Templates of expected measurement uncertainties for average prompt and total fission neutron multiplicities

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Received: 23 March 2023 / Received in final form: 18 July 2023 / Accepted: 30 August 2023

Abstract. In this paper, we provide templates of measurement uncertainty sources expected to appear for average prompt- and total-fission neutron multiplicities, \( \bar{v}_p \) and \( \bar{v}_t \), for the following measurement types: absolute manganese-bath experiments for \( \bar{v}_t \), absolute and ratio liquid-scintillator measurements for \( \bar{v}_p \). These templates also suggest a typical range of these uncertainties and their correlations based on a survey of available experimental data, associated literature, and feedback from experimentalists. In addition, the information needed to faithfully include the associated experimental data into the nuclear-data evaluation process is provided.

1 Introduction

The average prompt-fission neutron multiplicity, \( \bar{v}_p \), of an actinide is the average number of neutrons emitted promptly following fission of that nucleus and before the onset of \( \beta \)-decay. The average total-fission neutron multiplicity, \( \bar{v}_t \), differs from \( \bar{v}_p \) in that it also includes delayed neutrons. These observables can be measured for both spontaneous, e.g., the Neutron Data Standard (NDS) \( ^{252}\text{Cf(sf)} \) \( \bar{v}_t \), or neutron-induced fission, e.g., the \( ^{235}\text{U}(n,f) \) \( \bar{v}_p \) as a function of incident-neutron energy. Neutron multiplicities can also be measured as a function of other quantities, for instance, fission-fragment mass, charge, or total kinetic energy. Here, we focus on \( \bar{v}_p \) and \( \bar{v}_t \). The variable \( \bar{v} \) is also used as a short version encompassing both \( \bar{v}_p \) and \( \bar{v}_t \).

Observations of \( \bar{v} \) are among those that can be measured to the highest precision, i.e., lowest uncertainty, compared to other nuclear-reaction data, with uncertainties reported as low as 0.3–0.5% [1–8]. Given that many \( \bar{v} \) measurements are of high precision, these experiments must be carefully documented with regard to data analysis, corrections, and all pertinent uncertainties. Many high-precision measurements are given for \( ^{252}\text{Cf(sf)} \bar{v}_t \) [1–9], thermal constants \( ^{233,235}\text{U}(n,f) \bar{v}_t \) and \( ^{239,241}\text{Pu}(n,f) \bar{v}_t \) as described in [10–18]) and also for incident-neutron energy dependent \( \bar{v}_p \) and \( \bar{v}_t \) [14–16,19–26].

In Section 4, templates are provided for \( \bar{v} \) measurements using the absolute manganese-bath technique for \( \bar{v}_t \), and absolute and ratio liquid-scintillator technique for \( \bar{v}_p \); these techniques are described in Section 2 along with information needed to be provided on a particular experiment for nuclear data evaluations in Section 3. The templates on absolute \( \bar{v} \) measurements were established by analyzing \( ^{252}\text{Cf(sf)} \bar{v}_t \) data sets using information from their literature [2–9,15,27–30] and respective entries in the EXFOR (Experimental Nuclear Reaction Data) database [31–33]. The templates for ratio liquid-scintillator measurements were obtained using information from several \( ^{239}\text{Pu}(n,f) \bar{v}_t \) and \( ^{239}\text{Pu}(n,f) \bar{v}_p \) data sets from thermal to 20 MeV incident-neutron energy using their literature and EXFOR entries [12–26,34–42]. Most of these data sets were either measured at thermal energies or above 100 keV. Only scarce experimental information is available in the resonance range, where variations in \( \bar{v}_p \) have been investigated for \( ^{235}\text{U}(n,f) \) (negligible variations) and \( ^{239}\text{Pu}(n,f) \) (significant variations) [43,44]. The experimental information obtained from these analyses was supplemented by reviews on these experiments and associated measurement techniques [1,45–48], while references [11,43,49,50] provided input on \( \bar{v}_p \) and \( \bar{v}_t \) evaluations. Section 5 provides a short summary.
2 Measurement types

Three techniques were used for absolute $^{252}$Cf(sf) $\bar{\nu}$ measurements: manganese-sulfate baths (Mn baths) [3,5–7,9,11,30], boron piles [4,29] and doped liquid scintillators [1,2,8,15,27]. The differentiation is made according to the neutron-absorbing material, as each of these materials and associated measurement types carry distinct uncertainty sources. Boron-pile measurements will not be discussed here as the template would have been established on only two $^{252}$Cf(sf) $\bar{\nu}$ measurements [4,29], too small of a sample size for realistic estimates of typically appearing uncertainty sources. Mainly one ratio-type measurement was encountered in EXFOR entries for $^{239}$Pu $\bar{\nu}$ and $\gamma$ [12–26,34–42], namely, ratio liquid-scintillator measurements, which will be discussed below. Some $\gamma$, data have also been measured with setups that are traditionally used to determine prompt fission neutron spectra (PFNS), see e.g., [51–54]. To this end, the PFNS is measured via the techniques described in reference [55], extrapolated to span the entire outgoing-neutron energy, $E_{\text{out}}$, range, and then integrated over these energies to yield $\bar{\nu}_p$. Hence, the templates in Tables I and II of [55] apply to this measurement technique along with additional uncertainties for the extrapolation of the PFNS for $E_{\text{out}}$ and angular coverage of the neutron detectors. Given that and that only a few data sets use this technique, no separate template is provided.

2.1 Mn-bath measurements

In this type of experiment, $\bar{\nu}_f$ is absolutely determined by measuring the absolute neutron-emission rate, $c_n$, independently from the fission rate, $c_f$, (see Fig. 1), and then dividing them

$$\bar{\nu}_f = \frac{c_n}{c_f}. \quad (1)$$

The neutron-emission rate, $c_n$, is measured by a manganese-sulfate bath. To this end, the sample or detector enclosing it is placed in a re-entrant cavity such that it is located near the center of the bath. The Mn solution is very caustic and is never in contact with the sample or detector. When neutrons are emitted from the sample, some are captured by $^{55}$Mn forming radioactive $^{56}$Mn. For some measurements, the solution in the bath is continuously stirred and pumped through a tube and back into the bath. Detectors (often NaI scintillators) are located very near (or in a re-entrant well in) that tube, to detect the $\gamma$ rays from the decay of $^{56}$Mn, which yields an estimate of $c_n$ after applying necessary corrections. Non-pumped variants also were undertaken, where detectors were placed on the tank walls. The formal analysis procedure to obtain $c_n$ from this measurement was described in references [30,45] by:

$$c_n = \frac{c_\gamma}{\varepsilon_{\text{Mn}}} \frac{1 + \frac{\sigma_S}{\sigma_{\text{Mn}}(1 + \sigma_{\text{Mn}}) + \frac{N_{\text{Mn}}}{N_{\text{S}}} \sigma_H}}{(1 - L_n)(1 - S)(1 - O)}, \quad (2)$$

where $c_\gamma$ is the $\gamma$-counting rate measured by the Mn bath at saturation (i.e., activity equilibrium) [30,47] with an efficiency $\varepsilon_{\text{Mn}}$. Corrections have to be applied for thermalized neutrons captured by sulfur and hydrogen using their respective cross sections, $\sigma_S$ and $\sigma_H$, in ratio to that of Mn, $\sigma_{\text{Mn}}$, which can be determined independently. To this end, the ratio of the number of atoms of Mn to H, $\frac{N_{\text{Mn}}}{N_{\text{H}}}$, has to be considered. The ratio of Mn to S is always unity, due to the chemical composition of the bath and, hence, does not appear. No correction has to be applied for thermalized neutrons captured in $^{16}$O of the MnSO$_4$ in the bath, as the thermal neutron-capture cross-section of $^{16}$O is approximately a factor 10,000 smaller than the Mn cross-section. $^{16}$O is also present in the water of the aqueous solution; hence, the O/Mn also depends on the concentration of the solution. Different baths use different concentrations and this should be documented. The term $(1 + G \Gamma S)$ accounts for the resonance capture of neutrons in Mn, with G the resonance self-shielding factor, $\bar{\nu}$ the Wescott factor (epithermal flux parameter averaged over the Mn bath) and s being the resonance-activation integral of Mn. The term $(1 - L_n)$ accounts for neutrons leaking from the bath, $(1 - S)$ for neutrons recaptured by the source or material in the cavity, and $(1 - O)$ for those fast neutrons undergoing (n,α) and (n,p) reactions in oxygen in the water or sulfur present in the Mn-bath solution. Mn baths are often calibrated with respect to national standards [48]. The $\gamma$ detectors measuring decay from $^{56}$Mn can be calibrated by adding $^{56}$Mn to the solution. The activity can be determined absolutely very accurately making Mn-bath measurements a technique supported by basic radiation metrology. Thermal-neutron absorber impurities need to be known and corrected as described as part of the templates below. The temperature of the bath can also lead to a subtle effect if the pumping heats the water and it is not in thermal equilibrium with environmental temperatures.
A drawback of the Mn-bath technique is that it requires fairly strong neutron-emission rates. $^{252}$Cf naturally lends itself to such a technique, but it is a significant challenge to create deposits of strong enough fission rates that remain amenable to fission counting. However, it is a very important technique as it allows experimentalists to uncover unknown systematic biases in standard techniques, such as the liquid-scintillator technique. The fission rate, $c_f$, was determined, e.g., by solid-angle counting of fission fragments using Si-surface barrier detectors [9,30], fission-fission coincidence counting [3] or neutron-fission coincidence counting [7]. No explicit equation is given here to derive $c_f$ as it would be different for each of those three measurement techniques.

