cd-Polyethylene Bucket

A cadmium-polyethylene insert is used to tailor the neutron spectrum for this field. This Cd-poly "bucket" is designed to enhance the gamma ionization component in the mixed neutron/gamma reactor field by a factor of two. It also significantly reduces the thermal neutron component while leaving the epithermal component of the cavity spectrum. The "bucket" provides an inner experiment diameter of about 7.5-inches and accommodates packages with a height of 30-inches. The outer portion of the bucket is fabricated from high density polyethylene with a thickness of 0.55-inches and the inner cadmium sheet is 0.02-inches. The purpose of the bucket, depicted in Fig. 80, is to provide variation in the neutron-to-gamma radiation field when testing electronic components. The calculated neutron spectrum is shown in Fig. 75.

5. ACRR Pool-type Reactor Polyethylene-lead-graphite (PLG) Bucket

A polyethylene-lead-graphite (PLG) insert is used to tailor the neutron spectrum for this field. This PLG bucket is designed to fit within the 23.4-cm diameter

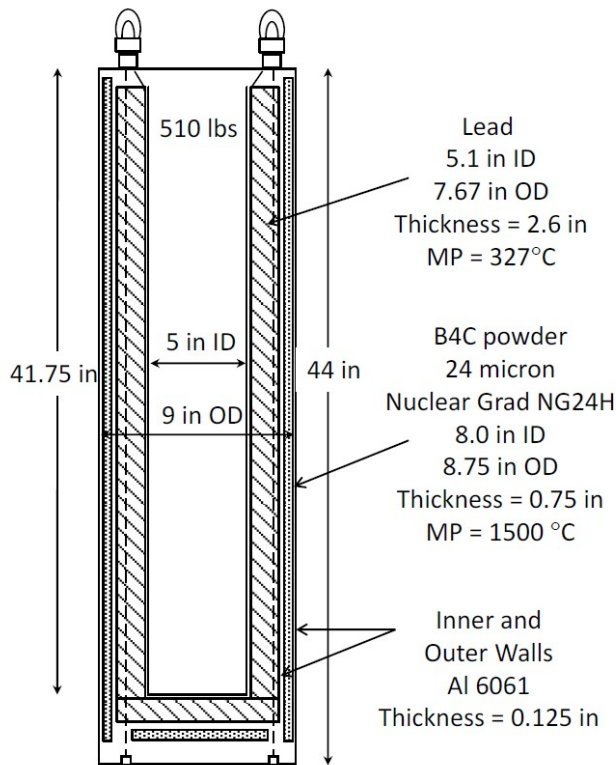


FIG. 79. "Bucket" used to tailor the ACRR-LB44 neutron spectrum.

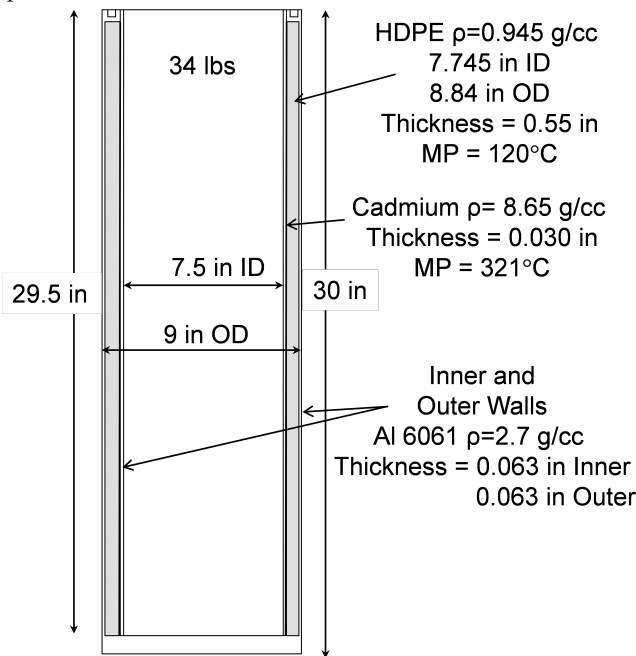


FIG. 80. "Bucket" used to tailor the ACRR-Cd-poly neutron spectrum.

ACRR central cavity. The purpose of the bucket, depicted in Fig. 81, is to provide some neutron moderation, reducing the fast neutron component relative to the total neutron fluence. The calculated neutron spectrum is shown in Fig. 75.

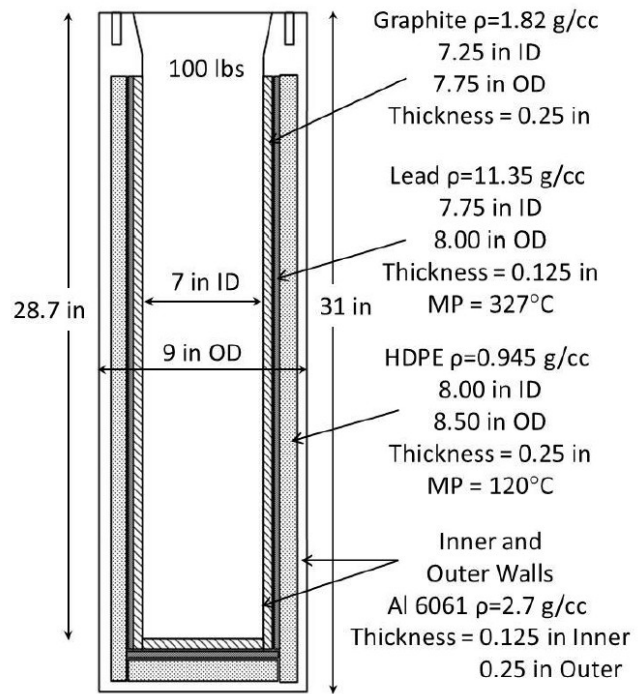


FIG. 81. "Bucket" used to tailor the ACRR-PLG neutron spectrum.

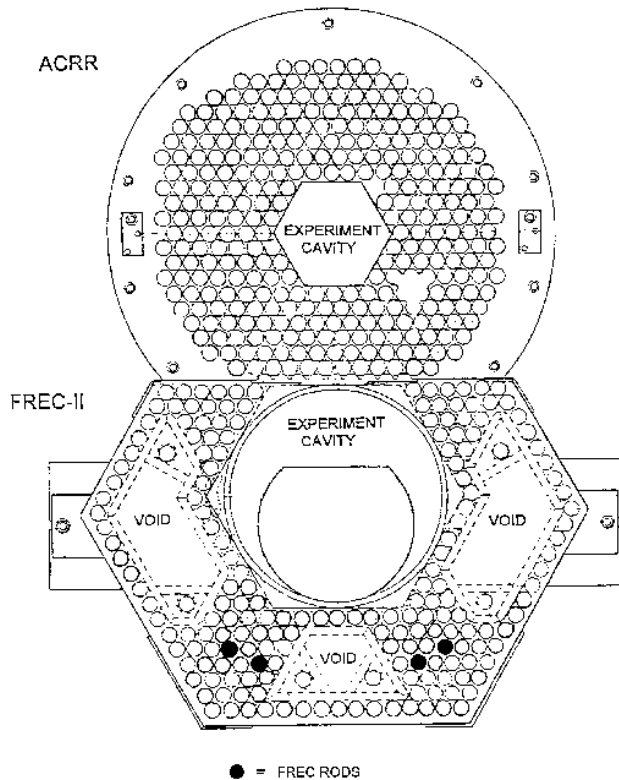


FIG. 82. ACRR fueled external cavity neutron spectrum.

6. ACRR Fueled Ring External Cavity (FREC-II)

FREC-II is an external cavity attached to the ACRR and it can be “coupled” or “de-coupled” to the ACRR core depending on the experimenter’s desires. In the “de-coupled” mode, FREC-II is tilted away from the core several degrees and the Ni plate is attached to the ACRR. In this configuration, the FREC-II has minimal effect on the neutronic behavior of the reactor. With the FREC-II “coupled,” a significant neutron and gamma-ray flux exists in the 20-inch, dry, FREC-II cavity.

Fig. 82 depicts the FREC-II cavity as it is coupled to the ACRR. The FREC-II uses uranium/zirconium-hydride (U-ZrH) TRIGA fuel. FREC-II is fueled with a 186-element array of $\text{UZrH}_{1.625}$ fuel elements. The fuel is 12 weight percent uranium enriched to 20 weight percent U-235. The hydrogen atomic ratio is 1.625 hydrogen atoms per Zr atom. The FREC-II fuel elements are stainless steel clad, 1.5 inches in diameter and 15 inches in fuel length. The UZrH fuel in each element is a single solid cylinder with a 0.25-inch diameter hole at its centre. The centre region maintains a Zr rod with the same diameter. FREC-II has four neutron poison elements that are fuel followed and can be positioned full-in (down position-poison in), full-out (up position-poison out), or any position in between. This feature allows for radial flux tilting in the 20-inch cavity. The FREC-II cavity allows for some advantages compared to the ACRR central cavity for consideration by experimenters: 1) the FREC-II can accommodate larger experiments, since its cavity is 20 inches in diameter compared to the 9-inch diameter central cavity; 2) the neutron and gamma-ray flux can be radially tilted by using the FREC rods and positioning the experiment within the cavity; and 3) the smaller neutron flux allows for a maximum pulse with a short pulse width (~ 8 ms) and smaller total neutron fluence. This same effect could be achieved in the central cavity by positioning the experiment package higher in the cavity.

The calculated neutron spectrum is shown in Fig. 75.

7. Mol-BR1 Cavity ^{235}U Fission Spectrum

A number of standard neutron fields were designed in the BR1 reactor at the Belgian Nuclear Research Centre SCK•CEN in Mol. They are designated as Mark-I, Mark-II, Mark-IIA or Mark-III. Many measurements of cross sections in these standard fields were reported [107–109]. In particular, most of the measurements of the $^{235}\text{U}(n_{th},f)$ PFNS spectrum-averaged cross sections were measured in the Mark-IIA neutron field, which is very similar to the current Mark-III. The standard fields are differentiated by enrichment, height and thickness of the ^{235}U cylinder and by the thickness of the Al claddings inside and outside of the uranium cylinder. The Mark-III standard neutron field was generated by means of a flexible ^{235}U source arrangement located at the centre of a spherical cavity in the graphite thermal column [107]. The large 100 cm diameter cavity has minimized the wall return corrections while permitting a large volume for active instruments. The access to the centre of the field is enabled through

a long 1 mm thick Cd tube, which is loaded with the irradiation samples. The source of fission neutrons is a ^{235}U cylinder wrapped around the cadmium tube. The Mark-III configuration was re-analysed recently [110] and the computational model of the core layout is shown in Fig. 83, with the irradiation facility at the top of the figure.

To refer the dosimetry reaction rates measured in the cavity centre to the ones in the $^{235}\text{U}(n_{th},f)$ PFNS, the neutron field perturbation corrections for the used facility configurations were calculated and applied by the authors of the measurements. The absolute neutron fluence was established relative to the ^{252}Cf source, or reactions of interest were measured as chosen ratios.

The neutron spectrum inside the MARK-III converter loaded in the BR1 cavity, computed by MCNP-6 with ENDF/B-VIII.0 neutron data, is depicted in Fig. 84. The standard fission neutron field at the BR-1 reactor has been made available for international cooperative efforts for neutron metrology. Thus the large fraction of the existing cross sections averaged over the reference $^{235}\text{U}(n_{th},f)$ prompt spectrum was measured there.

8. LR-0 Reactor at Řež

The LR-0 reactor of the Research Centre Řež near Prague, Czech Republic is an experimental zero-power light-water-moderated reactor. The core is placed in a 6.5 m high Al tank with diameter of 3.5 m. The large reactor vessel is a legacy of the former heavy water reactor project which was carried out back in the 60s. At the end of the 70s, the refurbishment was accomplished and the reactor core was converted to light-water-moderated to support research in this field. The first criticality of the light water version was reached in December 1982.

The main scope of the LR-0 research were some lattice studies of the VVER-440 and VVER-1000 reactor. Nominal thermal neutron flux density is $10^9 \text{ n.cm}^{-2}.\text{s}^{-1}$, which corresponds to the thermal power of 1 kW. Thanks to the low power, the fuel is considered to be physically fresh during all experiments. Later, the full-scale reflector segment of the VVER-1000 reflector was installed to the LR-0 vessel. Thus, the so called “VVER-1000 Mock-up” experiments were enabled. The only significant difference between a real VVER-1000 reactor core and the VVER-1000 Mock-up is the height, because the LR-0 does not need so much fuel as an energy-producing reactor. Despite the shorter fuel assemblies, the LR-0 reactor can be used for VVER-1000 and VVER-440 reactor experiments focused on criticality, neutron transport and dosimetry to support services for utilities. For reproducibility, the full-scale simulators of the VVER-1000 internals are permanently installed in the LR-0 tank (see Fig. 85, left).

The large reactor vessel offers space for the variety of experiments with cores physically and neutronicly decoupled from the VVER-1000 Mock-up. Based on this, smaller cores consisting of six hexagonal VVER-1000 fuel assemblies were designed to extend the experimental portfolio. The small core occupying only seven lattice positions is the simplest assembly which can be formed in the

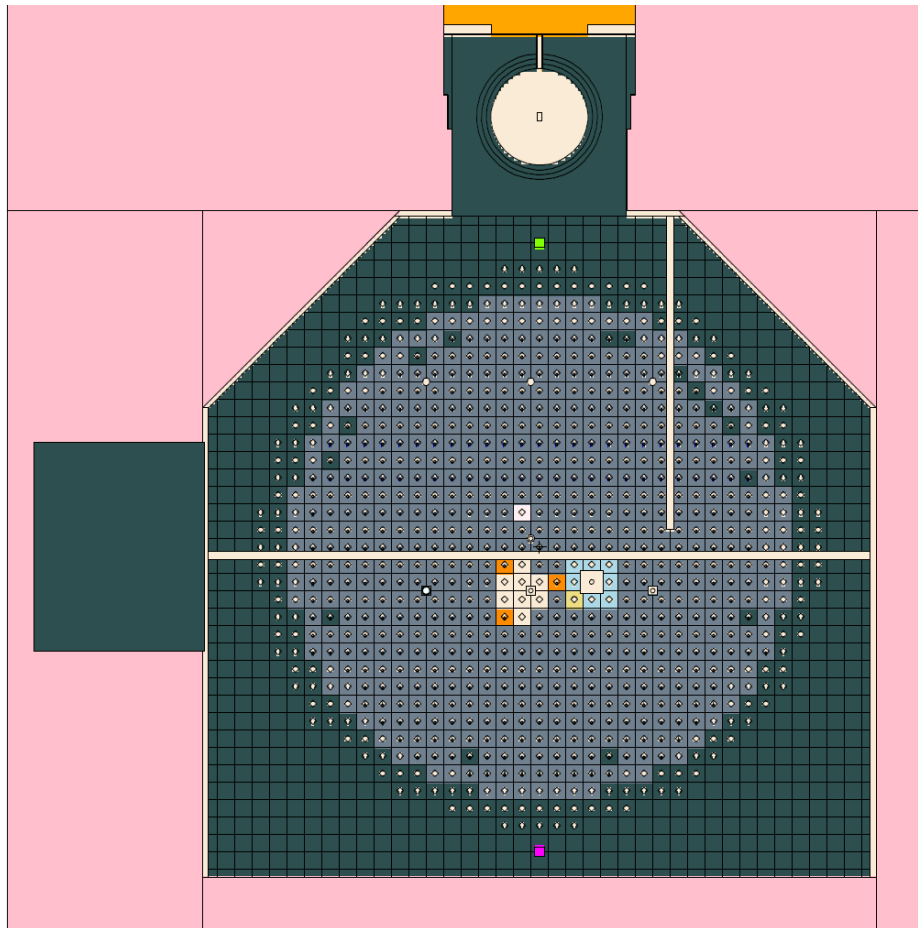


FIG. 83. (Color online) Computational model of the Mol-BR1 irradiation facility.

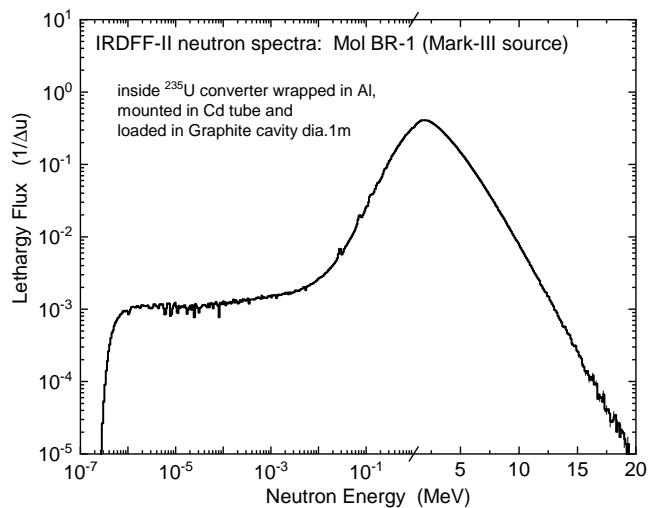


FIG. 84. The calculated fission neutron spectrum inside Mark-III U-235 converter in the Mol BR1 reactor graphite cavity. Note that the scale changes from log to linear above 1 MeV.

LR-0 reactor and it was also benchmarked. This benchmark core contains fuel assemblies with enrichment of 3.3 % surrounding one void hexagonal channel exactly fitting to the fuel lattice. As mentioned, the LR-0 reaches only negligible burn-up, thus redistribution of the fission source does not occur due to this effect and high repro-

ducibility of experiments can be guaranteed.

In the past, neutron spectra [111] and flux distribution [112] in the centre of the dry assembly have been validated as well as the criticality [113] and neutron emission distribution [114] of the driver core. Therefore, the field in the void assembly can be understood as the reference neutron field. Low flux experiments at 2×10^7 n.cm⁻².s⁻¹ in flux over 1 MeV with some larger samples have been performed. Only low activities of threshold reactions can be induced in this field, hence closer measuring geometry during gamma spectrometry was selected. This is possible thanks to a well-defined efficiency curve in the used HPGe semiconductor measurement arrangement. More details concerning the core description can be found in the IR-PhE Project Handbook [115]. The spectrum in the centre of the dry assembly of the Rež-LR0 reactor is shown in Fig. 86.

9. TRIGA-JSI Pneumatic Tube Irradiation Channel

The Jožef Stefan Institute (JSI) TRIGA reactor in Ljubljana, Slovenia is a 250 kW light-water pool-type TRIGA Mark II reactor with uranium zirconium hydride fuel elements and an annular graphite reflector, cooled by natural convection. The reactor achieved first criticality in 1966. Due to its very well characterized irradiation fields its main applications, at present, are Neutron Ac-

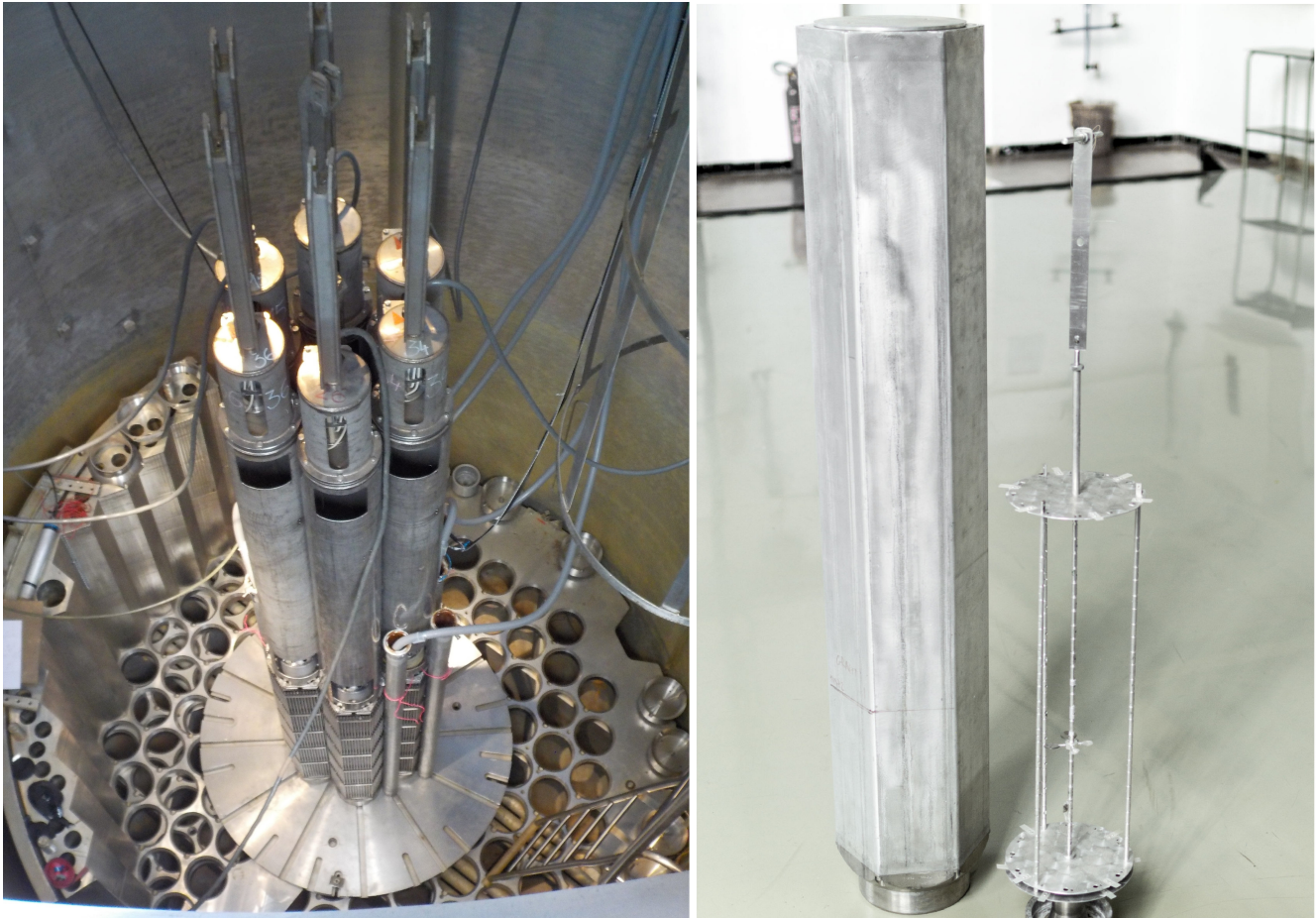


FIG. 85. (Color online) Core(left), special Al assembly (right) (from ANN. NUCL. EN. **112**, pp. 759–768 (2018)).

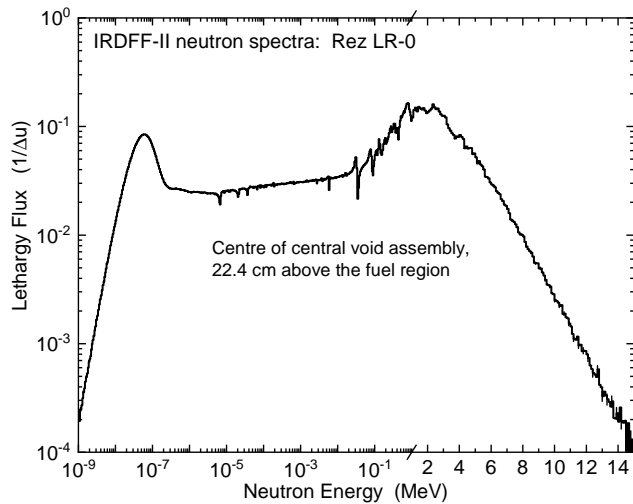


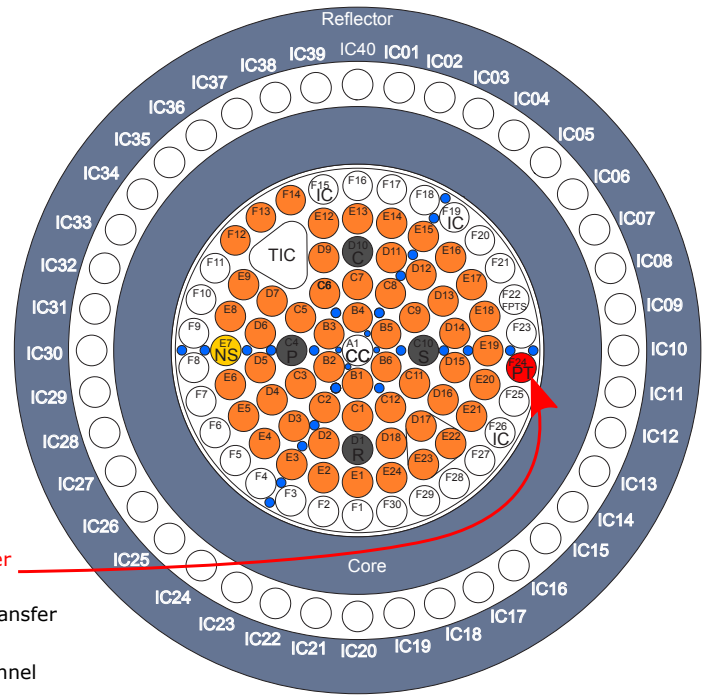
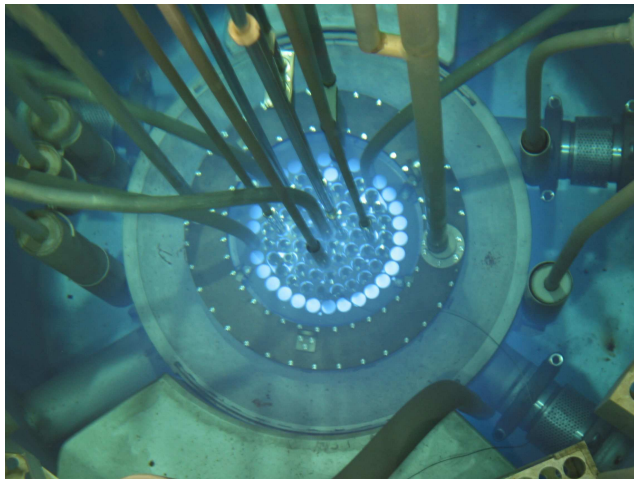
FIG. 86. Neutron field in the centre of the dry assembly of the Rež-LR0 reactor. Note that the scale changes from log to linear above 1 MeV.

tivation Analysis (NAA), radiation hardness testing of detectors and electronic components (*i.e.*, reference neutron radiation hardness irradiation facility for the CERN LHC accelerator and other accelerators within the EC H2020 project AIDA-II - Advanced European Infrastruc-

tures for Detectors at Accelerators), support for testing of neutron instrumentation detectors (fission chambers, self-powered neutron detectors) support for development of radiation-hard products, code testing and validation.

The JSI TRIGA reactor facility is described in detail in the publications [116, 117]. The reactor core is located at the bottom of a 6.25 m high open tank, 2 m in diameter. The core has a cylindrical configuration with 91 core locations which can be filled either by fuel elements or other components like control rods, a neutron source, irradiation channels, *etc.* The core lattice has an annular but not periodic structure. The fuel elements are arranged in six concentric rings: A, B, C, D, E and F, with 1, 6, 12, 18, 24 and 30 fuel locations per ring, respectively. The graphite reflector surrounding the reactor core is enclosed in aluminum casing and also houses a special irradiation facility, the rotary specimen rack, with 40 irradiation positions.

The reactor core is supported by a two 3/4-inch aluminium grid plates, which provide accurate axial and lateral positioning of the core components. The fuel elements are cylindrical with stainless steel (SS-304) cladding. Their total length is approximately 28 inches, their diameter is 1.478 inches. The fuel material in each element is 15 inches in length. The fuel elements are equipped with cylindrical graphite inserts at the top and bottom ends, 2.6 inches and 3.7 inches in length, respectively, which act as axial reflectors. In the centre of the fuel material there



- Fuel elements (20% U-235)
- Control rods
- Neutron source
- IC Irradiation channels
- In-core measurement positions
- CC Central channel
- PT Pneumatic sample transfer system
- FPTS Fast pneumatic sample transfer system
- TIC Triangular irradiation channel

FIG. 87. (Color online) Left: Photograph of the JSI TRIGA reactor core at 250 kW (full power). Right: schematic diagram of the reactor core configuration.

is a 0.25-inch-diameter hole which is filled by a zirconium rod. Between the fuel meat and the bottom graphite insert there is a 1/32-inch-thick molybdenum disc. The fuel itself is a homogeneous mixture of uranium and zirconium hydride with 12 wt.% uranium, enriched to 20 %.

Three control rods of the fueled-follower type are used in the reactor, *i.e.*, the Regulating (R), Shim (C), and Safety (S) control rods, identical in geometry and composition. A fourth control rod, the Transient Rod (T), is equipped with a pneumatic ejection system and enables reactor pulse operation. It is similar to the fueled-follower control rods, but with a so-called air follower, which replaces the fuel part in the fueled-follower control rods. The purpose of the air-follower is to reduce power peaking that could appear when the transient rod is in its fully withdrawn position during steady state operation.

During normal operation the reactor core is equipped with 7 irradiation channels. Six of the irradiation channels occupy one core position: the Central Channel (CC), located in the reactor core centre (A1), and five irradiation channels located in the outer F core ring (F15, F19, F22, F24 and F26). Two of these irradiation channels are equipped with pneumatic sample transfer systems, *i.e.*, the channel in position F22, also denoted as Fast Pneumatic Transfer System (FPTS), and in position F24, also denoted as Pneumatic Tube (PT). One channel, the Triangular Irradiation Channel (TIC) is larger and occupies three core positions (D8, E10 and E11), arranged in a triangle, and enables irradiations of objects with maximal lateral dimensions of approximately 50 mm. Fig. 87 displays a photograph of the TRIGA-JSI reactor core and a schematic diagram of the core configuration.

The measured quantities in the experiments were reaction rate ratios, either unfiltered relative to filtered, or

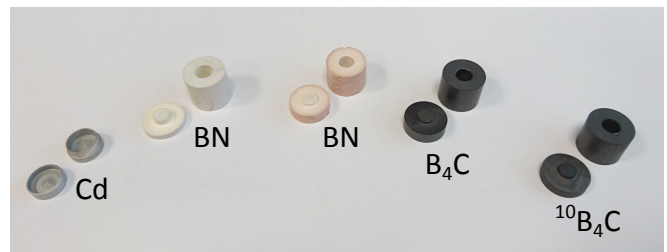


FIG. 88. (Color online) Neutron filters used in the experiments at the JSI TRIGA reactor. From left to right: Cd (1st campaign), BN (1st campaign), BN (2nd campaign), B₄C (2nd campaign), ¹⁰B₄C (2nd campaign).

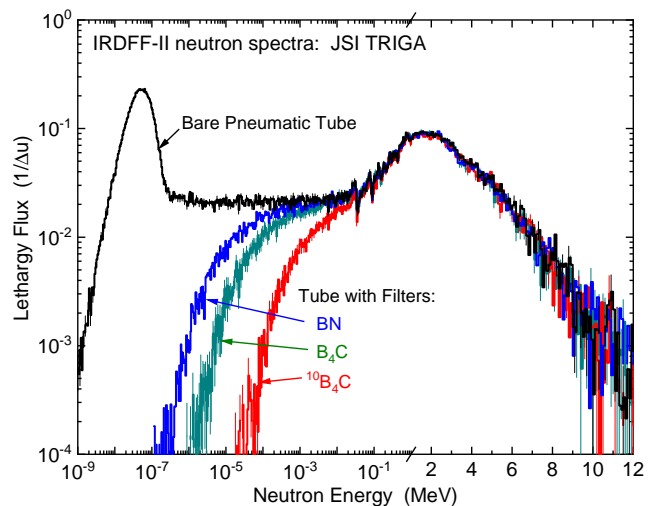


FIG. 89. (Color online) Unfiltered and filtered neutron fields in the TRIGA-JSI reactor. Note that the scale changes from log to linear above 1 MeV.

TABLE 16. Dimensions of the Cd, BN, B₄C and ¹⁰B₄C neutron filters used in the experiments at the JSI TRIGA reactor.

Filter	Outer diameter [mm]	Height [mm]	Inner diameter [mm]	Inner height [mm]
Cd (1st campaign)	10	6	8	4
BN (1st campaign)	13	14	5	6
BN (2nd campaign)	13	14	5	5
B ₄ C (2nd campaign)	13	15	5	5
¹⁰ B ₄ C (2nd campaign) (¹⁰ B enrichment > 96 %)	13	13	5	5

one reaction rate relative to another, derived from activation measurements with gamma spectrometry. Irradiations were performed in two experimental campaigns in the TRIGA-JSI reactor, both in the Pneumatic Tube (PT) irradiation channel. In the first campaign, cadmium (Cd) and boron nitride (BN) filters were employed. The objectives of the campaign were two: firstly the characterization of the neutron spectrum in the PT irradiation channel, and secondly, to investigate the suitability of boron nitride filters to shift the energy sensitivity of radiative capture reactions to the epithermal energy range, for cross-section validation [118]. In the second campaign, boron nitride (BN), boron carbide (B₄C) and enriched boron carbide (¹⁰B₄C) filters were employed. The objective of the campaign was to investigate new candidate radiative capture reactions, which are sensitive to the epithermal energy range (broadly between 1 keV and 1 MeV) under boron filters [119], which could be applicable for epithermal neutron dosimetry. The filter dimensions are reported in Table 16. Fig. 88 displays the filters. Fig. 89 shows the spectra with and without covers.

10. ISNF

The ISNF facility is set up in the spherical cavity inside the graphite column of the reactor. Eight enriched ²³⁵U fission discs, symmetrically mounted in the cavity, convert the thermal neutron flux with an average energy $\langle E \rangle = 0.0253$ eV to the fast fission spectrum. The summary flux is transmitted by the enriched ¹⁰B shell and it produces the standard' neutron field at the centre of the cavity.

The effective spectrum range $E_5\%$ - $E_{95\%}$, where the fraction of spectrum equals 5% and 95% of the total neutron fluence, lies between 8 keV and 4.1 MeV, an averaged energy $\langle E \rangle$ equals to 970 keV. The spectrum-averaged cross sections were measured at ISNF assembly operated at the research reactor of the U.S. National Bureau of Standards (Gaithersburg) [120–122].

The ISNF spectrum available in IRDF-2002, as seen in Figs. 90 and 91, has obviously unphysical irregularities in the vicinity of 0.4 MeV marked with question marks in Fig. 90. Failing to find the original data, we went back to the available description of the facility [120]. The detailed information documented there gave us an opportunity to conduct a re-analysis of the spectrum using the MCNP-6 code with neutron transport cross sections including ²³⁵U(*n_{th}*,*f*) PFNS from the ENDF/B-VIII.0 library. The

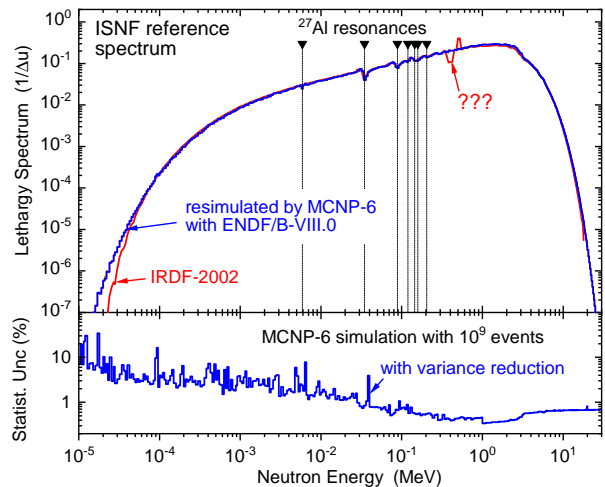


FIG. 90. (Color online) The ISNF reference spectrum: original from IRDF-2002 (red curve, the question marks point to unphysical oscillations) and recalculated in this work (blue). Arrows with dashed lines indicate the resonance energies of the Al total cross sections. The bottom part shows the statistical uncertainties obtained during the Monte Carlo simulation with variance reduction techniques.

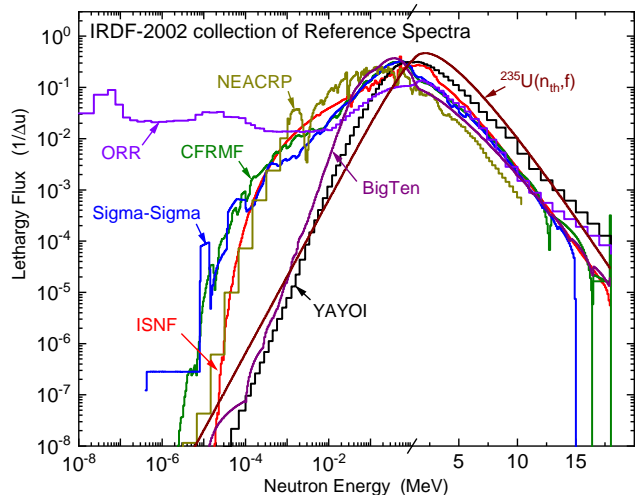


FIG. 91. (Color online) Neutron reference reactor spectra available in the IRDF-2002 collection [131]. Note that the scale changes from log to linear above 1 MeV.

calculated neutron energy distribution within the irradiation volume is shown in Fig. 90. This figure shows that the recalculated energy distribution is very close to the ISNF spectrum representation given in the IRDF-2002 library. It indeed exhibits resonances caused by the Al cap of the boron shell, but no longer exhibits the oscillations in the vicinity of 0.4 MeV.

This re-simulated ISNF spectrum, calculated in the 725-groups presentation, is used to compute the SACS and to support the validation of the IRDF-II cross sections (see Sec. VIII for details). The indicated numerical uncertainties only capture the statistical uncertainty of the Monte Carlo simulation and do not represent the complete spectrum uncertainty. To reach a similar numerical uncertainty over the whole neutron energy range, at the 1% level, variance reduction techniques were used. The

impact on statistical uncertainty of the variance reduction is shown in the bottom panel of Fig. 90.

11. Sigma-Sigma

Sigma-Sigma is a thermal-fast coupled spherical source assembly located within a conventional graphite thermal column of fission reactor. The column is a source of thermal neutrons that produce the fast fission spectrum in the spherical natural uranium shell (thermal-fast converter). The inner boron carbide shell shapes the neutron low-energy tail in such a way to increase the reaction rates within an appropriate energy range.

The effective spectrum range, $E_5\%$ - $E_{95\%}$ fluence metric, is 20 keV - 3.7 MeV, with an averaged energy $\langle E \rangle = 730$ keV. The Sigma-Sigma assemblies were operated at the Imperial College of Science and Technology (London) [123, 124], at the Belgian Nuclear Research Centre SCK•CEN, in Mol (Belgium) [125, 126], and at the Institute for Nuclear Technology (Bucharest) in cooperation with SCK•CEN [127, 128].

The Sigma-Sigma reference spectrum available in IRDF-2002 exhibits a suspicious spectrum shape below 10 keV, as seen in Fig. 91. The paper of A. Fabry and co-workers in 1975 [125] provided a tabulated spectrum which was based on the measurement and discrete ordinate calculation with nuclear data available at that time. Comparison of these two spectra shown in Fig. 92 confirms the unreliability of the IRDF-2002 data below ~ 10 keV.

Fortunately, several papers [125, 129, 130] have documented detailed specifications of the Sigma-Sigma assemblies. Using this information, we have constructed an MCNP model and re-simulated the neutron spectrum for this facility while employing the modern neutron fission and transport data from the ENDF/B-VIII.0 library. The computed spectrum is close to the original one by Fabry and the IRDF-2002 spectrum over most of the energy range. However, substantial differences are observed below 1 keV, the region where the resonances of the $^{238}\text{U}(n,\text{tot})$ cross sections produce a fine structure in the spectrum. The newly simulated Sigma-Sigma spectrum was used for the validation of IRDF-II cross sections, as reported in Sec. VIII. The indicated uncertainties only represent the numerical statistical contributions and have been reduced to a few percent by applying variance reduction techniques.

