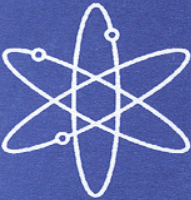


Sensitivity Analysis Applied to the Validation of the ^{10}B Capture Reaction in Nuclear Fuel Casks



Oak Ridge National Laboratory

**U.S. Nuclear Regulatory Commission
Office of Nuclear Material Safety and Safeguards
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Sensitivity Analysis Applied to the Validation of the ^{10}B Capture Reaction in Nuclear Fuel Casks

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ABSTRACT

Boron has commonly been used in nuclear fuel casks to ensure a sufficient margin of subcriticality. The amount of boron used in most casks far exceeds the amount of boron present in any of the available benchmark experiments. Such heavy loadings of boron in the casks may result in considerable spectral differences as compared to the benchmarks, resulting in boron sensitivities that are very different from those of the benchmarks. Before the calculations to determine the nuclear safety margin for various fuel loadings are deemed acceptable, as part of the safety basis, the computer code and cross sections must be validated against experimental benchmarks that cover the area of applicability of the proposed cask design. Therefore, this study was performed to determine if these available benchmarks can be used to validate a criticality code and neutron cross sections for the fuel casks. The sensitivity/uncertainty methodology has been applied to several application cask systems with different boron areal densities. Although, the sensitivities of the nuclear fuel cask applications are not completely covered by the set of benchmarks that were used in this study with regard to the ^{10}B capture cross section, the effect of this lack of coverage on the keff is minimal. Thus, the experimental biases are determined to be appropriate for the cask systems, and no additional bias (penalty) due to high boron loading need be imposed.

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EXECUTIVE SUMMARY

Boron has commonly been used in nuclear fuel casks to ensure a sufficient margin of subcriticality. The amount of boron used in most of the fuel casks far exceeds the amount of boron present in any of the available benchmark experiments. Such heavy loadings of boron in the casks may alter the spectrum considerably from the benchmarks or may result in boron sensitivities that are very different from those of the benchmarks. Therefore, this study was performed to determine if these available benchmarks can be used to validate a criticality code and neutron cross sections for the fuel casks. The natural question is this: are the experimental biases appropriate for the applications? In this study, sensitivity is defined as the percent change in the effective multiplication factor (k_{eff}) of the system for a 1% change in the parameter of interest (e.g., ^{10}B capture cross section). If it can be shown that the boron capture cross sections are adequately tested by the benchmarks using the sensitivity/uncertainty (S/U) method, it is judged that the existing benchmarks can be used to validate those applications, namely nuclear fuel casks that contain ^{10}B far in excess of benchmarks.

In this study, the S/U methodology has been applied to several application cask systems with different boron surface densities. The casks in this study have been shown to have very different boron capture sensitivities. Although, the sensitivities of the nuclear fuel cask applications are not completely covered by the set of benchmarks that were used in this study with regard to the ^{10}B capture cross section, the effect of this lack of coverage on the k_{eff} is minimal. Coverage is defined as having one or more benchmarks with sensitivities greater than the application's sensitivity for a specific nuclide, reaction, and energy group. Thus, the experimental biases are appropriate for the cask systems, and no additional bias (penalty) due to high boron loading need be imposed.

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1 INTRODUCTION

Many currently licensed or proposed storage and transportation casks for nuclear fuel employ sheets of borated material between fuel assemblies to enhance nuclear criticality safety. These borated materials include boral and borated steel. As part of the licensing process for these casks, calculations are performed to determine the subcritical margin for various fuel loadings. Before these calculations are deemed acceptable as part of the safety basis, the computer code and cross sections must be validated against experimental benchmarks that are applicable to the proposed cask design. There has been some question as to whether these calculations have been properly validated, given the limited set of experimental benchmarks available that utilize borated material between sets of fuel, as well as the difference between the experimental conditions and the new cask designs. In an effort to answer this question, the newly developed methods for sensitivity and uncertainty (S/U) analysis¹⁻³ at Oak Ridge National Laboratory were used to compare some nuclear fuel cask designs with experimental benchmarks, giving a measure of the similarity between the cask systems and the benchmark systems. In particular, the ¹⁰B capture reaction was studied to determine the extent to which the selected benchmarks are applicable for criticality code validation for selected fuel shipping casks containing both fresh and spent fuel with boron poison plates. KENO V.a and the 238-group ENDF/B-V SCALE cross-section library were used for all calculations.⁴