2.2 Absolute liquid-scintillator measurements

Absolute liquid-scintillator measurements belong to the class of delayed-coincidence measurements [1] and were used to report, e.g., $^{252}$Cf(sf) $\nu$ measurements [1,2,8,15,27] or also the neutron-multiplicity distribution, $P(\nu)$. In these measurements, a fission detector is placed within a through-tube running through the center of a large liquid scintillator, as shown in Figure 2.\(^1\) When the fission detector is triggered (often by fission fragments), a neutron-counting gate is opened for a defined time (between 30–100 $\mu$s across, e.g., the following ratio measurements [12–15,19–23,28,34,35]). During this time, some of the emitted neutrons are captured by the dopant in the scintillator material. The associated capture $\gamma$ rays cause scintillations that are subsequently detected by photomultipliers. This technique differs from Mn-bath measurements in as far as it is independent of the absolute fission rate as it measures $\nu$ per detected fission event. The liquid-scintillator material is loaded or doped with gadolinium [1,2,8,19–25] or cadmium [15,26–28]. Both have large neutron-capture cross sections at thermal energies and liberate approximately 6 MeV of $\gamma$ rays in the capture reaction. These $\gamma$ rays give an easily detected signal when converted to light in the liquid scintillator. Paraffin scintillators were also used for a few $^{239}$Pu $\nu$ ratio liquid-scintillator measurements [17,18,39,40]. The analysis of an absolute liquid-scintillator $\nu_p$ measurement

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\(^1\) To be exact, one does not need to use necessarily a liquid-scintillator neutron detector; one could, for instance, detect neutrons via plastic scintillator bars interlaced with thin Gd sheets, or poly assemblies with large $^3$He tubes.
can be described by the following equation:
\[
\nu_p = \frac{c - c_{\text{DG}} - b - c_{\text{ff}} - c_{\text{FE}}}{\varepsilon_n \tau} \omega. \tag{3}
\]

The photomultipliers convert the scintillator light produced by \( \gamma \) rays from the neutron-capture event in the scintillator into electrons and, hence, in a signal. These counts \( c \) are average counts per fission, as these \( \gamma \)s are only counted after fission triggered the start of the neutron-counting gate for a limited time. The fission count rate does not appear in this equation, contrary to Mn-bath measurements, because fission is used as a trigger and the desired quantity is the number of neutrons per fission. However, the fission rate must be low enough that neutrons from a not-detected fission event are not measured as part of the next fission-neutron coincidence. The total counts, \( c \), are corrected for (a) \( \gamma \) counts originating from delayed \( \gamma \)s due to radioactive decay of the fission fragment, \( c_{\text{DG}} \), (b) random background counts in the neutron detector, \( b \), produced by background neutrons not originating from fission, (c) false fission events, \( c_{\text{ff}} \), triggered by random coincidences, and (d) counts due to the so-called “French effect”, \( c_{\text{FE}} \) (described in detail as part of Sect. 4). The correction factor, \( \omega \), accounts for neutrons resulting from fission of impurity isotopes in the sample, while \( \tau \) is the dead-time correction. The remaining term is the average scintillator detector efficiency, \( \varepsilon_n \), which is obtained by:
\[
\varepsilon_n(E_{\text{inc}}) = c_\gamma \int dE_{\text{out}} d\Omega_{\text{out}} c(E_{\text{out}}, \theta) \chi(E_{\text{inc}}, E_{\text{out}})
\times L_n(E_{\text{out}}, \theta) a(E_{\text{inc}}, E_{\text{out}}, \theta) A_t(E_{\text{out}}, \theta), \tag{4}
\]
where \( c_\gamma \) is the efficiency for detecting \( \gamma \) rays produced in the neutron capture on gadolinium or cadmium. It is assumed to be independent of the neutron energy in reference [1]. The efficiency of the neutron-capture process, \( c(E_{\text{out}}, \theta) \), is usually calculated via Monte Carlo (MC) codes and depends on nuclear data [1] (cross sections, angular distributions, PFNS), as is the neutron-leakage correction, \( L_n \). Calculations of \( \varepsilon_n \) may also depend on a discrimination threshold to detect the \( \gamma \)s. Hence, neutrons that do not produce \( \gamma \)s are included in this calculation. The neutron-detector efficiency also requires knowledge of the PFNS, \( \chi \), and the angular distribution of fission neutrons, \( a \), which are either simulated or taken from nuclear data. The asymmetry introduced by the through-tube, \( A_t \), also has to be accounted for in the neutron-detector efficiency calculation. The neutron-detector efficiency, \( \varepsilon_n \), can also be absolutely measured by means of proton-recoil detectors located in the center of the scintillator, as was done in references [2,15]. These detectors measure the fraction of pulses, \( C_0 \), detected by the proton-recoil detector but not by the scintillator as a function of incident-neutron energy. One can then derive \( \varepsilon_n \) by (1 – \( C_0 \))/ (1 – \( \int dE_{\text{out}} d\Omega_{\text{out}} L_n(E_{\text{out}}, \theta) \chi(E_{\text{inc}}, E_{\text{out}}) a(E_{\text{inc}}, E_{\text{out}}, \theta) \)), where the denominator is simulated using nuclear data. The derived \( \varepsilon_n \) can then be compared to equation (4). One can also combine measurements and simulations, by measuring the efficiency for a few energies, while calculations, that were normalized to the measured \( \varepsilon_n \), fill in the missing energies [2,4]. This combined approach has the advantage of avoiding the need for absolute simulated values of \( \varepsilon_n \). Although equation (4) looks forbidding, in practice, it usually varies only gradually with mean prompt fission neutron energy.

### 2.3 Ratio liquid-scintillator measurements

These measurements differ from their absolute counterpart in that two measurements are performed. The first measurement determines \( \bar{\nu} \) of the desired isotope and is schematically the same as absolute measurements in Figure 2. The second experiment uses the same liquid scintillator and determines its efficiency by measuring a reference (or monitor) sample. This monitor is often \( ^{252}\text{Cf} \) and thus no incident-neutron beam is shown in Figure 2. The resulting experimental quantity is then a ratio of \( \bar{\nu} \) of the desired and monitor isotopes. The measured ratio relative to monitor \( m \) is determined by:
\[
\bar{\nu}_p/\bar{\nu}_p^m = \frac{c - c_{\text{DG}} - b - c_{\text{ff}}}{c^m - c_{\text{DG}}^m - b^m - c_{\text{ff}}^m} \omega \tau \varepsilon_d d, \tag{5}
\]
with an additional correction for the thickness of the sample, \( d \), leading to differences in the fission-fragment efficiency for specific fission-fragment pairs that may impact \( \nu \) [21,56]. This factor does not appear for absolute measurements of \( ^{252}\text{Cf} \), as the samples are usually sources of small diameter or thickness, or both. The efficiency, \( \varepsilon_n' \), changes in the ratio measurement to:
\[
\varepsilon_n'(E_{\text{inc}}) = d_{s/m} \int dE_{\text{out}} d\Omega_{\text{out}} \frac{\chi(E_{\text{inc}}, E_{\text{out}})}{\chi^m(E_{\text{inc}}, E_{\text{out}})}
\times \frac{a(E_{\text{inc}}, E_{\text{out}}, \theta)}{a^m(E_{\text{inc}}, E_{\text{out}}, \theta)}, \tag{6}
\]

Note that \( c(E_{\text{out}}, \theta) \), \( A_t \), and \( \epsilon_n \) from equation (4) dropped out, while ratios of PFNS and angular distributions between isotope of interest and monitor appear, thus reducing uncertainties associated with these terms. The difference in \( L_n \) between the monitor and isotope of interest is accounted for in the difference in PFNS, angular distribution, and the displacement of the monitor sample compared to the isotope of interest, \( d_{s/m} \). This effect is often measured by performing and analyzing the monitor measurement with various positions of the sample, e.g., in references [19–23]. In the case of induced-fission measurements, fragment detection can depend on \( E_{\text{inc}} \) because of the momentum brought into the system.