12. FNS-Graphite Neutron Field with a D-T Source

An experiment was performed at the JAEA-FNS facility in Tokai-mura, Ibaraki-ken, Japan to measure spectral indices in a graphite block irradiated with a D-T source. Reactor-grade graphite blocks were stacked to form a pseudo-cylindrical assembly of 31.4 cm in equivalent radius and 61.0 cm in thickness. The average density of the graphite blocks was 1.64 g/cm³. The graphite assembly was placed at a distance of 20.3 cm from the DT neutron source. Several foils for the reaction rate measurement of

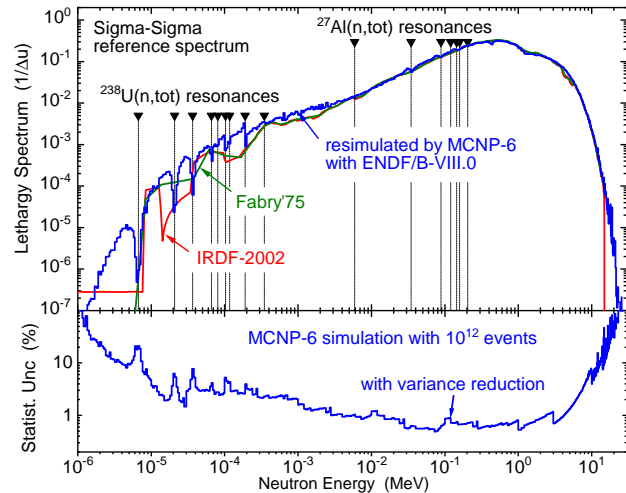


FIG. 92. (Color online) The Sigma-Sigma reference spectrum: original from IRDF-2002 (red curve), published by Fabry [125] (green) and recalculated in this work (blue). The bottom part shows the statistical uncertainties obtained in the Monte-Carlo simulations with variance reduction techniques.

forty dosimetry reactions were inserted into small spaces between the graphite blocks along the central axis of the assembly; the reaction rates of the $^{93}\text{Nb}(n,2n)^{92m}\text{Nb}$, $^{115}\text{In}(n,n')^{115m}\text{In}$ and $^{197}\text{Au}(n,\gamma)^{198}\text{Au}$ reactions were measured at four points (9.6, 19.3, 29.3 and 39.5 cm in depth) in the graphite assembly in order to check the reproducibility of the previous in situ experiment on graphite, while those of the other reactions were measured at two points (9.6 and 29.3 cm in depth). The plot of the spectra corresponding to the two depths in graphite are shown in Fig. 93.

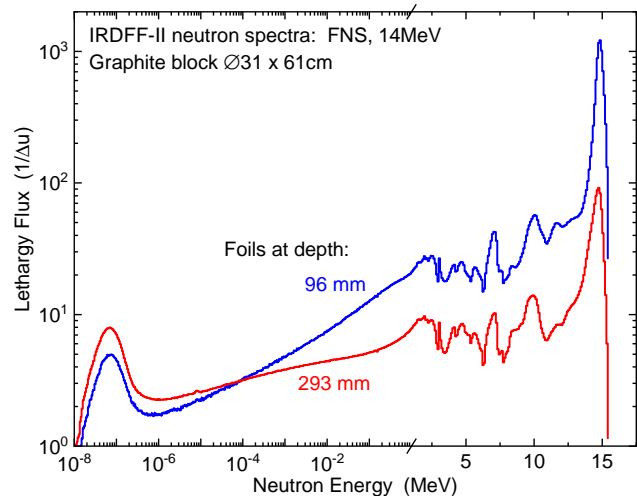


FIG. 93. (Color online) The FNS-Graphite reference spectrum from a D-T neutron source at 96 mm and 293 mm depth in the graphite block. Note that the scale changes from log to linear above 1 MeV.

13. Godiva Neutron Field from the ICSBEP Handbook

The Los Alamos National Laboratory HEU-MET-FAST-001 (*i.e.*, Godiva) critical assembly [132] consists of a number of discrete highly-enriched uranium metal pieces that produce a near spherical assembly when joined into a critical configuration. This assembly is modelled as a bare sphere with a 8.7404 cm radius. Reaction rate measurements were made at the assembly centre using a combination of fission counters and foils. Flux and reaction rate tallies were extracted from a detailed MCNP 6.2 [63] calculation. The tally region consists of a 0.5 cm diameter sphere centered within the critical assembly and the calculated flux and reaction rates result from a 10 billion neutron history job (50 warm-up cycles followed by 4 000 active cycles with 2 500 000 neutrons per cycle). Cross sections for the criticality transport calculation are from ENDF/B-VIII.0 [30] using 293.6 K ACE files generated by the MCNP Data Team [133]. Godiva's calculated flux spectrum for this central region is shown in Fig. 94.

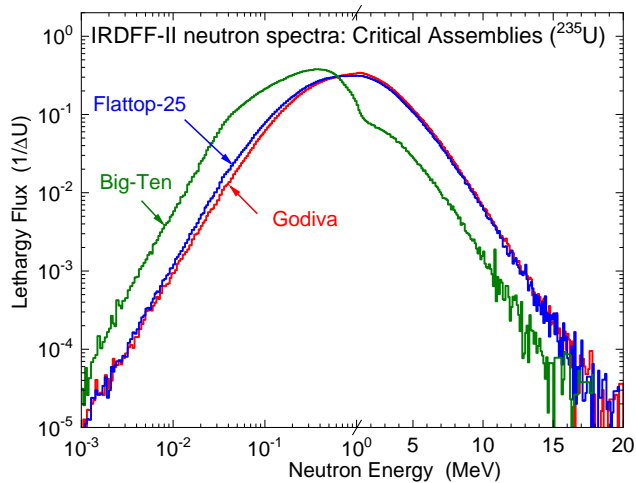


FIG. 94. (Color online) Neutron spectra for several critical assemblies containing ^{235}U from the ICSBEP Handbook [136]: Godiva, Flattop-25 and Big-Ten. The spectra are presented in the 725 energy groups and normalized to unity. Note that the scale changes from log to linear above 1 MeV.

14. Flattop-25 Neutron Field from the ICSBEP Handbook

The Los Alamos National Laboratory HEU-MET-FAST-028 (*i.e.*, Flattop-25) critical assembly [134] consists of a highly-enriched uranium metal central core surrounded by a large natural uranium metal reflector. When assembled in a critical configuration the core and reflector are nearly spherical in shape. Hence this assembly is modelled as a bare sphere of highly-enriched uranium surrounded by a large, spherical natural uranium reflector. The central core radius is 6.1156 cm while the spherical reflector radius is 18.0086 cm. There is no gap between the central core and the reflector in the computer model. Reaction rate measurements were made using fission counters and foils at or near the core centre with selected mea-

surements performed at various radial distances from the core centre extending into the reflector. In this report we focus on the central core. The tally region consists of a 0.5 cm diameter sphere centered within the critical assembly and the calculated flux and reaction rates result from a 5 billion neutron history MCNP 6.2 job (50 warm-up cycles followed by 2 000 active cycles with 2 500 000 neutrons per cycle). Cross sections for the kcode transport job are from ENDF/B-VIII.0 [30] using 293.6 K ACE files generated by the MCNP Data Team [133]. The Flattop-25 central region multigroup calculated flux spectrum is shown in Fig. 94.

15. Big-Ten Neutron Field from the ICSBEP Handbook

The Los Alamos National Laboratory IEU-MET-FAST-007 (*i.e.*, Big-10) critical assembly [135] is a large cylindrical assembly consisting of a heterogeneous mix of solid and annular uranium metal plates with enrichments vary from “depleted” to highly-enriched. The assembly has a central region consisting of 10 % enriched uranium where reaction rate and flux measurements were made. The calculational model incorporates the uranium metal plates so that the various “depleted”, normal uranium, 10 % enriched uranium and highly-enriched uranium regions are accurately portrayed. For calculated flux and reaction rate tallies a small, 0.5 cm diameter sphere is situated within the 10 % enriched region where the fission counters and foils were positioned. The MCNP 6.2 calculation was run for 5 billion active histories (50 warm-up cycles followed by 2 000 active cycles with 2 500 000 neutrons per cycle). Cross sections for the kcode transport job are from ENDF/B-VIII.0 [30] using 293.6 K ACE files generated by the MCNP Data Team [133]. The calculated multigroup flux spectrum within this tally sphere is shown in Fig. 94.

16. CFRMF

The Coupled Fast Reactivity Measurements Facility (CFRMF) is located at Idaho National Laboratory (INL) in Idaho Falls, Idaho, U.S.A. The CFRMF is a zoned core critical assembly with a fast-neutron spectrum zone in the centre of an enriched ^{235}U and ^{10}B annulus inside the uranium block [138–140]. The water moderated thermal driver zone are conventional plate-type fuel elements of fully enriched uranium clad in aluminum.

In the past there was a continuing effort to improve the characterization of the CFRMF spectrum both through calculation and spectrometry measurements. Different deterministic and Monte-Carlo radiation transport computer codes have been employed in the calculations. J.W. Rogers and co-workers [140] have published the results of a one-dimensional calculation with S6/P1 approximation and 71-group average cross sections from ENDF/B-IV covering the energy range 16 MeV to epithermal. This spectrum, as seen in Fig. 95, agrees with the spectrum assigned for CFRMF in the IRDF-2002 which is represented there at 461 energy points. Exceptions include

three points that deviate from the smooth shape above 10 MeV as seen in Fig. 91.

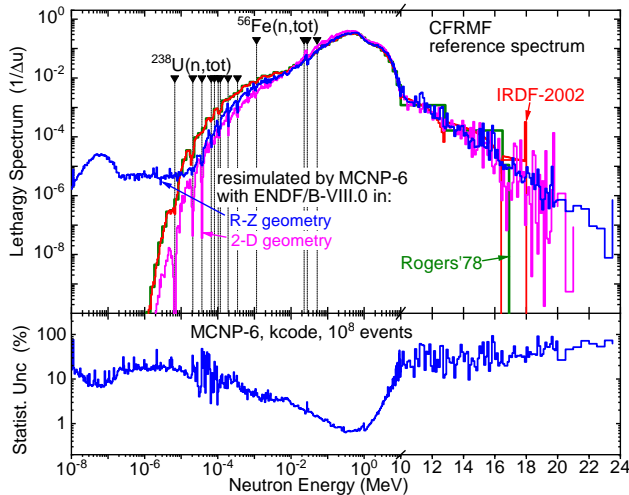


FIG. 95. (Color online) The CFRMF reference spectrum: original from IRDF-2002 (red curve), published by Rogers [140] (green) and recalculated in this work using 2-D (pink) and R-Z (blue) models. The bottom part shows the statistical uncertainties obtained in the Monte-Carlo criticality simulations with variance reduction. Note that the scale changes from log to linear above 1 MeV.

The one-dimensional model of the CFRMF facility is documented in a published paper [138]. We used this model to re-calculate the CFRMF critical assembly by the MCNP-6 code and employing up-to-date data from ENDF/B-VIII.0. As seen in Fig. 95, the newly calculated spectrum demonstrates more fine structure resulting from the strongest resonances in the neutron total cross section for ^{235}U and ^{56}Fe (component of stainless steel claddings). However, our calculation is lower than the IRDF-2002 and Roger [140] spectra for energies below ~ 10 keV. Another and more realistic model in the R-Z geometry was published in [140]. It resulted in a representation of the CFRMF spectrum that is very similar to the cylindrical representation, shown in Fig. 95, but additionally brings in the thermal and epithermal neutrons, which penetrate through the bottom and top ends of the access tube that accommodates the activation samples.

Since the reason for this systematic difference is not clear at this time, we are investigating refinements in the model geometry and have continued to use the IRDF-2002 spectrum (after correction of values at energies 12.7, 16.3 and 17.9 MeV) for the calculation of the spectrum-averaged cross sections. Calculation of the spectrum with this model predicts that the 5 % and 95 % limits are between 15 keV and 2.8 MeV and its average energy $\langle E \rangle$ equals 741 keV.

17. IPPE-BR1 Neutron Field from the ICSBEP Handbook

The Institute of Physics and Power Engineering FUND-IPPE-FR-MULT-RRR-001 is a Pu metal fuelled critical assembly [137] located in Obninsk, Russia. The core consists of (i) an empty centre tube where foils to be irra-

diated are located, (ii) 77 Pu metal rods are set on a hexagonal pitch, (iii) 31 peripheral copper metal rods to fill out the inner core region, (iv) annular uranium metal cylinders used for reactivity control and as radial reflectors, and (v) axial reflectors of uranium metal (lower) and copper metal (upper). The active fuel region is 13.1 cm tall. The MCNP 6.2 model includes a tally region at the core mid-plane in the voided centre rod position. This tally region is 1 cm tall. The MCNP 6.2 model was run for 5 billion active histories (50 warm-up cycles followed by 2 000 active cycles with 2 500 000 neutrons per cycle). Cross sections for the kcode transport job are from ENDF/B-VIII.0 [30] using 293.6 K ACE files generated by the MCNP Data Team [133]. The calculated multi-group flux spectrum within this tally sphere is shown in Fig. 96.

D. Potential Reference Neutron Fields from Detailed Computational Models

The neutron fields for several well-known Pu-fuelled critical assemblies are shown in this section; namely the LANL “Jezebel” (a bare, mostly ^{239}Pu , sphere), “Jezebel-240” (a bare Pu sphere with higher ^{240}Pu content), “Flat-top-Pu” (a Jezebel-like Pu spherical core reflected by natural Uranium) and “Thor” (a Jezebel-like Pu sphere reflected by Thorium). Criticality calculations yield accurate k_{eff} predictions but as will be shown in Sec. VIII K the reaction rate C/E results exhibit trends not seen in the Uranium fuelled assemblies. This suggests that some aspect of these Pu-fuelled assemblies, perhaps the ^{239}Pu PFNS, has unrecognized deficiencies. Hence while we present our calculations for these assemblies, the reader is cautioned that these neutron fields are not judged to be of benchmark quality at this time.

1. Jezebel Neutron Field from the ICSBEP Handbook

The Los Alamos National Laboratory PU-MET-FAST-001 (*i.e.*, Jezebel) critical assembly [141] consists of a number of discrete Pu metal pieces that, when assembled into a critical configuration, is nearly spherical in shape. The Pu metal is greater than 95 % ^{239}Pu with an approximate 1 % Ga alloy for phase stabilization. This benchmark evaluation has undergone significant revisions in recent years which allow for the development of complex computer models representing the individual Pu metal pieces. In addition, the simpler legacy model, a simple sphere with one material has been retained. This simple model, with its updated core radius and material data, is used here. Although the core model is 6.39157 cm in radius and the flux and reaction rate tallies are obtained for a 0.5 cm diameter spherical tally region located at the core centre. The MCNP 6.2 calculation was run for 25 billion active histories (50 warm-up cycles followed by 10 000 active cycles with 2 500 000 neutrons per cycle). Cross sections for the kcode transport job are from ENDF/B-VIII.0 [30] using 293.6 K ACE files generated by the MCNP Data Team [133]. The calculated multi-

group flux spectrum within this tally sphere is shown in Fig. 96.

2. Jezebel-240 Neutron Field from the ICSBEP Handbook

The Los Alamos National Laboratory PU-MET-FAST-002 (*i.e.*, Jezebel-240) critical assembly [142] is similar to the PMF001 assembly described above; the primary difference being in the Pu metal isotopic content. Whereas Jezebel is >95 % ^{239}Pu , the Jezebel-240 assembly is a little less than 80 % ^{239}Pu , approximately 20 % ^{240}Pu and includes smaller amounts of ^{241}Pu and ^{242}Pu . This assembly is modelled as a bare sphere 6.6595 cm in radius. Once again a small, 0.5 cm diameter, tally region is defined at the core centre where the various measurements were made. The MCNP 6.2 calculation was run for 10 billion active histories (50 warm-up cycles followed by 4 000 active cycles with 2 500 000 neutrons per cycle). Cross sections for the kcode transport job are from ENDF/B-VIII.0 [30] using 293.6 K ACE files generated by the MCNP Data Team [133]. The calculated multigroup flux spectrum within this tally sphere is very similar to the one of Jezebel in Fig. 96 and is not shown explicitly.

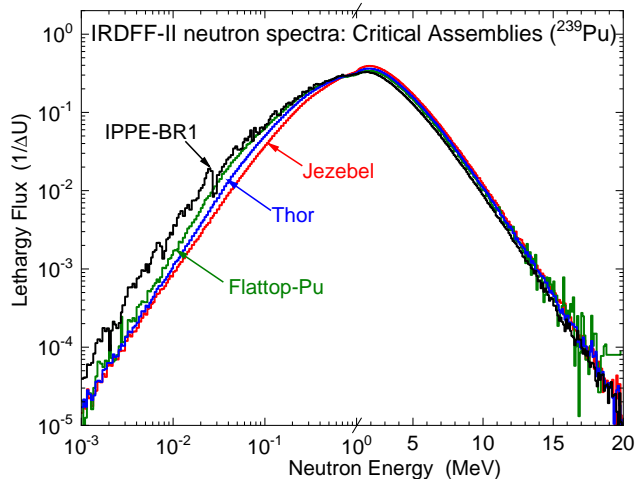


FIG. 96. (Color online) Neutron spectra for several critical assemblies containing ^{239}Pu from the ICSBEP Handbook [136]: Jezebel, Flattop-Pu, Thor and IPPE-BR-1. The spectra are presented in the 725 energy groups and normalized to unity. Note that the scale changes from log to linear above 1 MeV.

3. Flattop-Pu Neutron Field from the ICSBEP Handbook

The Los Alamos National Laboratory PU-MET-FAST-006 (*i.e.*, Flattop-Pu) critical assembly [143] is similar to the Flattop-25 assembly described previously with the exception that the central core is Pu metal rather than highly enriched uranium. The spherical Pu core radius is 4.5332 cm and the outer reflector radius is 24.1420 cm. As with Flattop-25 there is no gap between the central core and the reflector in the computer model. The central core flux and reaction rate tallies are obtained for a

0.5 cm diameter sphere at the located at the centre of the core. An MCNP 6.2 calculation was run for 1 billion histories (50 warm-up cycles followed by 400 active cycles with 2 500 000 neutrons per cycle). Cross sections for the kcode transport job are from ENDF/B-VIII.0 [30] using 293.6 K ACE files generated by the MCNP Data Team [133]. The calculated multigroup flux spectrum within this tally sphere is shown in Fig. 96.

4. THOR Neutron Field from the ICSBEP Handbook

The Los Alamos National Laboratory PU-MET-FAST-008 (*i.e.*, THOR) critical assembly [144] is similar to the Flattop-Pu assembly described previously with the exception that the heavy metal reflector is a cylinder of Thorium metal. The spherical Pu core radius is 5.310 cm and centered in the Thorium cylinder whose diameter and height are both 53.34 cm. Once again there is no gap between the core and reflector in the computer model. The MCNP 6.2 job includes a small, 0.5 cm diameter sphere in the centre of the core for flux and reaction rate tallies and was run for 5 billion active histories (50 warm-up cycles followed by 2 000 active cycles with 2 500 000 neutrons per cycle). Cross sections for the kcode transport job are from ENDF/B-VIII.0 [30] using 293.6 K ACE files generated by the MCNP Data Team [133]. The calculated multigroup flux spectrum within this tally sphere is very similar to the one of Jezebel in Fig. 96.

E. Legacy Reference Neutron Fields

The reference spectra incorporated within the IRDF-II library build upon the spectra collected in earlier dosimetry libraries. The reference neutrons fields generated at the research reactors, and frequently used for measurement of the spectrum-averaged dosimetry cross sections, were collected and documented in the IRDF-2002 database [131]. Some of these reference spectra were transferred over from the IRDF-90 [145]. The IRDF-2002 library included 9 spectra. The spectra, and associated references are:

- two ^{235}U thermal induced fission spectra, one from NBS (J. Grundl, private communication), second – from the ENDF/B-V evaluation;
- Intermediate-energy Standard Neutron Field (ISNF) [120];
- Coupled Fast Reactivity Measurement Facility (CFRMF) [120];
- 10 % enriched Uranium critical assembly (Big-Ten) [120];
- Coupled thermal/fast uranium and boron carbide spherical assembly (Sigma-Sigma) [120];
- Oak Ridge Research Reactor (ORR) [L. Greenwood, private communication];

- YAYOI reactor [L. Greenwood, private communication];
- Central zone flux of the NEACRP Benchmarks [B. Goal, private communication].

No uncertainties or associated correlation matrices for these spectra were provided in IRDF-2002. In 2005, the ^{252}Cf spontaneous fission neutron spectrum and its covariance matrix, evaluated by H. Mannhart [146], were added to IRDF-2002.

All compiled IRDF-2002 spectra were displayed in Fig. 91 above. It is worthwhile to stress that systematic measurements of SACS were carried out and consequently the corresponding data were compiled in the EXFOR database only for the following reference spectra: ISNF, CFRMF, Sigma-Sigma.

Only a subset of these spectra was used for the verification of the IRDF-II cross sections. Big-Ten critical assembly, ISNF and Sigma-Sigma reactor facilities were re-simulated using the latest ENDF/B-VIII.0 evaluation as described in Sects. VIC 10, VIC 11, and VIC 15, respectively. The CFRMF spectrum could not be reproduced in full, but because of the large number of measurements in this field the original spectrum provided by the authors for IRDF-2002 was adopted as described in Sec. VIC 16.

In ORR and YAYOI spectra unexplained structures are observed and the information describing the irradiation set-up is insufficient to develop new models. Therefore those IRDF-2002 spectra were not considered for validation. The Central zone flux of the NEACRP Benchmarks is not a real benchmark field and was discarded.

F. Analytical Reference Neutron Fields

1. Thermal Maxwellian

The thermal Maxwellian spectrum can be used to calculate spectrum-averaged cross sections $\bar{\sigma}$ in a well-thermalised spectrum. The spectrum corresponds to a temperature of 0.0253 eV and is normalised to $2/\sqrt{\pi}$ so that the averaged cross section can be compared to the σ_0 cross section value at 0.0253 eV, the ratio $\bar{\sigma}/\sigma_0$ being the Westcott g -factor.

2. $1/E$ Spectrum in the Energy Range 0.55 eV to 2 MeV

From neutron slowing-down theory it can be shown that high-energy neutrons in an ideal non-absorbing moderator with a constant cross section follow a $1/E$ distribution down to thermal energies. This is a reasonable approximation for a reactor spectrum. Cadmium has a very strong resonance near 0.2 eV. A cadmium cover of thickness 1 mm effectively filters thermal neutrons below 0.55 eV. At high energies the fission spectrum in a reactor falls off rapidly above 2 MeV. The $1/E$ spectrum defined in the energy range 0.55 eV to 2 MeV is a reasonable approximation for resonance integral calculations in a reactor spectrum under a cadmium cover, which can

be compared to measured values by the cadmium ratio method, for example.

3. $1/E$ Spectrum in the Energy Range 0.5 eV to 20 MeV

Some authors prefer different energy boundaries to define the resonance integral. In the Atlas of Resonance Parameters by Mughabghab [71], the integration boundaries are from 0.5 eV to 20 MeV. For convenience, both forms of the $1/E$ neutron spectrum are included in the spectra file of IRDF-II.

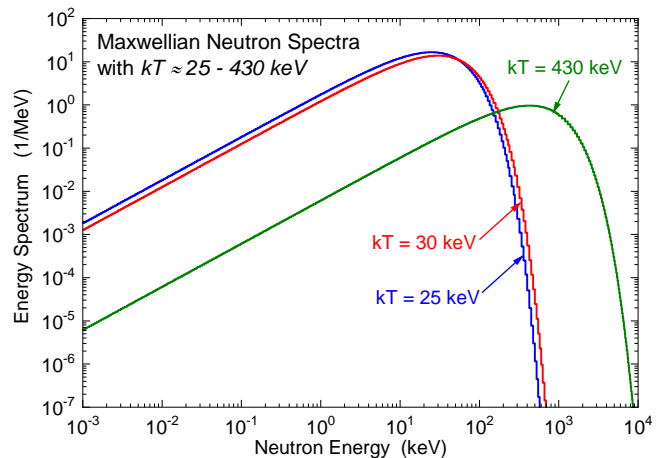


FIG. 97. (Color online) Maxwellian spectra with E_{Lab} ranging from 25 keV to 430 keV.

4. High Temperature Maxwellian

A set of high temperature quasi-Maxwellian neutron spectra, $kT = E \approx 25$ –60 keV, are frequently used for studying the neutron capture reactions along the astrophysical s-process path [147]. While the ideal spectra are defined by the analytic form, these neutron spectra can be approximately realized using the thick target $^7\text{Li}(p,n)$ reaction for incident proton energies near 2 MeV with modulation of the proton beam. The Maxwellian Averaged Cross Sections (MACS) for activation reactions are typically reported by the astrophysical community, but it must be remembered that the spectrum temperature is expressed in the centre-of-mass (CM) coordinate system, while the incident particle energy is normally given in the Lab system. Fortunately, the conversion is fairly straightforward. It can be shown that:

$$\bar{\sigma}(E_{\text{CM}}) = \bar{\sigma}(E_{\text{Lab}} \times (A + 1)/A), \quad (1)$$

where A is the ratio of the target mass to that of a neutron. For example, for hydrogen:

$$\bar{\sigma}(E_{\text{CM}} = 30\text{keV}) = \bar{\sigma}(E_{\text{Lab}} \approx 60\text{keV}) \quad (2)$$

A range of Maxwellian neutron spectra, with E_{Lab} ranging from thermal to 60 keV, are defined within the IRDF-II library and can be used to support validation activities. Fig. 97 shows a representative sample of these spectra.

5. Constant

The constant $\phi = 1$ is a trivial field that allows simple averages to be calculated with the same tool. Namely, it allows the calculation of flat-spectrum average cross sections or the spectrum integrals when applied to the IRDF-II library of neutron spectrum fields.

6. Linear

The linear spectrum $\phi = E$ is a trivial field that allows the calculation of the average spectrum energy when applied to a spectrum in the library of spectrum fields in ENDF-6 format.

G. Thermal Cross Sections and Resonance Integrals

For all of the non-threshold reactions, the thermal cross sections and resonance integrals in the IRDF-II library entries have been compared to the data from the 2016 Atlas of Thermal Neutron Capture Cross Sections Resonance Integrals and Thermal Cross Sections [71]. Since in many cases these data were considered in the development of the underlying nuclear data evaluations, this data comparison constitutes a verification step in the library development.

An independent comparison can be made to the values derived from the Kayzero data-base for neutron activation analysis (NAA) by the k_0 standardization method.

The Kayzero data-base is a library of nuclear constants that has been developed over many years and has been extensively tested in practical application for analytical work in certified laboratories. Most of the k_0 and Q_0 factors in the library have been measured independently of any differential cross section, so they provide a convenient source of information for the validation of thermal capture cross sections and resonance integrals. The k_0 factors are related to the cross sections:

$$k_0 = \frac{M_s \Theta_a \gamma_a \sigma_{0,a}}{M_a \Theta_s \gamma_s \sigma_{0,s}} \quad (3)$$

where M is the molar mass, Θ is the isotopic abundance, γ is the gamma-emission probability and σ is the cross section. Subscript a stands for the measured nuclide and s for the gold standard.

The Q_0 factors are simply the ratios of the resonance integral I_0 in a pure $1/E$ spectrum and the thermal cross section:

$$Q_0 = \frac{I_0}{\sigma_0} \quad (4)$$

Comparison of the cross sections and resonance integrals in the IRDF-II library to the values derived from the Kayzero data-base [69] provides an alternative source of validation for the thermal constants and the resonance integrals and augments the comparison with information provided in the Atlas of Resonance Parameters by Mughabghab [71].

In this implementation of k_0 -based metrics, the k_0 factors from the Kayzero library, the natural abundances and the gamma emission probabilities from the IRDF-II decay data library are combined and used to reconstruct the thermal cross sections and resonance integrals. In principle, the derived values correspond to idealized thermal reactor spectrum with a Maxwellian thermal component and a $1/E$ epithermal component. The deviation from a $1/v$ behaviour of the cross sections in the thermal range, deviation from the $1/E$ spectrum shape in the epithermal range and the non-ideal cadmium filter are taken into account (although not rigorously in all cases), but these are corrections that are usually smaller than the experimental uncertainties. The thermal cross sections and resonance integrals derived from the Kayzero library were taken as the reference for comparison with the Atlas values and the quantities as extracted from the IRDF-II library. Results from this validation step are presented in Sec. VIII.

VII. VALIDATION DATA – COMPARISON OF INTEGRAL METRICS IN STANDARD AND REFERENCE NEUTRON BENCHMARK FIELDS

One of the most powerful metrics employed in the validation of the IRDF-II cross sections is the comparison of calculated-to-experimental (C/E) spectrum-averaged cross sections in various standard and reference benchmark neutron fields. Since an absolute determination of the neutron fluence can be difficult to assess for some of the reactor fields, a spectral index (SI), or the ratio of the activity for a given reaction to that for a common monitor reaction, is often used. The C/E ratio for the spectral indices is then the relevant metric used for validation. Another metric that supports validation is a comparison of the consistency of the relative activity for a set of measurements gathered in a given well-characterized neutron field. The following sections address the consideration of these integral metrics in the benchmark neutron fields.

The criteria that were adopted to characterize the quality of the validation evidence based on the comparison of the measured and calculated of the spectrum-averaged cross section are shown in Table 17. A “good” validation status requires, as a baseline criterion, that the calculated-to-experimental (C/E) ratio of the spectrum-averaged cross section be within two standard deviations of unity and, in addition, that it lies within the interval of [0.9: 1.1]. In addition to this characteristic, in order to ensure the experimental measurement had adequate fidelity we required that the experimental uncertainty be less than 15 %. The total uncertainty in the C/E must reflect the root-mean-squared sum of the measurement uncertainty, the spectrum uncertainty, and the cross-section uncertainty. To be sure that the benchmark neutron field was adequately characterized, we also required that the total uncertainty be less than 25 %.

A “good” validation status is color-coded green. An “acceptable” validation status is color-coded yellow and reflects a C/E ratio within three standard deviations of unity, a value within the interval [0.8, 1.25], a measurement uncertainty less than 30 %, and a total uncertainty

TABLE 17. Criteria used to evaluate consistency validation evidence based on a least-square adjustment analysis for various reactions.

#	Metric	Status		
		Good	Acceptable	Poor
1	C/E	≤ 2 std. dev.	≤ 3 std. dev.	> 3 std. dev.
2	C/E Interval	within [0.9,1.1]	within [0.8,1.25]	outside [0.8,1.25]
3	Experimental Uncertainty	≤ 15 %	≤ 30 %	> 30 %
4	Total Uncertainty	≤ 25 %	≤ 40 %	> 40 %

less than 40 %. A “poor” validation status is color-coded as red and corresponds to a C/E that is more than three standard deviations from unity, or a value that is outside the interval [0.8, 1.25]. A measurement uncertainty greater than 30 % or a total uncertainty greater than 40 % does not necessarily reflect a “poor” validation state, but it does reflect an absence of a meaningful validation. These fields are left uncolored to reflect unknown status due to the poor quality of available information – either the measurement uncertainty or the neutron field characterization.

A. ^{252}Cf SFNS

The standard ^{252}Cf SFNS spectrum was evaluated purely from differential time-of-flight data and is independent of SACS. Therefore, the spectrum can be used to validate SACS (and vice versa). The SACS in the standard ^{252}Cf SFNS spectrum were evaluated mostly by Mannhart [149]. It is assumed that all acceptable SACS measurements available at the time were included in the evaluation. The measured values and the C/E values as a function of representative neutron energy $E_{50\%}$ (energy at which the reaction rate integral reaches 50 % of the integral over the full energy range) are shown in Table 18 and Fig. 98.

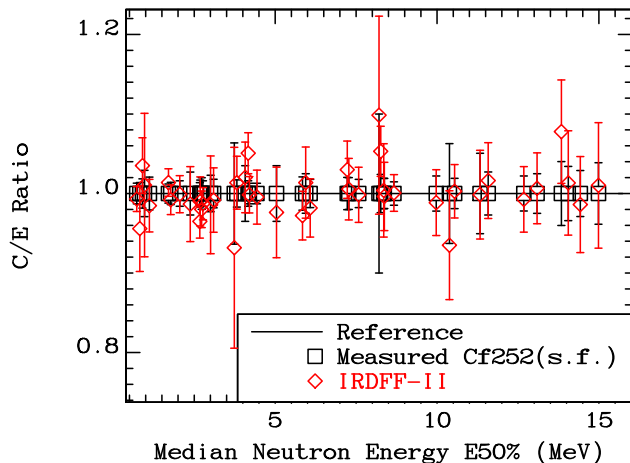


FIG. 98. (Color online) C/E relative to median energy $E_{50\%}$ % for the IRDF-II cross sections averaged in the ^{252}Cf (s.f.) neutron field. Plotted values are listed in Table 18.

More recently, new measurements were performed at Řež to confirm Mannhart evaluations. The intense ^{252}Cf source (about 10^9 n/cm²/s) at Řež is well characterized, and the activation measurements achieved low uncertain-

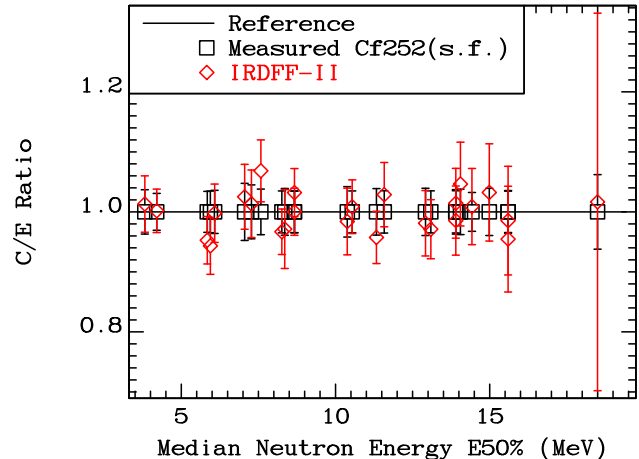


FIG. 99. (Color online) C/E relative to median energy $E_{50\%}$ for the IRDF-II cross sections averaged in the ^{252}Cf (s.f.) neutron field measured at Řež. Plotted values are listed in Table 19.

ties [159]. The results are shown in Table 19 and Fig. 99. The C/E uncertainties are colour-coded as defined in Table 17. The SACS in the standard ^{252}Cf SFNS spectrum have practically no outliers and no trend in energy is observed. A special case is the $^{169}\text{Tm}(n,3n)$ reaction (Tm1693), which has a very high threshold, with median energy >18 MeV, where the assigned uncertainty of the ^{252}Cf (s.f.) spectrum is very large due to the lack of measured data, and represents the dominant contribution to the large SACS uncertainty. Still, the C/E for that very special reaction is excellent.

TABLE 18. Measured and calculated spectrum averaged cross sections for $^{252}\text{Cf}(\text{s.f.})$ neutron standard field.

Reaction Notation	$E_{50\%}$ [MeV]	Measured SACS [mb]	Unc. [%]	Reference	Calculated SACS [mb]	Tot.Unc. [%]	X.S.Unc. [%]	C/E	Uncert. [%]	Diff. [%]
Au197g	0.724	75.5000	1.3	[148]	74.9780	0.90	0.52	0.9931	1.85	-0.69
Ta181g	0.818	87.3000	1.4	[148]	83.4430	5.47	5.41	0.9558	5.64	-4.42
Th232g	0.930	87.0000	1.8	[148]	93.4500	3.80	3.77	1.0741	4.22	7.41
Cu63g	0.964	10.3000	2.9	[148]	10.4090	8.44	8.41	1.0106	8.93	1.06
In115gm	1.116	125.6000	2.1	[149]	123.6800	2.64	2.61	0.9847	3.46	-1.53
U235f	1.703	1210.0000	1.2	[149]	1226.7000	1.21	1.21	1.0138	1.70	1.38
Pu239f	1.774	1812.0000	1.4	[149]	1797.8000	1.25	1.25	0.9922	1.85	-0.78
Np237f	2.053	1361.0000	1.6	[149]	1359.8000	1.70	1.69	0.9991	2.33	-0.09
Rh103m	2.380	734.3000	2.7	[146]	724.9200	3.95	3.94	0.9801	2.39	-1.99
In115nm	2.673	197.4000	1.4	[149]	190.4800	1.70	1.66	0.9649	2.18	-3.51
Nb93nm	2.685	147.5000	1.7	[87]	146.0300	2.61	2.59	1.0002	4.31	0.02
In113nm	2.730	161.2000	2.0	[146]	157.9900	1.24	1.18	0.9801	2.39	-1.99
U238f	2.768	325.7000	1.6	[149]	321.5400	1.29	1.22	0.9872	2.09	-1.28
Th232f	3.006	84.5500	2.3	[150]	83.3500	5.80	5.78	0.9864	8.20	-1.36
Hg199nm	3.098	298.4000	1.8	[149]	295.9400	3.66	3.63	0.9918	4.08	-0.82
BH3	3.737	11.0000	6.4	[151]	10.2500	11.97	11.96	0.9318	13.56	-6.82
Ti47p	3.817	19.2700	1.7	[149]	19.5330	2.80	2.73	1.0136	3.26	1.36
S32p	4.074	72.5400	3.5	[149]	74.0150	2.59	2.48	1.0203	4.35	2.03
Zn64p	4.167	40.5900	1.6	[149]	42.6470	1.86	1.69	1.0376	3.67	3.76
Ni58p	4.203	117.5000	1.3	[149]	117.3100	1.89	1.74	0.9984	2.29	-0.16
Fe54p	4.438	86.8400	1.3	[149]	86.4490	3.16	3.06	0.9955	3.43	-0.45
Pb204nm	5.041	20.8700	3.5	[149]	20.3740	4.67	4.57	0.9772	5.07	-2.28
Al27p	5.843	4.8800	2.1	[149]	4.7445	2.35	2.05	0.9722	3.18	-2.78
Co59p	5.943	1.6900	2.5	[149]	1.7132	3.65	3.46	1.0137	4.41	1.37
Ti46p	6.081	14.0700	1.8	[149]	13.8140	3.28	3.05	0.9818	3.73	-1.82
Si28p	7.226	6.9000	2.0	[152]	7.1069	2.85	2.47	1.0300	3.46	3.00
Cu63a	7.274	0.6887	2.0	[149]	0.6925	3.28	2.97	1.0055	3.82	0.55
Fe56p	7.579	1.4650	1.8	[149]	1.4626	2.98	2.60	0.9984	3.47	-0.16
U2382	8.208	19.2000	10.	[153]	21.0920	5.34	5.10	1.0442	12.13	4.42
Mg24p	8.260	1.9960	2.4	[149]	2.1020	1.78	0.80	1.0531	3.02	5.31
Ti48p	8.354	0.4247	1.9	[149]	0.4264	5.53	5.30	1.0039	5.84	0.39
Co59a	8.372	0.2218	1.9	[149]	0.2210	3.87	3.54	0.9963	4.30	-0.37
Al27a	8.668	1.0160	1.5	[149]	1.0167	1.77	0.71	1.0007	2.18	0.07
V51a	9.975	0.0390	2.2	[149]	0.0386	3.56	3.02	0.9885	4.19	-1.15
Tm1692	10.400	6.6900	6.3	[77]	6.2551	3.77	3.21	1.069	5.64	+6.90
Au1972	10.542	5.5060	1.9	[149]	5.5213	2.75	1.87	1.0028	3.30	0.28
Nb932m	11.328	0.7910	5.1	[149]	0.7899	2.38	0.84	0.9986	5.02	-0.14
I1272	11.580	2.0690	2.7	[149]	2.1027	3.83	3.03	1.0163	4.70	1.63
Cu652	12.680	0.6582	2.2	[149]	0.6533	3.52	1.89	0.9926	4.16	-0.74
Co592	13.090	0.4051	2.5	[149]	0.4078	3.66	1.52	1.0066	4.44	0.66
Cu632	13.840	0.1844	4.0	[149]	0.1987	4.51	1.38	1.0778	6.02	7.78
F192	14.052	0.0161	3.4	[149]	0.0163	5.50	2.90	1.0133	6.45	1.33
Zr902	14.424	0.2210	2.9	[149]	0.2179	5.39	0.91	0.9861	6.12	-1.39
Ni582	14.986	8.56E-3	3.6	[146]	8.65E-3	6.79	1.29	1.0102	7.81	1.02

B. $^{235}\text{U}(\text{n}_{\text{th}},\text{f})$ PFNS

The reference prompt fission neutron spectrum (PFNS) for $^{235}\text{U}(\text{n}_{\text{th}},\text{f})$ [98–100] was shown in Fig. 68 as ratio to the Maxwellian distribution with $kT = 1.32$ MeV. The uncertainties in that evaluated spectrum were constrained at high energy by the use of $^{90}\text{Zr}(\text{n},2\text{n})$ SACS point. Recommended SACS in the $^{235}\text{U}(\text{n}_{\text{th}},\text{f})$ PFNS are listed in Table 20. The C/E ratios of SACS in the reference $^{235}\text{U}(\text{n}_{\text{th}},\text{f})$ PFNS as a function of representative neutron energy $E_{50\%}$ are shown in Fig. 100. For the $^{235}\text{U}(\text{n}_{\text{th}},\text{f})$ PFNS field no trend in energy is observed. An outlier is the tritium production reaction on ^{10}B (B10H3) measured by Wyman in 1958. Measurements of the capture SACS are usually subject to large corrections for room-return and multiple scattering, which might be the cause of some discrepancy in the U238g reaction. Poorer

(but acceptable) agreement is also noted for $^{63}\text{Cu}(\text{n},2\text{n})$, $^{238}\text{U}(\text{n},2\text{n})$ and $^{204}\text{Pb}(\text{n},\text{n}')$ reactions. New extensive measurements of SACS in the reference $^{235}\text{U}(\text{n}_{\text{th}},\text{f})$ neutron field are on-going.