2 METHOD

The S/U tools are based on the first-order perturbation theory, in which the relative changes in the effective multiplication factor (k_{eff}) are calculated due to relative changes in the cross sections for each nuclide, reaction, and neutron energy group. Each nuclide, reaction (i.e., neutron capture, scattering, fission etc. as well as the parameters $\bar{\nu}$ and χ), and energy group are referred to as a “triplet. A “pair,” on the other hand, is defined as a specific nuclide and reaction combination (e.g., ^{10}B capture, ^{235}U fission). The triplets for a specific nuclide and reaction are summed over energy to obtain a pair. The relative change, or sensitivity, quantifies the importance of a nuclide-reaction-energy group triplet to the computed k_{eff} and, as such, is indicative of how important it is to know the reaction cross sections accurately. Sensitivity coefficients are defined such that they represent the percentage effect on k_{eff} (or more generally on some response) due to a percentage change in an input parameter, which is the cross section for fissionable material systems for which the S/U methodology has been implemented in SCALE. These sensitivity coefficients are typically presented as “profiles,” where the change in k_{eff} , due to cross sections, is given as a function of the energy of the cross section. These sensitivity profiles can be generated for each material in the system and may include various nuclear reactions (e.g., scatter, absorption, fission) as well as the neutron energy distribution from fission, χ , and average number of neutrons emitted per fission, $\bar{\nu}$.

The mathematical definition of a sensitivity “profile” is

$$S^{j,x,i} = \frac{\partial k_{eff} / k_{eff}}{\partial \Sigma^{j,x,i} / \Sigma^{j,x,i}}, \quad (1)$$

where i is the neutron energy group index, and x represents the reaction type (e.g., fission, scatter, capture) for nuclide j . The sensitivity coefficients (group value or sum over all groups) can be interpreted as percent change in the system k_{eff} for a 1% change in the cross section (corresponding group value or sum over all groups) for the reaction of interest.

The main issue that is addressed in this study is how to assess the computational penalty for an application with a significant number of triplets that are not covered by any combination of the benchmarks. Coverage is defined as having one or more benchmarks with sensitivities greater than the application’s sensitivity for a specific nuclide-reaction-energy group triplet.

The sensitivity of the system k_{eff} to each nuclide (for all reactions and all energy groups) is calculated with TSUNAMI-1D (formerly SEN1), which employs the one-dimensional discrete ordinates code XSDRNPM, or TSUNAMI-3D (formerly SEN3), which employs the Monte Carlo code KENO V.a, sensitivity analysis tools² for all applications and benchmarks for which similarity to the application is to be assessed. The benchmarks that exhibit greater sensitivities to the nuclide-reaction-energy group triplets than the application are considered to be appropriate for code and data validation for the application for those triplets. Consequently, the associated computational bias and uncertainty can be determined using an appropriate trending analysis based on integral parameters such as the E_{sum} value discussed below. In many cases, due to the limited number of benchmarks and the diverse variety of application systems, many benchmarks would have to be combined to achieve complete coverage of all triplets. Some benchmarks may

provide coverage for high-energy groups, while others may provide coverage for low-energy groups. If all triplets for an application are covered by the selected benchmarks (a single one or a combination of benchmarks), then the application is considered covered by the benchmarks that are included in the analysis and no computational penalty is considered necessary.

2.1 Integral Parameters

The sensitivity coefficients are used to derive integral parameters that quantitatively measure the applicability of an experiment to the validation of the criticality code and data to be used in the system (e.g., cask) assessment. The integral parameters that have been utilized in this study are briefly described in the following sections.

2.1.1 Integral Parameter E_{sum}

One such sensitivity-based parameter, denoted as E_{sum} , corresponds to the summation of the product of the sensitivity coefficients for two systems over energy groups and nuclides, normalized such that when summed over nuclides and reactions, an E_{sum} value of 0 indicates the systems are totally dissimilar and an E_{sum} value of 1 indicates that the two systems are precisely the same. The E_{sum} value is defined as

$$E_{sum} = M^{-1} \sum_j \sum_i S_a^{j,f,i} S_e^{j,f,i} + M^{-1} \sum_j \sum_i S_a^{j,c,i} S_e^{j,c,i} + M^{-1} \sum_j \sum_i S_a^{j,s,i} S_e^{j,s,i}, \quad (2)$$

where

$$M = \sum_x \left\{ \sum_j \sum_i (S_a^{j,x,i})^2 \sum_j \sum_i (S_e^{j,x,i})^2 \right\}^{1/2}, \quad (3)$$

and S is the sensitivity of k_{eff} for the design system application, a , or experimental configuration, e , to the fission, capture, and scattering cross sections (f , c , and s , respectively, or x as a generic name for a reaction) for energy group i and nuclide j . Mathematically, an E_{sum} value as low as -1 could be generated, but this would be the result of a rare combination of system sensitivities (i.e., for all reactions considered, the application's sensitivities are equal but opposite in sign to the benchmark's sensitivities). An E_{sum} value of 0 indicates that the systems are dissimilar. Thus, two systems exhibiting comparable integral responses in k_{eff} due to differential changes in the cross-section data are considered to be similar.