### 3 Information needed for evaluations

Sufficient experimental data on \( \bar{\nu} \), a short version encompassing both \( \bar{\nu}_p \) and \( \bar{\nu}_t \), are available for only a few isotopes, and only in restricted incident-neutron energy ranges, to evaluate \( \bar{\nu}_p \) or \( \bar{\nu}_t \) based exclusively on experimental data. A well-known evaluation that provides \( \bar{\nu}_t \)
values based on only experimental data is one of the thermal constants that is part of the Neutron Data Standard (NDS) project [10]. The thermal-constants effort is based on the work of Axton et al. [11]. Similar evaluation efforts were undertaken, e.g., [46,50], and all rely on a detailed analysis of experimental uncertainties including corrections of mean values and uncertainties. Nuclear data of $\overline{\nu}_p$ or $\overline{\nu}_t$ in the resolved resonance range are evaluated based on theory and available experimental data, e.g., undertaken as part of [49], but resonance data for $\overline{\nu}_p$ or $\overline{\nu}_t$ are not given for all isotopes. For instance, they are provided for $^{239}\text{Pu}$ $\overline{\nu}_p$ in ENDF/B-VIII.0 [57], but not for $^{238}\text{Pu}$ and $^{240–242}\text{Pu}$ $\overline{\nu}_p$. The evaluations in the fast range can be based on a Bayesian analysis of differential experimental data (e.g., $^{239}\text{Pu}$ and $^{240}\text{Pu}$ $\overline{\nu}_p$ in ENDF/B-VIII.0), thus following fluctuations observed in the experimental data, or can be based on a simple linear fit to data (e.g., $^{241}\text{Pu}$ and $^{242}\text{Pu}$ $\overline{\nu}_p$ in ENDF/B-VIII.0). Approximating $\overline{\nu}_p$ by a linear function is based on the assumption that the fission yields and total-kinetic energy of the fission fragments do not vary with excitation energy and that the total energy released through fission $\gamma_s$ follows a linear dependence [58]. However, measurements of the total kinetic energy and fission yields as a function of incident-neutron energy show an energy-dependent behavior, see, e.g., [59]. In line with these physics considerations, several $^{235,238}\text{U}$ and $^{239}\text{Pu}$ $\overline{\nu}$ measurements as a function of incident-neutron energy, e.g., [15,17–25,36,38,54], have shown that the experimental data cannot be described well by one linear function. In the absence of differential experimental data, evaluated data are also based on models with systematics of model parameters, as done, e.g., in reference [60].

The incident-neutron energy, $E_{\text{inc}}$, and either $\overline{\nu}_p$ or $\overline{\nu}_t$ are used as a bare-minimum input for the evaluation. If $\overline{\nu}_p$ is evaluated and the measurement is of $\overline{\nu}_t$, the evaluator needs to correct for the delayed component. If the data were measured in ratio to a monitor, it would be desirable if the ratio data were reported. Otherwise, the nuclear data of the monitor observable, often $^{252}\text{Cf(sf)}$ or $^{235}\text{U(n,f)}$ $\overline{\nu}$, or a reference should be provided. It would be desirable to explicitly state what PFNS was used either by reference or model parameters. Given the convolution of the PFNS with many other observables in the analysis of $\overline{\nu}$ measurements in equations (4) and (6), it is difficult to correct with a new PFNS. However, if one knows how close the used PFNS was to current nuclear data, one can estimate potential missing uncertainties due to limited knowledge of the PFNS at the time of the experiment. Partial uncertainties for all uncertainty sources listed in the templates should be provided, if applicable to a particular measurement. The $\overline{\nu}_p$ and $\overline{\nu}_t$ can be measured to high precision. However, even small variations in $\overline{\nu}$ of major actinides can impact the simulated neutrophotomultiplication factor, $k_{\text{eff}}$, of critical assemblies by a substantial amount. For instance, a change of 0.1% in a relevant energy range of $^{239}\text{Pu(n,f)}$ $\overline{\nu}_t$ can lead to a 100-pcm (i.e., a 0.1%) change in $k_{\text{eff}}$ of a Pu assembly, where approximately 210 pcm is the difference between a controlled critical assembly and an accident emitting lethal radiation doses [61]. Hence, reporting complete and realistic uncertainties for $\overline{\nu}_p$ and $\overline{\nu}_t$ measurements is central for realistic application simulations and their bounds. Along the same lines, the experimental set-up (e.g., time the gate is open, size and isotopic composition of the neutron detector, through-tube size, neutron-producing reaction, impurity level), all pertinent corrections (e.g., background, foil thickness, angular distribution of fission fragments, dead-time, impurities, geometry, spurious structures in neutron flux, delayed $\gamma$s, displacement of fission sample, false fissions, French effect) and analysis techniques should be documented in great detail, enabling the evaluator to judge the quality of the measurements and data reduction at a later time.

4 Template

A listing of all relevant uncertainty sources for $\overline{\nu}_p$ and $\overline{\nu}_t$ measurements is provided below, along with the proposed template values. All uncertainties here are given relative to $\overline{\nu}_p$ and $\overline{\nu}_t$ in percent (%) unless otherwise stated. Some uncertainty values seem very small, but one should consider that these measurements were built to reduce correction factors to a minimum and, hence, the associated uncertainties. Table 1 provides a template of uncertainties expected in absolute and ratio liquid-scintillator measurements, while Table 2 gives a template for Mn-bath measurements. Table 2 focuses mostly on uncertainties expected in measuring the neutron-count rate, $c_n$. Several different techniques were employed to measure the fission count rate, $c_f$, with differing uncertainty sources and values for each measurement type. Given that and the low number of $^{252}\text{Cf(sf)}$ $\overline{\nu}$ Mn-bath measurements, not enough statistics were available to give a template for specific uncertainty sources for $c_f$ for each measurement type. Hence, the uncertainty sources entering $\delta c_f$ were not split out in Table 2.

Counting-statistics uncertainties. No counting-statistic uncertainties, $\delta c$, are provided in both templates since these depend strongly on the measurement time, neutron flux (if applicable), detector response, etc., and are, thus, hard to estimate. We recommend rejecting a data set for evaluation purposes if this very basic uncertainty information cannot be, at least, roughly estimated (e.g., one-third of the total uncertainty), given that many $\overline{\nu}_p$ and $\overline{\nu}_t$ measurements are claimed to be of high precision. This high precision needs to be backed up by a reasonably informative uncertainty quantification.

Counting-statistics values depend strongly on the absolute measurement time, fission-detector response, and neutron flux for liquid-scintillator measurements. Hence, $\delta c$ varies significantly across different measurements. For instance, $\delta c$ values of 0.0012% to 3.9% can be found across $^{239}\text{Pu(n,f)}$ $\overline{\nu}$ measurements [12,14,19–23,26,35,38] and 0.08–0.8% can be found in EXFOR across $^{252}\text{Cf(sf)}$ $\overline{\nu}$ measurements [1,4,5,7–9,27,29,30]. The neutron-counting statistics uncertainty, $\delta c_n$, across various $^{252}\text{Cf(sf)}$ $\overline{\nu}$ Mn-bath measurements assume values from 0.1–0.2% [3,5,9,30,45], while the fission-counting uncertainty varies from 0.05–0.2% across references [5,7,30]. However, it
Table 1. Typical uncertainty sources encountered in absolute and ratio liquid-scintillator measurements of $\nu_t$ are listed, along with realistic ranges of estimates that can be assumed if none are provided for a particular measurement. Also, off-diagonal correlation coefficients for each uncertainty source (for the same and different experiments) are roughly estimated. We implicitly assume that the typical tanks have high or similar detector efficiencies of $\sim 80\%$ which is indeed often the case. The correlation functions are defined in reference [62].

<table>
<thead>
<tr>
<th>Unc.</th>
<th>Absolute (%)</th>
<th>Ratio (%)</th>
<th>Cor(Expᵢ)</th>
<th>Cor(Expᵢ,Expⱼ)</th>
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<tbody>
<tr>
<td>$\delta c$</td>
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<td>Must be provided ($\delta c$ &amp; $\delta c^m$)</td>
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<td>None</td>
</tr>
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<td>$\delta c_{DG}$</td>
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<td>Full</td>
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<td>0.5</td>
<td>Gaussian</td>
<td>0.2 for same n source</td>
</tr>
<tr>
<td>$\delta c_{ff}$</td>
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<td>0.22 (high $\alpha$-activity sample)</td>
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<td>0.2</td>
</tr>
<tr>
<td>$\delta c_{FE}$</td>
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<td>–</td>
<td>Gaussian</td>
<td>0.2</td>
</tr>
<tr>
<td>$\delta \omega$</td>
<td>see Table 3</td>
<td>see Table 3</td>
<td>0.9</td>
<td>0.9 (same method &amp; isot.)</td>
</tr>
<tr>
<td>$\delta \tau$</td>
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<td>0.08</td>
<td>Full</td>
<td>Low ($\sim 0.2$)</td>
</tr>
<tr>
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<td>Gaussian</td>
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<td>Full (same $E_{inc}$)</td>
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<td>Full</td>
<td>0.5</td>
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<tr>
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<td>Full</td>
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<tr>
<td>$\delta d$</td>
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<td>Full</td>
<td>0.8–0.9 (not corrected)</td>
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<td>Estimate from similar facilities</td>
<td>Full in</td>
<td>0</td>
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<td>holds that $\delta c_n$ should be provided for a high-precision measurement. The correlation matrix associated with $\delta c$ is usually diagonal for $\nu_p$ and $\nu_t$ measured within the same experiment, and not correlated to other measurements.</td>
<td></td>
<td></td>
<td>E_{inc} space</td>
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</tbody>
</table>

Delayed gamma-ray uncertainties. Delayed $\gamma$s arise mainly as part of the decay of isomeric states with half-lives on the order of 70 $\mu$s for gates that are open less than 500 $\mu$s [63]. Some delayed $\gamma$s from the $\beta$ decay of the fission fragments of another fission event could also be measured, but to a lesser extent. These delayed $\gamma$s are recorded in liquid-scintillator measurements along with those emitted after capturing prompt-fission neutrons in the scintillator material while the neutron-counting gate is open. This uncertainty source does not appear for Mn-bath measurements, as the $\gamma$ rays emitted from the Mn solution are measured in a re-entrant well separated from the source.