C. Ratio of ^{252}Cf SFNS to $^{235}\text{U}(\text{n}_{\text{th}},\text{f})$ PFNS Spectrum-averaged Cross Sections

The activation cross sections in the ^{252}Cf SFNS and $^{235}\text{U}(\text{n}_{\text{th}},\text{f})$ PFNS benchmark neutron fields discussed in Secs. VII A and VII B form the foundation for the IRDF-II validation. In addition to these two separate sets of validation data for a fission neutron spectrum, the smooth behaviour, *i.e.*, the systematics, of the ratios of these spectrum-averaged cross sections, as a function of the E_{50} response energy for a given reaction, provides

TABLE 19. New SACS for ^{252}Cf -252(s.f.) neutron benchmark field measured at Řež.

Reaction Notation	$E_{50\%}$ [MeV]	Measured SACS [mb]	Unc. [%]	Reference	Calculated SACS [mb]	Tot.Unc. [%]	X.S.Unc. [%]	C/E	Uncert. [%]	Diff. [%]
Ti47p	3.817	19.2800	3.70	[154]	19.5330	2.80	2.73	1.0131	4.64	1.31
Ni58p	4.203	117.1000	3.07	[154]	117.3100	1.89	1.74	1.0018	3.61	0.18
Al27p	5.843	4.9800	3.41	[155]	4.7445	2.35	2.05	0.9527	4.14	-4.73
Co59p	5.943	1.8160	3.47	[156]	1.7132	3.65	3.46	0.9434	5.04	-5.66
Ti46p	6.081	13.8450	3.60	[154]	13.8140	3.28	3.05	0.9978	4.87	-0.22
Ni60p	7.054	2.7300	4.76	[155]	2.7985	2.27	1.81	1.0251	5.28	2.51
Cu63a	7.274	0.6833	4.49	[154]	0.6925	3.28	2.97	1.0135	5.56	1.35
Fe56p	7.579	1.3690	3.80	[154]	1.4626	2.98	2.60	1.0684	4.83	6.84
Mg24p	8.260	2.1740	3.50	[154]	2.1020	1.78	0.80	0.9669	3.92	-3.31
Ti48p	8.354	0.4389	3.90	[154]	0.4264	5.53	5.30	0.9715	6.76	-2.85
Al27a	8.668	0.9851	3.45	[155]	1.0167	1.77	0.71	1.0321	3.88	3.21
Al27a	8.668	1.0165	3.49	[154]	1.0167	1.77	0.71	1.0002	3.92	0.02
Tm1692	10.400	6.3570	4.20	[156]	6.2551	3.77	3.21	0.9840	5.64	-1.60
Au1972	10.542	5.4740	3.49	[156]	5.5213	2.75	1.87	1.0086	4.44	0.86
Nb932m	11.328	0.8250	3.88	[156]	0.7899	2.38	0.84	0.9574	4.55	-4.26
I1272	11.580	2.0440	3.52	[157]	2.1027	3.83	3.03	1.0287	5.20	2.87
Mn552	12.900	0.4820	3.94	[156]	0.4729	3.92	2.32	0.9811	5.56	-1.89
Co592	13.090	0.4199	3.50	[156]	0.4078	3.66	1.52	0.9712	5.06	-2.88
Y892	13.902	0.3410	3.52	[155]	0.3459	4.53	1.24	1.0144	5.74	1.44
Y892	13.902	0.3510	3.70	[156]	0.3459	4.53	1.24	0.9855	5.85	-1.45
F192	14.053	0.0156	3.78	[155]	0.0163	5.50	2.90	1.0464	6.67	4.64
Zr902	14.424	0.2160	3.24	[155]	0.2179	5.39	0.91	1.0089	6.29	0.89
Ni582	14.987	8.37E-3	3.94	[154]	8.65E-3	6.79	1.29	1.0324	7.85	3.24
Na232	15.597	8.70E-3	3.45	[155]	8.57E-3	8.52	1.28	0.9854	9.19	-1.46
Na232	15.597	8.98E-3	3.56	[157]	8.57E-3	8.52	1.28	0.9547	9.24	-4.53
Tm1693	18.506	0.0145	6.21	[154]	0.0147	30.32	5.76	1.0166	30.95	1.66

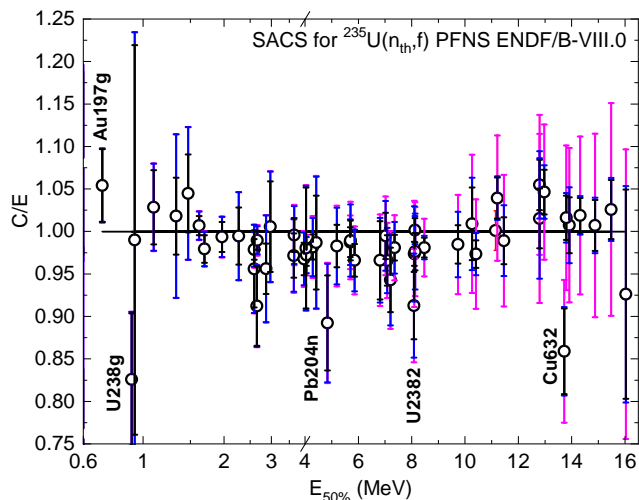


FIG. 100. (Color online) C/E as a function of median energy $E_{50\%}$ for the IRDF-II cross sections averaged in the $^{235}\text{U}(n_{th},f)$ PFNS field. Uncertainty bars reflect a r.m.s. combination of uncertainty contributions from the measurement (black), IRDF-II cross sections (blue) and spectra (pink). Plotted values are listed in Table 20.

another good indicator of the fidelity of the cross sections. This ratio is independent of the gamma emission probability, which is a significant source of measured SACS uncertainties. The application of this metric and a systematic of this ratio is addressed in Ref. [187]. The systematic can be used also to assess the quality of experimental data in one of the reference fields [187]. The actual status of C/E ratios computed with IRDF-II is depicted in Fig. 101. The ratio for the reaction $^{63}\text{Cu}(n,2n)$ is an

outlier indicating that one or both of the SACS measurements used to calculate the ratio are likely wrong. Indeed, it can be seen that the corresponding C/E of this particular reaction in the $^{235}\text{U}(n_{th},f)$ PFNS neutron field is 0.859(9.8%) pointing out to a possible experimental issue. A similar situation is found for the $^{238}\text{U}(n,2n)$ reaction, where the C/E in the $^{235}\text{U}(n_{th},f)$ PFNS neutron field is 0.913(7.3%), which also points to a potential experimental problem. Corresponding C/E in the ^{252}Cf SFNS field show good agreement for these two reactions.

D. Measurements in the SPR-III Fast Burst Reactor Central Cavity

The baseline measured quantities in this reference benchmark field are end-of-irradiation (EOI) spectrum-averaged cross sections measured in operation #13127, which was a 16-minute steady-state operation that took place on 1/27/2006 and represented a total facility-reported reactor integrated power of 125 MJ.

Although a baseline set of activation foils were used in this reactor operation #13127, it was desirable to add more reactions to the set of measured data, so additional reactor exposures of activation foils were conducted. In order to eliminate the effect of variations in the reactor power during the other exposures, a $^{58}\text{Ni}(n,p)^{58}\text{Co}$ monitor activity was used to facilitate a renormalization of data gathered in the other operations to the baseline exposure. When required, corrections were made in the measured activities of other foils to convert them to represent the end-of-irradiation from the baseline 16-minute uniform power operation. The other reactor operations

TABLE 20. Measured and calculated SACS for the IRDF-II cross sections in the $^{235}\text{U}(n_{th,f})$ neutron benchmark field.

Reaction	$E_{50\%}$	Measured SACS			Calculated SACS				C/E
Notation	MeV	Value, mb	Unc.,%	Reference	Value, mb	XS Unc.,%	Sp.Unc.,%	Tot.Unc.,%	
Au197g	0.705	7.400E+01	4.05	[173]	7.8011E+01	0.51	0.80	0.95	1.054 ± 4.16 %
U238g	0.907	8.500E+01	9.41	[173]	7.0197E+01	1.97	0.69	2.09	0.826 ± 9.64 %
Cu63g	0.932	1.080E+01	23.15	[173]	1.0693E+01	8.57	0.64	8.60	0.990 ± 24.70 %
In115gm	1.095	1.245E+02	4.25	[81]	1.2805E+02	2.61	0.58	2.67	1.029 ± 5.02 %
B10He4	1.329	5.410E+02	4.44	[130]	5.5077E+02	8.35	0.54	8.37	1.018 ± 9.47 %
Li6He4	1.472	4.560E+02	4.39	[130]	4.7641E+02	6.03	0.60	6.06	1.045 ± 7.48 %
U235f	1.618	1.217E+03	1.12	[146]	1.2253E+03	1.21	0.36	1.26	1.007 ± 1.69 %
Pu239f	1.691	1.831E+03	1.65	[146]	1.7933E+03	1.25	0.37	1.30	0.979 ± 2.10 %
Np237f	1.965	1.350E+03	1.78	[146]	1.3413E+03	1.58	0.47	1.75	0.994 ± 2.43 %
Rh103nm	2.267	7.103E+02	3.38	[74, 179, 180]	7.0643E+02	3.95	0.53	3.99	0.995 ± 5.23 %
Nb93n	2.589	1.476E+02	4.74	[174]	1.4109E+02	2.63	0.66	2.71	0.956 ± 5.46 %
In115nm	2.589	1.878E+02	1.23	[146]	1.8383E+02	1.68	0.69	1.82	0.979 ± 2.20 %
In113nm	2.651	1.668E+02	5.13	[81]	1.5217E+02	1.19	0.72	1.39	0.912 ± 5.31 %
U238f	2.659	3.094E+02	1.13	[146]	3.0618E+02	1.22	0.75	1.43	0.990 ± 1.82 %
Th232f	2.867	8.186E+01	3.10	[108]	7.8267E+01	5.79	0.78	5.84	0.956 ± 6.61 %
Hg199nm	2.972	2.787E+02	5.30	[79]	2.8024E+02	3.68	0.77	3.76	1.006 ± 6.50 %
B10H3	3.533	3.023E+01	6.70	[160]	4.7915E+01	12.37	0.85	12.40	1.585 ± 14.09 %
P31p	3.633	3.586E+01	2.68	[152]	3.4842E+01	3.47	1.06	3.63	0.972 ± 4.51 %
Ti47p	3.647	1.784E+01	1.99	[146]	1.7765E+01	2.79	0.99	2.96	0.996 ± 3.57 %
S32p	3.967	6.908E+01	1.97	[146]	6.6846E+01	2.53	1.14	2.78	0.968 ± 3.41 %
Zn64p	4.036	3.889E+01	7.25	[79]	3.8133E+01	1.74	1.20	2.12	0.981 ± 7.55 %
Ni58p	4.051	1.082E+02	1.30	[146]	1.0519E+02	1.75	1.14	2.09	0.972 ± 2.46 %
Fe54p	4.294	7.809E+01	1.50	[81]	7.6659E+01	3.16	1.24	3.39	0.982 ± 3.71 %
Zn67p	4.421	9.660E-01	5.59	[81]	9.5344E-01	5.52	1.19	5.64	0.987 ± 7.94 %
Pb204nm	4.839	1.946E+01	6.27	[182, 183]	1.7364E+01	4.66	1.42	4.87	0.892 ± 7.94 %
Mo92pm	5.190	6.687E+00	2.53	[81]	6.5723E+00	3.83	1.49	4.11	0.983 ± 4.83 %
Al27p	5.700	3.902E+00	1.77	[146]	3.8527E+00	3.49	12.23	2.66	0.987 ± 3.20 %
Co59p	5.703	1.396E+00	2.36	[146]	1.3812E+00	3.59	1.68	3.97	0.989 ± 4.62 %
Ti46p	5.862	1.151E+01	1.99	[146]	1.1118E+01	3.19	1.74	3.64	0.966 ± 4.15 %
Ni60p	6.811	2.180E+00	4.77	[148]	2.1062E+00	1.98	2.10	2.88	0.966 ± 5.57 %
Cu63a	7.019	5.173E-01	2.74	[146, 165]	5.1459E-01	3.09	2.16	3.77	0.995 ± 4.66 %
Si28p	7.079	5.470E+00	3.07	[152]	5.2831E+00	2.53	2.28	3.41	0.966 ± 4.59 %
Fe54a	7.205	8.650E-01	4.02	[74, 167, 168]	8.1560E-01	4.02	2.25	4.59	0.943 ± 6.10 %
Fe56p	7.362	1.079E+00	1.54	[146]	1.0583E+00	2.68	2.38	3.59	0.981 ± 3.91 %
U2382	8.087	1.600E+01	4.34	[184–186]	1.4604E+01	5.10	2.92	5.87	0.913 ± 7.30 %
Ti48p	8.103	3.014E-01	1.46	[146, 165]	2.9347E-01	5.57	2.84	6.25	0.974 ± 6.42 %
Mg24p	8.125	1.449E+00	1.43	[146, 165]	1.4512E+00	0.83	2.85	2.97	1.002 ± 3.30 %
Co59a	8.127	1.563E-01	2.25	[146]	1.5260E-01	3.99	2.81	4.88	0.976 ± 5.37 %
Al27a	8.471	7.005E-01	1.19	[146, 161]	6.8720E-01	0.74	3.16	3.24	0.981 ± 3.45 %
V51a	9.737	2.429E-02	2.29	[146]	2.3918E-02	3.17	4.46	5.48	0.985 ± 5.94 %
Tm1692	10.265	3.735E+00	4.23	[87]	3.7685E+00	3.34	6.01	6.88	1.009 ± 8.08 %
Au1972	10.414	3.387E+00	1.67	[146, 165]	3.2969E+00	1.93	6.11	6.41	0.973 ± 6.62 %
Nb932	11.210	4.348E-01	2.32	[81]	4.5189E-01	0.86	6.65	6.71	1.039 ± 7.10 %
I1272	11.459	1.199E+00	2.79	[79, 166]	1.1860E+00	3.16	6.60	7.32	0.989 ± 7.83 %
Mn552	12.796	2.356E-01	2.81	[146, 166]	2.4852E-01	2.48	6.84	7.28	1.055 ± 7.80 %
As752	12.797	3.207E-01	3.47	[165, 174, 175]	3.2561E-01	6.07	6.87	9.16	1.015 ± 9.80 %
Co592	12.974	2.028E-01	2.51	[146]	2.1220E-01	1.65	6.98	7.17	1.046 ± 7.60 %
Cu632	13.721	1.160E-01	5.88	[172]	9.9640E-02	1.44	7.69	7.82	0.859 ± 9.80 %
Y892	13.797	1.697E-01	2.34	[165, 175–178]	1.7274E-01	1.30	7.87	7.97	1.015 ± 8.37 %
F192	13.911	8.056E-03	3.26	[146, 161, 162]	8.1151E-03	2.99	7.86	8.41	1.007 ± 9.02 %
Zr902	14.320	1.041E-01	2.02	[146, 161, 175]	1.0607E-01	0.92	8.84	8.89	1.019 ± 9.12 %
Ni582	14.879	4.076E-03	2.98	[109, 169–171]	4.1047E-03	1.29	10.23	10.31	1.007 ± 10.73 %
Na232	15.483	3.860E-03	3.39	[162, 163]	3.9599E-03	1.27	11.65	11.72	1.026 ± 12.20 %
Al272	16.023	3.760E-03	13.30	[84]	3.4823E-03	2.06	1.69	12.72	0.926 ± 18.40 %

are detailed in Table 21.

In some of the cases covers were used to shield the activation foil and shift the sensitivity of the measured response to higher energies. These measurements used the following cover geometries:

1. “bare” indicating a bare foil with a thickness be-

tween 5 and 10 mils;

2. “Cd” indicating a cylindrical cadmium cover with thickness of 2.587×10^{-3} atoms/barn;
3. “Cdtk” indicating a cylindrical cadmium cover with thickness of 4.705×10^{-3} atoms/barn;

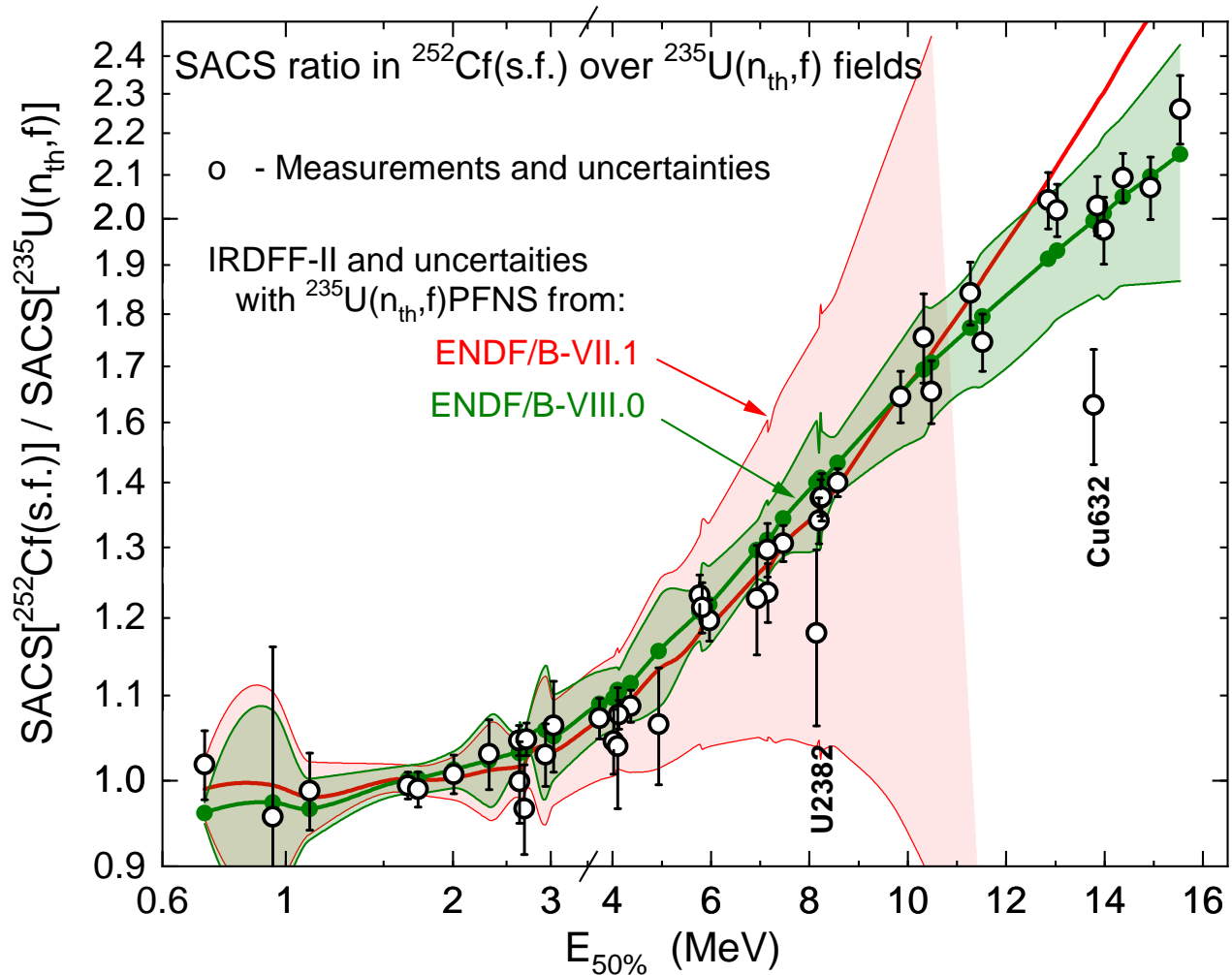


FIG. 101. (Color online) Ratio of SACS for the ^{252}Cf SFNS and $^{235}\text{U}(n_{th},f)$ PFNS fields as a function of median energy $E_{50\%}$: circles - measurements, crosses - computed with the IRDFF-II cross sections. Plotted values are calculated as ratios of SACS from Tables 18 and 19 to SACS listed in Table 20.

TABLE 21. Reactor operations used to support the SPR-III fast burst reactor central cavity spectral characterization.

Date [d/m/y]	Oper. No.	Reactor Power	Comment
19/01/2006	13090	10 kW, 250 s	3x3 grid Foils Set #1
19/01/2006	13091	10 kW,	^{235}U foil
19/01/2006	13092	10 kW,	^{239}Pu foil
26/01/2006	13117	5 kW, 275 s	^{237}Np foil
26/01/2006	13118	—	^{238}U foil
27/01/2006	13127	125 MJ, 16 min,	Foil Set #2
10/02/2006	13156	27.5 MJ, 40 min,	—
14/04/2006	13295	27.5 MJ, 138 min,	Sc in boron ball
14/04/2006	13296	127.5 MJ, 217 min,	Mn in boron ball
19/05/2006	13368	25 MJ, 32 min,	Fe in boron ball
19/05/2006	13369	—	Ag in boron ball
30/06/2006	13452	25 MJ, 190 min,	3x3 grid Foils Set #1

4. “Cdtk/B4C” indicates an inner thick cadmium cover with an areal thickness of 4.705×10^{-3} atoms/barn and an outer spherical boron ball. The boron ball is a spherical shell with an outer diameter of 4.76 cm and thickness of 1.03 cm. The B_4C has a density of 2.5 g/cm^2 and used isotopically enriched Boron (91.67 % ^{10}B and 8.33 % ^{11}B).

A detailed description of the covers and of how self-

shielding effects are rigorously addressed in their use in the activation measurements can be found in Reference [46]. Data were gathered for a total of 40 combinations of reactions and covers in this reference field.

The fission foils were in the form of a sintered oxide (uranium oxide-to-metal ratio of 1.995; plutonium ratio of 1.886) measuring approximately 0.5 inch by 0.03-inch thick, and having an approximate mass of 1 gram. The isotopic mixtures of the various fission foils are given in Table 22.

The baseline set of end-of-irradiation activity measurements, normalized using the $^{58}\text{Ni}(n,p)^{58}\text{Co}$ monitor reaction used in the same reactor operation, are shown in Table 23. We cannot compare these measured activities to the corresponding calculated dosimetry metrics, spectrum-averaged cross sections, in the absence of a known neutron fluence level for the reference operation. Since a spectral index (SI) is defined to be the ratio of the spectrum-averaged cross sections for two specified reactions – where one is generally considered to be a reference monitor reaction – this quantity is independent of the neutron fluence. Thus, we have adopted the C/E for spectral indices, using the $^{58}\text{Ni}(n,p)^{58}\text{Co}$ reaction as the reference reaction, as the relevant comparison metric.

TABLE 22. Isotopic composition of fission foils

Foil	Isotope	Atom Fraction*
Enriched ^{235}U	^{235}U	0.9300
	^{234}U	0.00981
	^{236}U	0.00359
	^{238}U	0.0566
Depleted ^{238}U	^{238}U	0.9979
	^{234}U	0.00001
	^{235}U	0.00205
	^{236}U	0.00004
^{239}Pu	^{239}Pu	0.869965
	^{238}Pu	0.0006798
	^{240}Pu	0.115968688
	^{241}Pu	0.010797
	^{242}Pu	0.00235936
	^{235}U	0.000199946
	^{237}Np	0.00002999

* Atom fraction sum of the number of atoms of the prime sensor element is unity. Due to contaminants, the total sum may be greater than 1.0.

The C/E spectral index metric, for the reaction “ α ” and the monitor “Ni58p”, is notated as the $\text{CoE}^{SI}_{\alpha/\text{Ni58p}}$.

Table 23 documents the data in this field for the 40 measured reactions. The entries in the table start with the $^{58}\text{Ni}(n,p)$ normalizing reaction but are then ordered with respect to the increasing E_{50} mean response energy for the reactions. The table includes:

1. Reaction identifiers;
2. Median response energy in this neutron field for the indicated dosimeter/cover combination;
3. measured end-of-irradiation (EOI) activities with measurement uncertainty;
4. uncertainty contribution in the calculated spectrum-averaged cross sections due to the *a priori* spectrum;
5. calculated spectral index and associated uncertainty, including the contributions from the cross section and spectrum;
6. calculated-over-experimental (C/E) ratio of the spectral index (SI) and associated uncertainty.

Column 5 of Table 23 shows the contribution from the *a priori* spectrum to the uncertainty in the spectrum-averaged cross sections. The *a priori* spectrum is based solely on a calculation and, while the statistical uncertainty in this Monte Carlo calculation was typically less than 1 % in a given energy bin, the uncertainty in the modeling and in the transport cross sections result is a very large overall spectrum uncertainty - as was seen in Fig. 102. Since the uncertainty in the calculated spectral index has the spectrum appearing in both the numerator and the denominator of the metric, a Monte Carlo approach was used to propagate the spectrum uncertainty through this ratio of integrals. This methodology is described in reference [188]. The use of spectral indices to make the C/E comparison means that the spectrum contribution to the SI uncertainty can be much smaller for

reactions that have the dominant response in the same spectral region as the reference $^{58}\text{Ni}(n,p)^{58}\text{Co}$ reaction. This is clearly seen in the column 7 spectrum contribution to the SI uncertainty. Here the spectrum-contribution to the SI uncertainty becomes zero for the reference reaction, and is much reduced for reactions where the dominant sensor response is similar to that from the reference reaction. For regions where the dosimeter dominant energy-dependent response differs from that of the reference reaction, the spectrum contribution to the SI uncertainty can be more than twice as large as that seen in the calculated ΔACS

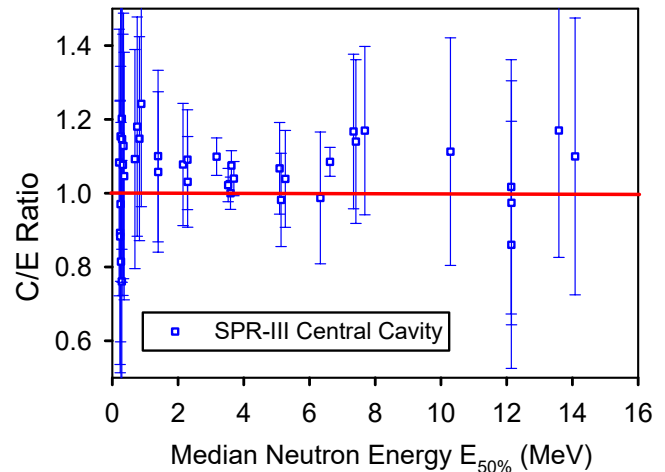


FIG. 102. (Color online) $\text{CoE}^{SI}_{\alpha/\text{Ni58p}}$ Ratios for the calculated spectrum in the SPR-III fast burst reactor central cavity reference benchmark field.

Columns 8 and 9 show the overall calculated-to-experimental (C/E) ratios, $\text{CoE}^{SI}_{\alpha/\text{Ni58p}}$, and associated uncertainties. The $\text{CoE}^{SI}_{\alpha/\text{Ni58p}}$ value is the primary metric reported in this benchmark field. The uncertainty shown for this C/E metric is taken to be the root-mean-square of the measurement uncertainty and the spectrum and cross section contributions to the calculated spectrum-averaged cross section uncertainty. The color-coding in the $\text{CoE}^{SI}_{\alpha/\text{Ni58p}}$ columns corresponds to the criteria appearing in Table 17. Good or acceptable validation evidence was seen for 39 of the 40 measured activities in this reference field. The only discrepant reaction among this set of data were for the $^{23}\text{Na}(n,\gamma)^{24}\text{Na}$ reaction and, even in this case, the $\text{CoE}^{SI}_{\alpha/\text{Ni58p}}$ was within one standard deviation of unity - but the ratio fell outside the criteria adopted in Table 17 and there was a large spectrum contribution to the $\text{CoE}^{SI}_{\alpha/\text{Ni58p}}$ uncertainty.

Fig. 102 shows the resulting $\text{CoE}^{SI}_{\alpha/\text{Ni58p}}$ ratios for this benchmark neutron field. The x-axis represents the E_{50} mean response energy for the foil/cover combination in that calculated reference field spectrum. The large uncertainties, due to the *a priori* spectrum uncertainty, are clearly seen here. A small positive bias in the C/E is also seen. The usefulness of this $\text{CoE}^{SI}_{\alpha/\text{Ni58p}}$ metric in the validation process is limited by the large uncertainty due to the spectrum in this neutron field. Sec. VIII will address another metric that can more rigorously address the uncertainty in the spectrum and thus provide more powerful validation evidence.

TABLE 23. Measured EOI activities and calculated quantities supporting the SI C/E for SPR-III central cavity neutron benchmark field.

Entry Label	$E_{50\%}$ [MeV]	Measured Act.		Calc. SACS [unc.]	Spectral		SI C/E	
		Bq/atom or Fiss./atom	Unc. [%]		Index (SI)	Unc. [%]	Value	Unc. [%]
Sc45g-Cd	0.2060	1.3720E-20	3.30	23.14	2.0093E-01	35.84	1.0832	36.12
Mn55g-Cd	0.2348	6.6180E-18	2.60	10.61	1.0265E-01	35.45	0.8926	35.68
Au197g-bare	0.2385	7.4140E-18	4.50	9.87	2.8463E+00	36.52	0.8827	36.92
Au197g-Cd	0.2505	6.5740E-18	4.50	10.11	2.7742E+00	36.93	0.9706	37.33
Sc45g-Cdtk/B4C	0.2515	1.1920E-20	3.50	11.26	1.8599E-01	39.01	1.1540	39.29
Na23g-bare	0.2628	7.0870E-20	2.10	4.21	5.8091E-03	37.54	0.8146	37.73
Fe58g-Cd	0.2883	7.9740E-21	2.80	9.63	6.8808E-02	35.13	1.2018	35.38
Na23g-Cdna	0.2887	7.1700E-20	2.10	3.41	5.4893E-03	32.59	0.7609	32.81
Mn55g-Cdtk/B4C	0.2958	4.5570E-18	2.70	9.18	9.0805E-02	38.60	1.1468	38.82
W186g-bare	0.3215	7.4543E-18	9.02	8.56	1.2806E+00	34.08	1.0769	35.39
Cu63g-Cd	0.3395	3.3020E-18	2.20	7.62	3.1820E-01	35.77	1.1281	35.97
Na23g-Cdtk/B4C	0.3589	5.9630E-20	2.10	5.28	6.2782E-03	33.33	1.0464	33.54
U235f-Cdtk	0.6877	1.7550E-11	3.00	1.27	2.4819E+01	29.38	1.0925	29.69
U235f-Cdtk/B4C	0.7568	1.5000E-11	3.00	0.37	2.2920E+01	29.39	1.1804	29.71
Pu239f-Cdtk	0.8244	2.2330E-11	2.70	1.44	3.3176E+01	27.33	1.1477	27.64
Pu239f-Cdtk/B4C	0.8812	1.9120E-11	2.70	2.33	3.0745E+01	27.56	1.2422	27.86
Np237f-Cdtk	1.3904	1.2340E-11	2.80	11.85	1.7581E+01	22.87	1.1006	23.25
Np237f-Cdtk/B4C	1.3952	1.1820E-11	2.80	12.22	1.6182E+01	21.34	1.0576	21.74
In115nm-bare	2.1498	6.2040E-17	4.70	18.12	2.0190E+00	15.57	1.0779	16.55
U238f-Cd	2.2862	2.3170E-12	3.20	19.29	3.2721E+00	12.77	1.0909	13.53
U238f-Cdtk/B4C	2.2909	2.2230E-12	3.20	19.53	2.9664E+00	11.44	1.0308	12.28
Ti47p-Cd	3.1728	2.8820E-19	2.70	21.61	1.7116E-01	2.97	1.0991	5.07
S32p-bare	3.5235	2.5080E-19	3.00	23.78	5.9089E-01	1.36	1.0221	4.52
Ni58p-Cd	3.5954	8.7520E-20	3.10	23.51	1.0000E+00	0.00	1.0000	4.38
Zn64p-Cd	3.6230	3.8820E-18	2.20	23.89	3.5650E-01	1.36	1.0749	4.04
Fe54p-Cd	3.6969	1.4000E-20	3.20	24.14	7.3304E-01	1.27	1.0396	4.63
Al27p-Cd	5.0886	2.8720E-17	3.00	27.07	3.3676E-02	11.64	1.0675	12.41
Co59p-bare	5.1337	1.8835E-21	5.18	26.70	1.3281E-02	11.12	0.9818	12.65
Ti46p-Cd	5.2634	7.3250E-21	3.40	27.59	1.0288E-01	12.29	1.0389	13.13
Cu63a-bare	6.3286	1.6723E-23	5.14	28.41	5.1287E-03	16.82	0.9873	17.86
Fe56p-Cd	6.6224	5.4080E-19	2.40	29.70	1.0197E-02	0.01	1.0851	3.92
Ti48p-Cd	7.3414	8.6840E-21	1.20	30.16	2.9767E-03	20.68	1.1673	20.94
Mg24p-Cd	7.4113	1.2420E-19	3.00	30.77	1.4247E-02	21.75	1.1400	22.18
Al27a-Cd	7.6839	6.1350E-20	1.90	30.83	7.2208E-03	22.52	1.1696	22.81
Nb932-bare	10.2890	2.4821E-21	4.72	32.61	4.5229E-03	30.32	1.1127	30.84
Co592-bare	12.1390	1.6686E-22	7.10	33.11	1.9395E-03	33.58	1.0172	34.46
Mn552-bare	12.1420	4.7562E-23	8.13	32.36	2.0609E-03	32.34	0.8601	33.49
Mn552-Cdtk/B4C	12.1450	4.0170E-23	5.01	32.40	1.9708E-03	32.53	0.9740	33.06
Zr902-Cd	13.5890	1.6160E-21	3.40	33.12	9.9648E-04	34.08	1.1699	34.39
Ni582-bare	14.0830	1.5485E-22	5.87	33.06	4.1111E-05	36.92	1.0997	37.51

E. ACRR Pool-type Reactor Central Cavity

The baseline measured quantities in this reference benchmark field are end-of-irradiation (EOI) spectrum-averaged cross sections measured in operation #10639, which was a 153.2 MJ pulsed operation that took place on 9/19/2013. In order to eliminate the effect of variations in the reactor power during the other exposures, a $^{58}\text{Ni}(n,p)^{58}\text{Co}$ monitor activity was used to facilitate a renormalization of data gathered in the other operations to the baseline exposure. When required, corrections were made in the measured activities of other foils to convert them to represent the end-of-irradiation from the baseline 16-minute uniform power operation. The other reactor operations are detailed in Table 24. As discussed in Sec. VIII D, in some of the cases covers were used to shield the activation foil and shift the sensitivity of the

measured response to higher energies.

Data were gathered for a total of 33 combinations of reactions and covers in this reference field. The baseline set of end-of-irradiation activity measurements, normalized using the $^{58}\text{Ni}(n,p)^{58}\text{Co}$ monitor reaction used in the same reactor operation, are shown in Table 25. As in Sec. VIII D, a spectral index is defined to be the ratio of the spectrum-averaged cross sections for two specified reactions, where one is generally considered to be a reference monitor reaction. This quantity is independent of the neutron fluence. Table 25 provides the calculated spectrum-averaged cross sections for the measured reactions, with covers, and reports both the spectrum and cross section uncertainty contributions to the calculated spectrum-averaged cross sections. The entries in the table are ordered with respect to the increasing E_{50} mean response energy for the reactions. The overall calculated-to-experimental (C/E) ratios, $\text{CoE}_{\alpha/Ni58p}^{SI}$, are also shown

TABLE 24. Reactor operations used to support the ACRR central cavity spectral characterization.

Date [d/m/y]	Oper. No.	Operation Type	Comment
18/09/2013	10636	Steady 172.4 MJ	Activation foil set 2: Co, Fe, Sc, Ni, and Ti
19/09/2013	10638	Pulse 157.7 MJ	Activation foil set 3: Zr, Mo, Ag, Au, Au, Au-dilute, and W
02/10/2013	10649	Pulse 148.0 MJ	Activation foil sets 4 and 5: Na, Mg, Mn, Fe, Cu, Zn, In, Nb, and Al
19/09/2013	10637	Pulse 151.5 MJ	Cd covered activation foil sets 6 and 7: Co, Fe, Sc, Ni, Mo, Ag, Au, and Au-dilute
03/10/2013	10650	Pulse 152.8 MJ	Cd covered activation foil sets 8 and 9: Na, Mg, Mn, Fe, In, Cu
08/04/2014	10870	Steady ≈150 MJ	Depleted U activation foil in Cd cover and B ₄ C ball
08/04/2014	10871	Steady ≈150 MJ	Enriched U activation foil in Cd cover and B ₄ C ball
08/04/2014	10872	Steady ≈150 MJ	Np activation foil in Cd cover and B ₄ C ball
08/04/2014	10873	Steady ≈150 MJ	Pu activation foil in Cd cover and B ₄ C ball

in the table. The uncertainty shown for this C/E metric is taken to be the root-mean-square of the measurement uncertainty and the spectrum and cross section contributions to the calculated spectrum-averaged cross section uncertainty. Several of the table entries have such a large uncertainty that they do not qualify as acceptable validation cases, but they have a good C/E value, so they are also not designated as discrepant. These table entries correspond to measurements that are not of validation quality, typically due to the spectrum uncertainty, and are left uncoloured.

Fig. 103 shows the resulting C/E ratios for this benchmark neutron field. The reactions with a dominant high energy response, similar to that of the $^{58}\text{Ni}(n,p)^{58}\text{Co}$ reference reaction, show smaller uncertainty bounds and general good agreement. The low energy dominant response reactions show much larger uncertainty bounds and considerable scatter in their central values.

F. Other “Bucket”-Modifying Environments in the ACRR Reference Reactor Fields

C/E ratios for the spectral indices in several other reference reactor fields have been considered. These neutron fields and activity measurements include:

1. ACRR LB44; 35 measured activities
2. ACRR PLG; 35 measured activities
3. ACRR Cd-poly; 31 measured activities
4. FREC-II; 33 measured activities

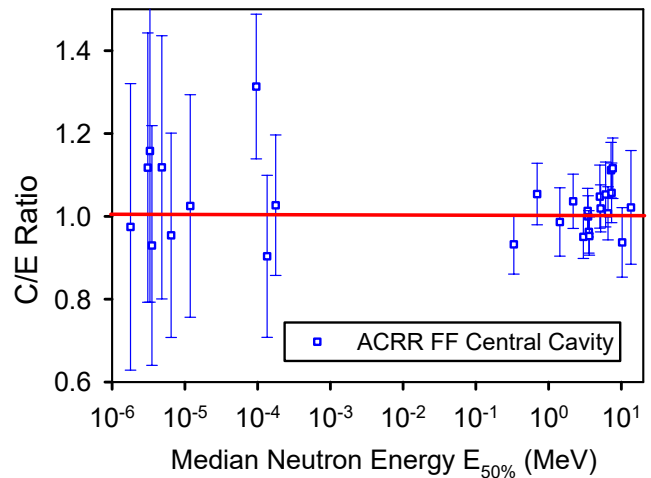


FIG. 103. (Color online) $\text{CoE}^{SI}_{\alpha/Ni58p}$ Ratios for the Calculated Spectrum in the ACRR Central Cavity Reference Benchmark Field.

The utility of these data is best addressed in the context of the consistency of the measured activities in Sec. VIII F rather than as isolated spectral indices.

G. Mol-BR1 Mark-III Irradiation Facility

New measurements from the BR1 reactor at the Belgian Nuclear Research Centre SCK•CEN in Mol were published recently [110]. The aim of the work was to make an update of the Mark-III spectrum and flux, using state of the art computing tools and certified neutron activation measurements. The IRDFF-v1.02 library, which was available at the time, was used. The measured flux calibration factors were scaled by the corresponding calculated cross sections listed in the same publication to recover the measured reaction rates, which in turn were normalized by the $^{58}\text{Ni}(n,p)$ reaction rate to remove the absolute flux dependence. The ratios were compared to the calculated values using the IRDFF-II cross sections, as shown in Table 26 and Fig. 104. The figure undoubtedly displays excellent agreement between measurements and calculations.

H. LR-0 Reactor at Řež

The measured quantities in the experiments are reaction rate ratios and neutron spectra. Irradiations were carried out at an atmospheric pressure and room temperature, using fairly large samples due to the low neutron flux. It was shown that the neutron energy spectrum above 6 MeV is indistinguishable from the ^{235}U PFNS [115]. Therefore, when corrected, the cross sections of reactions with the threshold over 6 MeV averaged in the LR-0 spectrum correspond to spectrum-averaged cross sections in ^{235}U PFNS [111]. Corrections are based on the ratio between the share of neutrons over 6 MeV in LR-0 spectrum and in the ^{235}U PFNS. Another correction, taking into

TABLE 25. Measured EOI activities and calculated quantities supporting the SI C/E for ACRR central cavity neutron benchmark field.