The E_{sum} parameter is considered “global” in nature, in that a single quantity identifies similarity between two systems based only on the magnitude and shape of the sensitivity profiles for fission, capture, and scatter.

2.1.2 Integral Parameter c_k

An alternative and complementary approach to exploring the similarity of systems based solely on the use of sensitivity data is the use of uncertainty analyses. The procedure involves the propagation of estimated cross-section uncertainty information to the calculated k_{eff} value of a

given system via the sensitivity coefficients. Mathematically, this is accomplished by a quadratic product of the group-wise sensitivity profile vectors by nuclide and reaction type with the cross-section uncertainty matrices by nuclide and reaction type. The result of this procedure is not only an estimate of the uncertainty in the system k_{eff} due to cross sections but also an estimate of the correlated uncertainty between systems. These correlated uncertainties can be represented by correlation coefficients, which represent the degree of correlation in the uncertainties between the two systems. This parameter, denoted as c_k , not only has the desirability of a single quantity relating the two systems but also measures the similarity of the systems in terms of uncertainty, not just sensitivity. The underlying assumption in this approach is that the cross-section uncertainty data for all nuclides and reactions of interest are tabulated and processed for use by these procedures. However, tabulated cross-section covariance data are not available for all nuclide-reaction pairs. Nuclide-reaction pairs without tabulated data are either omitted from the analysis or included with user-defined covariance values.

When the matrix of uncertainty information for all of the cross sections is denoted as $C_{\alpha\alpha}$, and the sensitivity matrix relating changes in each constituent material and process to the system k_{eff} is labeled as S_k , the uncertainty matrix for the system k_{eff} values, C_{kk} , is given as

$$C_{kk} = S_k C_{\alpha\alpha} S_k^{\dagger} , \quad (4)$$

where \dagger indicates a transpose.

The C_{kk} matrix consists of variance values σ_p^2 for each of the critical systems under consideration (the diagonal elements), as well as the so-called ‘‘covariance’’ between systems σ_{pq}^2 (the off-diagonal elements). These off-diagonal elements represent the shared or common variance, hence the term covariance, between any two systems. For presentation, these off-diagonal elements are typically divided by the square root of the corresponding (same row) diagonal elements (i.e., the respective standard deviations) to generate a correlation coefficient matrix. Thus, the c_k coefficients are defined as

$$c_k = \sigma_{pq}^2 / (\sigma_p \sigma_q) , \quad (5)$$

such that each c_k value represents the correlation coefficient between uncertainties in system p and system q .

These correlations arise due to the fact that the uncertainties in the k_{eff} values for two different systems are related, since they contain the same materials. Cross-section uncertainties will propagate to all systems containing these materials. Systems with the same materials and similar spectra would be correlated, while systems with different materials or differing spectra would not be correlated. The interpretation of the correlation matrix is the following: a value of 0 represents no correlation between the systems, a value of 1 represents full correlation between the systems, and a value of -1 represents a full anticorrelation.

2.1.3 Integral Parameter g

The sensitivity differences for an application's triplet is computed by taking the group value for the application and subtracting the group value for each benchmark—forming a vector of values, each element of which represents the sensitivity difference between the application and the corresponding benchmark:

$$Z_{ab}^{j,x,i} = \begin{cases} S_a^{j,x,i} - S_b^{j,x,i}, & \text{if } |S_a^{j,x,i}| > |S_b^{j,x,i}| \text{ and } S_a^{j,x,i} \cdot S_b^{j,x,i} > 0 \\ 0, & \text{if } |S_a^{j,x,i}| \leq |S_b^{j,x,i}| \\ 0, & \text{if } S_a^{j,x,i} \cdot S_b^{j,x,i} < 0 \end{cases}, \quad (6)$$

where:

i is the neutron energy group index,

j is the nuclide index,

x is the reaction index (e.g., fission, capture, etc.),

a represents the application,

b represents the benchmark,

$S_a^{j,x,i}$ is the application's sensitivity for neutron energy group i , nuclide j , and reaction x , and

$S_b^{j,x,i}$ is the benchmark's sensitivity for neutron energy group i , nuclide j , and reaction x

The difference is taken only when the application's sensitivity triplet $S_a^{j,x,i}$ and benchmark's sensitivity triplet $S_b^{j,x,i}$ have the same sign.

For each application triplet, there are as many sensitivity difference ($Z_{ab}^{j,x,i}$) values as there are benchmarks (i.e., b spans all benchmarks). The minimum of the sensitivity differences or the minimum noncoverage for the application is then the minimum of the $Z_{ab}^{j,x,i}$ values for each triplet calculated over all benchmarks:

$$Z_a^{j,x,i} = \min(Z_{ab}^{j,x,i}), \quad (7)$$

where b spans all benchmarks.