The resulting delayed-$\gamma$ background, $c_{DG}$, is corrected in absolute and ratio liquid-scintillator measurements by a combination of measurements and calculations of delayed $\gamma$-ray cascades [1,21–23,38,64]. For instance, a measurement with a low bias, i.e., low $E_\gamma$ threshold, (including delayed $\gamma$s) and a high bias (supposed to press most of the delayed $\gamma$s) gave an estimate of $c_{DG}$ for the experiment of Gwin et al. [21–23]. This measurement was counter-checked with simulations based on expected $\gamma$-ray cascades. Uncertainties, $\delta c_{DG}$, ranging from 0.1–0.2% were found for liquid-scintillator ratio data [21–23,38,64], with a median of 0.12% that is adopted for Table 1. $\delta c_{DG}$ for absolute data was estimated to be 0.1% based on the assessment of these uncertainties for Boldeman, Asplund-Nilsson, and Hopkins data in reference [1]. A full correlation is assumed for $\delta c_{DG}$ for the same and different experiments using the absolute and ratio liquid-scintillator technique, given that most likely the same decay data and assumptions for the simulation are used.

Random-background uncertainties. False-fission and delayed-$\gamma$ background events are not part of the random-background correction, $b$, for liquid-scintillator measurements as these two corrections are treated separately.

Two different types of backgrounds are encountered in ratio liquid-scintillator measurements: (a) neutron-beam-related backgrounds and (b) delayed neutrons. Delayed neutrons can be corrected with nuclear data. Neutron-beam-related background counts can be detected in the neutron counter either with their initial, or slightly degraded, incident-neutron energy, or can thermalize and
Table 2. Typical uncertainty sources encountered in Mn-bath measurements of $\tau_p$ and $\tau_f$ are listed along with realistic ranges of estimates if none are provided for a particular measurement. Also, correlation coefficients for each uncertainty source between different experiments are roughly estimated. No correlation coefficient is given between uncertainties of the same experiment at different incident-neutron energies as Mn-bath measurements are usually undertaken at one incident-neutron energy (thermal) or for spontaneous fission. A medium-range correlation would correspond to correlation coefficients of 0.5–0.8, while “Low” corresponds to approximately 0.2.

<table>
<thead>
<tr>
<th>Unc.</th>
<th>Unc. (%)</th>
<th>Cor(Expj,Expj)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\delta_{cs}$</td>
<td>$\delta_{cs}$</td>
<td>$c_f$</td>
</tr>
<tr>
<td>$\delta_{\Delta E_{mn}}^{252}$ &amp; $\delta_{\Delta E_{mn}}^{235}$</td>
<td>0.29</td>
<td>Full</td>
</tr>
<tr>
<td>$\delta_{\Delta N_{mn}}^{252}$</td>
<td>0.05</td>
<td>Medium</td>
</tr>
<tr>
<td>$\delta(1 + G7s)$</td>
<td>0.09</td>
<td>Full</td>
</tr>
<tr>
<td>$\delta_{\Delta E_{Mn}}^{252}$</td>
<td>0.2</td>
<td>Medium</td>
</tr>
<tr>
<td>$\delta(1 - L_n)$</td>
<td>0.02</td>
<td>Medium</td>
</tr>
<tr>
<td>$\delta(1 - S)$</td>
<td>0.05</td>
<td>Full</td>
</tr>
<tr>
<td>$\delta(1 - O)$</td>
<td>0.1</td>
<td>Full</td>
</tr>
<tr>
<td>Mn-bath impurities</td>
<td>0.1</td>
<td>0</td>
</tr>
<tr>
<td>$\delta_{cf}$</td>
<td>$\delta_{cf}$</td>
<td>$c_f$</td>
</tr>
<tr>
<td>$\delta_{\tau_f}$</td>
<td>0.32</td>
<td>Medium</td>
</tr>
<tr>
<td>$\delta_{\tau}$</td>
<td>0.05</td>
<td>Low</td>
</tr>
<tr>
<td>$\delta_{\omega}$</td>
<td>See Table 3</td>
<td>See Table 1</td>
</tr>
</tbody>
</table>

lead to a random room background. Some neutron-producing reactions can lead to groups of secondary neutrons of lower energy than the anticipated one. Thus, these secondary neutrons induce fission of lower incident-neutron energy than assumed to be measured, biasing $\tau$. All of these neutron beam-related backgrounds would not drop out in ratio measurements relative to $^{252}$Cf(sf) $\tau_f$, as the latter is measured without the beam.

The random, room-return, neutrons, and stray $\gamma$s were measured by opening a gate 200–250 $\mu$s after the main neutron-counting gate was closed, assuming that no more prompt-fission neutrons should be detected by then [15,17,19,20,34,36]. Also, measurements using random gates were employed to determine this part of $b$ [21–23]. The number of thermalized neutrons was often reduced by adequate shielding and also by measurements with and without this shielding [34,36]. Background originating from secondary neutron groups in the beam was usually determined by measuring the energy of the neutron flux directly [15,16,39]. Uncertainties are reported for experiments described in references [12,16,21–23,34,35,38] spanning from 0.08–3.5%, with a median of 0.5% that is suggested as a value for $\delta b$ if nothing else is known about the measurement. A Gaussian correlation is suggested for $\delta b$ across the same experiment, given that $b$ is often measured. Zero correlation is suggested for $\delta b$ across experiments, unless the same neutron-beam type was used (common second-energy groups). In this case, a correlation of 0.2 is suggested.

The delayed and incident-beam neutrons can also lead to $b$ for absolute liquid-scintillator experiments, which might be surprising as most of them are measuring spontaneously-fissioning sources. Even then, the neutron-detector efficiency is often measured with a neutron beam [2,15]. In these two experiments, a combination of shielding and filters reduced this background and measurements with random gates quantified $b$. The size of $b$ is often reduced as a triple coincidence (fission fragments in the fission detector, prompt-fission $\gamma$s in the scintillator, and fission neutrons in a proton-recoil detector) is used to open the gate. In addition to these above-mentioned contributions to $b$, the proton-recoil background may need to be quantified [1]. Asplund-Nilsson et al. addressed this background by pulse-shape discrimination [27]. Uncertainties of 0.1–0.2% are mentioned in reference [1], where we adopt the mean due to lack of statistics.

No $b$ needs to be corrected for Mn-bath measurements of $^{252}$Cf(sf) $\tau_f$, as no neutron beam is used for these measurements. However, if this technique is used for neutron-induced measurements of $\tau_f$, uncertainties due to incident-beam neutrons should be considered.

False-fission uncertainties. False-fission uncertainties, $\delta_{cf}$, are related to a random-coincidence background in ratio and absolute liquid-scintillator measurements that leads to an incorrect opening of the neutron-counting gate. As no gate is opened in Mn-bath measurements, this uncertainty source does not apply.

The background counts, $c_{ff}$, originate from the following effect: the fission detector is fired by $\alpha$ particles or noise and thus opens the neutron-counting gate. The neutron detector then registers background neutrons [21–23,64] assuming they are fission neutrons. This effect is usually measured and calculated. For instance, Gwin et al. [21–23] introduced a mock $\alpha$ pulse into the electronics of their experiment and measured the resulting background neutrons that can then be subtracted from all measured counts. However, one needs to know how often $\alpha$ particles are likely to fire the fission detector compared to the rate of fission. This rate of false-fission events compared to fission ones is usually estimated from the $\alpha$ activity of the sample. The resulting background-count rates can be counter-checked with a beam-off measurement [21–23,64].

The associated uncertainty, $\delta_{cf}$, is larger at low $E_{inc}$, as more false-fission events are registered relative to real ones [21–23]. Uncertainties are given for data of references [15,21–23,35,38] range from 0–5% with a relatively high median of 0.22% that is adopted here for samples with high $\alpha$ activity ($^{239}$Pu), while a value of 0.15% is adopted for lower-activity samples ($^{235,238}$U).

\(^2\) Some $\alpha$-particles can usually be discriminated from fission events on the signal amplitude by putting a threshold on the device. However, a pile-up of highly radioactive samples (e.g., $^{239}$Pu) can pose an issue as the $\alpha$-signal amplitude (not the shape) becomes similar to that of fission fragments. Then, a background measurement with the sample in place but without the beam can help identify the rate of false-fission events as mentioned below.
A medium-range uncertainty with Gaussian correlation is given as part of $c_{ff}$ corrections are measured and the associated uncertainties are random background-counting uncertainties. Some measurements take recourse to nuclear data and assumptions on the $\alpha$-activity of the sample to estimate the relative frequency of gates opened by $\alpha$ from the sample; then a Gaussian correlation is also adequate. These assumptions affect equally all $E_{\text{inc}}$ measured with the same sample. Correlations between $\delta c_{ff}$ of different measurements are assumed to be weak (with correlation coefficient 0.2), given that the only assumption common to both measurements, and thus introducing non-zero correlations, would be the $\alpha$ activity of the isotope in question.