Entry Label	$E_{50\%}$ [MeV]	Measured Act.		Calc. SACS Unc. [%]	Spectral Index (SI) Unc. [%]		SI C/E Value Unc. [%]	
		Bq/atom or Fiss./atom	Unc. [%]		Index (SI)	Unc. [%]	Value	Unc. [%]
Sc45g-bare	4.7701E-08	7.3161E-16	2.97	43.79	8.9065E+01	50.05	1.0063	50.26
Na23g-bare	6.0353E-08	1.8587E-15	2.97	38.98	1.7629E+00	45.73	1.0532	45.96
Mn55g-bare	6.8033E-08	3.4622E-13	2.76	36.16	5.1304E+01	42.67	0.9531	42.91
Fe58g-bare	7.2933E-08	8.3270E-17	2.69	33.19	5.4350E+00	39.79	1.0160	40.03
Co59g-bare	1.2520E-07	5.6707E-17	4.83	27.78	1.7960E+02	34.44	1.1392	25.58
Sc45g-Cd	1.7900E-06	1.1446E-16	2.97	28.59	1.3490E+01	34.29	0.9746	34.59
Au197g-bare	3.0900E-06	5.6538E-13	3.77	25.63	2.4586E+03	32.11	1.1177	32.52
Au197g-Cd	3.3100E-06	4.7342E-13	3.77	30.74	2.1328E+03	36.15	1.1580	36.51
Ag109g-bare	3.5260E-06	8.7524E-17	4.30	20.00	2.9347E+01	28.40	0.9297	28.94
Ag109g-Cd	4.8240E-06	5.7839E-17	4.29	25.61	2.3331E+01	31.29	1.1184	31.77
Na23g-Cd	6.4680E-06	4.2940E-16	2.97	18.52	3.6902E-01	24.23	0.9544	24.67
W186g-bare	1.1850E-05	2.2104E-13	2.50	18.33	3.2344E+02	26.52	1.0251	26.87
Co59g-Cd	9.5720E-05	1.8963E-17	4.83	13.66	6.9245E+01	16.41	1.3135	17.46
Mn55g-Cd	1.3490E-04	9.1556E-14	2.76	13.37	1.2863E+01	19.04	0.9036	19.56
Fe58g-Cd	1.7710E-04	2.5208E-17	2.69	11.82	1.6630E+00	16.38	1.0270	16.96
rmleu-Cdtk/B4C	3.3292E-01	3.1399E-09	4.03	8.11	3.3913E+01	4.79	0.9324	7.18
rmlpu-Cdtk/B4C	6.9648E-01	3.1557E-09	4.03	7.94	3.8531E+01	5.16	1.0540	7.43
Np237f-Cdtk/B4C	1.4277E+00	1.5815E-09	4.03	8.89	1.8067E+01	6.30	0.9865	8.26
rmldu-Cdtk/B4C	2.1768E+00	2.8531E-10	4.03	9.03	3.4254E+00	3.78	1.0365	6.55
Ti47p-bare	3.0036E+00	3.8055E-17	3.69	9.07	1.7495E-01	1.13	0.9505	5.22
S32p-bare	3.4419E+00	2.7894E-17	4.12	9.29	5.8302E-01	0.61	1.0133	5.45
Ni58p-bare	3.4929E+00	9.7808E-18	3.52	9.14	1.0000E+00	0.00	1.0000	4.98
Zn64p-bare	3.5683E+00	4.7963E-16	3.69	9.35	3.5292E-01	0.59	0.9626	5.13
Fe54p-bare	3.6245E+00	1.6954E-18	2.97	9.18	7.2790E-01	0.47	0.9526	4.63
Co59p-Cd	5.0698E+00	1.7965E-19	6.13	9.49	1.2094E-02	2.81	1.0478	7.61
Ti46p-bare	5.1988E+00	7.8515E-19	3.05	9.21	9.6822E-02	3.25	1.0193	5.68
Ni60p-bare	6.0778E+00	6.3203E-21	5.66	9.36	1.8502E-02	4.11	1.0529	7.83
Fe56p-bare	6.5505E+00	5.8136E-17	2.69	9.23	9.1059E-03	4.65	1.0074	6.42
Ti48p-bare	7.2201E+00	8.8424E-19	2.44	9.38	2.5831E-03	5.16	1.1116	6.71
Mg24p-bare	7.3367E+00	1.2890E-17	3.05	9.38	1.2267E-02	5.49	1.0569	7.20
Al27a-bare	7.6106E+00	6.1585E-18	3.05	9.57	6.1899E-03	5.62	1.1163	7.30
Nb932-bare	1.0266E+01	2.5526E-19	2.83	9.94	3.5065E-03	7.08	0.9372	8.40
Zr902-bare	1.3553E+01	1.7102E-19	5.48	13.77	8.2414E-04	12.09	1.0217	13.73

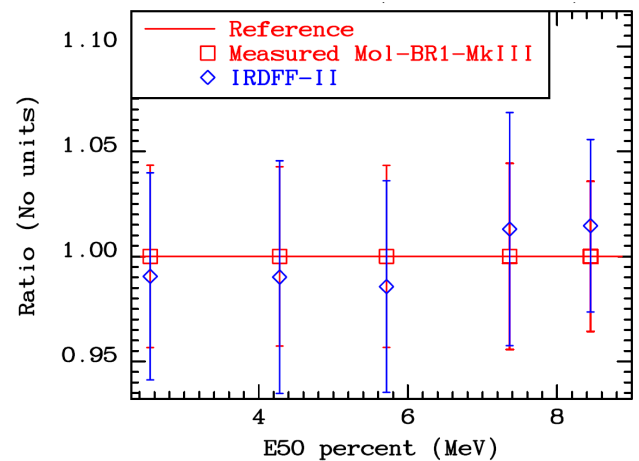
TABLE 26. Measured and calculated SI in the Mark-III facility of the BR1 reactor in Mol.

Entry Label	$E_{50\%}$ [MeV]	Measured SI		Calculated SI		SI C/E & and monitor	
		& Unc. [%]	& Unc. [%]	& Unc. [%]	& Unc. [%]	SI C/E	Unc. [%]
In115nm/Ni58p	2.54	1.816	4.34	1.798	2.43	0.990	5.0
Fe54p/Ni58p	4.28	0.733	4.27	0.726	3.61	0.990	5.6
Al27p/Ni58p	5.71	0.037	4.34	0.036	2.70	0.986	5.1
Fe56p/Ni58p	7.36	9.93E-3	4.43	0.010	3.22	1.013	5.5
Al27a/Ni58p	8.45	6.44E-3	3.57	6.53E-3	1.90	1.015	4.0

account the difference between LR-0 and PFNS, which is defined as a ratio between cross section averaged in reactor field and the ^{235}U PFNS field, is applied as well [189].

The neutron transport in large samples for correct evaluation of significant flux loss effect, as well as correction to flux level, are determined using well validated mathematical model of the irradiation assembly.

The efficiencies of the semiconductor detector were determined using a precise mathematical model of the used HPGe [189], which has been compiled using experimentally determined HPGe crystal parameters and its dead layer [190]. The same arrangement of the gamma spec-

FIG. 104. (Color online) C/E Spectral Index Ratios Relative to $^{58}\text{Ni}(n,p)$ for the Calculated Spectrum in the Mol-BR1-Mark-III Reference Benchmark Field.

trum measurement was used in case of ^{252}Cf irradiation experiments as well (see Table 19, [155]). Thanks to the

usage of the same arrangement in case of ^{235}U as well as ^{252}Cf experiments, reaction rates ratio obtained in the ^{235}U and ^{252}Cf irradiations can be obtained with low uncertainties. Results are presented in the section on the ^{235}U PFNS and ^{252}Cf SFNS spectrum-averaged cross sections.

I. TRIGA-JSI Pneumatic Tube Irradiation Channel

The measured quantities in the two measurement campaigns were the induced activities after irradiation of bare samples and samples with covers of cadmium, boron nitride, boron carbide and enriched boron carbide. The reported unfiltered relative to filtered reaction rate ratios were derived from gamma spectrometry measurements of samples with the same geometry and material composition, performed with the same detector at the same geometry, and of the same gamma line. Therefore, the uncertainties in the derived unfiltered relative to filtered reaction rate ratios are unaffected by uncertainties in the sample material composition, detection efficiency, coincidence correction factors, or gamma emission probabilities. The uncertainties in the unfiltered relative to filtered reaction rate ratios reflect the uncertainties in the measured peak areas, sample mass and the irradiation, cooling and measurement timing. The uncertainties in the ratios of one reaction rate relative to another, on the other hand, are affected by all the above mentioned sources of uncertainty and indeed are reflected in the magnitude of the uncertainties.

For the JSI TRIGA case, a neutron spectrum adjustment procedure was carried out using the GRUPINT code, developed at the JSI [191]. The code enables the parametrization of the spectrum with a 19-parameter analytical function, and subsequent fitting of selected function parameters to measured reaction-rate ratios. Fine structure present in the spectra due to neutron resonances is taken into account through a form function, which is defined as the ratio of the calculated and the fitted analytical function. Measured reaction rate ratios can be labelled as active in the spectrum parameter fitting, or excluded. The active reaction rate ratios in the JSI TRIGA case were cadmium ratios R_{Cd} (defined as the ratios of the bare relative to cadmium filtered reaction rates), for the $^{197}\text{Au}(n, \gamma)$ and $^{238}\text{U}(n, \gamma)$ reactions, and the ratios of the $^{27}\text{Al}(n, p)$ and $^{27}\text{Al}(n, \alpha)$ reaction rates relative to the $^{197}\text{Au}(n, \gamma)$ reaction rate. Neutron spectrum adjustment is required in order to improve the thermal to fission peak strength in the spectra resulting from the Monte Carlo calculations. In the present case the adjustment procedure resulted in an increase of the thermal spectrum by about 10 %.

In addition to cadmium filters, filters made from boron nitride (BN), boron carbide (B_4C) and enriched boron carbide ($^{10}\text{B}_4\text{C}$) were employed as well. The filter effect on the neutron spectrum, *i.e.*, the transmission function $t(E)$ is modelled in the GRUPINT code as a exponential

attenuation function governed by two parameters:

$$t(E) = e^{-\sigma_{tr}(E)nd_{eff}}, \quad (5)$$

where n is the atom density in the filter material and d_{eff} is the effective filter thickness, the first parameter, and $\sigma_{tr}(E)$ is the filter transport cross-section, defined as:

$$\sigma_{tr}(E) = \sigma_a(E) + \xi\sigma_s(E), \quad (6)$$

where $\sigma_a(E)$ is the filter absorption cross-section, $\sigma_s(E)$ is the filter scattering cross-section and ξ is the effective scattering fraction, the second parameter. Filtered reaction rates are calculated as:

$$R = \int_0^\infty \sigma(E)t(E)\varphi(E)dE, \quad (7)$$

where $\sigma(E)$ is the reaction cross-section, $t(E)$ is the filter transmission function and $\varphi(E)$ is the neutron spectrum, in the present case the adjusted neutron spectrum, resulting from the adjustment procedure described above.

The BN, B_4C and $^{10}\text{B}_4\text{C}$ filter cross sections were generated on the basis of ENDF/B-VIII.0 nuclear data, the filter chemical compositions and the measured filter densities, using the PREPRO-2018 nuclear data processing code package. The effective filter thicknesses d_{eff} were first adjusted so that the transmission function shapes approximately matched the shapes calculated using MCNP, and second, finely adjusted on the basis of the experimental BN, B_4C and $^{10}\text{B}_4\text{C}$ ratios for the $^{197}\text{Au}(n, \gamma)$ and $^{238}\text{U}(n, \gamma)$ reactions. The effective scattering fractions ξ , were adjusted on the basis of the filter transmission functions calculated with MCNP; however, the fractions ξ were seen to be very close to zero, implying that neutron scattering has a low impact on the overall filter transmission function. Fig. 105 displays the filter transmission functions calculated using MCNP and adjusted on the basis of experimental data.

It is very important to note, that since in the TRIGA-JSI case a subset of the experimental data is used for neutron spectrum adjustment or the adjustment of the parameters of the filter transmission functions, these data cannot be part of the TRIGA-JSI data validation case. The relevance of the TRIGA-JSI case is in the comparison of calculated and measured reaction rate ratios for other measured nuclear reactions in the experimental campaigns, which are included in the IRDF-II library. Additional measurements of a specific selection of (n, γ) reactions of potential interest for epithermal neutron dosimetry were made [119], but they are outside the scope of the present analysis. Table 27 reports the experimental and calculated BN, B_4C and $^{10}\text{B}_4\text{C}$ reaction rate ratios.

J. Uranium Fueled ICSBEP Assemblies from the ICSBEP Handbook

In this section the C/E results for the spectral indices (SI) for the uranium-fueled ICSBEP assemblies (HMF001,

TABLE 27. Comparison between experimental and calculated BN, B₄C and ¹⁰B₄C reaction rate ratios. The experimental results for the ¹⁹⁷Au(*n*, γ) and ²³⁸U(*n*, γ) reactions have been used for adjustment of the BN, B₄C and ¹⁰B₄C transmission function parameters.

React.	R_{BN} , exp.	R_{BN} , calc.	Diff. [%]	R_{B_4C} , exp.	R_{B_4C} , calc.	Diff. [%]	$R_{^{10}B_4C}$, exp.	$R_{^{10}B_4C}$, calc.	Diff. [%]
¹⁹⁷ Au	8.17±1.2%	8.28±8.52%	1.3	27.81±1.2%	28.76±13.8%	3.4	256.04±1.1%	258.65±4.52%	1.0
²³⁸ U	2.60±1.1%	2.53±7.01%	-2.7	5.44±1.2%	5.18±6.00%	-4.7	51.39±1.0%	48.28±4.77%	-6.0
²³⁸ U	2.63±1.4%	2.53±7.01%	-3.7	5.32±1.3%	5.18±6.00%	-2.5	45.47±1.3%	48.28±4.77%	6.2
²³⁸ U	2.46±1.1%	2.53±7.01%	3.1	5.33±1.1%	5.18±6.00%	-2.8	49.66±1.0%	48.28±4.77%	-2.8
⁵⁵ Mn	39.13±2.5%	35.83±5.17%	-8.4	48.26±2.6%	47.04±5.54%	-2.5	183.11±3.0%	147.43±6.64%	-19.5
²³² Th	4.07±2.2%	4.19±18.9%	2.7	6.50±2.2%	6.42±18.2%	-1.2	31.13±2.1%	30.39±11.3%	-2.4
²³ Na	92.77±2.3%	87.38±3.99%	-5.8	136.03±2.2%	122.64±4.07%	-9.8	283.94±2.2%	229.53±4.46%	-19.2

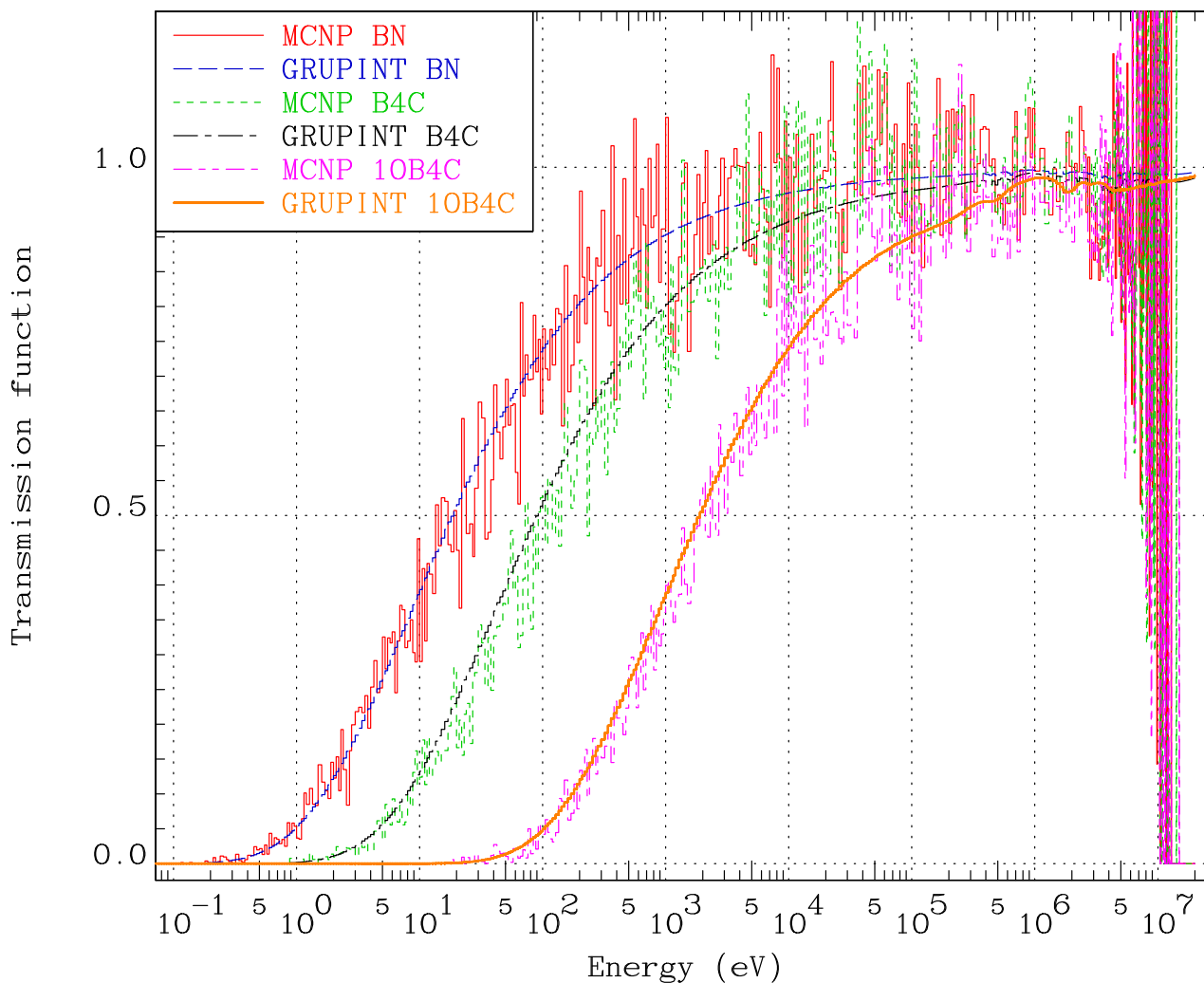


FIG. 105. (Color online) Comparison between the filter transmission functions $t(E)$ for the BN, B₄C and ¹⁰B₄C filters calculated with MCNP and as modelled in the GRUPINT code, with the function parameters adjusted to measured ¹⁹⁷Au(*n*, γ) and ²³⁸U(*n*, γ) reaction rate ratios.

HMF028 and IMF007) are summarized. As indicated earlier the simulation models are derived from the specifications provided in the ICSBEP Handbook. However the Handbook does not provide a summary of experimental reaction rate data. The reaction rate data cited here are derived from a variety of sources (*e.g.*, References [30, 39, 192–194]).

Table 28 provides results for HMF001, with the corresponding SI C/E values shown in Fig. 106. Aside from

some scatter in the low energy activation foil capture reactions, where C/E values oscillate between 0.80 and 1.30, the calculated SI C/E are generally in good agreement with experiment, with all values within 10 % of unity for $E_{50\%}$ values up to 8 MeV. Even SI C/E for the high energy ⁶³Cu(*n*,2*n*) reaction is in good agreement with experiment, given the large measurement and stochastic uncertainties for this reaction.

Table 29 provides results for HMF028, with the corre-

TABLE 28. Measured and calculated SI in the central region of HMF001 (Godiva).

Entry Label and monitor	$E_{50\%}$ MeV	Measured		Calculated		SI C/E		Diff [%]
		SI and Unc. [%]		SI and Unc. [%]		Value and Unc. [%]		
Mn55g/U5f	0.328	2.70E-3	7.41	3.42E-3	21.79	1.2673	23.02	26.73
Au197g/U5f	0.345	0.1000	2.00	0.0946	1.29	0.9458	2.38	-5.42
Ta181g/U5f	0.368	0.1230	9.76	0.1013	6.21	0.8238	11.56	-17.62
Nb93g/U5f	0.376	0.0300	10.00	0.0318	2.11	1.0588	10.22	5.88
Co59g/U5f	0.432	0.0380	7.89	5.48E-3	2.76	0.1442	8.36	-85.58
Cu63g/U5f	0.450	0.0117	5.13	0.0114	10.70	0.9780	11.86	-2.20
U8g/U8f	0.549	0.4760		0.4695	2.20	0.9863	2.20	-1.37
In115gm/U5f	0.594	0.1168	0.68	0.1266	2.48	1.0841	2.57	8.41
La139g/U5f	0.635	7.30E-3	8.22	6.38E-3	5.20	0.8736	9.73	-12.64
Pu9f/U5f	1.051	1.3650		1.3841	1.73	1.0140	1.73	1.40
Np7f/U5f	1.614	0.8516	1.41	0.8338	2.12	0.9791	2.53	-2.09
U8f/U5f	2.576	0.1629		0.1584	1.71	0.9726	1.71	-2.74
U8f/P31p	2.576	8.9460	4.61	9.1893	3.67	1.0272	5.89	2.72
Al27p/P31p	5.757	0.1126	4.88	0.1142	4.03	1.0143	6.33	1.43
Fe56p/P31p	7.397	0.0310	5.80	0.0327	4.39	1.0545	7.27	5.45
U82/U5f	8.082	7.73E-3	4.00	7.93E-3	5.25	1.0260	6.60	2.60
Al27a/P31p	8.461	0.0203	5.91	0.0215	3.55	1.0584	6.89	5.84
Cu632/P31p	13.619	2.94E-3	9.52	2.79E-3	4.49	0.9477	10.53	-5.23

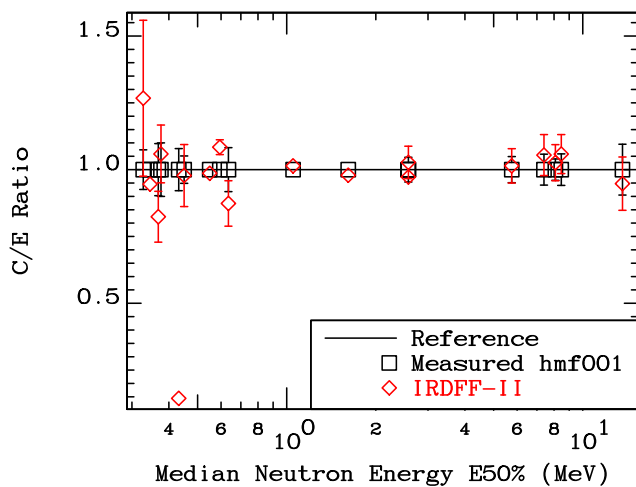


FIG. 106. (Color online) Ratio of Calculated and Measured SI in the central region of HMF001 (Godiva).

sponding SI C/E values shown in Fig. 107. Aside from the $^{169}\text{Tm}(n,2n)$ result (at $E_{50\%} \approx 10$ MeV), which seems anomalous given the excellent $^{197}\text{Au}(n,2n)$ result at nearly the same $E_{50\%}$ energy, these C/E results are excellent.

Table 30 provides results for IMF007, with the corresponding SI C/E values shown in Fig. 108. As with HMF001 there is a large scatter in the low energy activation foil capture reaction C/E values. This large variation, both below and above unity, suggests a deficiency in the underlying measurements rather than in the source spectrum. Unfortunately many of these measurements were made decades ago making it difficult to know whether details such as gamma transition intensity, decay half-life, possible capture branch to isomers, *etc* were properly accounted for. Once again the $^{169}\text{Tm}(n,2n)$ SI C/E, near 10.3 MeV, stands out for its marked deviation from unity. Also once again the $^{197}\text{Au}(n,2n)$ result is near unity suggesting the issue is with the measured ^{169}Tm SACS (or a decay data deficiency) rather than problems in the

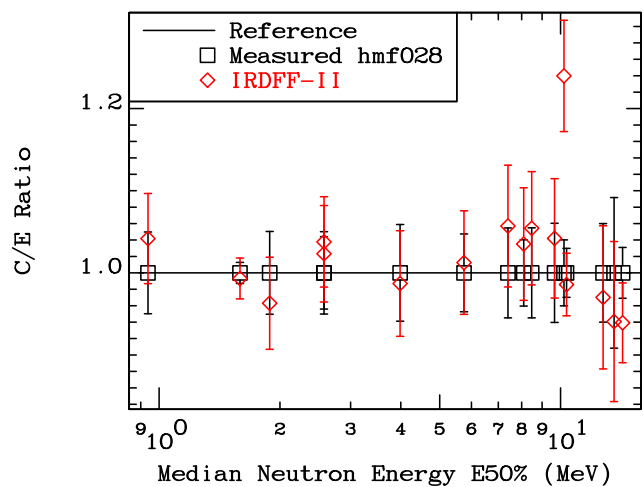


FIG. 107. (Color online) Ratio of Calculated and Measured SI in the central region of HMF028 (Flatop-25).

IMF007 spectrum. In general we observe good C/E results with most values being consistent with unity to within their respective uncertainties.

K. Plutonium Fueled ICSBEP Assemblies from the ICSBEP Handbook

In contrast to the good SI C/E results for the uranium fueled ICSBEP assemblies that were noted in Sec. VII J the results for plutonium fueled ICSBEP assemblies are poor. Measured reaction rates, our calculated values and the corresponding SI C/E results for the PMF001 (Jezebel), PMF006 (Flatop-Pu) and PMF008 (Thor) critical assemblies are given in Table 31.

The SI C/E values, plotted against $E_{50\%}$ as has been done with all other results, are shown in Figs. 109, 110 and 111. These figures display a clear trend of increasing C/E with increasing energy. With a similar trend seen

TABLE 29. Measured and calculated SI in the central region of HMF028 (Flattop-25).

Entry Label and monitor	$E_{50\%}$ MeV	Measured		Calculated		SI C/E		Diff [%]
		SI and Unc. [%]		SI and Unc. [%]		Value and Unc. [%]		
Pu9f/U5f	0.938	1.3070	4.97	1.3615	1.73	1.0417	5.27	4.17
Np7f/U5f	1.590	0.7804	1.28	0.7751	2.12	0.9932	2.48	-0.68
Am1f/U5f	1.886	0.7540	5.04	0.7261	2.92	0.9631	5.82	-3.69
U8f/U5f	2.575	0.1397	5.01	0.1450	1.72	1.0377	5.30	3.77
U8f/P31p	2.575	8.9810	4.41	9.1903	3.67	1.0233	5.74	2.33
S32p/U5f	3.985	0.0306	5.88	0.0302	2.79	0.9870	6.51	-1.30
Al27p/P31p	5.749	0.1128	4.74	0.1135	4.03	1.0125	6.22	1.25
Fe56p/P31p	7.398	0.0306	5.47	0.0324	4.39	1.0570	7.02	5.70
U82/U5f	8.097	6.94E-3	4.03	7.18E-3	5.25	1.0351	6.62	3.51
Al27a/P31p	8.476	0.0202	5.48	0.0213	3.56	1.0534	6.54	5.43
V51a/U5f	9.665	1.11E-5	6.04	1.16E-5	3.50	1.0420	6.98	4.20
Tm1692/U5f	10.200	1.47E-3	4.01	1.82E-3	3.71	1.2398	5.47	23.98
Au1972/U5f	10.351	1.61E-3	2.98	1.59E-3	2.46	0.9858	3.87	-1.42
As752/U5f	12.767	1.50E-4	6.00	1.46E-4	6.69	0.9702	8.99	-2.98
Cu632/P31p	13.597	2.94E-3	9.17	2.76E-3	4.82	0.9407	10.36	-5.93
Zr902/U5f	14.279	4.85E-5	3.09	4.56E-5	4.14	0.9392	5.17	-6.08

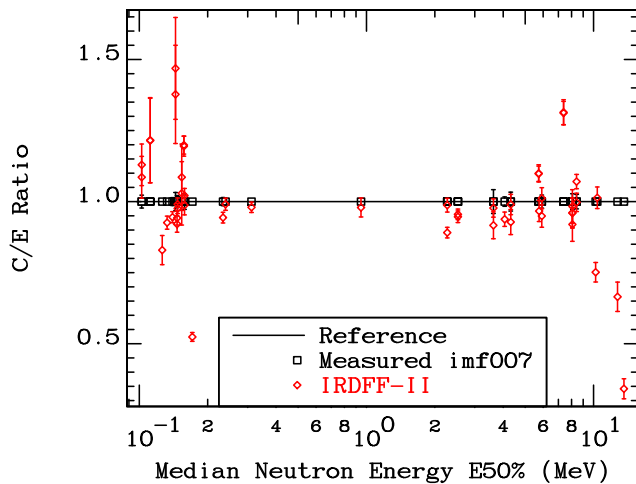


FIG. 108. (Color online) Ratio of Calculated and Measured SI in the central region of HMF028 (Big-Ten).

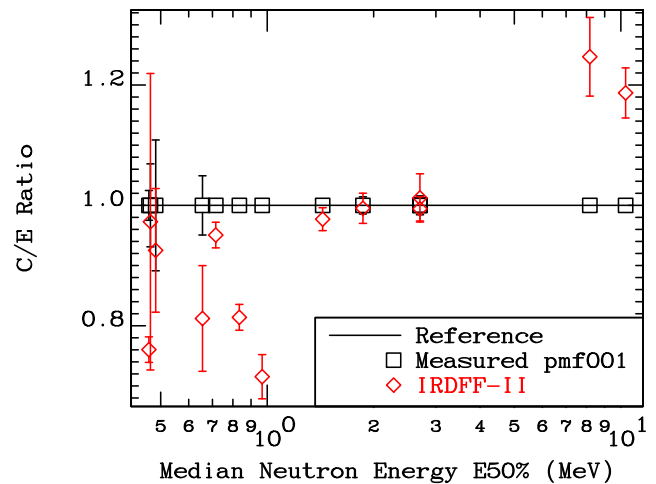


FIG. 109. (Color online) Ratio of Calculated and Measured SI in the central region of PMF001 (Jezebel) assembly.

for multiple assemblies it seems likely that the source for this trend is in the high energy tail of the ^{239}Pu prompt fission spectrum. However, we have also calculated the ICSBEP FMR001 assembly (more formally designated FUND-IPPE-FR-MULT-RRR-001 in the ICSBEP Handbook), a Pu metal rodded critical assembly with copper and uranium reflectors. The SI measured data, our calculations and the corresponding C/E values are given in Table 32 and the C/E values are shown in Fig. 112. In contrast to the other Pu fueled assemblies these results are quite good, with minimal variation in C/E out to the highest available energy. The large C/E variation seen at low energies is for a variety of capture reactions and mimics the pattern seen in the results for uranium fueled assemblies.

In view of these generally poorer, and certainly conflicting, Pu fueled system results the reader is cautioned that SI results for plutonium fueled systems obtained with current evaluated nuclear data files is subject to larger than expected biases.

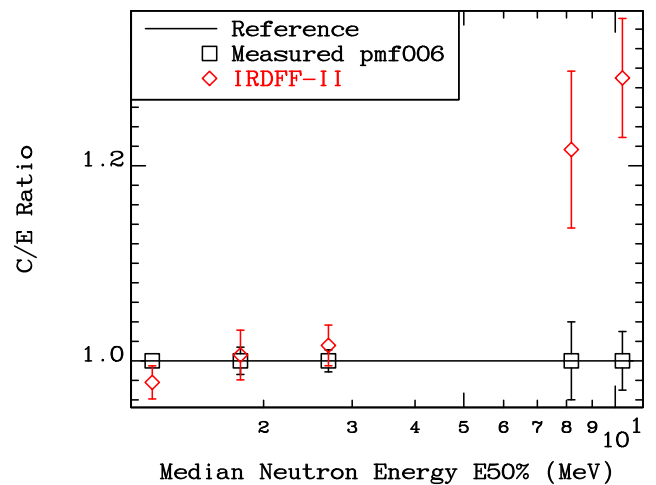


FIG. 110. (Color online) Ratio of Calculated and Measured SI in the central region of PMF006 (Flattop-Pu) assembly.

TABLE 30. Measured and calculated SI in the central region of IMF007 (Big-Ten).

Entry Label and monitor	$E_{50\%}$ MeV	Measured		Calculated		SI C/E		Diff [%]
		SI and Unc. [%]		SI and Unc. [%]		Value and Unc. [%]		
Sc45g/U5f	0.102	0.0132	2.27	0.0143	6.36	1.0857	6.75	8.57
Sc45g/Ni58p	0.102	1.1000		1.2424	6.49	1.1295	6.49	12.95
Mn55g/U5f	0.112	5.37E-3		6.52E-3	12.25	1.2147	12.25	21.47
Mn55g/Ni58p	0.112	0.4650		0.5655	12.32	1.2161	12.32	21.61
Ta181g/U5f	0.126	0.2160		0.1791	6.18	0.8293	6.18	-17.07
W186g/U5f	0.133	.05688		0.0527	2.48	0.9259	2.48	-7.41
B10a/U5f	0.136	1.0110	1.38	0.9557	1.50	0.9453	2.04	-5.47
Fe58g/U5f	0.144	3.10E-3	3.23	4.27E-3	12.13	1.3773	12.53	37.73
Fe58g/Ni58p	0.144	0.2520		0.3701	12.20	1.4688	12.20	46.88
Co59g/U5f	0.147	9.50E-3	2.11	8.75E-3	2.27	0.9209	3.10	-7.91
Co59g/Ni58p	0.147	0.8050		0.7585	2.61	0.9422	2.61	-5.78
Au197g/U5f	0.147	0.1670	1.80	0.1655	1.29	0.9907	2.21	-0.93
Au197g/Ni58p	0.147	14.700		14.3431	1.83	0.9757	1.83	-2.43
Cu63g/U5f	0.154	0.0164	0.61	0.0178	10.75	1.0858	10.77	8.58
Cu63g/Ni58p	0.154	1.5000		1.5437	10.83	1.0291	10.83	2.91
In115gm/U5f	0.157	0.1460		0.1750	2.64	1.1990	2.64	19.90
In115gm/Ni58p	0.157	12.700		15.1752	2.94	1.1949	2.94	19.49
U8g/U5f	0.158	0.1100	2.73	0.1086	2.09	0.9873	3.43	-1.27
U8g/Ni58p	0.158	9.2200		9.4151	2.45	1.0212	2.45	2.12
In113gm/U5f	0.171	0.4220		0.2210	2.95	0.5238	2.95	-47.62
Li6a/U5f	0.231	0.7100	1.41	0.6544	1.35	0.9217	1.95	-5.64
U5f/Ni58p	0.240	87.5000		86.6913	2.14	0.9908	2.14	-0.92
Pu9f/U5f	0.312	1.1936	0.70	1.1697	1.74	0.9799	1.84	-2.01
Np7f/U5f	0.946	0.3223	1.20	0.3156	3.14	0.9794	4.87	-2.06
In115nm/U5f	2.269	0.0271		0.0241	2.09	0.8909	2.09	-10.91
In115nm/Ni58p	2.269	2.1200		2.0931	2.46	0.9873	2.46	-1.27
U8f/U5f	2.528	0.0374	0.90	0.0357	1.73	0.9551	24.13	-4.49
U8f/Ni58p	2.528	3.2700		3.0957	2.16	0.9467	2.16	-5.33
Ti47p/U5f	3.622	2.15E-3	4.19	1.97E-3	3.01	0.9166	5.16	-8.34
Ti47p/Ni58p	3.622	0.1748		0.1708	3.28	0.9773	3.28	-2.27
Ni58p/U5f	4.061	0.0123	1.63	0.0115	2.14	0.9378	2.69	-6.22
Fe54p/U5f	4.318	9.00E-3	3.33	8.35E-3	3.38	0.9282	4.75	-7.18
Fe54p/Ni58p	4.318	0.7317		0.7242	3.62	0.9920	3.62	-1.03
Al27p/U5f	5.732	3.88E-4		4.26E-4	2.45	1.0981	2.45	9.81
Al27p/Ni58p	5.732	0.0336		0.0369	2.77	1.0993	2.77	9.93
Co59p/U5f	5.757	1.58E-4		1.53E-4	3.81	0.9667	3.81	-3.33
Ti46p/U5f	5.916	1.30E-3	2.31	1.23E-3	3.43	0.9491	4.13	-5.09
Ti46p/Ni58p	5.916	0.1057		0.1070	3.67	0.9813	3.67	1.20
Fe56p/U5f	7.378	9.12E-5		1.20E-4	3.11	1.3117	3.11	31.17
Fe56p/Ni58p	7.378	7.89E-3		0.0104	3.36	1.3144	3.36	31.44
U82/U5f	8.067	1.74E-3		1.67E-3	5.39	0.9607	5.39	-3.93
U82/Ni58p	8.067	0.1510		0.1449	5.54	0.9597	5.54	-4.03
Ti48p/U5f	8.068	3.60E-5	2.78	3.31E-5	5.87	0.9198	6.49	-8.02
Ti48p/Ni58p	8.068	2.92E-3		2.87E-3	6.01	0.9831	6.01	-1.69
Al27a/U5f	8.417	7.80E-5	2.56	7.78E-5	1.97	0.9974	3.24	-0.26
Al27a/Ni58p	8.417	6.30E-3		6.74E-3	2.36	1.0705	2.36	7.05
Tm1692/U5f	10.244	5.45E-4		4.10E-4	4.58	0.7515	4.58	-24.85
Au1972/U5f	10.380	3.52E-4		3.57E-4	3.74	1.0135	3.74	1.35
Co592/U5f	12.753	3.14E-5		2.09E-5	7.77	0.6650	7.77	-33.50
Y892/U5f	13.602	4.67E-5		1.59E-5	10.33	0.3414	10.33	-65.86

L. Legacy Reference Reactor Fields

We have revisited the older reference SACS data obtained at the following reactor facilities: coupled thermal/fast uranium and boron carbide spherical assembly (Sigma-Sigma), Coupled Fast Reactivity Measurement Facility (CFRMF), Intermediate-energy Standard Neutron Fields (ISNF) and Glory hole of the Tokyo University research reactor (YAYOI) [195]. Their neutron spectra are

available in the IRDF-2002 database [131] which in turn were adopted from the IRDF-90 library [145]; the latter was assembled in 1993. It is important to notice that the measurements in these neutron fields were not previously incorporated into the evidence package supporting validation of the IRDF cross section library.

For the present analysis, we selected the facilities and spectra for ISNF and Sigma-Sigma after their spectra were re-calculated based on the transport cross sections from the ENDF/B-VIII.0 library and represented in 725 groups

TABLE 31. Measured and calculated SI in the central region of PMF001 (Jezebel), PMF006 (Flattop-Pu) and PMF008 (Thor) assemblies.

Entry Label and monitor	$E_{50\%}$ MeV	Measured		Calculated		SI C/E		Diff [%]
		SI and Unc. [%]		SI and Unc. [%]		Value and Unc. [%]		
Jezebel								
Au197g/U5f	0.463	0.1012	2.47	0.0769	1.29	0.7603	2.79	-23.97
Mn55g/U5f	0.467	2.90E-3	6.90	2.82E-3	24.36	0.9728	25.32	-2.72
Nb93g/U5f	0.484	0.0276	10.87	0.0255	2.26	0.9252	11.10	-7.48
Cu63g/U5f	0.656	0.0122	4.92	9.91E-3	9.64	0.8120	10.82	-18.80
U8g/U5f	0.715	0.0677		0.0644	2.24	0.9506	2.24	-4.94
In115gm/U5f	0.834	0.1390		0.1131	2.63	0.8140	2.63	-18.60
La139g/U5f	0.968	8.20E-3		5.87E-3	5.13	0.7153	5.13	-28.47
Pu9f/U5f	1.435	1.4609	0.89	1.4274	1.73	0.9771	1.95	-2.29
Np7f/U5f	1.865	0.9835	1.42	0.9787	2.06	0.9951	2.50	-0.49
U8f/U5f	2.704	0.2133	1.08	0.2121	1.71	0.9942	2.02	-0.58
U82/U5f	8.172	0.0106		0.0132	5.24	1.2469	5.24	24.69
Tm1692/U5f	10.317	3.13E-3		3.71E-3	3.50	1.1868	3.50	18.68
Flattop-Pu								
Pu9f/U5f	1.202	1.4150		1.3837	1.73	0.9779	1.73	-2.21
Np7f/U5f	1.799	0.8561	1.40	0.8612	2.08	1.0060	2.51	0.60
U8f/U5f	2.691	0.1772	1.13	0.1800	1.72	1.0158	2.06	1.58
U82/U5f	8.178	9.02E-3	3.99	0.0110	5.27	1.2166	6.61	21.66
Tm1692/U5f	10.333	2.43E-3	3.00	3.13E-3	3.65	1.2900	4.73	29.00
Thor								
U8g/U5f	0.619	0.0830	3.61	0.0681	2.21	0.8209	4.24	-17.91
Th2g/U8g	0.651	1.2000		1.3586	3.81	1.1322	3.81	10.80
U5f/U8f	1.160	5.1300		5.1445	1.71	1.0028	1.71	0.28
Np7f/U5f	1.824	0.9419	1.17	0.9169	2.07	0.9735	2.38	-2.65
Np7f/U8f	1.824	4.6900		4.7172	2.08	1.0058	2.08	0.58
U8f/U5f	2.693	0.1962	1.10	0.1944	1.71	0.9907	2.03	-0.93
Th2f/U8f	2.931	0.2600		0.2575	5.91	0.9904	5.91	-0.96
U82/U8f	8.165	0.0530	5.66	0.0612	5.24	1.1543	7.72	15.43

TABLE 32. Measured and calculated SI in the central region of FMR001 assembly (formally designated FUND-IPPE-FR-MULT-RRR-001 in ICSBEP).