The sum of the minimum sensitivity differences over all energy groups ($Z_a^{j,x}$) is defined as the minimum of the sensitivity differences:

$$Z_a^{j,x} = \sum_i Z_a^{j,x,i}. \quad (8)$$

Coverage for a nuclide-reaction pair is assessed by analyzing the normalized differences in sensitivities between the application and each benchmark. This normalized difference for each nuclide and all reactions of interest is defined as:

$$g'_{j,x} = \frac{\sum_i Z_{ab}^{j,x,i}}{\sum_i S_a^{j,x,i}}, \quad (9)$$

where the difference ($Z_{ab}^{j,x,i}$) is calculated based on the conditional equation (6).

A g' value is calculated for an application against each benchmark for each nuclide-reaction pair. Hence, for N benchmarks there are N values of g' for an application for each nuclide-reaction pair. For a specific nuclide-reaction pair, the integral parameter g' is defined as the summed difference between the sensitivity coefficients for an application and a benchmark for all energy groups where the application's sensitivity is greater than the benchmark's sensitivity, normalized with respect to the application's total sensitivity. Total sensitivity is defined as the sensitivity for a specific nuclide-reaction pair that results from summation of the sensitivity triplets over all energy groups. Hence, throughout this document, total sensitivity should not be equated with the sensitivity to the total cross section.

The normalization of g' bounds its values between 0 and 1. A g' value of 0 indicates complete coverage (i.e., $|S_a^{j,x,i}| < |S_b^{j,x,i}|$ for all triplets), whereas a g' value of 1 indicates a complete noncoverage due to the benchmark's sensitivity being exactly zero. A large g' value indicates that the application's sensitivity for a specific nuclide-reaction pair is greater than the benchmark's sensitivity for the same nuclide-reaction pair in some or all neutron energy groups. A large g' value also indicates that the summed difference $Z_{ab}^{j,x}$ is a large fraction of the application's sensitivity.

Consider a hypothetical application that has high sensitivity only in the low neutron energy range (i.e., high below 1 eV and zero above 1 eV) for an arbitrary nuclide-reaction pair. Also consider a hypothetical benchmark that has high sensitivity only in neutron energies above 1 eV for the same nuclide-reaction pair (i.e., high above 1 eV and zero below 1 eV). The g' value for this case would be exactly 1. If the application's sensitivity above 1 eV is nonzero, then the g' value would be less than 1 and the actual value would depend on how large the application's sensitivity value is in this range (i.e., above 1 eV). Similarly, if the benchmark's sensitivity below 1 eV is nonzero, then the g' value would be less than 1 and the actual value would depend on how large the benchmark's sensitivity is in this range (i.e., below 1 eV).

Historically, the integral parameters that have been used in the sensitivity analysis have been defined such that a normalized integral parameter value of 1 indicates total agreement. Therefore, a slightly different version of the integral parameter g' , which can be thought of as the complement parameter, has been defined:

$$g_{j,x} = 1 - g'_{j,x} = 1 - \frac{\sum_i Z_{ab}^{j,x,i}}{\sum_i S_a^{j,x,i}}, \quad (10)$$

such that a g value of 1 indicates complete coverage, whereas a g value of 0 indicates a complete noncoverage due to benchmark's sensitivity being exactly zero.

The mathematical interpretation of the g value is illustrated in Figure 1 as differences between the application and benchmark sensitivities. In the energy range where the application's sensitivity is greater than the benchmark's sensitivity, the difference is calculated and is shown as the shaded area in Figure 1. The energy range of the shaded area indicates the range for which

the application's sensitivities are not covered by the benchmark. The g' value is the ratio of this shaded area, $\sum_i Z_{ab}^{j,x,i}$, to the area under the application's sensitivity curve, $\sum_i S_a^{j,x,i}$. The g value is then simply $1 - g'$. By definition, covered triplets do not contribute to the numerator of g' . Thus, the degree of coverage for a specific triplet does not influence the value of g . This feature offers an advantage over previous nuclide-reaction-specific integral parameters, which could be artificially inflated by triplets where the benchmark's sensitivity greatly exceeds that of the application.

A large g value indicates that the covered part of the application's sensitivity for a specific nuclide-reaction pair makes up the majority of the application's sensitivity for that nuclide-reaction pair. Though not necessary to consider in the analysis, if the value of the total sensitivity for a nuclide-reaction pair is "small," the application can be considered "not important" (or can even be considered adequately covered by the benchmark over all energy groups for that nuclide-reaction pair), even with a small g value. Here "small sensitivity" is arbitrarily defined as resulting in less than 0.001 change in k_{eff} for a 100% change in the cross sections. If an application is completely covered, there is no need to apply a computational penalty.