In absolute liquid-scintillator $^{252}\text{Cf}(sf)$ $\bar{\nu}$ measurements, $c_{ff}$ is usually negligible because a coincidence between the fission detector and prompt-fission $\gamma$ rays being detected in the scintillator was required to open the gate [2,8,15]. However, as they also used proton-recoil measurements to open the gate at the same time, this introduced the French effect into their results, which does not appear for ratio liquid-scintillator measurements.

**French effect uncertainties.** The French effect, found by Soleilhac and first reported at the IAEA Consultants meeting held at Studsvik, Sweden, in June 1970 [65], $c_{FE}$, only affects absolute liquid-scintillator measurements, as it originates from using a coincidence of several signals to open the neutron-counting gate. The bias results from a dependence on registering fission fragments in the fission chamber, scintillator pulses from prompt-fission $\gamma$s, and proton recoils on the number of neutrons detected per gate [65]. The physics mechanism behind this effect is not fully understood and the magnitude depends on the set-up and so can be difficult to estimate retrospectively, but for new measurements can be experimentally assessed. It affects the measurements of [1,2,8,15,27]. Reported uncertainties in the associated literature, EXFOR, and in references [1,46] cite uncertainties from 0.1–0.48%. The effect is measured and depends strongly on the set-up used. Boldeman states in reference [1] that this effect is usually small (except for the data of Ref. [27]), hence, a small uncertainty of 0.1% is recommended with a Gaussian correlation and low correlation between experiments.

**Impurity uncertainties.** Sample-impurity uncertainties, $\delta \omega$, apply to all measurement types. $\delta \omega$ can be effectively controlled by employing high-purity samples such as for instance used in [12,21–23,34,38] (they utilized $^{239}\text{Pu}$ samples with purity close to or larger than 99.9%). Spontaneously fissioning contaminants and their contribution to $\bar{\nu}_p$ and $\bar{\nu}_t$ were determined in incident-neutron energy-dependent $\bar{\nu}$ measurements [21–23,35] with a beam-off measurement. Non-spontaneously fissioning contaminants can be measured (e.g., by mass-spectroscopy [66]) and their contribution to $\bar{\nu}$ can then be corrected by using nuclear data [17,21–23,38]. The value of $\delta \omega$ depends thus on the accuracy of the contamination measurement, the level of impurity, and the nuclear data used. For $^{239}\text{Pu}(n,f)$ $\bar{\nu}$, $\delta \omega$ values between 0.03 and 0.1% can be found [17,21–23,35] with some experiments giving even lower values of 0.001% [12,38]. $\delta \omega$ values spanning from 0.02–0.1% can be found for $^{252}\text{Cf}(sf)$ $\bar{\nu}$ measurements. Also, EXFOR files were searched for $^{232}\text{Th}$, $^{237}\text{Np}$, $^{233,235,238}\text{U}$, $^{240,241}\text{Pu}$, and $^{241}\text{Am}$ $\bar{\nu}_p$ to obtain recommended values of $\delta \omega$ per isotope in Table 3. Unfortunately, $\delta \omega$ values were found for only a few isotopes. Due to the scarce statistics (i.e., a small number of documented cases), only a value of $\delta \omega$ could be given per isotope, not depending on impurity level or incident-neutron energy. Both factors play a role: for instance, a $^{238}\text{U}$ impurity in a $^{253}\text{U}$ sample could bias $\bar{\nu}$ in the fast range, but is negligible below the fission threshold of $^{238}\text{U}$. Also, as said above, $\delta \omega$ can be driven to be negligible by using very pure samples. The issue of isotopic impurity is an important one for samples that are not highly enriched, which could be the case in the earliest measurements of $\bar{\nu}$. The $\delta \omega$ values in Table 3 are recommended to be used if no information is given on the impurity levels in the sample. Like all other uncertainty values, $\delta \omega$ is understood to be given relative to $\bar{\nu}$.

It is interesting to note that the largest median value of $\delta \omega$ appears for $^{252}\text{Cf}(sf)$ $\bar{\nu}$ despite it being the standard. The reason for this finding might be due to considerable alpha pile-up in the impurity-level determination. Also, a significant $^{250}\text{Cf}$ component is inevitable with fresh sources having a mass fraction of ca. 10/85 compared to $^{252}\text{Cf}(sf)$ which increases with age and becomes prohibitively large for samples older than roughly 15 years [67]. For fresh sources, the $\delta \omega$ related to the $^{250}\text{Cf}$ content should be small.

It should also be mentioned that some small adjustments to older data might be used as $\bar{\nu}$ and half-lives of the impurities might be different enough to impact the estimate of $\omega$ correction factors. It is expected that this does not apply to $^{252}\text{Cf}$ sample as the spontaneous-fission branching ratio is very high.

A larger correlation coefficient is assumed for correlations between $\delta \omega$ of the same experiment as usually the same sample is used at all $E_{\text{inc}}$ and nuclear data used for the correction is strongly correlated. If the same method was employed across different experiments and the same isotope was measured, a strong correlation can be assumed. If a different isotope was measured, the correlation is low because different nuclear data were used for the correction of impurities in the sample.

<table>
<thead>
<tr>
<th>Isotope</th>
<th>$\delta \omega$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{233}\text{U}$</td>
<td>0.05</td>
</tr>
<tr>
<td>$^{235}\text{U}$</td>
<td>0.05</td>
</tr>
<tr>
<td>$^{239}\text{Pu}$</td>
<td>0.07</td>
</tr>
<tr>
<td>$^{241}\text{Pu}$</td>
<td>0.1</td>
</tr>
<tr>
<td>$^{252}\text{Cf}$</td>
<td>0.1</td>
</tr>
</tbody>
</table>
Dead-time uncertainties. Dead-time uncertainties, $\delta\tau$, apply to all measurement types. When neutron pulses pile up during the experiment, the system is dead to measuring other neutron events. Similarly, when $\alpha$ particles pile up, the fission-detection system (for $c_f$ in Mn-bath measurements) and for opening the gate in liquid-scintillator measurements) can be dead. The $\overline{\tau}_p$ and $\overline{\tau}_t$ need to be corrected for this dead-time effect. Descriptions of numerical correction procedures can be found in references [17, 21–23, 36, 38], while it was minimized in reference [36] by carefully selecting detectors and shielding to reduce dead time induced by background counts. For ratio measurements, the dead-time uncertainty reduces to the ratio of dead-time for the isotope of interest and the monitor sample. Related dead-time uncertainties in ratio-liquid-scintillator measurements, $\delta\tau$, for $\tau$ span values of 0.02–0.5% with a median of 0.08% that is recommended in Table 1. A median of 0.1% is found for absolute liquid-scintillator measurements, while a total $\delta\tau$ (for fission and neutron counting) of 0.05% is given for Mn-bath measurements.

This uncertainty source would be strongly correlated for the same experiment as functional forms are used for correcting $\tau$ while a weak correlation is assumed between measurements due to reliance on different equipment and different background-reduction techniques.

Uncertainty in detecting capture-$\gamma$ rays. The uncertainty in the detection of $\gamma$ rays resulting from neutron-capture in the scintillator, $\delta\varepsilon_{\gamma}$, are considered in $\delta\varepsilon_c$, as these two uncertainties are usually given in a combined manner. The reason for this is that usually $\varepsilon_{\gamma}$ is determined by measuring $\varepsilon_n$ at a specific $E_{inc}$ and angle. $\varepsilon_{\gamma}$ is then extracted by knowing $\varepsilon_c$ and $L_n$ from MC simulations and having a defined anisotropy introduced by the through-tube, $A_t$. Hence, there is a strong correlation between $\delta\varepsilon_{\gamma}$ and $\delta\varepsilon_c$. $\varepsilon_{\gamma}$ only appears for absolute liquid-scintillator measurements and drops out in ratio measurements.

Uncertainty of neutron-capture efficiency. The uncertainty related to the neutron-capture efficiency of the Gd or Cd-loaded scintillator material, $\delta\varepsilon_c$, only appears for absolute liquid-scintillator measurements while $\varepsilon_c$ cancels in ratio measurements in equation (6). The term $\varepsilon_c$ is often calculated with neutron-transport codes. To this end, the codes rely on nuclear data of the capturing material (e.g., hydrogen, gadolinium, cadmium) dependent on, both, energy and angle [1, 15]. $\varepsilon_c$ can also be measured in combination with $\varepsilon_{\gamma}$ [2, 15, 27]. Even if $\varepsilon_c$ is determined by simulations, usually validation measurements are performed [1, 15]. The neutron-detection efficiency is then measured at a specific $E_{inc}$ and angle. However, one still needs MC simulations for $L_n$ and nuclear data for $\chi$ and $a$ to obtain $\varepsilon_n$ [2, 15, 27].

Uncertainties of 0.1–0.2% are reported for the data of Boldeman and Asplund-Nilsson et al. in reference [1], while 0.388% is given in the EXFOR entry of [8]. Again, the median of 0.2% is assumed which applies to the combination of $\delta\varepsilon_c$ and $\delta\varepsilon_{\gamma}$. A Gaussian correlation is assumed as both measurements and nuclear data are used for determining $\varepsilon_c\varepsilon_{\gamma}$.