Entry Label and monitor	$E_{50\%}$ MeV	Measured		Calculated		SI C/E		Diff [%]
		SI and Unc. [%]		SI and Unc. [%]		Value and Unc. [%]		
Mn55g/U5f	0.230	2.97E-3	5.05	3.84E-3	18.46	1.2933	19.14	29.33
Au197g/U5f	0.283	0.1050	4.76	0.1000	1.29	0.9526	4.93	-4.74
Fe58g/U5f	0.296	2.28E-3	3.95	2.61E-3	10.53	1.1428	11.24	14.28
Co59g/U5f	0.349	6.40E-3	4.69	5.78E-3	2.63	0.9036	5.37	-9.64
Cu63g/U5f	0.367	0.0114	4.39	0.0121	9.76	1.0606	10.70	6.06
U8g/U5f	0.468	0.0770	3.90	0.0767	2.14	0.9967	4.44	-0.33
Th2g/U5f	0.518	0.1090	3.67	0.1012	1.94	0.9287	4.15	-7.13
Pu9f/U5f	1.093	1.3300	3.01	1.3642	1.73	1.0257	3.47	2.57
Np7f/U5f	1.714	0.7710	2.98	0.8191	2.11	1.0624	3.66	6.24
Am1f/U5f	1.982	0.8250	3.03	0.7871	2.96	0.9541	4.24	-4.59
In115nm/U5f	2.516	0.1020	5.88	0.1002	2.06	0.9821	6.23	-1.79
U8f/U5f	2.629	0.1650	3.03	0.1642	1.71	0.9953	3.48	-0.47
Th2f/U5f	2.857	0.0430	3.02	0.0420	5.91	0.9774	6.64	-2.26
Ti47p/U5f	3.702	9.70E-3	5.15	9.48E-3	2.99	0.9772	5.96	-2.28
Ni58p/U5f	4.129	0.0550	5.45	0.0560	2.11	1.0183	5.85	1.83
Fe54p/U5f	4.383	0.0447	3.36	0.0409	3.31	0.9145	4.72	-8.55
Al27p/U5f	5.806	2.21E-3	6.79	2.18E-3	2.38	0.9862	7.19	-1.38
Co59p/U5f	5.895	8.40E-4	4.76	7.85E-4	3.70	0.9343	6.03	-6.57
Ti46p/U5f	6.044	6.60E-3	4.55	6.33E-3	3.31	0.9596	5.63	-4.04
Fe54a/U5f	7.364	5.00E-4	4.00	4.99E-4	3.87	0.9984	5.57	-0.16
Fe56p/U5f	7.509	6.10E-4	3.28	6.55E-4	2.90	1.0743	4.38	7.43
U82/U5f	8.161	9.16E-3	5.46	9.39E-3	5.24	1.0246	7.57	2.46
Mg24p/U5f	8.207	9.00E-4	4.44	9.32E-4	1.45	1.0354	4.68	3.54
Ti48p/U5f	8.258	1.80E-4	4.44	1.88E-4	5.53	1.0442	7.10	4.42
Co59a/U5f	8.281	9.50E-5	4.21	9.75E-5	3.83	1.0263	5.69	2.63
Al27a/U5f	8.591	4.30E-4	4.65	4.47E-4	1.40	1.0391	4.86	3.91

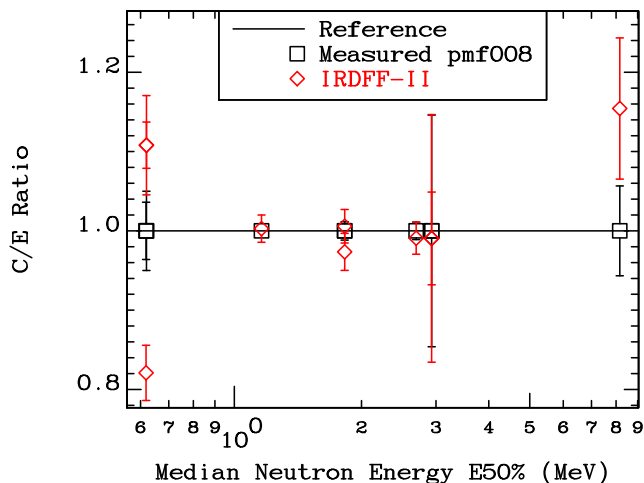


FIG. 111. (Color online) Ratio of Calculated and Measured SI in the central region of PMF008 (Thor) assembly.

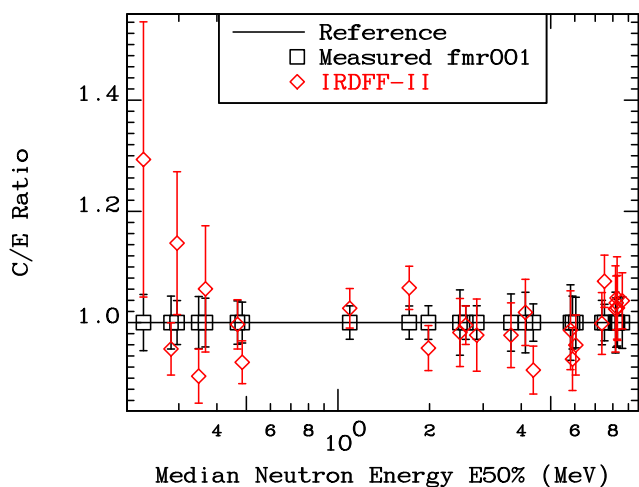


FIG. 112. (Color online) Ratio of Calculated and Measured SI in the central region of FMR001 assembly (formally designated FUND-IPPE-FR-MULT-RRR-001 in ICSBEP).

up to 60 MeV, as described in Secs. VIC 10 and VIC 11. The CFRMF spectrum in 460 energy groups was adopted directly from IRDF-2002 as we were not able to reproduce it, only the material number was changed from MAT=5 into MAT=9006.

We have examined the use of the YAYOI spectrum from IRDF-2002 (MAT= 9, in 100 energy groups) with the SACS measured by K. Kobayashi *et al.*[184, 196] for validation with this neutron environment. However practically all C/E ratios turned out to be below unity by 20-50 %. The difference is too large to be attributed to the rather coarse 100-groups representation of the energy spectrum. Most likely there are some unidentified differences in the interpretation of the published measured spectrum-averaged cross section values and the neutron field characterization data. Attempts to use an alternative YAYOI spectrum received through private communications did not improve the agreement, so work in this benchmark field remains for a future study.

The experimental spectrum-averaged reaction cross sections were taken from EXFOR and were also checked

against the original publications. Some experimental data are in the form of spectral indices, *i.e.*, the reaction rate ratios measured relative to $^{235}\text{U}(n,f)$, $^{238}\text{U}(n,f)$ or $^{197}\text{Au}(n,\gamma)$ – and these data were used in this form. The calculated energy-averaged cross sections were obtained from the IRDFF-II cross sections and facility spectra with the help of the RR_UNC code.

The C/E ratios for SACS or spectral indices are presented in Table 33 and Fig. 113 as a function of the mean response energy $E_{50\%}$. The error bars displayed in Fig. 113 correspond to the sum of the experimental, IRDFF-II cross sections and the spectra uncertainties when the latter were available. It is seen that deviation of C/E from unity lies within two to three-sigma for most reactions and facilities, however with several exceptions which are explicitly labelled in the figure. The agreement seems to be worse for $E_{50\%} < 0.1$ MeV, *i.e.*, for evaluated IRDFF-II cross sections in the epithermal region.

It is worth noting that the research reactor benchmarks have notably softer spectra than the fission sources ^{252}Cf SFNS or $^{235}\text{U}(n_{th},f)$ PFNS. Because of this, they are capable of delivering validation data for reactions that augments those obtained in the fission source neutron fields. This is particularly true for the (n,γ) reaction on targets, such as ^{45}Sc , ^{55}Mn , ^{58}Fe , and ^{109}Ag , where the observed C/E agreement is seen to be within 2–3 standard deviations of unity.

In the case of reactions sensitive to neutron energies above 10 MeV, we see agreement between the C/E values in the Sigma-Sigma and $^{235}\text{U}(n_{th},f)$ PFNS benchmark neutron fields for the $^{59}\text{Co}(n,2n)$ reaction ($E_{50\%} = 12.7$ MeV): $C/E = 0.900 \pm 0.049$ and $C/E = 0.995 \pm 0.044$, respectively.

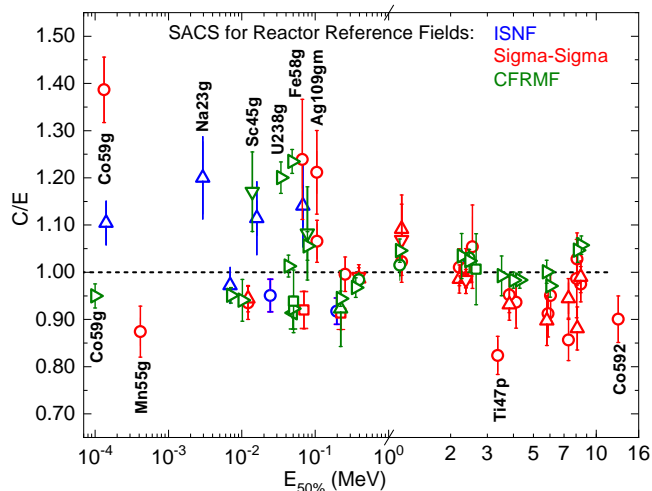


FIG. 113. (Color online) C/E ratios for SACS and spectral indices (SI) measured in the reference reactor fields: ISNF (blue symbols), CFRMF (green), Sigma-Sigma (red). Plotted C/E values are listed in Table 33.

TABLE 33: Measured and calculated SACS or SI ratio in the reference reactor neutron benchmark fields ISNF, Sigma-Sigma, and CFRMF. For the calculated SACS uncertainties the contributions from the dosimetry cross sections (Sigma Unc.) and neutron spectra (Spect. Unc.) are given separately. The C/E uncertainty is the sum of experimental and calculated ones.

Reaction or SI	E _{50%} [MeV]	Experimental SACS or Spectral Index (SI)			Calculated SACS or SI			C/E
		SACS [mb] or SI ratio	Exp. Unc. [%]	Ref.	SACS [mb] or SI ratio	Sigma		
						Unc. [%]	Spect. Unc. [%]	
Intermediate Energy Standard Neutron Field (ISNF)								
Co59g	1.416E-04	3.630E+01	4.13	[122]	4.009E+00	0.85	1.13	1.104 ± 4.22%
Na23g	2.944E-03	1.570E+00	6.40	[122]	1.884E+00	9.31	1.01	1.200 ± 11.2%
Au197g	6.946E-03	4.110E+02	2.68	[122]	3.995E+02	2.86	0.26	0.972 ± 4.91%
Sc45g	1.598E-02	2.440E+01	3.30	[122]	2.719E+01	6.13	0.26	1.115 ± 6.95%
B10He4	2.451E-02	1.831E+03	3.28	[121]	1.741E+03	1.60	0.16	0.951 ± 3.65%
Ag109gm	6.809E-02	2.380E+01	3.78	[122]	2.715E+01	6.67	0.12	1.141 ± 7.67%
Li6He4	1.963E-01	8.310E+02	3.01	[121]	7.625E+02	0.40	0.10	0.918 ± 3.03%
In115nm	2.262E+00	9.700E+01	2.58	[122]	9.846E+01	1.74	0.07	1.015 ± 3.11%
Secondary Intermediate-Energy Standard Neutron Field (Sigma-Sigma)								
Co59g	1.330E-04	3.640E+01	4.95	[128]	5.047E+01	0.78	0.10	1.387 ± 5.01%
Mn55g	4.153E-04	3.550E+01	3.38	[128]	3.103E+01	5.18	0.10	0.874 ± 6.19%
Au197g	1.219E-02	4.103E+02	3.41	[197]	3.837E+02	1.56	0.10	0.935 ± 3.75%
Au197g/U5f	1.325E-02	2.699E-01	2.50	[198]	2.548E-01	1.98	0.10	0.944 ± 3.15%
Fe58g	6.688E-02	5.890E+00	4.92	[128]	7.298E+00	9.03	0.10	1.239 ± 10.2%
B10He4	7.038E-02	1.695E+03	4.19	[130]	1.560E+03	1.06	0.20	0.920 ± 4.32%
Ag109gm	1.052E-01	2.430E+01	3.70	[128]	2.944E+01	6.31	0.10	1.212 ± 7.32%
In115gm	1.061E-01	2.602E+02	3.46	[197]	2.773E+02	2.35	0.10	1.066 ± 4.18%
Li6He4	2.236E-01	9.310E+02	3.76	[130]	8.503E+02	0.49	0.12	0.913 ± 3.79%
U235f	2.558E-01	1.512E+03	3.51	[197]	1.506E+03	1.11	0.10	0.996 ± 3.68%
Pu9f/U5f	3.930E-01	1.179E+00	2.00	[198]	1.164E+00	0.43	0.10	0.987 ± 2.57%
Pu9f/U5f	3.933E-01	1.175E+00	2.30	[126]	1.164E+00	0.43	0.10	0.993 ± 2.58%
Np237f	1.163E+00	6.030E+02	3.65	[197]	6.164E+02	0.00	0.10	1.022 ± 3.65%
Np7f/U5f	1.163E+00	3.830E-01	7.06	[126]	4.094E-01	1.67	0.10	1.071 ± 7.06%
Np7f/U5f	1.165E+00	3.751E-01	6.29	[198]	4.094E-01	1.67	0.10	1.091 ± 6.38%
In115m	2.201E+00	5.480E+01	3.47	[197]	5.537E+01	1.72	0.10	1.010 ± 3.87%
In115nm/U5f	2.202E+00	3.731E-02	2.50	[198]	3.677E-02	1.72	0.10	0.986 ± 3.23%
U238f	2.368E+00	8.346E+01	3.95	[197]	8.417E+01	1.22	0.10	1.009 ± 4.14%
U8f/U5f	2.368E+00	5.680E-02	2.70	[126]	5.590E-02	0.53	0.10	0.974 ± 2.96%
U8f/U5f	2.383E+00	5.664E-02	1.50	[198]	5.590E-02	0.53	0.10	0.987 ± 2.23%
Th232f	2.544E+00	1.980E+01	6.06	[197]	2.087E+01	5.80	0.10	1.054 ± 8.39%
Ti47p	3.363E+00	5.200E+00	4.04	[128]	4.284E+00	2.76	0.10	0.824 ± 4.89%
Ni58p	3.828E+00	2.532E+01	3.55	[197]	2.410E+01	1.72	0.10	0.952 ± 3.95%
Ni58p/U5f	3.828E+00	1.718E-02	3.10	[198]	1.601E-02	1.72	0.10	0.932 ± 3.71%
Fe54p	4.123E+00	1.820E+01	4.95	[128]	1.705E+01	3.19	0.10	0.937 ± 5.88%
Al27p/U5f	5.793E+00	6.148E-04	5.50	[198]	5.518E-04	2.07	0.10	0.898 ± 5.98%
Co59p	5.870E+00	3.250E-01	4.00	[128]	2.965E-01	3.64	0.10	0.912 ± 5.41%
Ti46p	6.040E+00	2.510E+00	4.78	[128]	2.387E+00	3.21	0.10	0.951 ± 5.76%
Fe56p	7.361E+00	2.800E-01	4.29	[128]	2.398E-01	2.72	0.10	0.856 ± 5.08%
Fe56p/U5f	7.361E+00	1.686E-04	3.50	[198]	1.593E-04	2.72	0.10	0.945 ± 4.57%
Ti48p	8.051E+00	6.700E-02	6.27	[128]	6.610E-02	5.59	0.10	0.987 ± 8.40%
Co59a	8.090E+00	3.340E-02	3.89	[128]	3.433E-02	3.71	0.10	1.028 ± 5.38%
Mg24p/U5f	8.112E+00	2.472E-04	5.10	[198]	2.178E-04	0.84	0.10	0.881 ± 5.28%
Al27a	8.433E+00	1.600E-01	3.75	[128]	1.552E-01	0.74	0.10	0.975 ± 3.82%
Al27a/U5f	8.433E+00	1.041E-04	3.30	[198]	1.031E-04	0.74	0.10	0.990 ± 3.56%
Co592	1.275E+01	4.330E-02	5.08	[128]	3.899E-02	2.01	0.10	0.900 ± 5.46%
Coupled Fast Reactivity Measurement Facility (CFRMF)								
Co59g/U5f	1.000E-04	5.890E-02	2.30	[140]	5.594E-02	0.75	0.10	0.950 ± 2.80%
Au197g/U5f	7.100E-03	2.730E-01	1.00	[140]	2.596E-01	2.52	0.10	0.951 ± 3.05%
Cu63g/U5f	1.020E-02	2.920E-02	4.50	[140]	2.747E-02	6.70	0.10	0.941 ± 8.19%
La139g/Au197ng	1.391E-02	4.600E-02	0.00	[201]	5.384E-02	7.25	0.10	1.170 ± 7.20%
U8g/U5f	3.400E-02	1.190E-01	2.40	[140]	1.429E-01	1.13	0.10	1.201 ± 3.00%
Sc45g/U5f	4.320E-02	1.510E-02	1.80	[140]	1.530E-02	5.79	0.10	1.013 ± 6.22%
Fe58g/U5f	4.850E-02	3.940E-03	1.50	[140]	4.865E-03	8.71	0.10	1.235 ± 8.95%
B10He4	4.963E-02	1.900E+03	3.68	[200]	1.737E+03	0.91	0.10	0.914 ± 3.79%
B10He4a/U5f	4.963E-02	1.102E+00	1.60	[140]	1.102E+00	0.91	0.10	0.923 ± 2.31%
Th232g	5.080E-02	2.910E+02	3.10	[203]	2.731E+02	6.37	0.10	0.938 ± 7.08%
In115gm/U5f	7.810E-02	1.810E-01	2.50	[140]	1.911E-01	2.36	0.10	1.056 ± 3.71%
In115gm/Au197g	7.810E-02	6.800E-01	8.82	[201]	7.360E-01	2.36	0.10	1.082 ± 9.12%
Li6He4	2.007E-01	1.004E+03	8.67	[199]	9.268E+02	0.84	0.10	0.923 ± 8.71%

TABLE 33: continued

Reaction or SI	$E_{50\%}$ [MeV]	Experimental SACS or Spectral Index (SI)			Calculated SACS or SI			C/E
		SACS [mb] or SI ratio	Exp. Unc. [%]	Ref.	SACS [mb] or SI ratio	Sigma Unc. [%]	Spect. Unc. [%]	
Li6He4/U5f	2.222E-01	5.881E-01	1.00	[140]	5.679E-01	0.84	0.10	0.944 ± 1.91%
Pu9f/U5f	3.563E-01	1.165E+00	1.60	[140]	1.127E+00	1.14	0.10	0.968 ± 2.41%
Pu9f/U5f	3.930E-01	1.145E+00	2.30	[202]	1.127E+00	0.42	0.10	0.984 ± 1.56%
Np7f/U5f	1.139E+00	3.560E-01	1.90	[140]	3.724E-01	2.23	0.10	1.046 ± 2.36%
Np7f/U5f	1.139E+00	7.340E+00	7.06	[202]	7.451E+00	2.23	0.10	1.015 ± 3.20%
In115nm/U5f	2.258E+00	3.190E-02	4.40	[140]	3.300E-02	1.70	0.10	1.034 ± 4.92%
U8f/U5f	2.451E+00	4.880E-02	1.90	[140]	4.997E-02	1.22	0.10	1.024 ± 2.66%
U8f/U5f	2.451E+00	4.850E-02	2.70	[202]	4.997E-02	1.22	0.10	1.030 ± 1.86%
Th232f	2.651E+00	1.960E+01	4.70	[203]	1.973E+01	5.79	0.10	1.006 ± 7.46%
Ti47p/U5f	3.512E+00	2.690E-03	4.00	[140]	2.670E-03	2.74	0.10	0.992 ± 5.05%
B10H3	3.641E+00	7.780E+00	8.61	[199]	1.707E+01	16.5	0.10	2.194 ± 18.6%
Ni58p/U5f	4.006E+00	1.560E-02	1.00	[140]	1.534E-02	1.73	0.10	0.983 ± 2.44%
Fe54p/U5f	4.297E+00	1.120E-02	1.00	[140]	1.101E-02	3.13	0.10	0.983 ± 3.57%
Al27p/U5f	5.782E+00	5.630E-04	2.00	[140]	5.635E-04	2.07	0.10	1.001 ± 3.20%
Ti46p/U5f	6.009E+00	1.680E-03	2.00	[140]	1.631E-03	3.13	0.10	0.971 ± 3.97%
Ti48p/U5f	8.153E+00	4.430E-05	2.00	[140]	4.638E-05	5.54	0.10	1.047 ± 6.05%
Al27a/U5f	8.488E+00	1.040E-04	1.20	[140]	1.100E-04	0.73	0.10	1.057 ± 1.98%

TABLE 34. Comparison of the thermal cross sections by Mughabghab [71] and IRDF-II to the values derived from the Kayzero library [69]. Relative uncertainties and σ_0 differences are given in %.

Target	Prod.	Kayzero		Mughabghab			IRDF-II		
		σ_0 [b]	Unc. [%]	σ_0 [b]	Unc. [%]	Diff. [%]	σ_0 [b]	Unc. [%]	Diff. [%]
²³ Na	²⁴ Na	0.513	0.6	0.525	1.0	2.3	0.536	1.6	4.4
⁴⁵ Sc	⁴⁶ Sc	26.3	0.4	27.2	0.7	3.5	27.2	6.9	3.6
⁵⁰ Cr	⁵¹ Cr	15.18	0.6	14.7	2.7	-3.2	14.4	2.0	-5.0
⁵⁵ Mn	⁵⁶ Mn	13.21	0.5	13.36	0.4	1.2	13.28	1.1	0.5
⁵⁸ Fe	⁵⁹ Fe	1.29	1.5	1.32	2.3	1.9	1.31	5.1	1.5
⁵⁹ Co	⁶⁰ Co	37.92	0.3	37.18	0.2	-2.0	37.18	0.7	-2.0
⁶³ Cu	⁶⁴ Cu	4.6	0.8	4.5	0.4	-2.6	4.47	4.1	-3.3
⁹³ Nb	⁹⁴ Nb	1.150	22.	1.15	4.3	0.0	1.149	4.0	1.1
⁹³ Nb	^{94m} Nb	0.864	12.	0.853	0.9	0.0	0.862	0.9	0.0
¹⁰⁹ Ag	^{110m} Ag	3.909	1.3	3.95	1.3	1.1	4.211	5.1	7.7
¹¹³ In	^{114m} In	8.733	1.7	8.1	3.7	-7.2	8.134	4.6	-6.9
¹¹⁵ In	^{116m} In	157.5	0.7	162.3	0.4	3.1	159.8	1.0	1.5
¹³⁹ La	¹⁴⁰ La	9.29	0.6	9.21	0.4	-0.8	9.04	3.9	-2.7
¹⁸¹ Ta	¹⁸² Ta	20.3	4.7	20.4	1.5	0.3	20.7	3.0	1.6
¹⁸⁶ W	¹⁸⁷ W	34.7	3.0	38.1	1.3	9.9	38.1	5.0	9.9
¹⁹⁷ Au	¹⁹⁸ Au	98.65	0.1	98.65	0.1	0.0	98.70	0.5	0.0
²³² Th	²³³ Th	7.34	0.5	7.35	0.4	0.1	7.34	5.4	0.0
²³⁸ U	²³⁹ U	2.67	0.6	2.682	0.7	0.5	2.68	1.0	0.5

M. Thermal Reactor Neutron Fields - Thermal Cross Sections and Resonance Integrals

There is a direct relation between average cross sections with $1/v$ shape and the 2200 m/s value in a Maxwellian thermal reactor spectrum. The resonance integral also represents an idealized epithermal reactor spectrum. These quantities can be compared to compilations such as the Atlas of Resonance Parameters [71]. The values of the thermal capture cross sections reported in the Atlas are based on decades of experience with the interpretation of different kinds of experimental data, resonance theory and systematics. There could also be an element of subjective

judgement involved, in the case of inconsistent or lacking data. Since the values in the Atlas were often used in the evaluation process, such comparison should be considered as data verification, rather than validation.

The k_0 and Q_0 constants for neutron activation analysis by the k_0 standardization method in the Kayzero library were measured independently over the years. The thermal cross sections and the resonance integrals derived from them are representative of an integral measurement in an idealized thermal reactor system. They can be used for data validation purposes.

The k_0 factors can be measured quite accurately in irradiation facilities with strongly enhanced thermal spectra. They provide the ratio to the gold standard of the product of the cross section and the gamma-emission probability. In some cases the uncertainty in the gamma-emission probabilities introduces a dominant contribution to the uncertainty in the derived cross section. Other sources of uncertainty originate from detector calibration, measured masses and other details related to the analysis of the measured gamma spectra. The methodology was intended mainly for use with $1/v$ absorbers. The Westcott g -factors and the corrections for the non-ideal cadmium filter were seldom used in the interpretation of the measured data, but for the nuclides relevant for dosimetry these corrections are generally smaller than the overall experimental uncertainties.

In Table 34 the thermal cross sections from the compilation by Mughabghab and in the IRDF-II library are compared to the reference values derived from the Kayzero library. Generally, the values from IRDF-II lie close to the reference values, except for the ¹⁸⁶W capture cross section, which differs by nearly 10 %. However, according to the consistency criteria defined in Table 17, the thermal capture cross sections of ²³Na and ⁵⁹Co also show poor agreement, mainly because the uncertainties assigned to the cross sections are small. Comparison of the cross sections in the Atlas by Mughabghab shows poor agreement for ⁴⁵Sc, ⁵⁹Co and ¹¹⁵In and marginally acceptable agreement for ²³Na and ⁶³Cu for the same rea-

TABLE 35. Comparison of the resonance integrals I_0 from IRDF-II to the values derived from the Kayzero library [69] and the values by Mughabghab [71]. I_0 is calculated by (8) using $E1 = 0.55$ eV and $E2 = 2$ MeV for the Kayzero comparison and $E1 = 0.5$ eV and $E2 = 20$ MeV for the Mughabghab comparison. Relative uncertainties and I_0 differences are given in %.

Target	Prod.	Kayzero		IRDF-II		Diff. [%]	Mughabghab-2016		IRDF-II		
		I_0 [b]	Unc. [%]	I_0 [b]	Unc. [%]		I_0 [b]	Unc. [%]	I_0 [b]	Unc. [%]	Diff. [%]
²³ Na	²⁴ Na	0.3029	4.7	0.3041	4.5	0.4	0.3120	3.2	0.3158	4.5	1.2
⁴⁵ Sc	⁴⁶ Sc	11.3	0.4	11.32	7.6	0.2	12.1	4.1	11.89	7.6	-1.8
⁵⁰ Cr	⁵¹ Cr	8.047	2.5	6.528	3.9	-19.	7.2	2.8	6.834	3.9	-5.1
⁵⁵ Mn	⁵⁶ Mn	13.91	3.0	13.25	3.8	-4.7	13.3	3.8	13.53	3.8	1.7
⁵⁸ Fe	⁵⁹ Fe	1.26	1.8	1.246	4.9	-1.3	1.34	12.	1.275	4.9	-4.8
⁵⁹ Co	⁶⁰ Co	75.58	3.0	75.01	0.8	-0.8	75.8	2.6	75.80	0.8	0.0
⁶³ Cu	⁶⁴ Cu	5.3	0.8	4.885	4.2	-7.3	4.97	1.6	4.988	4.2	0.4
⁹³ Nb	⁹⁴ Nb	8.45	12.	8.725	3.4	3.2	8.37	4.4	8.753	3.4	4.6
⁹³ Nb	^{94^m} Nb	6.35	12.	6.546	3.4	3.1	6.21	6.4	6.567	3.4	5.8
¹⁰⁹ Ag	^{110^m} Ag	65.27	4.4	68.39	7.0	4.8	65.1	4.5	68.50	7.0	5.3
¹¹³ In	^{114^m} In	211.33	2.4	224.7	8.7	6.3	182.9	5.5	225.1	8.7	23.0
¹¹⁵ In	^{116^m} In	2645.4	2.0	2539.	3.1	-4.0	2650	3.8	2647.	3.1	-0.1
¹³⁹ La	¹⁴⁰ La	11.52	0.6	11.92	5.5	3.5	12.1	5.0	12.11	5.5	0.1
¹⁸¹ Ta	¹⁸² Ta	677.6	4.7	659.5	3.8	-2.7	655	3.1	660.0	3.8	0.8
¹⁸⁶ W	¹⁸⁷ W	474.9	3.5	483.4	4.5	1.8	480	3.1	484.2	4.5	0.9
¹⁹⁷ Au	¹⁹⁸ Au	1550.	1.8	1542.	2.1	-0.5	1550	1.8	1545.	2.1	-0.3
²³² Th	²³³ Th	84.42	3.6	84.14	19.	-0.3	83.3	1.8	84.29	19.	1.2
²³⁸ U	²³⁹ U	276.0	1.4	274.9	1.1	-0.4	277	1.1	275.0	1.1	-0.7

son, even though the values are comparably close to the reference. It must be mentioned that the main contribution to the uncertainties of the thermal cross sections of ⁹³Nb come from the gamma-emission probabilities.

The case of ¹⁸⁶W is an interesting one, where we believe the gamma-emission probabilities are mainly responsible for the difference. In the recent re-evaluation of the gamma-emission probabilities the values changed by about 20 %, but the resulting derived thermal capture cross section could not be fitted in a rigorous resonance analysis performed at JRC in Geel [88]. The case of the thermal capture cross section of ¹⁸⁶W remains an open issue.

Resonance integrals can be derived from the Q_0 factors, which are usually measured by the cadmium ratio method. The method is convenient because it practically eliminates the dependence on the detector efficiency and other features related to gamma-spectrum analysis. Its drawback is that it requires two sample irradiations, bare and under Cd cover, where the presence of Cd may distort the local spectrum in the irradiation facility close to the Cd-covered sample. The other source of ambiguity is the definition of the resonance integral. The common definition:

$$I_0 = \int_{E1}^{E2} \sigma(E) \frac{dE}{E}, \quad (8)$$

where $E1 = 0.5$ eV and $E2$ goes to infinity is not what is measured. Mughabghab used $E1 = 0.5$ eV and $E2 = 20$ MeV. The effective Cd cutoff energy for a 1 mm thick Cd cover is around 0.55 eV and it is not sharp. Secondly, measurements are made in thermal-reactor spectra, which fall off rapidly above 2 MeV. A better approximation to compare with measured data is to use $E1 = 0.55$ eV and $E2 = 2$ MeV. An additional refinement is to define the Cd transmission correction factor F_{cd} to correct for the

difference between the idealized sharp cutoff at 0.55 eV and the actual cutoff by the rising Cd cross section with decreasing neutron energy. The Cd factor is defined as

$$F_{cd} = \frac{\int_0^\infty t(E)\sigma(E)\phi(E)dE}{\int_{E_{cd}}^{E_3} \sigma(E)\phi(E)dE} \quad (9)$$

From the definition it follows that measured values by the Cd ratio method must be divided by F_{cd} for comparison with the idealized values. A compilation of cadmium factors is available in the IAEA document **INDC(NDS)-0693** for all nuclides of interest for NAA.

The F_{cd} factors were seldom used in the determination of the Q_0 factors for the Kayzero library. In the current Kayzero library an attempt was made to measure the F_{cd} factor of ¹⁸⁶W by making measurements in three irradiation channels with different spectral characteristics, but no uncertainty was specified. We believe that the value 0.908 adopted by DeCorte [204] is unrealistic and that good agreement of the resonance integral with Atlas and with IRDF-II in Table 35 is a coincidence, stemming from error compensation between F_{cd} and the gamma-emission probabilities in deriving the thermal capture cross section.

Another possible source of discrepancy in the measured resonance integrals stems from the fast-fission spectrum contribution. This depends on the irradiation facility. It may become significant in the Cd-ratio measurements, since these have to be made in conditions, where the epithermal contribution to the reaction rate is not so small. By a minor extension to the governing equations the fast fission effect can be treated explicitly [70]. The effect is usually small, but it has been demonstrated that in some cases it can amount to a few percent in the measured Q_0 [205]. However, this methodology is not generally used by the NAA community.

TABLE 36. Measured and calculated SACS for the IRDF-II cross sections in High Temperature Maxwellian Neutron Fields. Data from KADoNiS 1.0 [206] are the stellar Maxwellian averaged cross sections in the centre-of-mass system.

Reaction Notation	kT_{lab} [keV]	$E_{50\%}$ [keV]	Measured SACS		Reference	Calculated SACS			C/E
			Value [mb]	Unc. [%]		Value [mb]	Unc. [%]	Spec. Unc. [%]	
Mn55g	30.55	25.78	3.315E+01	9.6	[206]	3.283E+01	6.05	0.00	0.990 ± 11.3%
Cu63g	30.48	25.84	6.093E+01	10.4	[206]	7.144E+01	9.70	0.00	1.173 ± 14.2%
Au197g	25.00	26.64	6.450E+02	3.9	[209]	6.899E+02	0.69	0.00	1.070 ± 4.1%
Co59g	30.51	27.05	4.379E+01	4.6	[206]	3.444E+01	2.57	0.00	0.787 ± 5.3%
U238g	25.30	29.00	3.910E+02	4.35	[210]	3.899E+02	1.44	0.00	0.997 ± 4.6%
Nb93g	30.32	29.46	2.807E+02	2.4	[206]	2.638E+02	0.78	0.00	0.940 ± 2.5%
La139g	30.22	29.52	3.284E+01	9.7	[206]	4.159E+01	10.8	0.00	1.266 ± 14.5%
Au197g	28.00	30.25	6.160E+02	2.8	[208]	6.451E+02	0.64	0.00	1.047 ± 2.8%
Au197g	28.00	30.25	6.200E+02	2.6	[208]	6.451E+02	0.64	0.00	1.040 ± 2.7%
Ta181g	30.17	31.71	7.765E+02	2.5	[206]	7.506E+02	3.87	0.00	0.967 ± 4.6%
Sc45g	30.67	32.58	6.234E+01	3.9	[206]	6.797E+01	6.99	0.00	1.090 ± 8.0%
Au197g	30.15	32.65	6.200E+02	1.8	[206]	6.181E+02	0.61	0.00	0.997 ± 1.9%
In113gm	30.26	35.03	1.561E+02	29.9	[206]	1.616E+02	4.05	0.00	1.035 ± 30.2%
In115gm	30.26	35.04	6.305E+02	30.1	[206]	6.080E+02	4.05	0.00	0.964 ± 30.4%
In113gg	30.27	35.04	7.867E+02	10.7	[206]	7.696E+02	0.00	0.00	0.978 ± 10.7%
Na23g	31.30	35.27	1.825E+00	3.1	[206]	2.023E+00	8.49	0.00	1.109 ± 9.1%
In113g	30.27	35.91	8.201E+02	14.0	[206]	9.539E+02	0.00	0.00	1.163 ± 14.0%
Fe58g	30.52	37.38	1.450E+01	6.5	[206]	1.506E+01	16.3	0.00	1.039 ± 17.5%
W186g	30.16	37.46	2.291E+02	5.5	[206]	1.920E+02	1.93	0.00	0.838 ± 5.8%
Ag109g	30.28	42.23	4.075E+01	3.8	[206]	5.785E+01	6.76	0.00	1.420 ± 7.7%
U238g	426.0	632.2	1.080E+02	3.70	[210]	3.899E+02	1.82	0.00	1.011 ± 4.1%

The Atlas value of the resonance integrals of ^{45}Sc and ^{55}Mn are within two-sigma uncertainty compared to the values derived from the Q_0 factors in the Kayzero library, but the resonance integrals of ^{50}Cr , ^{63}Cu and ^{113}In disagree with the values derived from the Kayzero library, as seen from Table 35. The value for ^{113}In was derived from the total capture resonance integral, assuming the same branching ratio as for the thermal cross section, which might be inappropriate. The values calculated from the IRDF-II library generally agree to within two-sigma with Kayzero, except ^{50}Cr . The neutron-induced resonance data of these nuclides and particularly for $n+^{50}\text{Cr}$ remain an open issue that requires further investigation.

N. High Temperature Maxwellian ($kT \approx 25\text{--}430$ keV)

Most of the experimental Maxwellian spectrum-averaged cross sections, MACS, were taken from the latest version 1.0 of KADoNiS [206] based on the gold standard $^{197}\text{Au}(n,\gamma) = 611.6 \pm 6.0$ mb. This database presents the so-called stellar SACS which were derived from the experimental by applying the factor $2/\sqrt{\pi}$ and converting them in the centre-of-mass system to the temperature $kT_{cm} = 30$ keV [207]. For comparison with these data we have computed the IRDF-II SACS for the corresponding $kT_{lab} = kT_{cm} * (1 + A)/A$, where A is the ratio of the mass of the target nucleus to that of a neutron. Additionally, all KADoNiS cross sections and their uncertainties were renormalized to the newly introduced SACS gold standard $^{197}\text{Au}(n,\gamma) = 620 \pm 11$ mb [32] which adds a +1.4% correction.

Several other measurements [208–210] were carried out, but were not included in KADoNiS yet. In particular, A. Wallner and co-workers [210] have measured SACS,

not by the traditional activation technique, but by the highly-accurate Accelerator Mass Spectroscopy technique (AMS). In addition to the measurement at typical neutron spectrum average energy of 25 keV they have measured SACS at higher $kT = 426$ keV. These experiments delivered their results in the laboratory system corresponding to different kT values, so the IRDF-II SACS were calculated at those temperatures.

The resulting C/E ratios with IRDF-II for the 17 dosimetry reactions addressed within the KADoNiS database and several new experiments, along with the associated experimental and IRDF-II cross section uncertainties, are listed in Table 36. The trends in the C/E ratios are displayed in Fig. 114. There the uncertainties of measurements and of the IRDF-II cross sections are displayed separately allowing to observe their contributions to the total C/E uncertainty.

As one can see, most of the IRDF-II Maxwellian averaged cross sections agree with measurements to within 1–2 sigma. However we do not observe a good agreement for (n,γ) reactions on ^{59}Co , ^{186}W and ^{109}Ag targets. Whether this is a problem in IRDF-II evaluations or it is an experimental problem remains an open question. The reaction $^{232}\text{Th}(n,\gamma)$ is a rare case among the IRDF-II (n,γ) reactions that is not validated in the Maxwellian-30 keV benchmark field yet.