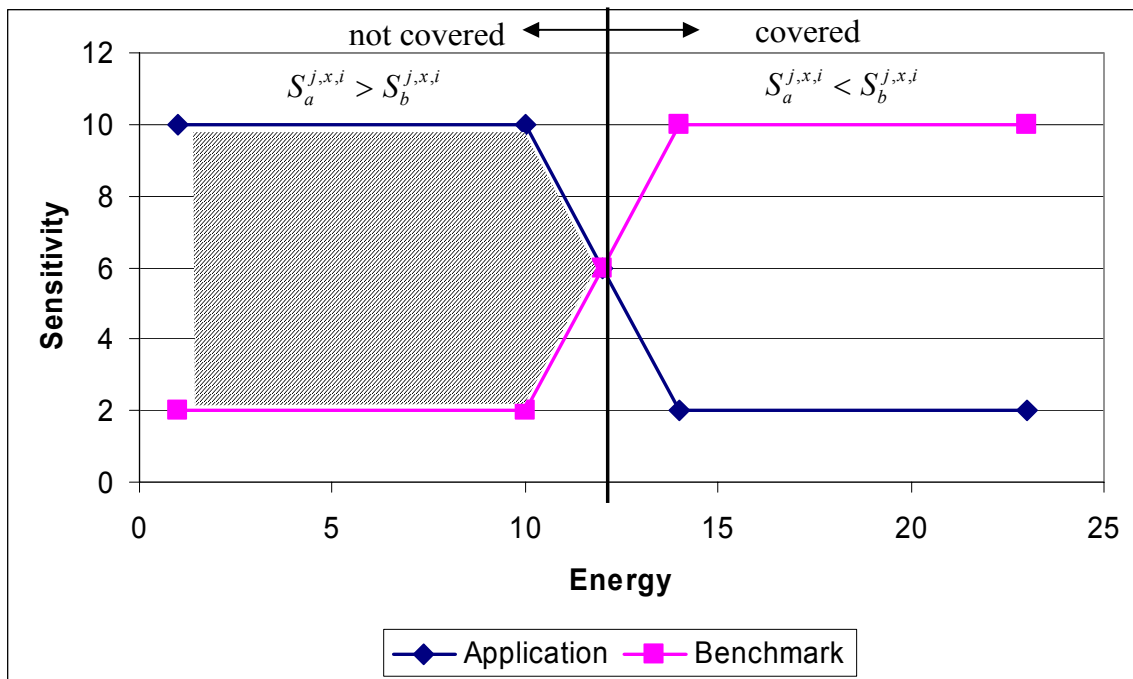


Figure 1 Illustration of g value

If the E_{sum} or c_k values between an application and an experiment indicate little similarity, the g value can be used to determine the underlying reason. Consider an application system with large sensitivity to a specific nuclide-reaction pair. If this application is analyzed with a set of

benchmarks that yield a small g value for the same nuclide-reaction pair, then the corresponding E_{sum} and c_k values will be small. Therefore, by calculating the g values for the nuclide-reaction pairs that yield large sensitivities, one can determine the effect of those pairs on the E_{sum} or c_k values.

The differences between the application and benchmark sensitivities are indicative of how well the benchmark covers the application. If these differences are small, then the benchmarks are considered adequate for the application for validation purposes and the effect of the uncertainties in the cross-section data would be quantified by trending analysis. If the differences are large, then the benchmarks are not adequate for the application for validation purposes and the computational penalty that is generated is not appropriate. The sensitivity difference value above which these differences are considered large has not been established in this study. That task is beyond the scope of this study and will be concluded in the future as part of the development of a guidance document. The guidance will discuss the use of the g parameter methodology for assessment of the degree of applicability of benchmarks and the corresponding computational penalty due to noncoverage.

2.2 Estimated Penalty for Noncoverage

The penalty assessment methodology presented here is based on the assumption that a benchmark with a greater sensitivity for the nuclide, reaction, and energy group triplet of interest sufficiently covers the triplet in the application. The approach that is used in this method is to determine the differences between the application and benchmark sensitivities for all triplets that are not covered, and to quantify the importance of this noncoverage in terms of its final effect on the k_{eff} value of the application using the cross-section uncertainties.

In the penalty assessment, the application's sensitivity for a nuclide-reaction pair is compared with all benchmarks that are included in the analysis on a group-wise basis. All nuclide-reaction pairs are processed sequentially (one at a time). For each pair being processed, the number of benchmarks that have greater sensitivities than the application is tallied for each group for adequacy of coverage. If some of the groups have no benchmarks that provide coverage, then the minimum of the sensitivity differences (noncoverage) between all benchmarks and the application for those noncovered groups is calculated (i.e., shaded area in Figure 1).

Before a sensitivity difference between the application and a benchmark can be used in the penalty assessment, the application and each considered benchmark must pass the similarity test based either on the integral parameter c_k or the integral parameter E_{sum} . These parameters are described in detail in Ref. 2. The sensitivity differences for the benchmarks that do not pass the similarity test between the benchmark and the application are eliminated from the vector. Similarity of benchmarks to the application is determined by using one of the integral parameters. Subsequent trending analysis yields the corresponding bias and uncertainty, which are incorporated into the subcritical limit. The method by which this bias is determined is beyond the scope of this study.