PFNS uncertainty. The PFNS, $\chi$, enters as an important factor in the analysis of each measurement technique presented here. It enters in equations (4) and (6) for absolute and ratio liquid-scintillator. $\chi$ does not appear explicitly in equation (2). However, it has to be considered implicitly when simulating the terms $(1 - L_n)$ (neutrons leaked), $(1 - S)$ (neutrons recaptured by material in the cavity) and $(1 - O)$ (neutrons lost by charged-particle reactions with oxygen or sulfur) via neutron-transport codes as one needs to know the energy distribution of neutrons being released as part of the fission process. To this end, representative values need to be selected for $\chi$, either from nuclear-data libraries or by assuming model-parameter values (e.g., Maxwellian temperature or Watt parameter values) entering a functional form. However, given the convoluted nature of $\chi$ with other observables to yield $\varepsilon_n$ or $(1 - L_n)(1 - S)(1 - O)$ in equation (2), it is neither straightforward to correct with more recent $\chi$ values nor to estimate $\delta\chi$ based on our present-day understanding of $\chi$. One would need to know all factors entering equations (4), (6), and $(1 - L_n)(1 - S)(1 - O)$. In addition, the absolute value of $\delta\chi$ depends not only on how well the chosen numerical values for $\chi$ reproduce actual physics properties, but also on other factors appearing in equations (4), (6), and (2). For instance, if one employs a highly efficient neutron detector, $\delta\chi$ relative to $\tau$ will be smaller than a less efficient one for the same values of $\chi$.

In absolute $\tau_p$ and $\tau_t$ measurements, $\chi$ enters absolutely as well, while the ratio of $\chi$ of the isotope of interest and the monitor isotope is considered in ratio measurements. Quantifying only the difference between two prompt-fission neutron spectra leads usually to smaller $\delta\chi$ than if $\chi$ has to be given absolutely. Conversely, $\delta\chi$ values of absolute $^{252}\text{Cf}(sf)$ $\tau$ liquid-scintillator measurements are overall larger than those found for ratio measurements of $^{239}\text{Pu}(n,f)$ $\tau_p$: $\delta\chi$s found for the $^{252}\text{Cf}(sf)$ $\tau$ data of [1, 2, 8, 27, 29] assume values in the range of 0.04–0.5% with a median of 0.23% that is used for the template for absolute liquid-scintillator measurements. Values ranging from 0.01–1% are found across several $^{239}\text{Pu}(n,f)$ $\tau_p$ measurements [12, 16, 17, 19–23, 26, 38] with a median of 0.16%. An additional complication with the PFNS of ratio measurements poses the incident-neutron energy dependence of $\chi$. If the monitor value is measured at one defined incident-neutron energy, as is the case for using the monitor $^{252}\text{Cf}(sf)$ $\tau$ ($E_{inc} = 0$ as spontaneously fissioning), the overall size of the difference between $\chi$ of interest and monitor depends on $E_{inc}$. The $\chi$ of an isotope of interest usually differs from the $\chi$ of $^{252}\text{Cf}(sf)$. Even greater differences are expected for neutron-induced fission near the second-chance-fission threshold. At these $E_{inc}$, distinct structures can be observed in $\chi$ for $E_{out}$ around a few hundred keV across many isotopes, see e.g., references [53, 68–71]. While $\chi$ also hardens with $E_{inc}$ for the isotope of interest, the mean energy might actually be closer to that of the $\chi$ of $^{252}\text{Cf}(sf)$ reducing the difference between the two $\chi$ which is counter-balanced by structures such as ≥2nd chance fission and the pre-equilibrium component for the $\chi$ of interest. Hence, the median of 0.16% is suggested for ratio liquid-scintillator measurements in Table 1 for all $E_{inc}$ except for the second-chance-fission.
threshold, where 0.5% is suggested. It should be mentioned that these values are isotope-dependent. Some isotopes, e.g., $^{238}$U, show larger second-chance fission structures than others leading to increased $\delta \chi$ for the same neutron-detection efficiency. No $\delta \chi$ is given in Table 2 for Mn-bath measurements given the convoluted nature of $\chi$ with the term $(1-L_n)(1-S)(1-O)$. Instead, uncertainties are provided for the individual terms—$(1-L_n)$, $(1-S)$, and $(1-O)$—which contain $\delta \chi$.

If the same values for $\chi/\chi^m$ are used for one (liquid-scintillator) experiment, $\delta \chi$ is fully correlated. If an energy-dependent $\chi/\chi^m$ is used, a Gaussian correlation is recommended. The reasoning behind this choice is that evaluations of $\chi$ are usually based on models leading to strong correlations between $\chi$ at different $E_{inc}$. If the same or similar assumptions on $\chi$ are made across different experiments at the same $E_{inc}$, $\delta \chi$ is fully correlated across these experiments. If the same $\chi$ is chosen across experiments at different $E_{inc}$, a Gaussian correlation would be recommended in line with the discussion above. It would be best practice for experimental results to be reported along with sufficient detail to allow the PFNS uncertainty to be re-calculated at a later time or when a new evaluation of $\chi$ becomes available.

**Neutron-leakage uncertainty.** The leakage of neutrons, $L_n$, out of the through-tube of liquid-scintillator measurements is convoluted in equation (4) with the PFNS, $\chi$, the angular distribution of prompt-fission neutrons, $a$, and the geometrical anisotropy of the experimental configuration $A_i$. All these factors, $\chi$, $a$, and $A_i$, enter in MC simulations of $L_n$ [1] that in turn enter the determination of $\varepsilon_A$. In ratio measurements, the geometrical anisotropy for monitor and isotope of interest should be nearly the same except for the effect of the displacement of the sample with uncertainties considered in $\delta L_n/\chi$. The difference in neutron leakage can be traced back to the difference in $\chi$ and $a$ for the monitor and isotope in question. Hence, $L_n$ does not appear in equation (6) and $\delta L_n$ is not applicable.

$\delta L_n$ is also given rarely for absolute-liquid scintillator measurements as it is either considered in $\delta A_i$ as in reference [2] or in the neutron-detector efficiency [15]. In a review by Boldeman [1] a value of 0.2% was given for his own $^{252}$Cf($sf$) $\pi_p$ measurement and 0.3% for the measurement of Asplund-Nilsson [27], while in the original publication, no $\delta L_n$ is reported separately and $L_n$ is discussed as part of $\delta \varepsilon_A$. A value of 0.2% is recommended for absolute liquid-scintillator measurements for the sake of completeness if not considered as part of other uncertainty sources. A full correlation is assumed for $\delta L_n$ of the same experiment as $L_n$ is usually calculated with MC codes relying on geometrical assumptions and nuclear data, while a strong correlation factor of 0.5 is assumed between experiments.

The leakage of neutrons out of the Mn bath, $(1-L_n)$, is considered explicitly in equation (2), and associated uncertainties, $\delta(1-L_n)$, ranging from 0.01–0.04% (about 10% of the correction factor itself) are provided by references [3,5,6,30,45] with a more recent MC study [47] confirming simulated uncertainties as low as 0.01%. We recommend a slightly larger median value of 0.02% (still negligibly small) which reflects the fact that most measurements were undertaken decades ago where neutron-transport or diffusion calculations were less accurate and precise than today’s counterparts [3,45]. These simulations rely on assuming a $\chi$ of the source as well as knowing the concentration of the Mn-bath solution, geometry of the set-up itself, and shielding [3]. $(1-L_n)$ can also be measured [5,30,45], e.g., by using two baths of varying size. Given that $\delta(1-L_n)$ relies on very similar nuclear data to be calculated but also assumptions that are unique to an experiment campaign (geometry) and explicit measurements [45], a medium-range correlation is assumed between $\delta(1-L_n)$ of different experiments.

**Uncertainty of angular distribution of fission neutrons.** A correction for the angular distribution of fission neutrons, $a$, is needed for $\pi$ measurements in the fast range. As measurements in the fast range are usually undertaken with the liquid-scintillator technique, $a$ appears here for these types of measurements. The incident-neutron energy beam leads to a preferred direction of the fission fragments with respect to the angle of incident neutrons, see e.g., [66], because of (a) the forward boost of fission fragments that becomes significant for $E_{inc} > 10$ MeV, and (b) the inherent anisotropy of fission fragments. The outgoing prompt-fission neutrons tend to be emitted into the direction of the moving fragments in the reference system of the laboratory [38] leading to a non-negligible angular dependence of fission neutrons. The resulting angular distribution of prompt-fission neutrons is both dependent on the incident-neutron energy and the isotope [72]. At thermal, the prompt-fission neutrons can be assumed to be emitted isotropically on average, while an angular distribution of fission neutrons becomes noticeable for incident-neutron energies as low as 1 MeV for $^{235,238}$U and $^{239}$Pu [72] and is isotropic only below approximately 200 keV. For spontaneous fission, the fission fragments are emitted isotropically and so are the neutrons. Hence, this effect does not appear if all of the fissions of $^{252}$Cf are detected. If only some part of the angular range of the fragments of $^{252}$Cf($sf$) is detected, as it will be for any fragment detector, then the associated neutrons will not be isotropic, but the effect will be less pronounced compared to neutron-induced fission in the fast range. Due to the same reasoning, $a/a^m$ does not cancel in ratio measurements where the isotope of interest is measured relative to $^{252}$Cf($sf$) $\pi$.