VIII. CONSISTENCY OF CROSS SECTIONS IN STANDARD AND REFERENCE FIELDS

Because of the uncertainty in the neutron fluence for a given reactor operation, a better metric than a spectral index for a fission reactor field is the consistency of the set of measured activities in a given reactor field. The consistency is best captured by the overall chi-squared

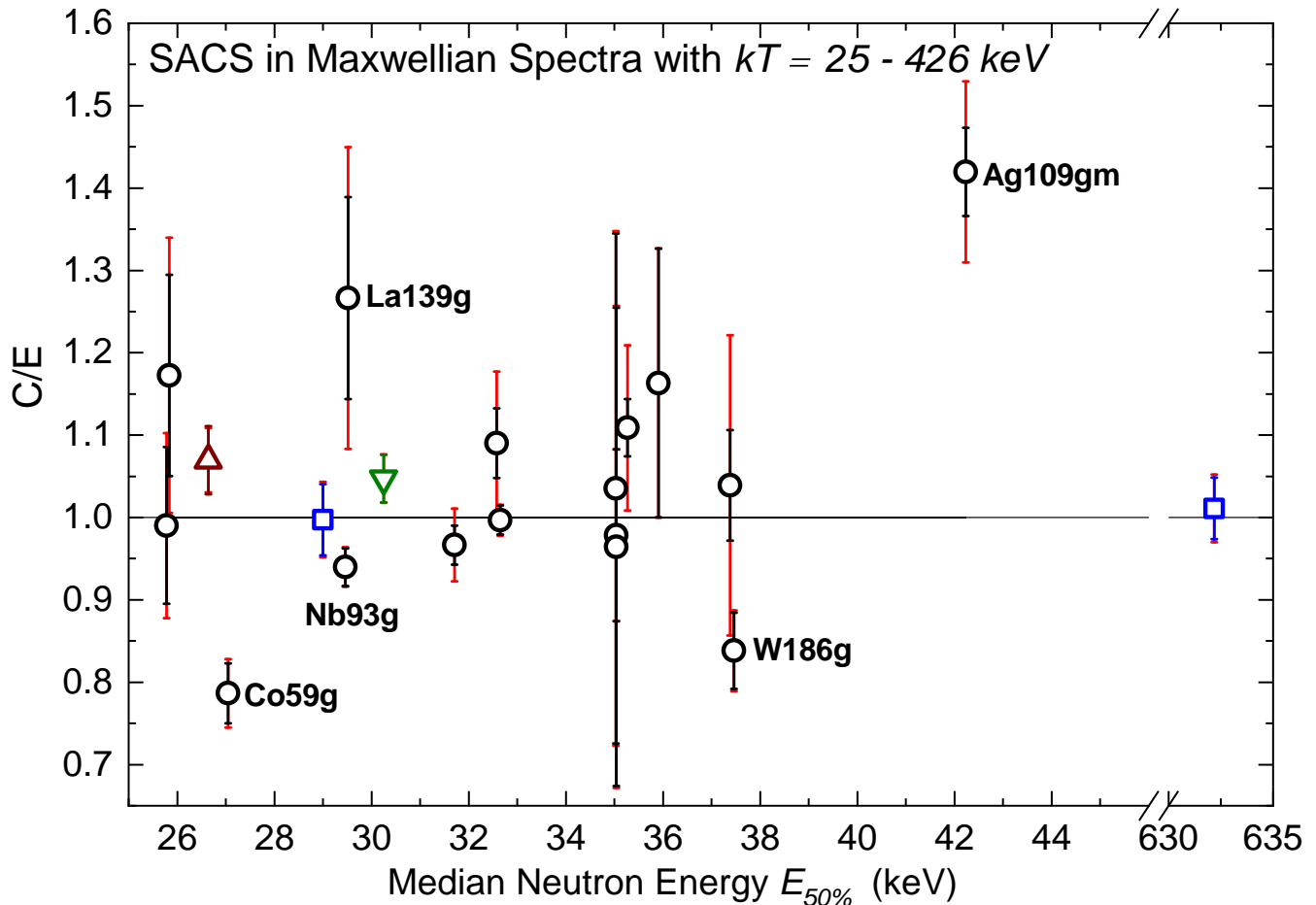


FIG. 114. (Color online) C/E ratios for SACS measured and calculated in the (25–426) keV Maxwellian spectra as a function of the median energy. Experimental data: \circ - KADoNiS-1.0 [206], other symbols – G. Feinberg *et al.* [208], A. Wallner *et al.* [210], P. Jimenez-Bonilla *et al.* [209]. Uncertainties resulting from measurements and IRDF-II cross sections are shown by black and red bars. Plotted values are listed in Table 36.

per degree of freedom (χ^2/dof) metric as obtained from a least-squares-based spectrum adjustment in neutron benchmark fields. The relevant reaction-specific metric is then the resulting least-squares-based activity adjustment and the C/E for the adjusted activity.

The following subsections address this least-squares metric for specific neutron benchmark fields.

A. $^{252}\text{Cf}(\text{s.f.})$

Sec. VII A reports on the validation evidence using individual measured values for the spectrum-averaged cross section in this standard neutron benchmark field. A more exacting validation metric, based on the consistency of the set of validation data gathered within this neutron field, is depicted in Table 37.

The LSL-M2 code was used to perform the least-squares analysis. The inputs included the measurements, with uncertainties, the IRDF-II cross sections, with covariance matrices, and the *a priori* spectrum, in this case as provided by time-of-flight measurements, along with the corresponding covariance matrix. This approach permitted all of the input data to be adjusted, consistent with the input uncertainties, but the major result was to

adjust the spectrum and to provide a high-fidelity test of the consistency of the data for the neutron field. A chi-squared per degree of freedom, χ^2/dof , is reported and provides a metric on the consistency of the data.

The least-squares calculation in this standard neutron benchmark field used 48 measured reactions. As expected from the consideration of the separate spectrum-averaged cross sections addressed in Sect. VI A, the measured data were very consistent with calculations and the overall least-squares spectrum adjustment had a very small χ^2/dof of 0.7042. A least-squares process using data for which there was an accurate identification of the *a priori* uncertainties should have a χ^2/dof metric of 1.00. This χ^2/dof less than unity suggests that some of the input data (spectrum, cross sections, or measured activities) actually may have had smaller uncertainties than were assigned to the input quantities.

The resulting adjustment in the spectrum-averaged activity, the reduced spectrum uncertainty contributions to the C/E, and the C/E values are tabulated in Table 37. The C/E column is color-coded in a manner consistent with the discussion in Sec. VI A. The discrepant reactions are shown in “red”, the “acceptable” reactions in “yellow”, and the “good” reactions in “green”. While a least-squares analysis can adjust all of the input data

TABLE 37. Least-squares-based consistency of data for the $^{252}\text{Cf}(\text{s.f.})$ neutron benchmark field

Reaction Notation	Act. [%]	Adj. Spectrum [%]	Unc. [%]	C/E
Au197g	-0.33%	0.42%		$0.991 \pm 1.79\%$
Ta181g	-0.35%	0.75%		$0.935 \pm 5.65\%$
Th232g	0.43%	0.43%		$1.032 \pm 5.53\%$
Cu63g	0.20%	0.61%		$1.010 \pm 8.64\%$
In115g	-1.20%	0.37%		$0.979 \pm 2.45\%$
U235f	1.94%	0.07%		$1.013 \pm 1.21\%$
Pu239f	-0.12%	0.07%		$0.992 \pm 1.38\%$
Np237f	0.66%	0.23%		$1.000 \pm 1.69\%$
In115n	-2.28%	0.38%		$0.964 \pm 1.57\%$
Nb93n	1.47%	0.36%		$1.010 \pm 3.73\%$
In113n	-1.39%	0.40%		$0.978 \pm 2.15\%$
U238f	-1.45%	0.40%		$0.978 \pm 1.70\%$
Th232f	-5.66%	0.43%		$0.935 \pm 5.08\%$
Hg199n	-0.54%	0.44%		$0.985 \pm 2.22\%$
BH3	-1.95%	0.51%		$0.931 \pm 11.58\%$
Ti47p	1.31%	0.62%		$1.015 \pm 2.16\%$
S32p	2.68%	0.73%		$1.023 \pm 3.65\%$
Zn64p	1.27%	0.78%		$1.009 \pm 3.31\%$
Ni58p	0.35%	0.75%		$1.000 \pm 1.66\%$
Fe54p	1.36%	0.78%		$1.015 \pm 1.79\%$
Pb204n	-1.58%	0.97%		$0.971 \pm 2.74\%$
Al27p	-4.51%	1.14%		$0.950 \pm 2.51\%$
Co59p	1.12%	1.15%		$1.015 \pm 3.16\%$
Ti46p	-1.44%	1.18%		$0.974 \pm 2.63\%$
Si28p	2.18%	1.41%		$1.031 \pm 2.67\%$
Cu63a	0.22%	1.37%		$1.006 \pm 2.93\%$
Fe56p	-0.23%	1.45%		$0.998 \pm 2.91\%$
Mg24p	4.59%	1.56%		$1.050 \pm 2.92\%$
U2382	3.92%	1.56%		$1.045 \pm 11.23\%$
Ti48p	0.07%	1.55%		$1.005 \pm 3.53\%$
Co59a	-0.14%	1.54%		$1.000 \pm 3.39\%$
Al27a	-0.11%	1.59%		$1.003 \pm 2.07\%$
U51a	-0.98%	1.85%		$0.989 \pm 3.23\%$
Au1972	0.04%	1.95%		$1.005 \pm 2.85\%$
Nb932	-0.44%	2.20%		$1.000 \pm 4.95\%$
I1272	1.10%	2.30%		$1.018 \pm 3.87\%$
Cu652	-0.84%	2.90%		$0.993 \pm 3.70\%$
Co592	0.36%	3.24%		$1.006 \pm 4.27\%$
Cu632	3.32%	4.10%		$1.032 \pm 5.74\%$
F192	1.09%	4.39%		$1.007 \pm 5.66\%$
Zr902	-1.39%	4.99%		$0.981 \pm 5.77\%$
Ni582	0.87%	5.90%		$0.997 \pm 7.07\%$
Ni60p	2.41%	1.35%		$1.027 \pm 5.03\%$
Tm1692	-1.75%	1.91%		$0.986 \pm 4.74\%$
Mn552	-1.49%	3.04%		$0.958 \pm 15.63\%$
Y892	1.21%	4.19%		$0.984 \pm 5.64\%$
Na232	-4.46%	6.79%		$0.990 \pm 7.63\%$
Tm1693	-0.33%	14.75%		$0.795 \pm 16.24\%$

that have identified associated uncertainties, changing their contribution to the uncertainty in the resulting C/E value, this table does not provide the details of the resulting cross section and measurement uncertainty contributions since these uncertainties are not changed significantly from those presented in the spectrum-averaged cross section comparison in Table 37. While the spectrum contribution to the overall C/E uncertainty is reduced here relative to that seen in the spectrum-averaged cross section comparison, the difference is not large because of the very-well-characterized *a priori* $^{252}\text{Cf}(\text{s.f.})$ spec-

trum, which was measured using time-of-flight techniques, used in the analysis. In less-well-characterized neutron fields, the least-squares analysis will significantly reduce the spectrum contribution to the resulting uncertainty in the C/E value.

Fig. 115 shows the C/E values in this $^{252}\text{Cf}(\text{s.f.})$ neutron benchmark field plotted versus the median response energy for the 48 reactions used in the least-squares adjustment. There is no significant trend seen in the C/E ratios versus the median response energy. The C/E uncertainties are clearly much larger in the low and high energy portion of the spectrum. The adjustment produced a 1.0% adjusted scale factor. The scale factor adjustment is taken out of the C/E values in this figure.

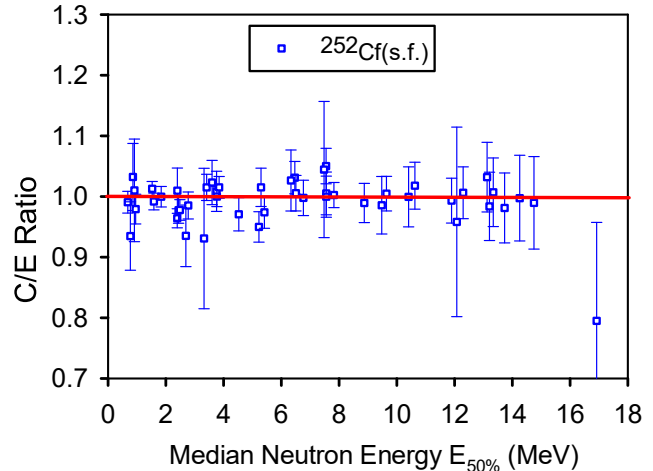


FIG. 115. (Color online) Least-squares-based C/E as a function of median energy $E_{50\%}$ for the activity measurements in the $^{252}\text{Cf}(\text{s.f.})$ standard neutron benchmark Field. Uncertainties bars reflect the combined rms uncertainty contributions from the measurement, cross sections, and spectrum. Plotted values are listed in Table 37.

The largest contributor to the overall χ^2/dof came from the Al27p reaction, which had an individual χ^2 contribution of 4.02, but, based on the acceptance criteria presented in Table 17, the Al27p reaction, with a C/E of $0.950 \pm 2.51\%$, still showed a “good” validation status in this neutron field. The next largest contributor to the overall χ^2/dof came from the In115n reaction, which had an individual χ^2 contribution of 3.23, with a C/E of $0.964 \pm 1.57\%$. The color coding in Table 37 shows that this reaction, because of its small overall C/E uncertainty, was just outside of the required two standard deviations and has a status of “acceptable” rather than “good” as validation evidence. The only “poor” validation status seen in the data available in this benchmark field came from the Tm1693 reaction. This reaction, which had an individual χ^2 contribution of 2.37, had a C/E of $0.795 \pm 16.24\%$, which, even though its large uncertainty gave it a C/E that was well within even the two standard deviations of unity required by Table 17 criterion #1 for a “good” status, it failed to meet the Table 17 C/E interval criterion #2 for even an “acceptable” status.

Note that the uncertainty of the TOF measured *a priori* spectrum is larger than 15% at the reaction threshold of 16 MeV and quickly rises up to 75% at 20 MeV, there-

fore the least-squares fit is not well constrained in the energy region where this particular dosimeter is sensitive. The least-squares analysis showed a very strong upward adjustment (+19.3%) in the high energy (> 17 MeV) portion of the spectrum and a downward shift in the reaction rate (-4.5%), that is, the least-squares process made adjustments in both the reaction probability and the spectrum in order to minimize the resulting disagreement in this high energy region.

B. $^{235}\text{U}(n_{th},f)$

Sect. VII B reports on the validation evidence using individual measured values for the spectrum-averaged cross sections in this reference neutron benchmark field. The more exacting validation metric, based on the consistency of the set of validation data gathered within this neutron field as assessed by the LSL-M2 least square code, is depicted in Table 38. The least square inputs included the measurements, with uncertainties, the IRDF-II cross sections, with covariance matrices, and the *a priori* spectrum, in this case as provided from the GMA analysis and incorporated into the ENDF/B-VIII.0 evaluation as addressed in Sec. VII B, along with the corresponding covariance matrix. The chi-squared per degree of freedom, χ^2/dof , is reported from the analysis and provides a metric on the consistency of the input data.

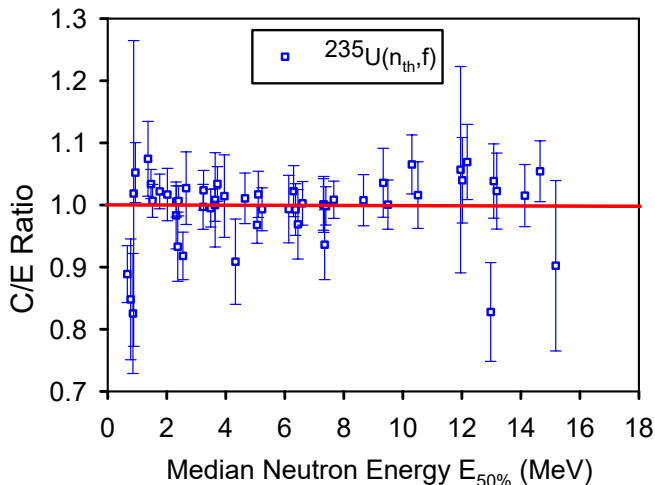


FIG. 116. (Color online) Least-squares-based C/E as a function of median energy $E_{50\%}$ for the activity measurements in the $^{235}\text{U}(n_{th},f)$ reference neutron benchmark Field. Uncertainties bars reflect the combined rms uncertainty contributions from the measurement, cross sections, and spectrum. Plotted values are listed in Table 38.

The least-squares calculation used 52 measured reactions. Two reactions that were included in the SACS analysis were not included in this least-squares analysis: the B10t reaction was not addressed in the analysis because it was correlated with the B10a reaction that was used and the LSL-M2 code implementation used here did not handle strongly correlated cross sections; the Zr902 reaction was not addressed in the analysis because it was used in support of the determination of the *a priori* $^{235}\text{U}(n_{th},f)$

TABLE 38. Least-squares-based consistency of data for the $^{235}\text{U}(n_{th},f)$ neutron reference benchmark field.

Reaction Notation	Act. Adj. [%]	Spectrum Unc. [%]	C/E
Li6He4	4.69%	0.65%	1.074 ± 6.02%
Au197g	-11.27%	0.67%	0.889 ± 4.58%
B10a	-3.82%	0.64%	0.848 ± 9.72%
U238g	-17.62%	0.64%	0.825 ± 9.66%
Cu63g	1.46%	0.61%	1.019 ± 24.61%
In115gm	4.78%	0.60%	1.052 ± 4.83%
U235f	3.11%	0.52%	1.034 ± 2.35%
Pu239f	0.4%	0.52%	1.007 ± 2.65%
Np237f	1.82%	0.56%	1.022 ± 2.79%
Rh103n	1.24%	0.57%	1.017 ± 4.22%
Nb93nm	-1.81%	0.62%	0.983 ± 5.38%
In115nm	0.28%	0.63%	1.006 ± 2.52%
In113nm	-6.94%	0.65%	0.933 ± 5.57%
U238f	0.4%	0.65%	1.007 ± 2.40%
Th232f	-8.05%	0.65%	0.918 ± 3.83%
Hg199n	2.34%	0.65%	1.027 ± 5.84%
P31p	-0.49%	0.78%	0.997 ± 3.62%
Ti47p	1.78%	0.73%	1.024 ± 3.19%
S32p	-0.67%	0.82%	0.995 ± 3.04%
Ni58p	-0.19%	0.80%	1.000 ± 2.62%
Zn64p	0.6%	0.85%	1.008 ± 7.59%
Fe54p	2.84%	0.84%	1.034 ± 2.78%
Zn67p	1.01%	0.80%	1.014 ± 6.64%
Pb204n	-8.62%	0.93%	0.909 ± 6.87%
Mo92p	0.67%	0.97%	1.011 ± 4.07%
Al27p	-3.01%	1.08%	0.968 ± 2.96%
Co59p	1.25%	1.07%	1.017 ± 3.72%
Ti46p	-0.5%	1.13%	0.993 ± 3.45%
Ni60p	-0.6%	1.34%	0.994 ± 5.43%
Cu63a	1.72%	1.36%	1.022 ± 4.11%
Si28p	-0.45%	1.56%	0.993 ± 4.16%
Fe54a	-2.1%	1.44%	0.969 ± 5.60%
Fe56p	0.21%	1.54%	1.003 ± 3.51%
Ti48p	-0.04%	1.70%	1.001 ± 4.15%
Co59a	-0.03%	1.68%	1.001 ± 4.50%
U2382	-5.03%	1.87%	0.936 ± 5.63%
Mg24p	-0.08%	1.83%	0.999 ± 3.08%
Al27a	0.65%	1.86%	1.009 ± 3.00%
V51a	1.79%	2.28%	1.008 ± 4.12%
Tm1692	-1.5%	2.74%	1.036 ± 5.55%
Au1972	3.55%	2.79%	1.001 ± 3.96%
Nb932m	-1.25%	3.61%	1.065 ± 4.75%
I1272	0.12%	3.79%	1.016 ± 5.36%
Mn552	0.55%	4.72%	1.057 ± 16.61%
As752	3.25%	4.81%	1.040 ± 6.88%
Co592	-19.81%	4.95%	1.069 ± 6.05%
Cu632	1.11%	4.94%	0.828 ± 7.95%
Y892	-0.29%	5.06%	1.039 ± 5.98%
F192	-0.31%	4.60%	1.023 ± 6.12%
Ni582	3.92%	3.42%	1.015 ± 5.00%
Na232	-10.7%	2.86%	1.054 ± 4.89%
Al272g	4.69%	2.30%	0.902 ± 13.72%

spectrum and this would have introduced a strong correlation that the LSL-M2 code could not address. As expected from the consideration of the separate spectrum-averaged cross sections addressed in Sect. VIB 1, most of the reactions included in the least-squares adjustment proved to provide good or acceptable validation evidence, and the least-squares analysis yielded a very good χ^2/dof of

0.943.

There were no “poor” or discrepant reactions. Four reactions (Au197g with an individual χ^2 contribution of 6.14 and a C/E of $0.889 \pm 4.58\%$; Th232f with a χ^2 contribution of 4.84 and a C/E of $0.918 \pm 3.83\%$; U238g with a χ^2 contribution of 3.33 and a C/E of $0.825 \pm 9.66\%$; and B10a with an χ^2 contribution of 2.48 and a C/E of $0.848 \pm 9.72\%$) were labeled as “acceptable” validation evidence, and the remaining reactions met the criteria for “good” validation evidence. The U238g reaction was seen to be labeled as “acceptable” in both the SACS analysis shown in Table 20 and in this least-squares analysis. The Cu632 and Pb204n reactions were only labeled as “acceptable” in the SACS analysis but were both “good” in this least squares analysis. The B10a, Au197g, and Th232f reactions are only labeled as “acceptable” in this least square analysis but met the criteria for “good” validation evidence in the SACS analysis. The fact that these three reactions also corresponded to the reactions with the lowest median response energy might indicate that it is the low energy portion of the *a priori* $^{235}\text{U}(n_{th},f)$ spectrum, and not the cross sections or measurements, that has an issue.

The resulting adjustment in the spectrum-averaged activities, the reduced spectrum uncertainty contributions to the C/E, and C/E values are tabulated in Table 38. The C/E column is again color-coded in a manner consistent with the discussion in Sec. VIB 1. Overall good or acceptable validation evidence, using these least-squares validation criteria, is found for all 52 IRDF-II reactions in the $^{235}\text{U}(n_{th},f)$ reference neutron benchmark field. It should be noted that, while the *a priori* $^{252}\text{Cf}(s.f.)$ neutron spectrum showed a high uncertainty and a large least-squares adjustment in the high energy (15–20 MeV) region, neither this large uncertainty nor the large least-squares adjustment were seen for the *a priori* $^{235}\text{U}(n_{th},f)$ spectrum. This observation is related to the role that the Zr902 reaction played in the determination of the *a priori* $^{235}\text{U}(n_{th},f)$ spectrum by significantly reducing the uncertainty of the evaluated spectra due to the low uncertainty of the employed Zr902 SACS (7%). Al272 was the reaction in this $^{235}\text{U}(n_{th},f)$ field that had the highest median response energy. While this reaction, with a median response energy of 15.18 MeV, had a significant measurement uncertainty (13.3%), a very small individual χ^2 component, 0.58, and a good C/E of $0.902 \pm 13.72\%$, the least-squares analysis resulted in only a 0.3% adjustment in the neutron spectrum above 15 MeV. This least-squares analysis is a very strong indication of the fidelity of the > 10 MeV portion of the *a priori* $^{235}\text{U}(n_{th},f)$ spectrum as defined in original PFNS evaluations [98–100] adopted for the ENDF/B-VIII.0 library [30, 40, 41].

Fig. 116 shows the C/E values in this $^{235}\text{U}(n_{th},f)$ neutron benchmark field plotted versus the median response energy for the 52 reactions used in the final least-squares adjustment. The adjustment produced a 0.3% adjusted scale factor. The scale factor adjustment is taken out of the C/E values in this figure. There is no significant trend seen in the C/E ratios versus the median response energy. The C/E uncertainties are larger for several of the reactions in the low and high energy portion of the spectrum

that reflects uncertainties of the *a priori* spectrum in those energy regions.

TABLE 39. Least-squares-based consistency of data for the SPR-III central cavity neutron reference benchmark field.

Reaction Notation	Act. Adj. Spectrum Unc.		C/E
	[%]	[%]	
Ni58p-Cd	-4.89	10.48	1.000 ± 10.96%
Sc45g-Cd	0.2	5.93	0.994 ± 8.90%
Mn55g-Cd	-3.43	5.64	0.802 ± 8.31%
Au197g-bare	-15.58	5.58	0.839 ± 7.17%
Au197g-Cd	-6.78	5.72	0.929 ± 7.28%
Sc45g-fiss	1.71	6.05	1.051 ± 8.57%
Na23g-bare	0.53	6.37	1.129 ± 11.26%
Fe58g-Cd	0.66	5.86	1.105 ± 12.04%
Na23g-Cdna	0.21	6.62	1.075 ± 11.78%
Mn55g-fiss	1.14	5.69	1.082 ± 8.69%
W186g-bare	-20.28	5.11	0.818 ± 10.41%
Cu63g-Cd	0.04	5.52	1.041 ± 11.99%
Na23g-fiss	0.59	6.24	1.178 ± 11.94%
U235f-Cdtk	-1.63	5.43	1.045 ± 6.21%
U235f-fiss	2.07	5.51	1.089 ± 6.27%
Pu239f-Cdtk	3.36	5.6	1.109 ± 6.21%
Pu239f-fiss	8.33	5.68	1.165 ± 6.29%
Np237f-Cdtk	-0.01	6.99	1.118 ± 7.55%
Np237f-fiss	-1.46	7.01	1.103 ± 7.57%
In115nm-bare	-0.84	9.03	1.068 ± 10.20%
U238f-Cd	-1.54	9.87	1.046 ± 10.37%
U238f-fiss	-2.6	9.85	1.036 ± 10.36%
Ti47p-Cd	4.05	9.77	1.102 ± 10.22%
S32p-bare	-5.13	11.07	0.990 ± 11.50%
Zn64p-Cd	2.35	11.19	1.077 ± 11.42%
Fe54p-Cd	-0.22	10.91	1.049 ± 11.40%
Al27p-Cd	6.4	12.06	1.138 ± 12.45%
Co59p-bare	-7.44	11.83	0.983 ± 13.04%
Ti46p-Cd	-0.9	12.4	1.054 ± 12.96%
Cu63a-bare	-10.01	12.86	0.966 ± 13.97%
Fe56p-Cd	-0.35	14.11	1.076 ± 14.43%
Ti48p-Cd	0.31	13.88	1.105 ± 14.22%
Mg24p-Cd	3.48	15.41	1.122 ± 15.71%
Al27a-Cd	-1.46	14.78	1.075 ± 14.90%
Nb932-bare	3.25	16.47	1.113 ± 17.14%
Co592-bare	4.16	14.94	1.087 ± 16.60%
Mn552-bare	-16.09	15.06	0.878 ± 17.15%
Mn552-fiss	-3.18	15.05	1.009 ± 15.90%
Zr902-Cd	5.45	15.17	1.142 ± 15.55%
Ni582-bare	-8.87	15.62	1.014 ± 16.69%

C. SPR-III Fast Burst Reactor Central Cavity

The least-squares analysis used 40 reaction/cover combinations and had a χ^2/dof of 2.409. Although this chi-squares is high, none of the resulting dosimeters are found to be discrepant when we use the acceptance criteria adopted in Table 17. Table 39 shows the resulting C/E along with the consistent least-squares spectrum uncertainty and adjustment in the activity measurement. Fig. 117 shows the C/E values in the SPR-III reference neutron benchmark field plotted versus the median response energy for the 40 reactions used in the final least-squares adjustment.

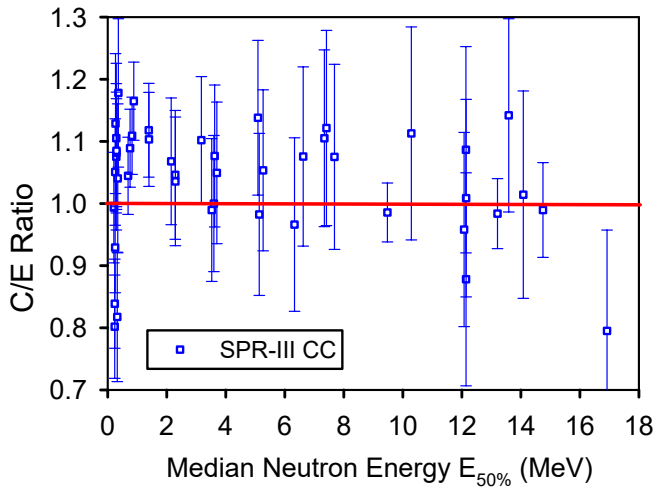


FIG. 117. (Color online) Least-squares-based C/E as a function of median energy $E_{50\%}$ for the activity measurements in the SPR-III fast-burst reactor central cavity reference neutron benchmark field. Uncertainties bars reflect the combined rms uncertainty contributions from the measurement, cross sections, and spectrum (remove bias). Plotted values are listed in Table 39.

D. ACRR Pool-type Reactor Central Cavity

The least-squares analysis for this neutron field used 33 reaction/cover combinations and had a χ^2/dof of 0.8984. This reflects an excellent consistency of the data (and associated input uncertainties). Table 40 shows the resulting C/E along with the consistent least-squares spectrum uncertainty and adjustment in the activity measurement. Two of the 33 reactions, Ag109g-bare and Co59g-Cd, were labeled as discrepant based on the criteria in Table 17 – due to the large C/E value even if this value was within two standard deviations of unity.

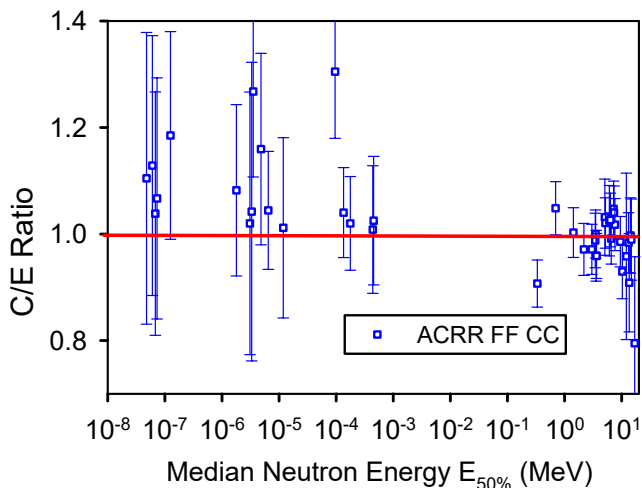


FIG. 118. (Color online) Least-squares-based C/E as a function of median energy $E_{50\%}$ for the activity measurements in the ACRR pool-type reactor central cavity reference neutron benchmark field. Uncertainties bars reflect the combined rms uncertainty contributions from the measurement, cross sections, and spectrum. Plotted values are listed in Table 40.

TABLE 40. Least-squares-based consistency of data for the ACRR central cavity neutron reference benchmark field.

Reaction	Act. Adj.	Spectrum	
Notation	[%]	Unc. [%]	C/E
Ni58p-bare	1.67	2.75	$1.000 \pm 4.53\%$
Sc45g-bare	0.16	26.17	$1.105 \pm 27.39\%$
Na23g-bare	3.27	24.15	$1.129 \pm 24.40\%$
Mn55g-bare	-3.83	22.67	$1.038 \pm 22.85\%$
Fe58g-bare	-0.17	22.09	$1.067 \pm 22.64\%$
Co59g-bare	1.87	18.88	$1.185 \pm 19.50\%$
Sc45g-Cd	0.07	14.32	$1.082 \pm 16.09\%$
Au197g-bare	-1.92	24.32	$1.020 \pm 24.66\%$
Au197g-Cd	0.94	27.71	$1.042 \pm 28.04\%$
Ag109g-bare	4.66	14.74	$1.268 \pm 16.04\%$
Ag109g-Cd	-0.8	16.34	$1.160 \pm 17.98\%$
Na23g-Cd	-1.05	10.17	$1.045 \pm 11.08\%$
W186g-bare	-0.12	16.5	$1.012 \pm 16.93\%$
Co59g-Cd	4.26	11.53	$1.305 \pm 12.53\%$
Mn55g-Cd	0.66	7.73	$1.040 \pm 8.45\%$
Fe58g-Cd	-0.18	7.31	$1.020 \pm 8.80\%$
rmleu-fiss	-7.44	1.8	$0.907 \pm 4.42\%$
rmlpu-fiss	5.04	1.87	$1.049 \pm 4.99\%$
Np237f-fiss	1.86	2.29	$1.003 \pm 4.67\%$
rmldu-fiss	-0.89	2.71	$0.971 \pm 4.89\%$
Ti47p-bare	-0.87	2.59	$0.971 \pm 4.71\%$
S32p-bare	0.73	2.93	$0.988 \pm 5.12\%$
Zn64p-bare	-2.15	2.95	$0.959 \pm 4.77\%$
Fe54p-bare	-2.18	2.87	$0.959 \pm 4.23\%$
Co59p-Cd	2.95	3.12	$1.032 \pm 7.16\%$
Ti46p-bare	1.43	3.23	$1.021 \pm 4.74\%$
Ni60p-bare	1.39	3.28	$1.025 \pm 6.62\%$
Fe56p-bare	-1.65	3.49	$0.992 \pm 4.81\%$
Ti48p-bare	1.35	3.5	$1.048 \pm 5.20\%$
Nb932-bare	-2.17	4.23	$0.930 \pm 5.10\%$
Mg24p-bare	1.93	3.91	$1.041 \pm 4.98\%$
Al27a-bare	0.19	3.8	$1.018 \pm 4.89\%$
Zr902-bare	-1.89	7.43	$0.909 \pm 9.23\%$

The table reflects a large spectrum contribution to the uncertainty, and there is also seen to be a correlation in the C/E between the two configurations, bare and cover, – both configurations showing a positive C/E value. It is very possible that the discrepancy reflects an issue with the *a priori* neutron spectrum, or the associated uncertainty, rather than with the dosimetry cross section. It should also be noted that both of these basic reactions appear in the set of sensors in both, a bare configuration, and with a cadmium cover; only one of the sensor configurations was labelled as discrepant. It is important that we also note a limitation in our current least-squares treatment. While a least-squares analysis should capture any correlation between input quantities, the LSL code used here did not include any correlation between the dosimetry responses. While this is not an issue for most dosimetry reactions, it could be an issue when for the response functions that address two different configurations for the same basic reaction. This is an area for later tool development. Fig. 118 shows the C/E values in the ACRR central cavity reference neutron benchmark field plotted versus the median response energy for the 33 reactions used in the final least-squares adjustment.

TABLE 41. Least-squares-based consistency of data for the ACRR LB44 neutron reference benchmark field.

Reaction Notation	Measured Activity [Bq/at or Fiss/at.]	Act. Unc. [%]	Adj. Unc. [%]	Spectrum Unc. [%]	C/E \pm Unc. [%]
Co59g-bare	6.2470E-18	0.8	0.84	14.15	1.485 \pm 14.33
Au197g-bare	2.4293E-14	2.8	-0.87	9.67	1.039 \pm 10.59
Mn55g-bare	4.0558E-14	2.0	2.08	4.81	1.195 \pm 6.37
Fe58g-bare	1.2442E-17	2.3	-0.81	4.54	1.095 \pm 8.08
Mo98g-bare	1.6478E-15	1.8	0.18	4.02	1.110 \pm 9.93
Cu63g-bare	6.6548E-15	5.7	-11.28	4.34	0.859 \pm 8.91
Sc45g-bare	1.6744E-17	0.7	0.11	2.8	1.171 \pm 15.18
Co59g-Cdtk/B4C	2.0990E-19	0.8	0.27	3.65	1.167 \pm 4.38
Au197g- Cdtk/B4C	2.7153E-15	4.2	-1.53	2.15	1.056 \pm 6.37
Fe58g-Cdtk/B4C	2.7876E-18	7.2	3.06	2.9	1.180 \pm 13.44
Mo98g-Cdtk/B4C	3.9782E-16	2.2	3.27	2.82	1.286 \pm 8.07
rmleu-Cdtk/B4C	3.8036E-09	2.1	-1.51	4.39	1.046 \pm 5.27
rmlpu-Cdtk/B4C	4.5468E-09	2.0	-2.73	5.69	1.007 \pm 6.76
Np237f-Cdtk/B4C	1.9753E-09	3.0	0.7	10.2	1.032 \pm 10.83
In115n-bare	9.8199E-15	5.6	-7.47	13.09	0.956 \pm 14.39
rmldu-Cdtk/B4C	2.8865E-10	2.0	2.44	14.44	1.072 \pm 14.72
Ti47p-bare	3.6124E-17	2.6	0.99	13.07	1.046 \pm 13.55
S32p-bare	2.6391E-17	2.9	-0.76	14.54	1.016 \pm 14.98
Ni58p-bare	9.5960E-18	1.6	-2.78	13.63	1.000 \pm 13.89
Fe54p-bare	1.4959E-18	1.1	1.64	14.18	1.048 \pm 14.39
Co59p-bare	1.6482E-19	2.4	-1.49	16.66	1.019 \pm 17.08
Ti46p-bare	6.7205E-19	0.8	0.93	17.24	1.057 \pm 17.45
Ni60p-bare	5.4018E-21	6.0	-0.2	17.11	1.053 \pm 18.27
Fe56p-bare	4.7933E-17	0.9	-0.22	17.89	1.056 \pm 18.13
Ti48p-bare	7.7602E-19	0.6	0.01	18.16	1.061 \pm 18.51
Al27a-bare	5.2690E-18	3.7	0.14	20.11	1.059 \pm 20.54
Nb932-bare	2.0258E-19	1.0	0	16.1	1.097 \pm 16.26
Co592-bare	1.2528E-20	12.0	8.36	20.27	1.206 \pm 23.68
Mn552-bare	3.5010E-21	16.0	-12.24	20.46	0.999 \pm 26.07

E. Other “Bucket”-Modifying Environments in Reference Reactor Fields

The least-squares analysis for the SPR-III fast burst reactor central cavity and the pool-type ACRR central cavity reactor were good validation cases for the range of activities that are easily measured in research reactors, as shown in the previous section. The consistency of activity measurements for other reactor spectra produced using spectrum modifying buckets has also been considered in our validation. Tables 41, 42, 43 and 44 present the measured activities and the C/E ratios obtained from the least-squares adjustment. The C/E ratios are color-coded to indicate the status of the consistency.

Inspection of these tables shows that, while providing a larger quantity of validation evidence, the consideration of these other Bucket-Modifying Environments did not significantly increase the scope of reactions for which we have validation evidence over what is seen in the SPR-III Cen-

tral Cavity and ACRR Central Cavity least-squares analyses. Consideration of the consistency of dosimeters in the least-squares analysis for these other research reactor configurations, addressing the following neutron benchmark fields, produced the following results:

1. ACRR LB44 with 29 sensors, a $\chi^2/dof = 1.279$, and 2 discrepant dosimeter/cover configurations (Co59g-bare, Mo98g-Cdtk-B4C);
2. ACRR PLG with 35 sensors, a $\chi^2/dof = 1.025$, and 1 discrepant dosimeter/cover configurations (Mn552-bare);
3. ACRR Cd-poly with 31 sensors, a $\chi^2/dof = 1.183$, and 1 discrepant dosimeter/cover configuration (Np237f-Cdtkj-B4C);
4. FREC-II with 33 sensors, a $\chi^2/dof = 1.711$, and 0 discrepant dosimeter/cover configurations.

TABLE 42. Least-squares-based consistency of data for the ACRR PLG neutron reference benchmark field.