The minimum of the sensitivity differences can be viewed as a measure of the inadequacy of the benchmark suite to replicate the physics of the application. This minimum sensitivity difference can be converted into an artificial computational penalty by multiplying the difference

by the cross-section covariance data for that nuclide-reaction pair. If the covariance data for the desired nuclide-reaction pair do not exist, a hypothetical matrix corresponding to 10, 20, 100%, etc., uncertainty in the microscopic cross-section data can be used to quantify possible effects from the nuclide-reaction pair assuming linearity of the k_{eff} response to the cross section. One should keep in mind, however, that first-order linear perturbation theory is valid for small changes in the variable (i.e., in this case, the uncertainty that is propagated to the final result). As such, it would be appropriate to choose a hypothetical matrix value that bounds the expected maximum uncertainty effect on the penalty. This approach, though not theoretically appropriate, is very useful for examining and estimating cross-section impacts on the computed results. If the analyst has *a priori* knowledge regarding the cross section uncertainties (at least the magnitude), a corresponding conservative matrix value can be selected. For example, if the uncertainties are not known but are expected to be less than 20%, then the penalty value from a 20% or larger uncertainty value can be used in determining the penalty. In this study, a 100% uncertainty in the cross sections for which there are no covariance data has been assumed.

The cross-section uncertainties and the minimum sensitivity differences are used to calculate an adjusted k_{eff} , which is interpreted to be the application's calculated k_{eff} increased for consideration of noncovered sensitivities. The adjusted k_{eff} (k'_{eff}) is the k_{eff} value after accounting for the noncoverage:

$$k'_{eff} = k_{eff} + k_{eff} \sqrt{Z_a C_{\alpha\alpha} Z_a^T} . \quad (11)$$

For N triplets in the problem, Z_a is an $N \times 1$ vector of the minimum of the sensitivity differences comprised of $Z_a^{j,x,i}$ values for all nuclides, reactions, and energy groups; Z_a^T is the transpose; and $C_{\alpha\alpha}$ is the $N \times N$ cross section covariance matrix. Hence, N is the number of nuclide-reaction pairs times the number of neutron energy groups (i.e., the number of triplets). In the above equation, the unit for Z_a is $(\Delta k_{eff} / k_{eff}) / (\Delta \Sigma / \Sigma)$, and the unit for $C_{\alpha\alpha}$ is $(\Delta \Sigma / \Sigma)^2$. Therefore the final penalty has the unit (Δk_{eff}) . The cross-section uncertainty is $(\Delta \Sigma / \Sigma)$ [e.g., 10% uncertainty in the cross sections implies $(\Delta \Sigma / \Sigma)$ is 0.1].

Hence, the adjusted k_{eff} represents the calculated effective multiplication factor after applying a penalty for the noncoverage of the sensitivities. The noncoverage is weighted by the uncertainties in the measured cross sections thereby creating a penalty that is the result of the cross-section uncertainties. If the cross-section uncertainties were zero (implies cross sections are known exactly), then the penalty would be zero even if the application's sensitivities were not covered completely. This result is reasonable and expected because the cross sections are known exactly, there would not be any concern about validating the cross sections or determining the degree of applicability of the benchmarks based on the cross sections. Therefore, no penalty would have to be assessed due to cross-section uncertainties as a result of noncoverage. The penalty would also be zero if the application is fully covered.

Currently, a single benchmark can provide coverage for an application. The adequacy of coverage by one benchmark has not yet been investigated. An alternate use of the g value may be developed in the future to increase the adjusted k_{eff} penalty when an insufficient number of benchmarks (e.g., one) provide coverage.

2.3 Implementation in SCALE

The new S/U analytical sequences TSUNAMI-1D and TSUNAMI-3D will be publicly available with the release of SCALE 5. These sequences compute the relative change in the system k_{eff} that would be observed for perturbations in the group-wise neutron cross-section data for each reaction of each nuclide in the system. The TSUNAMI-IP (formerly CANDE) code within SCALE 5 uses this sensitivity data to calculate integral parameters, which were described in previous sections. These integral parameters give a measure of the similarity between a given application and an experimental benchmark. A near-unity value for the integral parameter indicates that the experiment demonstrates similar properties to the application, based on the S/U criteria of the particular integral parameter evaluated, and thus the experiment is applicable for the criticality code validation of the application.

If two systems exhibit an E_{sum} or c_k value of 0.9 or greater, the systems may be considered to be applicable for the criticality code validation of one another. However, the values of E_{sum} or c_k may be dominated by some nuclide-reaction pairs in the system (e.g., ^{235}U fission and ^1H scattering reactions as opposed to boron capture). Therefore, one must be careful in using the E_{sum} or c_k values to derive conclusions on validation of all nuclides in the system.