The correction for this effect is usually calculated by taking into account the geometry of the experiment (especially the through-tube) and assuming an angular distribution of fission fragments [17,21–23,38,66]. It is strongly intertwined with the neutron-leakage, $L_n$, and asymmetry of the measurement configuration, $A_i$. If the neutron detector fully enclosed the fission detector (i.e., a $4\pi$ environment), no correction for $a$ would apply. However, the through-tube is needed for housing the detector and giving way to the neutron beam. This reduces the geometry to be less than $4\pi$. $\delta a$ is one of the least understood uncertainty sources for $\pi$, given that the angular distribution of fission neutrons is not very well-known today and even less so in the mid-60s to 80s, when many $\pi$ measurements were...
performed. Uncertainty values related to the angular distribution of fission neutrons, $\delta a$, range for $^{235}\text{Pu}(n,f)$ measurements from 0.002–0.3% [12,16,17,21–23,35,36,38]. The lowest uncertainties (0.002–0.1%) are reported for data below 2 MeV [12,17,35,38], while values of approximately 0.3% are reported for data at higher $E_{\text{inc}}$ [21–23,36]. Exceptions are data of [38,54] that have low $\delta a$ values despite being measured to $E_{\text{inc}}>2$ MeV, where the contribution of $a$ is non-negligible [72]. In the template, 0.1% is proposed for $\delta a$ from 0.2 up to 1 MeV going up to 0.3% above 2 MeV and 0.5% at the second-chance-fission threshold and at $E_{\text{inc}}>10$ MeV. $\delta a$ is assumed to be zero for $E_{\text{inc}}<0.2$ MeV. At the second-chance-fission threshold, the angular distribution of FF is very large in general, leading to a larger $a$. Also, the forward boost of fission fragments induced by incident neutrons should be considered for $E_{\text{inc}}>10$ MeV, leading to overall larger uncertainties in this energy range. It should be stressed that $\delta a$ and $a$ are not well understood and both values should be carefully examined within new experiments.

Off-diagonal correlation coefficients between 0.8–0.9 are assumed between $\delta a$s at different $E_{\text{inc}}$ for the same experiment, as nuclear data and functional relationships are usually used for corrections. A correlation of 0.6 is assumed between different experiments, as the same functional relationships would have been used and similar nuclear data, but the accuracy of geometrical considerations differs between measurements. Zero uncertainty is assumed for absolute measurements that are usually taken of spontaneously fissioning systems, where $a$ related to the detector is not applicable.

Asymmetry uncertainty. The asymmetry uncertainty, $\delta A_t$, only occurs for absolute liquid-scintillator measurements and has to be considered for a correct calculation of neutron leakage. For only one $^{252}\text{Cf}(sf)\,\pi$ measurement, i.e., [2], a separate uncertainty is given for $\delta A_t$ of 0.03%. Otherwise, $\delta A_t$ is part of either $\delta L_n$ or $\delta a$. Hence, we do not have an entry for $\delta A_t$ in Table 1, considering it part of $\delta L_n$ or $\delta a$. $\delta A_t$ drops out in ratio liquid-scintillator measurements, as the same asymmetry applies to the isotope of interest and monitor measurements. Any difference in $\delta A_t$ stemming from the variability in the location of the sample of interest and the monitor one is considered in $\delta d_{s/m}$. It can be estimated experimentally by shifting the sample.

Monitor uncertainty. The monitor uncertainty, $\delta \pi^m$, is usually straightforward to estimate if one knows what data were used for $\pi^m$ in equation (6) or if ratio data are explicitly provided. The latter case is preferred, as it gives evaluators the possibility to use the currently best available nuclear data for $\pi^m$. Usually, $^{252}\text{Cf}(sf)\,\pi_t$ is used as monitor. The current value recommended by the NDS project is 3.7637 ± 0.42 % [10]. This uncertainty only applies to ratio measurements.

If the same monitor applies to all data across incident-neutron energies, as is the case for measurements relative to $^{252}\text{Cf}(sf)\,\pi_t$, this would lead to a fully correlated $\delta \pi^m$ across one experiment. The same nuclear data should be used to represent the same a monitor observable throughout an evaluation, leading to a full correlation of $\delta \pi^m$ across all experiments measured relative to the same monitor observable.

Uncertainty due to thickness of sample. The uncertainty due to the thickness of the sample, $\delta d$, only appears for measurements that do not employ a point-source sample. Usually, small and thin samples are used for $^{252}\text{Cf}$ measurements. Hence, a template value is only given for ratio measurements. This correction factor applies because fission fragments do not leave the thick foil or lose too much of their kinetic energy before escaping the foil. Thus, they fail to trigger the fission detector that opens the gate to measure neutrons. The size of $d$ depends on the thickness of the sample, but also on how the sample was manufactured. If the sample material is sprayed or evaporated onto a foil, the resulting samples are usually the most uniform ones. Thus fewer fission fragments are lost than when a sample of the same average thickness is electroplated or, in the worst case, rolled or painted. This effect can be counter-balanced by a fission detector with good efficiency that is also able to register fission fragments of low kinetic energy.

This effect is discussed in great detail in references [21,23,56], where reference [56] provides a brief overview on previous work. Correction factors for different thicknesses were obtained in reference [56] by measuring a $^{252}\text{Cf}$ point source with increasing layers of lead. The results show that $d$ is linear up to a thickness of 2500 µg/cm² (correction factors are provided in Table I of this reference). Gwin [21–23] measured $\pi$ with two samples of different thicknesses and simulated the effect. These investigations led to an uncertainty of 0.05%, which is a lower bound, as Gwin employed a thin sample (100 µg/cm²), thus minimizing the effect of $d$. An uncertainty of 0.1% is assumed for a high fission-detection efficiency and a sample thicker than that of Gwin. $\delta d$ can rise to 0.3% for thick samples (700 µg/cm²).

A full correlation matrix is assumed for $\delta d$ of each individual experiment as $d$ is a multiplicative correction factor unless samples of different thicknesses were employed at different $E_{\text{inc}}$. It is assumed to be strongly correlated across different measurements, given that this effect was not frequently corrected. However, if an independent measurement was undertaken for an individual experimental effort, no correlation should apply.

Sample-displacement uncertainty. The uncertainty due to the displacement of the sample, $\delta d_{s/m}$, appears only for ratio measurements, because it accounts for possible biases introduced in the measured quantity by small deviations between the position of isotope of interest and monitor samples. This displacement would lead to a difference in the neutron leakage between the sample and monitor, and thus lead to a bias in $\pi$. The possible bias was quantified in references [12,21–23,38] for $^{239}\text{Pu}(n,f)\,\pi$ with very low uncertainty values $\delta d_{s/m}$ of 0.01–0.05% uncertainties. Gwin et al. [21–23] obtained these values by first determining a possible bias by measuring the $^{252}\text{Cf}$ sample in different positions in the through-tube. The authors then
assumed one-third to one-fifth of the measured correction as uncertainty. The upper bound of 0.05% is adopted for the template as a conservative estimate if this uncertainty source was not quantified.

As $\delta s/m$ is an energy-independent multiplicative correction factor, a full correlation is assumed for $\delta s/m$ across data of the same experiment. A zero correlation between $\delta s/m$ of different experiments is assumed, as a displacement of the sample would be random between these experiments, thus leading to an independent bias across measurements.

**Energy uncertainty.** The incident-neutron energy uncertainty, $\delta E_{\text{inc}}$, or resolution, $\Delta E_{\text{inc}}$, depends on the neutron source which can be a white or mono-energetic neutron source. For TOF measurements, it depends on the TOF-path length, energy, and uncertainty in resolving the exact timing of the fission pulse, while secondary-neutron groups can lead to a smearing of $E_{\text{inc}}$ in mono-energetic measurements. Given that, it would be best to estimate $\delta E_{\text{inc}}$, if missing, based on existing uncertainty values from the same or similar facilities at the same $E_{\text{inc}}$ rather than assuming one blanket value. To support this statement, it should be mentioned that $\delta E_{\text{inc}}$ ranging from 0.034% to 27.5% are reported across [12–26,34–38].

A full correlation is assumed for the $\delta E_{\text{inc}}$ of the same measurement in energy space, while no correlation is assumed between measurements, as the defining factors for uncertainties vary randomly between measurements.