Reaction Notation	Measured Activity		Act. Adj. Spectrum		C/E \pm Unc. [%]
	[Bq/at. or Fiss/at.]	Unc. [%]	Unc. [%]	Unc. [%]	
Sc45g-bare	1.6579E-15	2.38	0.06	30.63	1.216 \pm 31.62
Na23g-bare	3.9324E-15	2.38	1.7	29.74	1.245 \pm 29.90
Fe58g-bare	1.6831E-16	2.90	-0.1	28.23	1.205 \pm 28.68
Cu63g-bare	5.0078E-14	3.69	-2.61	28.17	1.164 \pm 28.52
Co59g-bare	1.1350E-16	2.50	-0.56	26.6	1.205 \pm 26.72
Sc45g-Cd	1.2948E-16	2.38	-0.65	14.74	1.054 \pm 16.43
Au197g-bare	7.1560E-13	2.69	-0.35	22.13	1.078 \pm 22.33
Au197g-Cd	5.3238E-13	2.69	0.21	28.07	1.026 \pm 28.26
Na23g-Cd	4.1746E-16	2.38	0.71	10.97	1.114 \pm 11.60
Co59g-Cd	2.2206E-17	2.50	0.86	11.24	1.169 \pm 11.54
Fe58g-Cd	2.6925E-17	2.90	0.89	7.34	1.072 \pm 8.90
Mn55g-Cd	1.1075E-13	2.62	-1.39	7.57	1.022 \pm 8.32
Cu63g-Cd	7.5389E-15	3.69	2.37	7.1	1.095 \pm 8.48
rmleu-Cdtk/B4C	2.5730E-09	2.83	-6.39	1.77	0.927 \pm 3.35
rmlpu-Cdtk/B4C	2.5696E-09	2.83	4.06	1.84	1.059 \pm 4.08
In115n-bare	7.0790E-15	5.95	-7.32	2.57	0.921 \pm 6.51
rmldu-Cdtk/B4C	2.1936E-10	2.90	1.78	2.72	1.014 \pm 4.01
Ti47p-bare	2.1537E-17	4.12	-2.23	2.58	0.974 \pm 4.71
S32p-bare	7.7936E-18	2.62	1.49	2.92	1.017 \pm 5.11
Ni58p-bare	3.7146E-16	3.61	-0.1	2.74	1.000 \pm 3.87
Zn64p-bare	1.2772E-18	3.36	-1.68	2.94	0.985 \pm 4.70
Fe54p-bare	1.4707E-19	3.36	1.13	2.87	1.014 \pm 4.51
Co59p-Cd	6.1277E-19	2.56	-0.04	3.12	1.013 \pm 4.98
Ti46p-bare	5.2239E-21	3.69	2.57	3.23	1.051 \pm 4.44
Ni60p-Cd	1.4983E-21	8.15	-3.24	3.28	0.989 \pm 5.04
Cu63a-bare	4.6190E-17	2.69	-17.86	3.22	0.839 \pm 8.96
Fe56p-bare	7.1657E-19	2.24	-1.64	3.49	1.009 \pm 4.81
Ti48p-bare	9.8561E-18	2.69	0.23	3.51	1.046 \pm 5.11
Mg24p-bare	4.8547E-18	2.97	4.87	3.92	1.098 \pm 4.77
Al27a-bare	1.9054E-19	2.33	-0.28	3.8	1.044 \pm 4.84
Nb932-bare	3.5325E-21	20.3	0.26	4.23	1.005 \pm 4.85
Mn552-bare	1.2657E-20	18.2	-18.09	5.4	0.764 \pm 21.04
Co592-Cd	1.4115E-19	3.61	-0.84	5.38	0.937 \pm 19.04
Zr902-bare	1.1537E-20	6.51	-1.98	7.49	0.882 \pm 8.32
Ni582-Cd	1.6579E-15	2.38	-0.36	8.73	0.885 \pm 10.91

There are also several reaction/cover combinations that are newly addressed in these neutron fields, such as:

1. Fe58g-Cdtk-B4C (configuration not used in ACRR-FF or SPR-III)
2. Co59g-Cdtk-B4C (configuration not used in ACRR-FF or SPR-III)
3. Co59g-Cd (which was discrepant in ACRR-FF)
4. Cu63g-bare (Cd cover was good in SPR-III)
5. Ag109g-bare (which was discrepant in ACRR-FF)
6. W186g-Cd

F. $^9\text{Be}(d,n)$ Reaction Neutron Fields

The IRDFF-II library contains cross sections up to 60 MeV for neutron dosimetry at accelerator-based neutron sources which require measurements for validation in high-energy neutron fields. Above 20 MeV there are many additional reaction channels that must be included to calculate the production of an activation product from elements that contain multiple isotopes. For example, the production of ^{54}Mn from Fe must include $^{56}\text{Fe}(n,X)^{54}\text{Mn}$ that contains reactions from other Fe isotopes that create more ^{54}Mn than the traditional $^{54}\text{Fe}(n,p)^{54}\text{Mn}$ reaction cross section used below 20 MeV. The impact of this ef-

TABLE 43. Least-squares-based consistency of data for the ACRR CdPoly neutron reference benchmark field.

Reaction Notation	Measured Activity [Bq/at. or Fiss/at.]	Act. Adj. Unc. [%]	Spectrum Unc. [%]	C/E \pm Unc. [%]	
Sc45g-bare	9.8273E-17	1.30	-0.63	14.76	0.940 \pm 16.44
Au197g-bare	3.7488E-13	1.70	-0.05	28.43	1.060 \pm 28.62
Na23g-bare	3.0951E-16	1.80	1.31	11.12	1.039 \pm 11.78
Ag109g-bare	4.8737E-17	0.60	0.13	17.07	1.088 \pm 18.33
W186g-bare	1.0211E-13	2.10	-0.02	19.55	1.014 \pm 20.09
Co59g-bare	1.3482E-17	1.40	1.4	11.22	1.274 \pm 11.51
Fe58g-bare	2.0046E-17	1.90	-0.91	7.39	0.965 \pm 8.88
Mn55g-bare	7.9671E-14	1.60	-1.38	7.58	0.968 \pm 8.32
Mo98g-bare	1.4967E-15	1.00	-0.01	7.94	1.017 \pm 12.50
Cu63g-bare	5.3525E-15	2.80	0.56	7.22	1.033 \pm 8.45
Nb93g-Cd	6.0332E-22	17.50	11.49	3.17	1.169 \pm 17.96
rmleu-Cdtk/B4C	1.5452E-09	2.20	0.8	1.74	1.047 \pm 3.46
Np237f-Cdtk/B4C	7.9845E-10	3.60	10.64	2.28	1.148 \pm 4.74
In115n-bare	5.1109E-15	5.60	-7.17	2.56	0.954 \pm 6.51
Ti47p-bare	2.3439E-17	3.10	-2.8	2.58	0.976 \pm 4.70
S32p-bare	1.7162E-17	3.60	0.59	2.93	1.004 \pm 5.12
Ni58p-bare	6.1122E-18	3.00	0.05	2.76	1.000 \pm 4.60
Zn64p-bare	2.9304E-16	2.80	-1.02	2.95	0.986 \pm 4.58
Fe54p-bare	1.0366E-18	1.70	-1.01	2.88	0.985 \pm 4.00
Al27p-bare	2.3141E-15	2.20	-4.11	3.17	0.951 \pm 4.40
Co59p-bare	1.2296E-19	3.10	-0.19	3.1	0.993 \pm 5.17
Ti46p-bare	4.9967E-19	1.50	3.28	3.21	1.043 \pm 4.38
Cu63a-Cd	1.1497E-21	4.40	-10.22	3.19	0.894 \pm 6.08
Fe56p-bare	3.8669E-17	1.60	-1.57	3.46	0.995 \pm 4.71
Ti48p-bare	6.0515E-19	1.00	0.45	3.47	1.040 \pm 5.02
Mg24p-bare	8.6405E-18	1.90	1.55	3.88	1.053 \pm 4.78
Al27a-bare	4.0228E-18	1.80	2.65	3.77	1.062 \pm 4.65
Nb932-bare	1.7793E-19	1.90	-1.88	4.19	0.986 \pm 5.04
Mn552-bare	2.8166E-21	4.50	-2.34	5.46	0.990 \pm 7.43
Co592-bare	1.1389E-20	18.30	3.94	5.44	1.054 \pm 19.24
Zr902-bare	1.2630E-19	2.20	0.97	7.59	1.050 \pm 8.15

fect on high energy neutron dosimetry was discussed in Ref. [211]. The IAEA Nuclear Data Section addressed this issue in the newly released IRDF-II library.

New evaluated cross section files for natural Al, Ti, Fe, Ni, and Zr activation to various activation products were especially developed for IRDF-II. While this file could be processed with the PREPRO codes, the format of the new file was not compatible with processing by the current version of NJOY. While interim steps have been taken to support processing this file with the NJOY code, we expect that NJOY compatibility with a legitimate ENDF-6 format in IRDF-II will soon be addressed.

The new natural element cross sections and covariances were processed into multi-group cross section libraries used by the STAYSL-PNNL computer code [212]. Integral testing was then performed using activation data measured in several $^9\text{Be}(d,n)$ neutron fields that have been

characterized by neutron time-of-flight (TOF) measurements at deuteron energies of 16 and 40 MeV [101, 213, 214]. Measurements at $E_d = 30$ MeV and $E_d = 40$ MeV were presented in INDC(NDS)-0731 [27], prior to the creation of the new Fe activation file. The plot of the spectra is shown in Fig. 72.

Eleven of the new natural element activation cross sections were tested in this report using previously published TOF neutron spectra to calculate the activation rates for direct comparison with the measured activation rates. These integral tests are independent of prior work since the new cross sections and covariances were not used in any spectral adjustments of the TOF neutron spectra using STAYSL-PNNL. The calculations were performed using 725-group cross section and covariance libraries. The measured activation rates were updated using the recommended gamma branching ratios listed in Table 8. Inte-

TABLE 44. Least-squares-based consistency of data for the ACRF FREC-II neutron reference benchmark field.

Reaction Notation	Measured Activity		Act. Adj. Spectrum		C/E \pm Unc. [%]
	[Bq/at. or Fiss/at.]	Unc. [%]	Unc. [%]	Unc. [%]	
Sc45g-bare	4.6441E-16	2.20	2.67	3.79	1.148 \pm 4.66
Mn55g-bare	1.9858E-13	1.70	0.98	28.06	0.894 \pm 28.37
Fe58g-bare	4.7655E-17	2.00	-1.83	21.94	0.854 \pm 22.29
Cu63g-bare	1.3579E-14	2.90	-0.64	28.15	0.886 \pm 28.26
Co59g-bare	3.1311E-17	2.20	0.72	11.26	1.119 \pm 11.54
Sc45g-Cd	2.4468E-17	2.20	-4.65	3.07	0.975 \pm 5.32
Au197g-bare	1.6471E-13	3.10	0.29	29.56	0.872 \pm 29.87
Au197g-Cd	1.0643E-13	1.80	0.18	6.93	1.002 \pm 8.25
Na23g-Cd	7.8216E-17	1.80	-1.2	2.88	0.980 \pm 4.13
Co59g-Cd	4.1486E-18	1.30	-3.48	3.46	1.041 \pm 5.08
Fe58g-Cd	5.4352E-18	1.90	0.6	29.64	0.869 \pm 30.07
Mo98g-bare	5.1417E-16	1.00	-1.22	7.22	0.955 \pm 8.76
Mo98g-Cd	4.6865E-16	1.00	-0.62	2.52	0.960 \pm 6.49
Cu63g-Cd	1.4968E-15	2.80	7.43	3.9	1.196 \pm 5.01
rmle-Cdtk/B4C	5.7944E-10	2.10	-0.64	29.81	0.858 \pm 29.93
rmlpu-Cdtk/B4C	5.8279E-10	2.10	-4.68	5.4	0.948 \pm 9.83
Np237f- Cdtk/B4C	3.1382E-10	2.10	2.3	7.66	0.920 \pm 12.31
In115n-bare	1.6796E-15	5.60	-2.46	7.31	0.883 \pm 12.02
rmldu-Cdtk/B4C	5.8963E-11	2.20	1.62	10.81	1.063 \pm 11.54
Ti47p-bare	8.0144E-18	3.20	2.4	4.22	1.098 \pm 5.02
S32p-bare	6.3767E-18	3.60	0.87	2.76	1.000 \pm 4.47
Ni58p-bare	2.0926E-18	2.80	0.67	2.25	0.974 \pm 3.71
Zn64p-bare	9.8155E-17	4.80	-3.16	2.65	0.936 \pm 4.02
Fe54p-bare	3.5797E-19	2.00	-9.91	1.74	0.864 \pm 3.39
Co59p-bare	4.3896E-20	3.40	3.95	1.83	1.026 \pm 4.09
Ti46p-bare	1.7181E-19	1.70	5.27	2.92	1.044 \pm 5.11
Fe56p-bare	1.3315E-17	2.50	0.56	31.89	0.899 \pm 32.97
Ti48p-bare	2.0869E-19	1.00	-0.89	14.65	0.985 \pm 16.44
Mg24p-bare	2.7379E-18	2.40	2.02	3.19	1.063 \pm 4.43
Al27a-bare	1.3531E-18	1.80	-1.74	2.58	0.966 \pm 4.76
Nb932-bare	5.8125E-20	1.80	-0.44	3.49	1.094 \pm 5.00
Mn552-bare	1.0289E-21	7.90	2.14	2.95	1.015 \pm 6.01
Zr902-bare	4.9566E-20	3.00	-1.54	7.62	0.929 \pm 8.44

gral testing results are presented in Table 45 - Table 46 below. The neutron spectrum for ${}^9\text{Be}(d,n)$ at $E_d = 16$ MeV does not exceed 20 MeV so it is a good check on the lower energy range. The spectra at $E_d = 40$ MeV provides a good test of the cross sections at progressively higher energies up to about 50 MeV. Tables 45 and 46 show generally good agreement between the measured activation rates and the calculated activation rates using the IRDF-II natural element cross sections within the uncertainties of the neutron time-of-flight spectra as well as the activation rate measurements.

The new cross section and cross section libraries were also tested by performing neutron spectral adjustments with the STAYSL-PNNL computer code using the 725-group cross section and covariance libraries. At 16 MeV,

we used 24 reactions for the small spectral adjustment and 25 reactions at 40 MeV. Both the TOF and adjusted spectral activation rate calculations are shown in Fig. 72. In both cases we achieved acceptable χ^2 values and overall standard deviations less than 10 % attesting to the high quality of the new IRDF-II file for neutron spectral adjustment up to 50 MeV using the new natural element cross sections.

IX. OTHER VALIDATION EVIDENCE

A large volume of additional data exists that supports the validation of the IRDF-II library, as has been reported at other venues [27]. This additional data includes

TABLE 45. Be(d,n) Integral Testing of Natural Element Cross Sections in IRDFE-II (725 group calculations), $E_d = 16$ MeV

Elem. Prod.	Measured at/at-s	\pm [%]	Calculated at/at-s	Diff. [%]	E-Range MeV	
					5%	95%
Ti	^{46}Sc 9.74E-18	3	1.04E-17	6.8	5.0	13.8
	^{47}Sc 5.77E-18	10	5.86E-18	1.6	3.1	17.7
	^{48}Sc 8.33E-18	10	7.90E-18	-5.2	7.1	15.2
Fe	^{54}Mn 1.92E-17	3	1.94E-17	0.8	3.5	12.4
	^{56}Mn 2.51E-17	3	2.43E-17	-3.2	6.4	14.4
	^{51}Cr 1.17E-18	10	1.12E-18	-3.9	6.3	14.9
Ni	^{58}Co 2.77E-16	3	2.81E-16	1.5	3.3	12.2
	^{57}Ni 7.90E-19	3	8.28E-19	4.8	13.2	19.6
	^{60}Co 1.40E-17	7	1.08E-17	-23	6.0	14.0
Zr	^{89}Zr 1.36E-17	3	1.41E-17	4.1	12.8	19.3

TABLE 46. Be(d,n) Integral Testing of Natural Element Cross Sections in IRDFE-II (725 group calculations), $E_d = 40$ MeV

Elem. Prod.	Measured at/at-s	\pm [%]	Calculated at/at-s	Diff. [%]	E-Range MeV	
					5%	95%
Al	^{24}Na 3.14e-15	3	3.06E-15	-2.5	8.9	22.0
Ti	^{46}Sc 1.42E-15	3	1.56E-15	9.7	8.3	37.0
	^{47}Sc 3.02E-15	3	2.94E-15	-2.8	12.9	33.0
	^{48}Sc 1.35E-15	3	1.49E-15	10.7	9.7	28.5
Fe	^{54}Mn 1.68E-15	3	1.38E-15	-17.9	5.3	38.0
	^{56}Mn 2.80E-15	3	2.98E-15	6.4	8.5	26.0
	^{51}Cr 2.30E-16	3	1.83E-16	-20.4	9.1	39.0
Ni	^{58}Co 1.12E-14	3	1.09E-14	-2.2	3.8	23.0
	^{57}Ni 1.34E-15	10	1.46E-15	8.7	14.8	29.5
	^{60}Co 1.04E-15	10	1.07E-15	2.5	8.1	26.0
Zr	^{89}Zr 1.70E-14	3	1.78E-14	4.7	14.2	28.5

measurements such as:

- JAEA/FNS benchmark testing with Li2O assembly and a DT source;
- ENEA Frascati copper block assembly with 14-MeV neutrons;
- CEA reactor data measured at MINERVE, EOLE, MASURCA, and CALIBAN reactor facilities;
- ASPIS-IRON88 benchmark described in the SINBAD package NEA-1517/75;
- activation experiments at iThemba Laboratory for Accelerator Based Sciences using 90 and 140 MeV quasi-monoenergetic neutron beams;
- Kyoto University Research Reactor 80 and 140 MeV p-Li quasi-monoenergetic neutron beams;

This evidence is not explicitly captured in this paper because the results, typically, exhibited a strong influence from other uncertainty contributors or they did not significantly expand the strength of the validation evidence for the IRDFE-II library.

X. VALIDATION STATUS

Secs. VII, VIII, and IX detail the validation evidence that has been assembled to support the IRDFE-II recommended cross sections in integral neutron fields. It is not sufficient for users of the library to just ask if a reaction cross section has been validated, users must also consider if the reaction cross section has been validated within the relevant energy response region addressed by their application. Table 47 summarizes the validation data in a format where users can more readily assess the validation status for a given reaction for a given class of neutron field. Results are presented for selected integral benchmarks that cover the complete range of average spectrum energies. For a more detailed information the reader is referred to Secs. VII, VIII, and IX.

Whereas all of the IRDFE-II metrology reactions have nuclear data evaluations that are based on some experimental data and trends informed by nuclear model calculations, some of the metrology reactions within the library are seen in Table 47 to lack any additional validation evidence in integral benchmark neutron fields. Many of the validation deficiencies are for high threshold energy reactions where measurements are not easily acquired in fission neutron fields.

TABLE 47: Validation evidence for IRDF-II metrology reactions from integral benchmark neutron fields.

Entry Label	Thermal x.s.	Resonance Integral	Reactor ACRR-CC	Reactor SPR-III	²³⁵ U(n,f) PFNS	²⁵² Cf(sf) neutrons	Be(d,n) $E_d=40$ MeV
	E_{av} 0.0506 eV	132.4 keV	0.575 MeV	1.251 MeV	2.000 MeV	2.121 MeV	15.58 MeV
	Unc. Dif.	Unc. Dif.	C/E	C/E	C/E	C/E	Unc. Dif.
LiH3							
LiHe4					1.074 ± 6.02%		
Li6H3							
Li6t							
Li6He4							
Li7H3							
Li7He4							
BH3 13						0.931 ± 11.58%	
BHe4							
B10H3							
B10a					0.848 ± 9.72%		
B10He4							
B11H3							
B11He4							
F192					1.023 ± 6.12%	1.007 ± 5.66%	
Na232					1.054 ± 4.89%	0.990 ± 7.63%	
Na23g	1.6% 4.4%	4.5% 0.4%	1.045 ± 11.08%	1.178 ± 11.94%			
MgNa24			1.041 ± 4.98%	1.122 ± 15.71%	0.999 ± 3.08%	1.050 ± 2.92%	
Mg24p			1.041 ± 4.98%	1.122 ± 15.71%	0.999 ± 3.08%	1.050 ± 2.92%	
Al27p				1.138 ± 12.45%	0.968 ± 2.96%	0.950 ± 2.51%	
Al27a			1.018 ± 4.89%	1.075 ± 14.90%	1.009 ± 3.00%	1.003 ± 2.07%	
Al27Na24			1.018 ± 4.89%	1.075 ± 14.90%	1.009 ± 3.00%		3% -2.5%
Al272g					0.902 ± 13.72%		
SiAl28							
Si28p					0.993 ± 4.16%	1.031 ± 2.67%	
Si29p							
P31p					0.997 ± 3.62%		
SP32			0.988 ± 5.12%	0.990 ± 11.50%	0.995 ± 3.04%	1.023 ± 3.65%	
S32p			0.988 ± 5.12%	0.990 ± 11.50%	0.995 ± 3.04%	1.023 ± 3.65%	
Sc45g	6.9% 3.6%	7.6% 0.2%	1.082 ± 16.09%	1.051 ± 8.57%			
TiTi45							
TiSc46				1.054 ± 12.96%	0.993 ± 3.45%	0.974 ± 2.63%	3% 9.7%
TiSc47					1.024 ± 3.19%	1.015 ± 2.16%	3% -2.8%
TiSc48					1.001 ± 4.15%	1.005 ± 3.53%	3% 10.7%
Ti462							
Ti46p			1.021 ± 4.74%	1.054 ± 12.96%	0.993 ± 3.45%		
Ti47p			0.971 ± 4.71%	1.102 ± 10.22%	1.024 ± 3.19%	1.015 ± 2.16%	
Ti48p			1.048 ± 5.20%	1.105 ± 14.22%	1.001 ± 4.15%		
V51a					1.008 ± 4.12%	0.989 ± 3.23%	
V51Sc48							
CrCr51							
Mn552				1.009 ± 15.90%	1.057 ± 16.61%	0.958 ± 15.63%	
Mn55g	1.1% 0.5%	3.8% -4.7%	1.040 ± 8.45%	1.082 ± 8.69%			
FeCr51					0.969 ± 5.60%		3% -20.4%
FeMn54			0.959 ± 4.23%				3% -17.9%
FeMn56			0.992 ± 4.81%				3% 6.4%
FeMn53							
Fe542							
Fe54p			0.959 ± 4.23%	1.049 ± 11.40%	1.034 ± 2.78%	1.015 ± 1.79%	
Fe54a					0.969 ± 5.60%		
Fe56p			0.992 ± 4.81%	1.076 ± 14.43%		0.998 ± 2.91%	
Fe58g	5.1% 1.5%	4.8% 1.3%	1.020 ± 8.80%	1.105 ± 12.04%			
Co592				1.087 ± 16.60%	1.069 ± 6.05%	1.006 ± 4.27%	
Co593							
Co59g	0.7% -2.0%	0.8% -0.8%	1.305 ± 12.53%				
Co59p			1.032 ± 7.16%	0.983 ± 13.04%	1.017 ± 3.72%	1.015 ± 3.16%	
Co59a					1.001 ± 4.50%	1.000 ± 3.39%	
Co59Mn56					1.001 ± 4.50%	1.000 ± 3.39%	
NiNi57				1.014 ± 16.69%	1.015 ± 5.00%	0.997 ± 7.07%	
NiCo58					1.000 ± 2.62%	1.000 ± 1.66%	3% -2.2%
NiCo60			1.025 ± 6.62%		0.994 ± 5.43%		10% 2.5%
Ni582				1.014 ± 16.69%	1.015 ± 5.00%	0.997 ± 7.07%	10% 8.7%

TABLE 47: (continued) Validation evidence for IRDF-II metrology reactions from integral benchmark neutron fields.

Entry Label	Thermal x.s.	Resonance Integral	Reactor ACRR-CC	Reactor SPR-III	$^{235}\text{U}(n,f)$ PFNS	$^{252}\text{Cf}(sf)$ neutrons	$\text{Be}(d,n)$ $E_d=40$ MeV
	E_{av} 0.0506 eV	132.4 keV	0.575 MeV	1.251 MeV	2.000 MeV	2.121 MeV	15.58 MeV
	Unc. Dif.	Unc. Dif.	C/E	C/E	C/E	C/E	Unc. Dif.
Ni58p			1.000 ± 4.53%	1.000 ± 10.96%	1.000 ± 2.62%	1.000 ± 1.66%	
Ni60p			1.025 ± 6.62%		0.994 ± 5.43%	1.027 ± 5.03%	
CuCo60				0.966 ± 13.97%	1.022 ± 4.11%	1.010 ± 8.64%	
CuCu62					0.828 ± 7.95%		
CuCu64				1.041 ± 11.99%		0.993 ± 3.70%	
Cu632					0.828 ± 7.95%	1.032 ± 5.74%	
Cu63g	4.1% -3.3%	4.2% -7.3%		1.041 ± 11.99%	1.019 ± 24.61%	1.010 ± 8.64%	
Cu63a				0.966 ± 13.97%	1.022 ± 4.11%	1.006 ± 2.93%	
Cu652						0.993 ± 3.70%	
ZnCu64			0.959 ± 4.77%		1.008 ± 7.59%	1.009 ± 3.31%	
ZnCu67					1.014 ± 6.64%		
Zn64p			0.959 ± 4.77%	1.077 ± 11.42%	1.008 ± 7.59%		
Zn67p					1.014 ± 6.64%		
Zn68Cu67							
As752					1.040 ± 6.88%		
Y892					1.039 ± 5.98%	0.984 ± 5.64%	
ZrZr89							3% 4.7%
Zr902			0.909 ± 9.23%	1.142 ± 15.55%		0.981 ± 5.77%	
Nb932m			0.930 ± 5.10%		1.065 ± 4.75%	1.000 ± 4.95%	
Nb93nm				1.113 ± 17.14%	0.983 ± 5.38%	1.010 ± 3.73%	
Nb93g	4.0% 1.1%	3.4% 3.2%					
Nb93gm	4.0% 0.9%	3.4% 3.1%					
MoNb92m							
Mo92pm					1.011 ± 4.07%		
Rh103nm					1.017 ± 4.22%		
Ag109g	5.1% 7.7%	7.0% 4.8%	1.160 ± 17.98%				
InIn114m	4.6% -6.9%	8.7% 6.3%					
In113gm	4.6% -6.9%	8.7% 6.3%					
In113nm					0.933 ± 5.57%	0.978 ± 2.15%	
In1152m							
In115nm				1.068 ± 10.20%	1.006 ± 2.52%	0.964 ± 1.57%	
In115gm	1.0% 1.5%	3.1% -4.0%			1.052 ± 4.83%	0.979 ± 2.45%	
I1272					1.016 ± 5.36%	1.018 ± 3.87%	
La139g	3.9% -2.7%	5.5% 3.5%					
Pr1412							
Tm1692					1.036 ± 5.55%	0.986 ± 4.74%	
Tm1593						0.795 ± 16.24%	
Ta181g	3.0% 1.6%	3.8% -2.7%				0.935 ± 5.65%	
W186g	5.0% 9.9%	4.5% 1.8%	1.012 ± 16.93%	0.818 ± 10.41%			
Au1972					1.001 ± 3.96%	1.005 ± 2.85%	
Au197g	0.5% 0.0%	2.1% -0.5%	1.042 ± 28.04%	0.929 ± 7.28%	0.889 ± 4.58%	0.991 ± 1.79%	
Hg199nm					1.027 ± 5.84%	0.985 ± 2.22%	
Pb204nm					0.909 ± 6.87%	0.971 ± 2.74%	
Bi2092							
Bi2093							
Bi2094							
Bi2095							
Bi2096							
Th232f					0.918 ± 3.83%	0.935 ± 5.08%	
Th232g	5.4% 0.0%	10% -0.3%				1.032 ± 5.53%	
U235f				1.089 ± 6.27%	1.034 ± 2.35%	1.013 ± 1.21%	
U2382					0.936 ± 5.63%	1.045 ± 11.23%	
U238f				1.036 ± 10.36%	1.007 ± 2.40%	0.978 ± 1.70%	
U238g	1.0% 0.5%	1.1% -0.4%			0.825 ± 9.66%		
Np237f			1.003 ± 4.67%	1.103 ± 7.57%	1.022 ± 2.79%	1.000 ± 1.69%	
Pu239f				1.165 ± 6.29%	1.007 ± 2.65%	0.992 ± 1.38%	
Am241f							

XI. LIBRARY DISTRIBUTIONS

The development of the IRDF-II library was the result of an international collaboration, organized through

a series of Research Projects at the International Atomic Energy Agency Nuclear Data Section (NDS). The library can be obtained from www-nds.iaea.org/IRDF/. It has

been incorporated into the IAEA nuclear data viewers, www-nds.iaea.org/exfor/endl.htm. Using that online system, comparisons can be readily made between different nuclear data libraries, measured data available in EXFOR [72], and the IRDF-II recommended cross section with uncertainties.

XII. CONCLUSIONS

This paper reports on the assembly of a new neutron metrology library - the IRDF-II - that represents the recommendations of an international collaboration and supports neutron metrology application in fission and fusion neutron fields. The elements of the IRDF-II library are described and the basis for the recommended nuclear data values is provided. The IRDF-II library, which now extends up to incident neutron energies of 60 MeV, contains a comprehensive, complete and consistent set of nuclear data that can be used for the characterization of neutron radiation fields and to support the neutron metrology used with experiments conducted in these neutron fields. In addition to providing the recommended nuclear data, this document summarizes the set of validation evidence gathered from integral neutron benchmark fields that confirms the consistency of the recommended nuclear data.

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- [1] International Reactor Dosimetry File 2002 (IRDF-2002), TECHNICAL REPORT SERIES **452**, IAEA, (2006); numerical data: <https://www-nds.iaea.org/irdf2002/>.
 - [2] R. Capote, K.I. Zolotarev, V.G. Pronyaev and A. Trkov, "Updating and Extending the IRDF-2002 Dosimetry Library", *J. ASTM INT.*, **9**, 1 (2012).
 - [3] D. Smith, *Probability, Statistics, and Data Uncertainties in Nuclear Science and Technology*, American Nuclear Society, LaGrange Park, Illinois (1991).
 - [4] R. Capote, D.L. Smith and A. Trkov, "Nuclear data evaluation methodology including estimates of covariances", *EPJ WEB CONF.* **8**, 04001 (2010).
 - [5] D.L. Smith and N. Otuka, *Nucl. Data Sheets* **113**, 3006-3053 (2012).
 - [6] E. Bauge, R. Capote, U. Fisher *et al.*, "Covariance Data in the Fast Neutron Region". Final report of the *NEA/WPEC-24*, Coord. M. Herman, Report **NEA/NSC/WPEC/DOC(2010)427** (Nuclear Energy Agency, OECD, Paris, 2011).
 - [7] W. P. Poenitz, "Evaluation Methods for Neutron Cross Section Standards", in *Proc. Conf. Nucl. Data Evaluation Methods and Procedures*, BNL report **BNL-NCS-51363**, Vol.1, p. 249 (1981). Codes and documentation available online at <https://www-nds.iaea.org/standards/codes.html>.
 - [8] D.L. Smith, "A Least-Squares Computational "Tool Kit"", Report **ANL/NDM-128**, Argonne National Laboratory, IL, USA (1993).
 - [9] D.M. Hetrick and C.Y. Fu, "GLUCS: A generalized least-squares program for updating cross section evaluations with correlated data sets," Oak Ridge National Laboratory, TN, USA (1980).
 - [10] T. Kawano and K. Shibata, "KALMAN," Report **JAERI-Data/Code 97-037**, JAERI, Tokai, Japan (1997).
 - [11] D.W. Muir, "Global Assessment of Nuclear Data Requirements (GANDR project)," IAEA report (6 volumes), Vienna, Austria (2007). Available online at <https://www-nds.iaea.org/gandr/docs.html>.
 - [12] V.N. Vinogradov, E.V. Gai and N.S. Rabotnov, "Analytical Approximation of Data in Nuclear and Neutron Physics," *Energoatomizdat*, Moscow (1987); in Russian.
 - [13] S.A. Badikov, E.V. Gai, M.A. Guseynov and N.S. Rabotnov, "Nuclear data processing, evaluation, transforma-

- tion and storage with Padé-approximants,” Proc. Int. Conf. Nuclear Data for Science and Technology, Jülich, Germany, 1991, ed: S.M. Qaim, 182–187 (1992) Springer-Verlag, Berlin.
- [14] A. Hermanne, A.V. Ignatyuk, R. Capote *et al.* “Reference cross sections for charged-particle monitor reactions,” *NUCL. DATA SHEETS* **148**, 338–382 (2018).
- [15] H.E. Padé, “Sur la Représentation Approchée d’une Fonction par des Fractions Rationnelles,” Supplement to ANN. SCI. L’ÉCOLE NORM. SUP., Series 3, Vol. **9**, 3–93 (1892).
- [16] P.R. Graves-Morris (Ed.), “Padé Approximants and their Applications,” Academic Press, New York (1973).
- [17] G.A. Baker Jr., “Essentials of Padé Approximants,” Academic Press, New York (1975).
- [18] E. Bauge and P. Dossantos-Uzarralde, “Evaluation of the covariance matrix of Pu-239 neutronic cross sections in the continuum using a Backward-Forward Monte Carlo method,” *J. KOR. PHYS. SOC.* **59**, 1218 (2011).
- [19] R. Capote, D.L. Smith, A. Trkov, M. Meghzifene, “A New Formulation of the Unified Monte Carlo Approach (UMC-B) and Cross-Section Evaluation for the Dosimetry Reaction $^{55}\text{Mn}(n,\gamma)^{56}\text{Mn}$,” *J. ASTM INT.* **9**, 104115 (2012).
- [20] D.L. Smith, “A Unified Monte Carlo Approach to Fast Neutron Cross Section Data Evaluation,” in PROC. 8TH INT. TOPICAL MEETING ON NUCL. APPL. AND UTILIZATION OF ACCELERATORS, Pocatello, ID, USA, 29 July 29–2 August, p. 736 (2007).
- [21] R. Capote and D.L. Smith, “An Investigation of the Performance of the Unified Monte Carlo Method of Neutron Cross Section Data Evaluation,” *NUCL. DATA SHEETS* **109**, 2768 (2008).
- [22] A.J. Koning, “Bayesian Monte Carlo method for nuclear data evaluation,” *EUR. PHYS. J.* **A51**, 184 (2015).
- [23] S. Goriely and R. Capote, “Uncertainties of mass extrapolations in Hartree-Fock-Bogoliubov mass models,” *PHYS. REV.* **C89**, 054318 (2014).
- [24] A. J. Koning, D. Rochman, J.-Ch. Sublet, N. Dzysiuk, M. Fleming, and S. van der Marck, “TENDLE722: Complete Nuclear Data Library for Innovative Nuclear Science and Technology,” *NUCL. DATA SHEETS* **155**, 1–55 (2019).
- [25] A.J. Koning, D. Rochman, J. Kopecky, J. Ch. Sublet, E. Bauge, S. Hilaire, P. Romain, B. Morillon, H. Duarte, S. van der Marck, S. Pomp, H. Sjostrand, R. Forrest, H. Henriksson, O. Cabellos, S. Goriely, J. Leppanen, H. Leeb, A. Plompen and R. Mills, “TENDL-2015: TALYS-based evaluated nuclear data library.” Available online at https://tendl.web.psi.ch/tendl_2015/tendl2015.html.
- [26] J. Knaster, F. Arbeiter, P. Cara, S. Chel, A. Facco, R. Heidinger, A. Ibarra, A. Kasugai, H. Kondo, G. Micciche, K. Ochiai, S. Ohira, Y. Okumura, K. Sakamoto, E. Wakai, “IFMIF, the European/Japanese efforts under the Broader Approach agreement towards a Li(d,xn) neutron source: Current status and future options”, *NUCL. MAT. AND ENERGY* **9**, 46–54 (2016).
- [27] L.R. Greenwood, M. Kostal, S.P. Simakov, A. Trkov, “Summary Report of the Third Research Coordination Meeting on Testing and Improving the International Reactor Dosimetry and Fusion File (IRDFF)”, March 20–24, report **INDC-NDS-0731**, IAEA, Vienna, Austria (2017).
- [28] J.-C. Sublet, L. Packer, J. Kopecky, R.A. Forrest, “Recent Progress in Neutron-, Proton- and Deuteron Induced Reaction Nuclear Data for EAF-2010 and the European Activation System,” in Proc. Int. Conf. on Nucl. Data for Science and Technology (ND2010), Jeju Island, Korea
- [29] K. Shibata *et al.*, “JENDL-4.0: A New Library for Nuclear Science and Engineering,” *J. NUCL. SCI. TECHN.* **48**, 1–30 (2011).
- [30] D. Brown, *et al.*, “ENDF/B-VIII.0: The 8th Major Release of the Nuclear Reaction Data Library with CIELO-project Cross Sections, New Standards and Thermal Scattering Data” *NUCL. DATA SHEETS* **148**, 1–142 (2018).
- [31] JEFF Scientific Working Group, “JOINT EVALUATED FISSION AND FUSION FILE (JEFF) RELEASE 3.3,” OECD, NEA (2018).
- [32] A.D. Carlson, V.G. Pronyaev, R. Capote *et al.*, “Evaluation of the Neutron Data Standards,” *NUCL. DATA SHEETS* **148**, 143 (2018)
- [33] M.B. Chadwick, M.W. Herman, P. Obložinský *et al.*, “ENDF/B-VII.1 nuclear data for science and technology: cross sections, covariances, fission product yields and decay data,” *NUCL. DATA SHEETS* **112**, 2887–2996 (2011).
- [34] A. Trkov, R. Capote, E. Sh. Soukhovitskii, L. C. Leal, M. Sin, I. Kodeli, D. W. Muir, “Covariances of Evaluated Nuclear Cross Section Data for ^{232}Th , $^{180,182,183,184,186}\text{W}$ and ^{55}Mn ,” *NUCL. DATA SHEETS* **112**, 3098–3119 (2011).
- [35] A. Koning *et al.*, “The JEFF-3.1 Nuclear Data Library”, *JEFF REPORT* **21**, OECD 2006, NEA No. 6190.
- [36] A. Trkov, R. Capote, I. Kodeli, L. Leal, “Evaluation of tungsten nuclear reaction data with covariances,” *NUCL. DATA SHEETS* **109**, 2905–2909 (2008).
- [37] R. Capote, L. Leal, Liu Ping *et al.*, “Evaluated nuclear data for nuclides within the thorium-uranium fuel cycle,” Technical report **STI/PUB/1435** (Vienna, International Atomic Energy Agency, 2010), ISBN 9789201010100.
- [38] I. Sirakov, R. Capote, F. Gunsing, P. Schillebeeckx, and A. Trkov, “An ENDF-6 compatible evaluation for neutron induced reactions of ^{232}Th in the unresolved resonance region,” *ANN. NUCL. EN.* **35**, 1223–1231 (2008).
- [39] M. B. Chadwick, E. Dupont, E. Bauge *et al.*, “The CIELO Collaboration: Neutron Reactions on ^1H , ^{16}O , ^{56}Fe , $^{235,238}\text{U}$, and ^{239}Pu ,” *NUCL. DATA SHEETS* **118**, 1–25 (2014).
- [40] M. B. Chadwick, R. Capote, A. Trkov, *et al.*, “CIELO collaboration summary results: International evaluations of neutron reactions on uranium, plutonium, iron, oxygen and hydrogen,” *NUCL. DATA SHEETS* **148**, 189–213 (2018).
- [41] R. Capote, A. Trkov, *et al.*, “IAEA CIELO evaluation of neutron-induced reactions on ^{235}U and ^{238}U targets for incident energies up to 30 MeV,” *NUCL. DATA SHEETS* **148**, 254–292 (2018).
- [42] **ASTM E722-14**, “Standard Practice for Characterizing Neutron Fluence Spectra in Terms of an Equivalent Monoenergetic Neutron Fluence for Radiation-Hardness Testing of Electronics,” *ASTM INTERNATIONAL*, West Conshohocken, PA, 2014.
- [43] **ASTM E693-17**, “Standard Practice for Characterizing Neutron Exposures in Iron and Low Alloy Steels in Terms of Displacement Per Atom (DPA),” *ASTM INTERNATIONAL*, West Conshohocken, PA, 2017.
- [44] W. N. McElroy, S. Berg, T. Crockett, and R. Hawkins, “A Computer-Automated Iterative Method for Neutron Flux Spectral Determination by Foil Activation,” **AFWL-TR-67-41**, Vol. 1, Air Force Weapons Laboratory, Kirtland AFB, New Mexico, July 1967.