The g parameter, however, indicates how well a benchmark provides coverage for the nuclide-reaction pair. Therefore, the g parameter is used to assess the validity of the computational penalty due to noncoverage. The $Z_a^{j,x}$, sum of sensitivity differences for noncovered groups for the nuclide-reaction pair, is used to calculate the computational penalty.

3 NUCLEAR FUEL CASK MODELS

Three nuclear fuel casks and two computational benchmark casks were selected to evaluate validation coverage by the experimental benchmarks identified for this study. In many of the existing fuel casks, the amount of boron far exceeds the amount of boron in most of the available benchmark experiments.^{5,6} A list and description of these available benchmark experiments are given in Appendix A. Of the five fuel cask models that were used in this study, the three existing cask models were loaded with fresh fuel of varying enrichment and the two computational benchmarks were loaded with burned fuel. Brief descriptions of the cask models are provided below.

1. Holtec HI-STAR MPC-24 cask,⁷ which holds 24 pressurized water reactor (PWR) assemblies and is designed for fresh fuel storage and transportation, contains 0.055-in.-thick Boral plates. The model used in this study contained 4.2% enriched fuel.
2. Holtec HI-STAR MPC-68 cask,⁷ which holds 68 boiling water reactor (BWR) assemblies and is also designed for fresh fuel storage and transportation, contains 0.08-in.-thick Boral plates. The model used in this study contained 4.2% enriched fuel.
3. General Atomics GA-4 cask,⁸ holds four PWR assemblies. In this model, the cask is loaded with fresh fuel assemblies containing 3% enriched fuel. The boron is present as B₄C pellets inserted into horizontal tubes within a cruciform stainless steel fuel support structure.
4. GBC-32 cask,⁹ which is designed as a computational benchmark problem holding 32 PWR assemblies, contains Boral plates that are 0.101 in. thick. The fuel in the model has been burned for 45 GWd/MTU and cooled for 5 years.
5. Organisation for Economic Co-operation and Development (OECD) Benchmark cask,¹⁰ which holds 21 PWR assemblies and is also a computational benchmark problem, contains borated steel with 1 wt % boron, which is 0.08 in. thick. The fuel in the model has been burned for 30 GWd/MTU and cooled for 1 year.

The nuclear fuel cask models, including their calculated energy of average lethargy causing fission (EALF) and their calculated ¹⁰B capture sensitivity coefficients, are summarized in Table 1.

The ¹⁰B surface number density (density per unit thickness) and ¹⁰B capture sensitivities in these application fuel casks and all benchmarks that were considered in this study are shown in Figures 2 and 3, respectively. The ¹⁰B capture sensitivity profiles for the application casks are shown in Figures 4 through 9. The value “a” in the legend indicates the total sensitivity for the reaction that is being plotted. If the sensitivity profile had a mixture of positive and negative sensitivity values, then the values that are opposite in sign to the total sensitivity (“a=”) would be shown with “osc=” in the legend.

Table 1 Nuclear fuel cask model parameters

Cask	Calculated $k_{eff} \pm \sigma$	EALF (eV)	^{10}B Capture sensitivity	^{10}B Form	Total ^{10}B (kg)	^{10}B Surface density (at/cm ²)
MPC-24	0.9458 ± 0.0005	2.257E-01	-2.62E-02	Boral	~12	1.216E+21
MPC-68	0.9349 ± 0.0005	2.775E-01	-5.05E-02	Boral	~15	1.658E+21
GA-4	0.9221 ± 0.0005	4.572E-01	-2.38E-02	B ₄ C	~8	4.750E+22
GBC-32	0.8941 ± 0.0004	2.474E-01	-2.76E-02	Boral	~12	1.688E+21
OECD	1.1303 ± 0.0005	6.311E-02	-4.45E-02	Borated steel	~6	3.918E+20