Uncertainties stemming from those of thermalized hydrogen and sulfur cross sections as ratios to manganese. The fraction of neutrons captured by Mn in the Mn-baths depends on the ratio of cross sections of neutron-capture in Mn relative to that of those captured in hydrogen and sulfur at thermal energies, $\sigma_{\text{H}}/\sigma_{\text{Mn}}$ and $\sigma_{\text{S}}/\sigma_{\text{Mn}}$, respectively. These cross-sections are usually represented by nuclear data and their uncertainties, $\delta\sigma_{\text{H,Mn}}$ and $\delta\sigma_{\text{S,Mn}}$, are propagated to uncertainties in $\chi$ [3,5]. The ratio can also be determined by varying the concentration of, e.g., hydrogen ($N_{\text{H}}/N_{\text{Mn}}$) in the solution [30,45]. $\delta\sigma_{\text{H,Mn}}$ and $\delta\sigma_{\text{S,Mn}}$ are usually reported in a combined manner (both contain $\sigma_{\text{Mn}}$), with uncertainties range from 0.2–0.29% [3,30,45]. Hence, additional measurements with varying $N_{\text{H}}/N_{\text{Mn}}$ were performed to determine $\sigma_{\text{H,Mn}}/\sigma_{\text{Mn}}$ [45]. A full correlation is assumed between $\delta\sigma_{\text{H,Mn}}$ and $\delta\sigma_{\text{S,Mn}}$ of different experiments, given that usually the same or similar nuclear data were employed. This uncertainty source naturally only applies to Mn-bath measurements.

Uncertainties due to number of hydrogen versus manganese atoms. The uncertainties in the ratio of the number of hydrogen vs Mn atoms, $\delta N_{\text{H}}/N_{\text{Mn}}$, can be determined very precisely by various methods (volumetric, gravimetric, and solution-density determination) [3,45]. It is rarely reported and Axton states in his review of Mn-bath measurements [45] that it is negligibly small, with a reproducibility uncertainty of 0.05% which is adopted in Table 2. A medium-range correlation is recommended between $\delta N_{\text{H}}/N_{\text{Mn}}$ of different experiments, given that similar methods are used to determine $N_{\text{H}}/N_{\text{Mn}}$ across experiments.

Uncertainties due to resonance capture of neutrons in manganese. The uncertainties in the resonance-capture term of neutrons in Mn, $\delta(1 + G7S)$, are usually given in a combined manner, given that all three parameters are used to derive the effective cross-section. This factor is either obtained by theoretical or MC calculations, depending on the Mn-activation integral and nuclear data [5,30,45]. Values for $\delta(1 + G7S)$ of 0.05–0.1% are reported for references [5,30,45] and the median of 0.09% is adopted in Table 2. Given that usually recourse to the same assumptions and data are taken, a full correlation is assumed between $\delta(1 + G7S)$ of different experiments. This uncertainty source naturally only applies to Mn-bath measurements.

Mn-bath efficiency uncertainties. Calibrating the Mn bath is usually undertaken with respect to well-known (standard) sources and by cross-comparison of various baths and sources in the international community [3,7,30,45]. Axton highlights in reference [45] that the efficiency, $\varepsilon_{\text{Mn}}$, derived from this calibration will deteriorate invariably over time and recommends repetition of calibrations.

The associated uncertainty, $\delta\varepsilon_{\text{Mn}}$, is a major uncertainty source for Mn-bath measurements of $\chi$, with values ranging from 0.11–0.3% across [3,5,6,30,45], where the median, 0.2%, is recommended for the template. Medium-sized correlations are suggested for $\delta\varepsilon_{\text{Mn}}$ between different measurements, given that common standards are used as well as cross-comparisons are undertaken across experiments.

Uncertainties due to neutrons captured by cavity material. In Mn-bath measurements, one has to account for neutrons lost as they enter the sample cavity or shell which creates the air space at the center of the solution and are recaptured by either the source or materials present in the cavity. The associated term, $(1 − S)$ in equation (2), is usually simulated by MC neutron-transport or diffusion codes, taking into account the concentration of the Mn solution, the PFNS, $\chi$, the geometry of the Mn bath, and diameter, thickness, and materials in the cavity [3,30,45,47]. The flux that is then calculated at the cavity boundary can be validated by measurements of this flux using various foils in the cavity. For instance, gold foils can lead to a smearing of $\chi$ and calibration will deteriorate invariably over time and recommends repetition of calibrations.
Uncertainties due to neutrons captured by oxygen and sulfur. Fast neutrons are captured in Mn baths by oxygen and sulfur in (n,α) and (n,p) reactions. The associated term, (1−O), in equation (2) is usually simulated with MC codes that track the position of neutrons from the source leading to (n,α) and (n,p) reactions [3,5,30,45,47]. To this end, one needs to know the outgoing-neutron energy distribution of prompt neutrons, χ, the concentration of the bath and the geometry of the set-up [45,47]. Validating experiments can be performed using known sources as described in references [7,45].

Associated uncertainties δ(1−O) of 0.06–0.1% are found across references [3,5,6,30,45], with a median of 0.1% that is recommended in Table 2. A recent report [47] obtained a value of 0.07% using modern codes and nuclear data. Also, a recent study by [73] showed that newly evaluated data (e.g., ENDF/B-VIII.1 cross sections which are 30–50% lower than those of ENDF/B-VIII.0) can have a non-negligible impact, for instance, on the 252Cf(sf) τ_f data of Spiegel et al. A full correlation is assumed for δ(1−O) between different experiments, as the underlying simulations usually depend on the same nuclear data and a similar χ.

Mn-bath impurity uncertainties. Impurities in the Mn-bath solution can lead to thermalized neutrons being captured in additional reactions than discussed above [5,45]. The associated uncertainty depends on the isotopic composition and abundance of the impurities. Aleksandrov et al. estimated that 0.1% of cα are recorded from events where neutrons are captured by impurities and gave this as a bounding uncertainty. This value is adopted in Table 2 and coincides with the uncertainty reported for [6]. It should be mentioned that this uncertainty estimate is highly dependent on the actual contamination level in the bath, and, hence, this uncertainty should only be used if no other information can be derived from other sources. One should be able to study this effect by gamma spectroscopy and half-life studies as well as losses due to competitive capture compared to Mn. Zero correlation is assumed for Mn-bath impurity uncertainties of different experiments, as two distinct baths are expected to have different impurities necessitating dissimilar corrections.

Fission-rate uncertainties. The fission-rate uncertainties, δc_f, encompass several uncertainty sources appearing for measuring the fission rate, c_f. As mentioned above, several techniques were employed to determine c_f in various 252Cf(sf) τ_f measurements: solid-angle counting of fission fragments using Si-surface barrier detectors [9,30], fission-fission coincidence counting [3] or neutron-fission coincidence counting [7]. We have for each measurement type too few individual measurements available to reasonably split out uncertainty sources and generate a template for each of those measurement techniques. If one compares combined uncertainties of δc_f across the measurements of [3,5,7,9,30], one ends up with a median value of 0.32%, which is recommended in Table 2. This value should be seen as a guide to compare if given total uncertainties related to δc_f are realistic, and as a rough estimate if nothing else is given. However, this is a major uncertainty source, and if it is not provided for a particular data set, we would recommend not to include the data in the evaluation, unless there are no other measurements of the observable available. Usually, δc_f contains counting statistics uncertainties (0.05–0.2% [5,7,30]), solid-angle uncertainties (0.05–0.6% [5,7,30]), back-scattering from sample (0.1–0.5% [6,30]), self-transfer of 252Cf onto the fission chamber (0.05%–0.4% [5,7,30]) and the half-life of the isotope in question (0.01–0.15% [5–7]). A medium-range correlation of δc_f between different experiments is assumed in Table II of [55], given its composite nature encompassing purely random as well as strongly correlated uncertainty sources.

5 Summary

Nuclear data measurements of average prompt- and total-fission neutron multiplicities fall into a few highly developed types. For each type, the sources of uncertainty have been identified and quantified by various practitioners over many years. Individual experiments, especially older ones, are not always described in as much detail on all individual uncertainty sources or corrections as needed for the nuclear-data evaluation process. Nonetheless, it is expected that measurements of a given type will be impacted by these known inaccuracies or missing uncertainties, whether reported or not. In the present work, templates of expected measurement uncertainties were established for average prompt- and total-fission neutron multiplicities. These templates list expected uncertainty sources for the manganese-bath, liquid-scintillator absolute, and ratio measurement types. They also give ranges of uncertainties for most sources and estimates of correlation coefficients between uncertainties of each source in case that some of them were not reported for a particular measurement and cannot be estimated in any other way. The uncertainty values were estimated conservatively, based on information found for 252Cf, 235,238U, and 239Pu multiplicity measurements in their literature or respective EXFOR entries, as well as on expert judgment from experimenters. In addition to these templates, a section lists what information on the experiments is needed to include its data faithfully in the evaluation process. Future experiments need to be carefully planned, executed, and reported in order to improve nuclear data in this area; many of the corrections can be assessed experimentally and confirmed by simulation as a matter of good practice. The templates presented here serve as a reminder of what uncertainty sources and input for the evaluation should be provided.

Conflict of interests

The authors declare that they have no competing interests to report.
Funding
We gratefully acknowledge the partial support of the Advanced Simulation and Computing program at LANL. This work was supported by the US Department of Energy through the Los Alamos National Laboratory. Los Alamos National Laboratory is operated by Triad National Security, LLC, for the National Nuclear Security Administration of the US Department of Energy under Contract No. 89233218CNA000001.

Data availability statement
The data that were created associated with this manuscript are all within its main text and tables.

Author contribution statement
DN wrote the original draft, AC and SC contributed text to that. All authors reviewed and proposed edits to the manuscripts. All authors were involved in investigations and data curation associated with the article and discussions on uncertainty quantification for $\mathcal{P}$.

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