- [45] F.W. Stallmann, "LSL-M2: A Computer Program for Least-Squares Logarithmic Adjustment of Neutron Spectra," **NUREG/CR-4349**, ORNL/TM-9933, March 1985.
- [46] P.J. Griffin, "A rigorous treatment of self-shielding and covers in neutron spectra determinations," **IEEE TNS** **42**, 1878–1885 (1999).
- [47] M.J. Norgett, M.T. Robinson, I.M. Torrens, "A Proposed Method of Calculating Displacement Dose Rates," **Nucl. Eng. Des.**, Vol. **33**, pp. 50–54, 1975.
- [48] **ASTM E1855-15**, "Standard Test Method for Use of 2N2222A Silicon Bipolar Transistors as Neutron Spectrum Sensors and Displacement Damage Monitors," ASTM INTERNATIONAL, West Conshohocken, PA, 2015.
- [49] **Decay Data Evaluation Project**. Data available online at www.nucleide.org/DDEP.htm.
- [50] **Evaluated Nuclear Structure Data File (ENSDF)**. Available online at www.nndc.bnl.gov/ensdf/. Developed and maintained by the International Network Of Nuclear Structure and Decay Data Evaluators (NSDD) (see www.nds.iaea.org/nsdd/).
- [51] **LiveChart of Nuclides**, IAEA decay data retrieval code available online at www.nds.iaea.org/medical/monitor_reactions.html.
- [52] **NuDat**, Brookhaven National Laboratory, USA. Decay data retrieval code available online at www.nndc.bnl.gov/nudat2/.
- [53] **Atomic weights of the elements 2017**. Available online at www.ciaaw.org.
- [54] M. Wang, G. Audi, F.G. Kondev, W.J. Huang, S. Naimi, X. Xu, "The AME2016 Atomic Mass Evaluation, (II) Tables, Graphs, and References", **CHIN. PHYS.** **C41**, Number 3 (2017); <https://www.nds.iaea.org/amdc/>.
- [55] J. Meija, T.B. Coplen, M. Berglund, W. A. Brand, P. De Bièvre, M. Gröning, N. E. Holden, J. Irrgeher, R. D. Loss, Thomas Walczyk, T. Prohaska, "Isotopic compositions of the elements 2013" (IUPAC Technical Report), **PURE APPL. CHEM.** **88**, 293–306 (2016). Published Online 2016-02-20 |; <https://doi.org/10.1515/pac-2015-0503>.
- [56] **ASTM E704-13**, "Standard Test Method for Measuring Reaction Rates by Radioactivation of Uranium-238," ASTM INTERNATIONAL, West Conshohocken, PA, 2013.
- [57] **ASTM E1005-16**, "Standard Test method for Application and Analysis of Radiometric Monitors for Reactor Vessel Surveillance," ASTM INTERNATIONAL, West Conshohocken, PA, 2016.
- [58] National Research Council 2005. "Monitoring Nuclear Weapons and Nuclear-Explosive Materials: An Assessment of Methods and Capabilities" (2005), Committee on International Security and Arms Control, National Academy of Sciences, The National Academies Press, Washington D.C., DOI 10.17226/11265. Available at: <http://nap.edu/11265>.
- [59] A.L. Nichols, D.L. Aldama, M. Verpelli, "Handbook of Nuclear Data for Safeguards," report **INDC(NDS)-0502**, International Atomic Energy Agency, Vienna, Austria (2007).
- [60] A. Trkov, M. Herman and D. A. Brown (Eds.), "ENDF-6 Formats Manual," CSEWG Document **ENDF-102**, Report **BNL-203218-2018-INRE**, Brookhaven National Laboratory, Upton, NY 11973-5000 (2018).
- [61] "Nuclear Data Processing System NJOY21 and NJOY2016," Los Alamos National Laboratory, <https://github.com/njoy>.
- [62] **PREPRO**
- [63] "MCNP-A General Monte Carlo Code for Neutron and Photon Transport, Version 3A", J. Briesmeister (Editor), report **LA-7396-M**, Los Alamos National Laboratory, Los Alamos, NM (1986).
- [64] **SERPENT**
- [65] C. Dunford, "ENDF Utility Codes," Brookhaven National Laboratory, Upton, NY 11973-5000.
- [66] **COVEIG**
- [67] A. Trkov, "Guidelines for Nuclear Data Verification and Validation," **INDC(SEC)-0107**, International Atomic Energy Agency, Vienna, Austria (2005). Available online at <https://www.nds.iaea.org/publications/indc/indc-sec-0107/>.
- [68] M.A. Kellett, O Bersillon, RW Mills; **JEFF report 20** (2009).
- [69] "The Kayzero database," http://www.kayzero.com/k0naa/k0naaorg/Nuclear_Data_SC/Nuclear_Data_SC.html.
- [70] A. Trkov, V. Radulović, "Nuclear reactions and physical models for neutron activation analysis," **J. RADIOAN. NUCL. CHEM.** **304**, 763-778, (2015)
- [71] S. F. Mughabghab, "Atlas of Neutron Resonances: Resonance Parameters and Thermal Cross Sections Z=1–100," 6th Edition, Elsevier, Amsterdam The Netherlands (2016).
- [72] N. Otuka, E. Dupont, V. Semkova, B. Pritychenko *et al.*, "Towards a More Complete and Accurate Experimental Nuclear Reaction Data Library (EXFOR): International Collaboration Between Nuclear Reaction Data Centres (NRDC)," **NUCL. DATA SHEETS** **120**, 272–276 (2014). Data available online (e.g., at www.nds.iaea.org/exfor/).
- [73] W. Mannhart, "Recent Experiments on ²⁵²Cf(sf) Spectrum-Averaged Neutron Cross Sections," Proc. 5th ASTM-EURATOM Symp. on Reactor Dosimetry, 1984, Geesthacht, Germany, Vol. 2, D. Reidel, Dordrecht, The Netherlands, 1985, p. 801–812.
- [74] O. Horibe *et al.*, Proc. of Conf. 50 Years with Nuclear Fission, Washington D.C., 25–28 April 1989, v.2, p.923.
- [75] W. Mannhart, Prog. Report **INDC(GER)-045**, pp.40–43, June 1999.
- [76] W. Mannhart, "Handbook on Nuclear Activation Cross Sections", IAEA Technical Report Ser. No. **273**, p.413 (1987).
- [77] W. Mannhart, "Validation of Differential Cross Sections with Integral Data", IAEA Techn. Meet. on "International Reactor Dosimetry File: IRDF-2002", Vienna, 27–29 Aug 2002, Report **INDC(NDS)-0435**, p.59 (2002).
- [78] ASTM E264, "Standard Test Method for Measuring Fast-Neutron Reaction Rates by Radioactivation of Nickel", ASTM INTERNATIONAL, West Conshohocken, PA, 2019.
- [79] K.I. Zolotarev, "Re-evaluation of microscopic and integral cross section data for important dosimetry reactions", Report **INDC(NDS)-0526**, IAEA, Vienna, Austria (2004).
- [80] **ASTM E265-15**, "Standard Test Method for Measuring Reaction Rates and Fast-Neutron Fluences by Radioactivation of Sulfur-32," ASTM International, West Conshohocken, PA, 2013.
- [81] K.I. Zolotarev and P.K. Zolotarev, "Evaluation of some (n,n'), (n,γ), (n,p), (n,2n) and (n,3n) reaction excitation functions for fission and fusion reactor dosimetry applications," Report **INDC(NDS)-0657**, IAEA, Vienna, Austria (2013).
- [82] M.C. Moxon, "Evaluation of the resonance region for

- 58Fe,” International Atomic Energy Agency, Vienna, **INDC(UK)-089**, 2004.
- [83] K.I. Zolotarev, “Evaluation of cross section data from threshold to 40-60 MeV for specific neutron reactions important for neutron dosimetry applications. Part I,” Report **INDC(NDS)-0546**, IAEA, Vienna, Austria (2009).
- [84] K.I. Zolotarev, “Evaluation of $^{23}\text{Na}(n,\gamma)^{24}\text{Na}$, $^{23}\text{Na}(n,2n)^{22}\text{Na}$ and $^{27}\text{Al}(n,2n)^{26}\text{Al}$ reaction cross sections for the IRDF library”, Report **INDC(NDS)-0705**, IAEA, Vienna, Austria (2017).
- [85] E. Simeckova and M. Majerle, private communication, 2019.
- [86] K.I. Zolotarev, “Evaluation and improvement of cross section accuracy for most important dosimetry reactions for $^{27}\text{Al}(n,p)$, $^{56}\text{Fe}(n,p)$ and $^{237}\text{Np}(n,f)$ including covariance data”, Report **INDC(CCP)-0438**, IAEA, Vienna, Austria (2004).
- [87] K.I. Zolotarev, “Evaluation of cross section data from threshold to 40-60 MeV for specific neutron reactions important for neutron dosimetry applications. Part II.”, Report **INDC(NDS)-0584**, IAEA, Vienna, Austria (2010).
- [88] P. Schillebeeckx *et al.*, “Evaluation of Neutron Resonance Cross Section Data at GELINA,” *NUCL. DATA SHEETS* **119**, 94–97 (2014), DOI: 10.1016/j.nds.2014.08.027
- [89] D.W. Kneff, B.M. Oliver, H. Farrar IV, L.R. Greenwood, “Helium production in pure elements, isotopes, and alloy steels by 14.8-MeV neutrons,” *NUCL. SC. & ENG.* **92**, 491 (1986).
- [90] Neutron Cross Sections for Reactor Dosimetry, International Atomic Energy Agency, Laboratory Activities, Vienna, Vol 1, 1978, p. 62 and M. Vlasov, IAEA Program on Benchmark Neutron Fields Applications for Reactor Dosimetry, Report **INDC(SEC)-54/L+Dos**, IAEA, Vienna, Austria (1976).
- [91] W. Mannhart, “Status of the Cf-252 Fission Neutron Spectrum Evaluation with Regard to Recent Experiments,” IAEA Consult. Meet. on “Physics of Neutron Emission in Fission”, Mito, Japan, 24–27 May 1988, Report **INDC(NDS)-0220**, pp. 305–336, IAEA, Vienna, Austria (1989).
- [92] **ASTM E170-17**, “Standard Terminology Relating to Radiation Measurements and Dosimetry,” ASTM International, West Conshohocken, PA, 2017.
- [93] W. Mannhart, “Evaluation of the Cf-252 Fission Neutron Spectrum between 0 MeV and 20 MeV,” in Properties of Neutron Sources, Report **IAEA-TECDOC-410**, pp. 158–171, IAEA, Vienna, Austria (1987).
- [94] N.V. Kornilov, “Verification of the ^{252}Cf standard in the energy range 2 - 20 MeV,” , Report **INDC(USA)-108**, IAEA, Vienna, Austria (2015); EXFOR 14431.
- [95] E. Blain, A. Daskalakis, R. C. Block and Y. Danon, “Measurement of prompt fission neutron spectrum for spontaneous fission of ^{252}Cf using γ multiplicity tagging,” *PHYS. REV.* **C95**, 064615 (2017); EXFOR 14477.
- [96] Nuclear Energy Agency, Collaborative International Evaluated Library Organization (CIELO) Pilot Project, WPEC Subgroup 40 (SG40), <https://www.oecd-nea.org/science/wpec/sg40-cielo/>.
- [97] R. Capote and A. Trkov (coordinators), “IAEA CIELO Data Development Project,” within the International Pilot Project of the OECD/NEA (see <https://www.nds.iaea.org/CIELO/>).
- [98] A. Trkov, R. Capote, “Evaluation of the Prompt Fission Neutron Spectrum of Thermal-neutron Induced Fission in U-235,” *PHYS. PROC.* **64**, 48–54 (2015).
- [99] R. Capote, A. Trkov, V.G. Pronyaev, “Current Issues in Nuclear Data Evaluation Methodology: ^{235}U Prompt Fission Neutron Spectra and Multiplicity for Thermal Neutrons,” *NUCL. DATA SHEETS* **123**, 8–15 (2015).
- [100] R. Capote, Y.-J. Chen, F.-J. Hamsch, N. V. Kornilov, J. P. Lestone, O. Litaize, B. Morillon, D. Neudecker, S. Oberstedt, T. Ohsawa, N. Otuka, V. G. Pronyaev, A. Saxena, O. Serot, O. A. Shcherbakov, N.-C. Shu, D. L. Smith, P. Talou, A. Trkov, A. C. Tudora, R. Vogt, A. C. Vorobyev, “Prompt fission neutron spectra of actinides,” *NUCL. DATA SHEETS* **131**, 1–106 (2016).
- [101] L.R. Greenwood, R.R. Heinrich, R.J. Kennerley, and R. Medrzychowski, “Development and Testing of Neutron Dosimetry Techniques for Accelerator-Based Neutron Irradiation Facilities,” *NUCL. TECHNOL.* **41**, 109 (1978) <https://doi.org/10.13182/NT78-A32137>.
- [102] M.J. Saltmarsh, C.A. Ludemann, C.B. Fulmer, R.C. Styles, “Characteristics of an Intense Neutron Source Based on the d+Be reaction,” *NUCL. INSTR. METH.* **143**, 81–90 (1977).
- [103] P.J. Griffin, K.R. DePriest, D.W. Vehar, E.J. Parma, Neutron Reference Benchmark Field Specification: SPR-III Central Cavity, Sandia National Laboratories, report **SAND2015-47190**, Albuquerque, NM, 2015.
- [104] J. G. Williams, D. W. Vehar, P. J. Griffin, D. B. King, “Covariance Matrices for Calculated Neutron Spectra and Measured Dosimeter Response,” *REACTOR DOSIMETRY STATE OF THE ART 2008. Proc. 13th Int. Symp. Held 25-30 May 2008 in Akersloot, The Netherlands. Edited by Wim Voorbraak (Institute for Energy, The Netherlands), Luigi Debarberis (Institute for Energy, The Netherlands), Pierre D’hondt (SCK-CEN, Belgium) & Jan Wagemans (SCK-CEN, Belgium)*. World Scientific Publishing Co. Pte. Ltd. (2009). ISBN 9789814271110, pp. 343–351.
- [105] K.R. DePriest, P.J. Cooper, and E.J. Parma, “MCNP/MCNPX model of the Annular Core Research Reactor,” report **SAND2006-3067**, Sandia National Laboratories, Albuquerque, NM (2006).
- [106] E.J. Parma *et al.*, “Radiation Characterization Summary: ACRR Central Cavity Free-Field Environment with the 32-Inch Pedestal at the Core Centerline (ACRR-FF-CC-32-cl),” report **SAND2015-6483**, Sandia National Laboratories, Albuquerque, NM (2015).
- [107] A. Fabry, G. Minsart *et al.*, “The Mol cavity fission spectrum standard neutron field and its applications,” in *Proc. 4th ASTM-EURATOM Symp. on Reactor Dosimetry*, NUREG/CP-0029 **Vol. 2**, 665–687 (1982).
- [108] D.M. Gilliam, J.A. Grundl, G.P. Lamaze *et al.*, in *Proc. 5th ASTM-EURATOM Symp. on Reactor Dosimetry*, Geesthacht, 1984, published by D. Reidel, Dordrecht, The Netherlands, **Vol. 2**, 867 (1985), EXFOR 23096.
- [109] W. Mannhart, “Spectrum-averaged neutron cross sections measured in the U-235 fission neutron field in Mol,” in *Proc. 5th ASTM-EURATOM Symp. on Reactor Dosimetry*, Geesthacht, 1984, published by D. Reidel, Dordrecht, The Netherlands (1985), p. 813–825.
- [110] J. Wagemans, E. Malambu, L. Borms, L. Fiorito, “The ^{235}U Prompt Fission Neutron Spectrum in the BR1 Reactor at SCK•CEN,” *EPJ WEB OF CONF.* **106**, 06003 (2016).
- [111] M. Košťál *et al.*, “On similarity of various reactor spectra and ^{235}U prompt fission neutron spectrum,” *APP. RAD. ISOT.* **135**, 83–91 (2018).
- [112] M. Košťál *et al.*, “Measurement of various monitors reaction rate in a special core at LR-0 reactor,” *ANN.*

- NUCL. EN. **112**, 759-768 (2018).
- [113] M. Košťál *et al.*, "Study of graphite reactivity worth on well-defined cores assembled on LR-0 reactor," ANN. NUCL. EN. **87**, 601-611 (2016).
- [114] M. Košťál *et al.*, "Determining the axial power profile of partly flooded fuel in a compact core assembled in reactor LR-0," ANN. NUCL. EN. **90**, 450-458 (2016).
- [115] M. Košťál *et al.*, "VVER-1000 physics experiments hexagonal lattices (1.275 cm Pitch) of Low Enriched U (3.3 wt.% $^{235}\text{UO}_2$) fuel assemblies in light water: $^{75}\text{As}(n,2n)$, $^{23}\text{Na}(n,2n)$, $^{90}\text{Zr}(n,2n)$, $^{89}\text{Y}(n,2n)$ reaction rates," **LR0-RESR-004**, submitted to IRPhE, 2018; EXFOR 31771.006.
- [116] R. Jeraj, M. Ravnik, "TRIGA Mark II reactor U(20) - zirconium hydride fuel rods in water with graphite reflector". in *International Handbook of Evaluated Criticality Safety Benchmarks Experiments*, NEA/NSC/DOC **95 03** (1999).
- [117] Ž. Štancar, *et al.*, "Reaction Rate Distribution Experiments at the Slovenian JSI TRIGA Mark II Research Reactor, TRIGA-FUND-RESR-002". In: *International Handbook of Evaluated Reactor Physics Benchmark Experiments [DVD]*, Nuclear Energy Agency, Paris OECD Nuclear Energy Agency (2017), p. 251.
- [118] V. Radulović, *et al.*, "Use of boron nitride for neutron spectrum characterization and cross-section validation in the epithermal range through integral activation measurements," NUCL. INSTR. METH. **A840**, 5-14 (2016).
- [119] V. Radulović, *et al.*, "Nuclear reactions suitable for epithermal neutron dosimetry using boron-based filters: preparation of an experimental campaign at the JSI TRIGA reactor," in *textitProc. 27th Int. Conf. Nucl. Energy for New Europe - NENE 2018*, Portorož, Slovenia, September 10-13, Nuclear Society of Slovenia (2018).
- [120] J. Grundl and D.C. Eisenhauer, "Benchmark neutron fields for reactor dosimetry," in *Neutron Cross Sections for Reactor Dosimetry*, report **IAEA-208**, v.1, 53-104, International Atomic Energy Agency, Vienna, Austria (1978).
- [121] B.M. Oliver, H. Farrar IV, D.M. Gilliam, E.P. Lippencott, "Spectrum-integrated helium generation cross sections for ^6Li and ^{10}B in the intermediate-energy field," in *Proc. 5th ASTM-EURATOM Symp. on Reactor Dosimetry*, Geesthacht, 1984, published by D. Reidel, Dordrecht, The Netherlands, **Vol. 2**, 877 (1985), EXFOR 13752.
- [122] G.P. Lamaze, C.M. Eisenhauer *et al.*, in *Int. Conf. Nucl. Data Sci. & Techn.*, Mito, 1988, 1033 (1988).
- [123] C.B. Besant, J. Emmett, C.G. Campbell *et al.*, "Design and construction of a fast reactor neutron spectrum generator - NISUS," NUCL. ENG. INT. **18**, 425-428 (1973).
- [124] A.H.M.A. Hannan and J.G. Williams, "Activation Foil Data for Nisus, MOL-SS and ^{235}U Fission Spectrum," in *Neutron Cross Sections for Reactor Dosimetry*, report **IAEA-208**, v.2, 209-225, International Atomic Energy Agency, Vienna, Austria (1978).
- [125] A. Fabry, G. De Leeuw and S. De Leeuw, "The Secondary Intermediate-Energy Standard Neutron Field at the MOL- Σ - Σ Facility," NUCL. TECH. **25**, 349-375 (1975).
- [126] A. Fabry, J.G. Williams *et al.*, "Benchmark neutron fields for reactor dosimetry," in *Neutron Cross Sections for Reactor Dosimetry*, report **IAEA-208**, v.2, 191, International Atomic Energy Agency, Vienna, Austria (1978).
- [127] I. Garlea, C. Miron, M. Lupu *et al.*, "Measuring of a few integral data in the Sigma-Sigma neutron field," REV. ROUM. PHYS. **23**, 409 (1978); EXFOR 30452.
- [128] I. Garlea and C. Miron, "Sigma Sigma-ITN Facility - Intermediate energy neutron source for reactor dosimetry," REV. ROUM. PHYS. **26**, 643 (1981); EXFOR 30568.
- [129] A. Fabry, "Role of Standard Fields", in *Fast Reactor Spectrum Measurements and their Interpretation*, Report **IAEA-138**, p. 89, International Atomic Energy Agency, Vienna, Austria (1971).
- [130] B.M. Oliver, H. Farrar IV, E.P. Lippencott and A. Fabry, "Spectrum-integrated helium generation cross sections for ^6Li and ^{10}B in the sigma-sigma and fission cavity standard neutron fields," in *4th Symp. on Reactor Dosimetry*, USA Washington D.C. 1982, Vol. 2, p. 889 (1982); EXFOR 23226.
- [131] Standard Spectra collection on the IAEA web page "International Reactor Dosimetry File: IRDF-2002" [1]. Online at <https://www-nds.iaea.org/irdf2002/index.html>.
- [132] R.J. LaBauve, "Bare, Highly Enriched Uranium Sphere (GODIVA)". HEU-MET-FAST-001 Revision 2 dated September 30, 2002 in the International Handbook of Evaluated Criticality Safety Benchmark Experiments, **NEA/NSC/DOC(95)03/II**.
- [133] J.L. Conlin *et al.*, "Release of ENDF/B-VIII.0 Based ACE Data Files," **LA-UR-18-24034** dated May 9, 2018. The data files may be downloaded from <https://nucleardata.lanl.gov>.
- [134] R.W. Brewer, "Uranium-235 Sphere Reflected by Normal Uranium using Flattop". HEU-MET-FAST-028 Revision 1 dated September 30, 1999 in the International Handbook of Evaluated Criticality Safety Benchmark Experiments, **NEA/NSC/DOC(95)03/II**.
- [135] A.L. Sapir, B.J. Krohn and R.D. Mosteller, "Big-Ten: A Large Mixed-Uranium Metal Cylindrical Core with 10% Average ^{235}U Enrichment, Surrounded by a Thick ^{238}U Reflector". IEU-MET-FAST-007 Revision 1 dated September 30, 1999 in the International Handbook of Evaluated Criticality Safety Benchmark Experiments, **NEA/NSC/DOC(95)03/II**.
- [136] ICSBEP 2016: International Handbook of Evaluated Criticality Safety Benchmark Experiments, Nuclear Energy Agency, OECD, Paris 2016. See <http://www.oecd-nea.org/science/wpncs/icsbep/handbook.html>.
- [137] Y.Khomyakov, "Cross Section Ratios Measured in the Core Center of the BR-1 Fast Spectrum Assembly". **FUND-IPPE-FR-MULT-RRR-001** Revision-0, September 30, 2009, International Handbook of Evaluated Criticality Safety Benchmark Experiments, **NEA/NSC/DOC(95)03/II**.
- [138] J.W. Rogers, D.A. Millsap and Y.D. Harker, "CFRM Neutron Field Flux Spectral Characterization," NUCL. TECH. **25**, 330-348 (1975).
- [139] J.W. Rogers, Y.D. Harker and D.A. Millsap, "Fast Neutron Spectrum and Dosimetry Studies in the Coupled Fast Reactivity Measurements Facility," Proc. of First ASTM-EUROATOM Symp. on Reactor Dosimetry, 22-26 Sep 1975 Petten, in Report EUR 5667 e/f, part II, p. 277-295, Luxembourg 1977.
- [140] J.W. Rogers, Y.D. Harker and D.A. Millsap, "The Coupled Fast Reactivity Measurement Facility (CFRMF)," in Report **IAEA-208**, Vienna 1976, v. II, p. 117-176; SACS will be compiled in EXFOR.
- [141] J.A. Favorite, "Bare Sphere of Plutonium-239 Metal (4.5 at.% ^{240}Pu , 1.02wt.% Ga)". PU-MET-FAST-001 Revision 4 dated September 30, 2016 in the International Handbook of Evaluated Criticality Safety Benchmark

- Experiments, **NEA/NSC/DOC(95)03/II**.
- [142] R.D. O'Dell, "240Pu Jezebel: Bare Sphere of Plutonium-239 Metal (20.1 at.% 240Pu, 1.01wt.% Ga)". PU-MET-FAST-002 Revision 1 dated September 30, 1999 in the International Handbook of Evaluated Criticality Safety Benchmark Experiments, **NEA/NSC/DOC(95)03/II**.
- [143] R.W. Brewer, "Plutonium Sphere Reflected by Normal Uranium using Flattop". PU-MET-FAST-006 Revision 1 dated September 30, 1999 in the International Handbook of Evaluated Criticality Safety Benchmark Experiments, **NEA/NSC/DOC(95)03/II**.
- [144] R.W. Brewer, "Benchmark Critical Experiment of Thorium Reflected Plutonium Sphere". PU-MET-FAST-008 Revision 2 dated September 30, 1999 in the International Handbook of Evaluated Criticality Safety Benchmark Experiments, **NEA/NSC/DOC(95)03/II**.
- [145] "The International Reactor Dosimetry File (IRDF-90 Version 2)," report **IAEA-NDS-141**, IAEA, Vienna (1999). Online at <https://www-nds.iaea.org/public/libraries/irdf/>.
- [146] W. Mannhart, "Status of the evaluation of the neutron spectrum of ^{252}Cf (s.f.)," IAEA Consult. Meet. on *International Neutron Cross-Section Standards: Measurements and Evaluation Techniques*, Vienna, 13–15 Oct 2008, Report **INDC(NDS)-0540**, Vienna, 2008. Available online.
- [147] M. Paul, M. Tessler, M. Friedman *et al.*, "Reactions along the astrophysical s-process path and prospects for neutron radiotherapy with the Liquid-Lithium Target (LiLiT) at the Soreq Applied Research Accelerator Facility (SARAF)," **EUR. PHYS. JOURN. A55**, 44 (2019).
- [148] S. Manojlovic, A. Trkov, "Nuclear Cross Section Measurement Analysis in the Californium-252 Spectrum with the Monte Carlo Method", Conf. "Nuclear Energy for New Europe 2011", Ljubljana, contribution 307.
- [149] W. Mannhart, "Response of Activation Reactions in the Neutron Field of ^{252}Cf (s.f.)," IAEA **Technical Report Series No. 452**, IAEA, Vienna, Austria, 2006, pp. 30–45.
- [150] J. Csikai, Z. Dezsó, "Average cross sections for the Cf-252 neutron spectrum. II. (n,f) reactions," in *VIIIth Gaussig Symp.*, Nov 1977, Report **ZFK-376**, Rossendorf, pp. 44–46 (1978); EXFOR 30415.004.
- [151] F. Cserpak, T. Biro, J. Csikai, "Measurement of cross-sections for (n,t) reactions on light nuclei," in *Int. Conf. on Nucl. Phys. and Nucl. Data*, Harwell 1978, p. 761 (1978); EXFOR 30474.002.
- [152] K.I. Zolotarev, "Evaluation of excitation functions for $^{28}\text{Si}(n,p)^{28}\text{Al}$, $^{31}\text{P}(n,p)^{31}\text{Si}$, and $^{113}\text{In}(n,\gamma)^{114m}\text{In}$ reactions," Report **INDC(NDS)-0668**, International Atomic Energy Agency, Vienna, Austria (2014).
- [153] M.V. Blinov, E.A. Gromova, Kovalenko *et al.*, "Cross Section of the $^{238}\text{U}(n,2n)^{237}\text{U}$ Reaction induced by Neutrons from ^{252}Cf Spontaneous Fission," **SOV. AT. EN. 65**, 761–764 (1988); EXFOR 40996.002.
- [154] M. Schulc *et al.*, "Validation of IRDF-II library by means of ^{252}Cf Spectral Averaged Cross Sections", **APP. RAD. ISOT.**, submitted for publication 8 Aug 2019.
- [155] M. Schulc, M. Kostal, S. Simakov *et al.*, "Validation of differential cross sections by means of ^{252}Cf spectral averaged cross sections," **APP. RAD. ISOT. 132**, 29–37 (2018); EXFOR 31786.002.
- [156] M. Schulc, M. Kostal, R. Capote *et al.*, "Validation of selected (n,2n) dosimetry reactions in IRDF-1.05 library," **APP. RAD. ISOT. 143**, 132–140 (2019).
- [157] M. Schulc, M. Kostal, E. Novak *et al.*, "Investigation of $^{127}\text{I}(n,2n)^{126}\text{I}$ and $^{23}\text{Na}(n,2n)^{22}\text{Na}$ reactions using ^{252}Cf neutron source," **ASME J. NUCL. RAD. SC. 5**, 030918 (2018).
- [158] A. Trkov, "Program RRUNC - calculation of uncertainties in reaction rates and cross sections," Jozef Stefan Institute, Ljubljana, Slovenia, https://www-nds.iaea.org/IRDF/rr_unc.for.
- [159] M. Schulc, *et al.*, "Application of ^{252}Cf neutron source for precise nuclear data experiments," **APP. RAD. ISOT. 151**, 187–195 (2019).
- [160] M.E. Wyman, E.M. Fryer and M.M. Thorpe, "(n,t) Cross Sections for B^{10} , B^{11} and Be^9 ," **PHYS. REV. 112**, 1264 (1958); EXFOR 11231.005.
- [161] M. Košťál *et al.*, "A reference neutron field for for measurement of spectrum averaged cross sections," **ANN. NUCL. ENERGY** (2019), to be published.
- [162] M. Košťál, M. Svadlenkova, P. Baron *et al.*, "Measurement of $^{23}\text{Na}(n,2n)$ cross section in well-defined reactor spectra," **APP. RAD. ISOT. 111**, 1–7 (2016).
- [163] M. Kostal, "VVER-1000 physics experiments hexagonal lattices (1.275 cm Pitch) of Low Enriched U (3.3 wt.% $^{235}\text{UO}_2$) fuel assemblies in light water: $^{75}\text{As}(n,2n)$, $^{23}\text{Na}(n,2n)$, $^{90}\text{Zr}(n,2n)$, $^{89}\text{Y}(n,2n)$ reaction rates," **LR0-RESR-004**, submitted to IRPhE (2018); EXFOR 31771.006.
- [164] D.J. Hughes, R.C. Garth, J.S. Levin, "Fast Neutron Cross Sections and Nuclear Level Densities," **PHYS. REV. 91**, 1423–1429 (1953); EXFOR 13860.014.
- [165] M. Košťál *et al.*, "Validation of $^{14}\text{N}(n,p)$ cross section in LR-0 reference field," **APP. RAD. ISOT.**, 2019, to be published.
- [166] N. Burianova, M. Kostal, M. Schulc *et al.*, "Measurement of selected differential cross sections in ^{235}U spectrum," **ASME J. NUCL. RAD. SC. 5**, 030906 (2019).
- [167] P.M. Suarez, M.A. Arribere *et al.*, "Fission Neutron Spectrum Averaged Cross-Section of the $^{54}\text{Fe}(n,f)^{51}\text{Cr}$ Reaction," **RAD. ACTA 78**, 11–13 (1997); EXFOR 31508.002.
- [168] A.V. Bushuev, A.V. Zvonarev *et al.*, Determination of the cross-sections for some neutron reactions with structural materials in the BN-350 reactor core, **Sov. J. of Atom Energy**, V. 63, p. 721 (1987); EXFOR 40956.012.
- [169] K. Kobayashi and I. Kimura, "The average cross section for the (n,2n) reactions of Mn-55, Ni-58, Co-59 and I-127 to the fission neutrons of U-235t," Report INDC(JAP)-54, p. 42 (1980), IAEA; EXFOR 21693.003.
- [170] O. Horibe and H. Chantani, "Cross sections of the reactions $^{55}\text{Mn}(n,2n)^{54}\text{Mn}$, $^{58}\text{Ni}(n,2n)^{57}\text{Ni}$ and $^{58}\text{Ni}(n,np)^{57}\text{Co}$ averaged over the U-235 fission neutron spectrum," *Conf. on Nucl. Data for Sci. and Technol., Julich 1991*, p.68 (1991); EXFOR23223.004.
- [171] T. Sekine and H. Baba, Cross sections of the (n,2n) reaction of ^{59}Co , ^{58}Ni , ^{70}Ge , ^{90}Zr and ^{203}Tl with fission neutrons, **J. Inorg. Nucl. Chem. 43**, 1427 (1981); EXFOR 21613.003.
- [172] J.A. Grundl, "A study of fission neutron spectra with high-energy activation detectors. Part II Fission Spectra," **NUCL. SCI. TECH. 31**, 191 (1968); EXFOR 12193.008.
- [173] A. Fabry, M. DE Coster, "Integral test of capture cross-sections in the energy range 0.1 to 2 MeV," *Proc. Int. Conf. on Nuclear Cross-Sections and Techn.*, Washington 1968, Vol. 2, pp. 1263–1267 (1968); EXFOR 20264.
- [174] K.I. Zolotarev, A.V. Ignatyuk, V.N. Manokhin, and A.B. Pashchenko, "RRDF-98 Russian Reactor Dosimetry File," Report **IAEA(NDS)-193**, IAEA, Vienna, Austria (1999).

- [175] M. Košťál, M. Schulc, J. Simon *et al.*, “Measurement of various monitors reaction rate in a special core at LR-0 reactor,” *ANN. NUCL. EN.* **112**, 759–768 (2018).
- [176] G. Rau, Averaged Cross Section of the Reaction $Y-89(n,2n)Y-88$, in: Progress Report EANDC(E)-89, February 1968, p. 41.
- [177] E. Steinnes, “Cross-Sections of Some (n,2n) Reactions Induced by Reactor Fast Neutrons,” *RAD. ACTA* **13**, 169 (1970).
- [178] A. A. Bojtcov, E.I. Grigoriev, V.S. Troshin, V.P. Yaryna, “Measurements and Evaluations of the Averaged Cross-Section for $89Y(n,2n)88Y$ Reaction for the Fission $235U$ Neutrons”, *VOPROCY ATOMNOJ NAUKI I TEHNIKI*, Ser.Yadernye Konstanty, 1 (1992) 55-56 (in Russian).
- [179] K. Kobayashi and T. Kobayashi, “Measurements of U-235 fission spectrum-averaged cross sections and neutron spectrum adjusted with the activation data,” Report **KURRI-AR-26**, p. 21 (1992), Kyoto University; EXFOR 22216.
- [180] E.I. Grigoriev, G.B. Tarnovskij, V.P. Jaryna, “Measurement of the neutron threshold reactions cross sections for U-235 fission spectrum,” *Proc. 6th All-Union Conf. on Neutron Physics, Kiev 2-6 Oct 1983*, Vol. 3, p. 187 (1983); EXFOR 40848.002.
- [181] K.I. Zolotarev, “Evaluation and improvement of cross section accuracy for most important dosimetry reactions including covariance data”, Report **INDC(NDS)-0431**, IAEA, Vienna, Austria (2002).
- [182] I. Kimura, K. Kobayashi and T. Shibata, “Measurements of Activation Cross Sections for Some Threshold Reactions for Neutrons with Fission-Type Reactor Spectrum,” *J. NUCL. SC. TECH.* **8**, 59–69 (1971); EXFOR <http://www-nds.iaea.org/EXFOR/20317.018>.
- [183] A.K. Brodskaya *et al.*, “Fission neutron spectra averaged cross-sections of (n,a), (n,p), (n,2n), (n,n) reactions for some elements,” *YAD.-FIZ. ISSLED.* **24**, 15 (1977) (in Russian).
- [184] K. Kobayashi, I. Kimura *et al.*, *J. NUCL. SC. TECH.* **13**, 531 (1976); EXFOR 20693.
- [185] T. Hashimoto, T. Sotobayashi and K. Kobayashi, “Measurement of Average Cross Section for $238U(n, 2n)237U$ Reaction,” *J. NUCL. SC. TECH.* **15**, 626–628 (1978); EXFOR 20810.002.
- [186] G. Shani, “(n,2n) cross-section measurement of Nb-93, Au-197 and U-238 with fission-neutron spectrum”, *ANN. NUCL. EN.* **10**, 473 (1983); EXFOR 30658.004.
- [187] R. Capote, A. Trkov, “Predicting Spectrum Averaged Cross Sections in Prompt Fission Neutron Fields,” *16th International Symposium on Reactor Dosimetry*, **ASTM STP1608**, M. H. Sparks, K. R. DePriest, and D. W. Vehear, Eds., ASTM International, West Conshohocken, PA, 2018, pp.117–123, <http://dx.doi.org/10.1520/STP160820170114>.
- [188] P.J. Griffin, D.W. Vehear, E.J. Parma, K.D. Hahn, “Rigorous Uncertainty Propagation Using a Dosimeter Transfer Calibration,” *16th International Symposium on Reactor Dosimetry*, **ASTM STP1608**, M. H. Sparks, K. R. DePriest, and D. W. Vehear, Eds., ASTM International, West Conshohocken, PA, 2018, pp. 124–134, 2018, <https://doi.org/10.1520/STP1608-EB>.
- [189] M. Košťál, M. Schulc, V. Rypar, *et al.*, “Validation of zirconium isotopes (n, γ) and (n,2n) cross sections in a comprehensive LR-0 reactor operative parameters set,” *APPL. RAD. ISOT.* **128**, 92–100 (2017).
- [190] P. Dryak, P. Kovar, “Experimental and MC determination of HPGe detector efficiency in the 40-2754 keV energy range for measuring point source geometry with the source-to-detector distance of 25 cm,” *APPL. RAD. ISOT.* **64**, 1346–1349 (2006).
- [191] Trkov, A., Radulovic \acute{c} , V., Snoj, L., “The GRUPINT neutron spectrum adjustment code - general features and characterization of the spectra in three irradiation channels of the JSI TRIGA reactor,” *16th International Symposium on Reactor Dosimetry*, May 7–12 (2017).
- [192] P.G.Young *et al.*, “Evaluation of Neutron Reactions for ENDF/B-VII: $^{232-241}U$ and ^{239}Pu ,” *NUCL. DATA SHEETS* **108**, 2589–2654 (2007).
- [193] S.C.Frankle and J.F.Briesmeister, “Spectral Measurements in Critical Assemblies: MCNP Specifications and Calculated Results,” Report **LA-13675**, December 1999.
- [194] Cross Section Evaluation Working Group Benchmark Specifications, **BNL 19302 (ENDF-202)**. Originally published November 1974 with updates to selected benchmarks or new additions in May 1975, October 1976, February 1978, October 1978, May 1981, November 1981, September 1982, December 1983, September 1986 and September 1991.
- [195] J.A. Grundl, C.M. Eisenhauer, “Compendium of Benchmark Neutron Fields for Reactor Dosimetry,” **Report NSIR 85-03151**, National Bureau of Standards, 1986.
- [196] K. Kobayashi, I. Kimura, Report **EUR-6813**, 1004, 1980; EXFOR 21589;
- [197] I. Garlea, C. Miron, *REV. ROUM. PHYS.* **23**, 409 (1978)
- [198] A.H.M.A. Hannan, J.G. Williams, Report **NEANDC(E)-172**, p. 8, France, 1976.
- [199] E.P. Lippencott, A.L. Pitner and L.S. Kellogg, “Measurement of $^{10}B(n,t)$ cross section in a fast-neutron spectrum”, *TRANS. AMER. NUCL. SOC.* **17**, p. 494 (1973); EXFOR 13756.
- [200] H. Farrar IV, W.N. McElroy and E.P. Lippencott, “Helium production cross sections of Boron for fast reactor neutron Spectra”, *NUCL. TECHN.* **25**, 305 (1975); EXFOR - to be compiled.
- [201] Y.D. Harker, E.H. Turk, Conf. on Develop. in Reactor Phys., Kiamesha Lake 1972, v. 2, p. 614
- [202] J.A. Grundl, D.M. Gilliam *et al.*, *NUCL. TECHN.* **25**, 237 (1975).
- [203] R.A. Anderl and Y.D. Harker, “Measurement of the Integral capture and fission cross sections for ^{232}Th in the CFRMF,” *Proc. Int. Conf. on Nucl. Cross Sections For Techn.*, Oct 1979, Knoxville, p. 475; EXFOR 14354.
- [204] F. De Corte, “The KO-standardization method,” Rijksuniversiteit, Gent, Thesis (1987).
- [205] V. Radulović, A. Trkov, R. Jaćimović, R. Jeraj, “Measurement of the neutron activation constants Q_0 and k_0 for the $^{27}Al(n,\gamma)^{28}Al$ reaction at the JSI TRIGA Mark II reactor,” *J. RADIOAN. NUCL. CHEM.* **298**, 1791–1800(2013).
- [206] *Karlsruhe Astrophysical Database of Nucleosynthesis in Stars KADoNiS version 1.0*; <https://exp-astro.de/kadonis1.0/>.
- [207] Z.Y. Bao, H. Beer, F. Kappeler *et al.*, “Neutron cross section for nucleosynthesis studies,” *AT. DATA NUCL. DATA TABLES* **76**, 70-154 (2000).
- [208] G. Feinberg, M. Friedman, A. Krasa *et al.*, “Quasi-stellar neutrons from the $^7Li(p,n)^7Be$ reaction with an energy-broadened proton beam,” *PHYS. REV.* **C85**, 055810 (2012); EXFOR 23284.
- [209] P. Jiménez-Bonilla and J. Praena, “Measurement of the $^{197}Au(n,\gamma)$ stellar cross section at $kT=30$ keV by activation”, 13th Symp. on Nuclei in the Cosmos, Debrecen, 2014, p. 102 (2014); EXFOR 23282.

- [210] A. Wallner, T. Belgya, M. Bichler *et al.*, “Novel Method to Study Neutron Capture of ^{235}U and ^{238}U Simultaneously at keV Energies”, *PHYS. REV LETT.* **112**, 192501 (2014); EXFOR 23170.
- [211] L.R. Greenwood *et al.*, Testing and Improving of the International Reactor Dosimetry and Fusion File (IRDF), International Atomic Energy Agency, Vienna, Austria, Report **INDC(NDS)-0731**, Vienna, Austria (2017).
- [212] L.R. Greenwood, B.P. Pierson, and C.D. Johnson, “Enhancement of STAYSL-PNNL with IRDF to 60 MeV,” *Reactor Dosimetry: 16th International Symposium*, **ASTM STP1608**, M. H. Sparks, K. R. DePriest, and D. W. Vehar, Eds., ASTM International, West Conshohocken, PA, 2018, pp. 265–275, <http://dx.doi.org/10.1520/STP1608201700542>.
- [213] D.R. Nethaway, R.A. Van Konynenberg, M. W. Guinan, and L. R. Greenwood, “Neutron Spectra from 30-MeV Deuterons on a Thick Beryllium Target,” *Proc. Symp. on Neutron Cross Sections from 10–40 MeV*,” Report **BNL-NCS-50681**, p.135 (1977).
- [214] L.R. Greenwood, R. R. Heinrich, M. J. Saltmarsh, and C. B. Fulmer, “Integral Tests of Neutron Activation Cross Sections in a $^9\text{Be}(d,n)$ Field at $E_d = 40$ MeV,” *NUCL. SCI. ENG.* **72**, 175 (1979) <https://doi.org/10.13182/NSE79-A19462>.