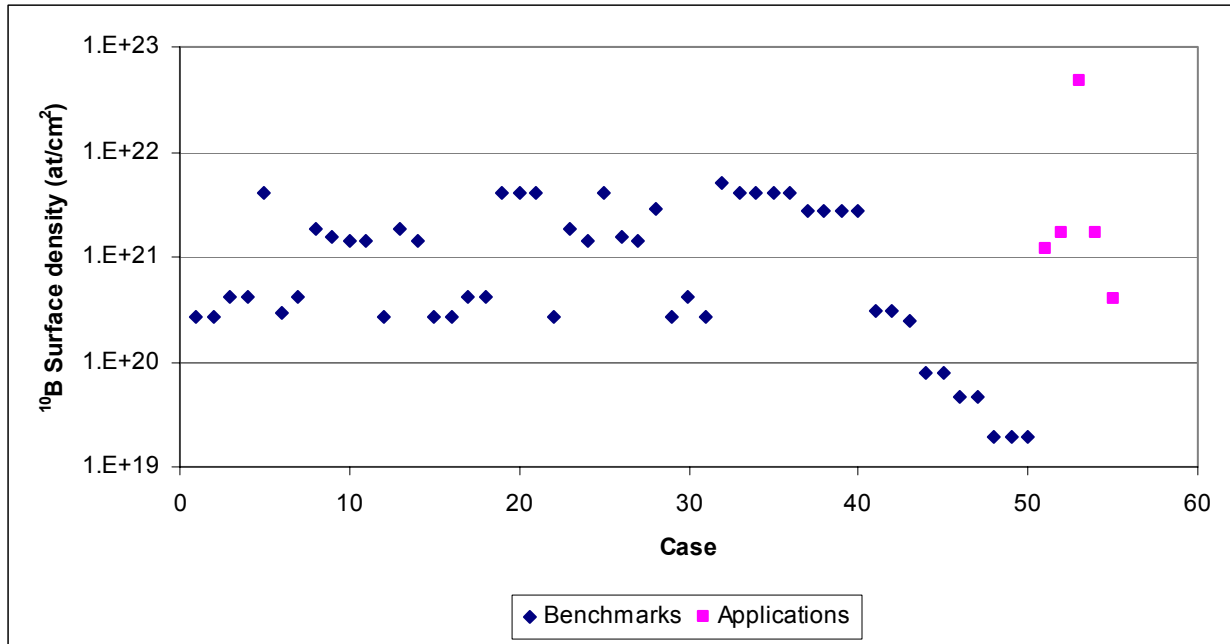


Figure 2 Maximum ^{10}B surface number density for all benchmarks and applications

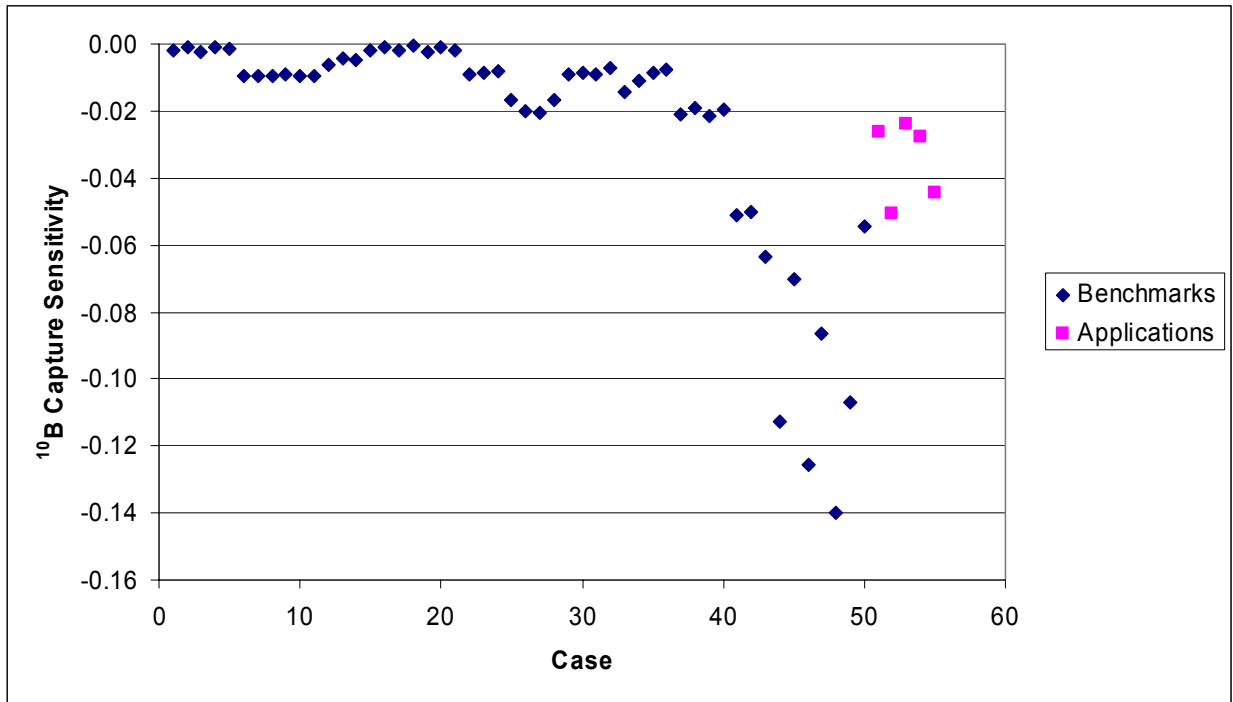


Figure 3 Capture sensitivities of ¹⁰B for all benchmarks and applications

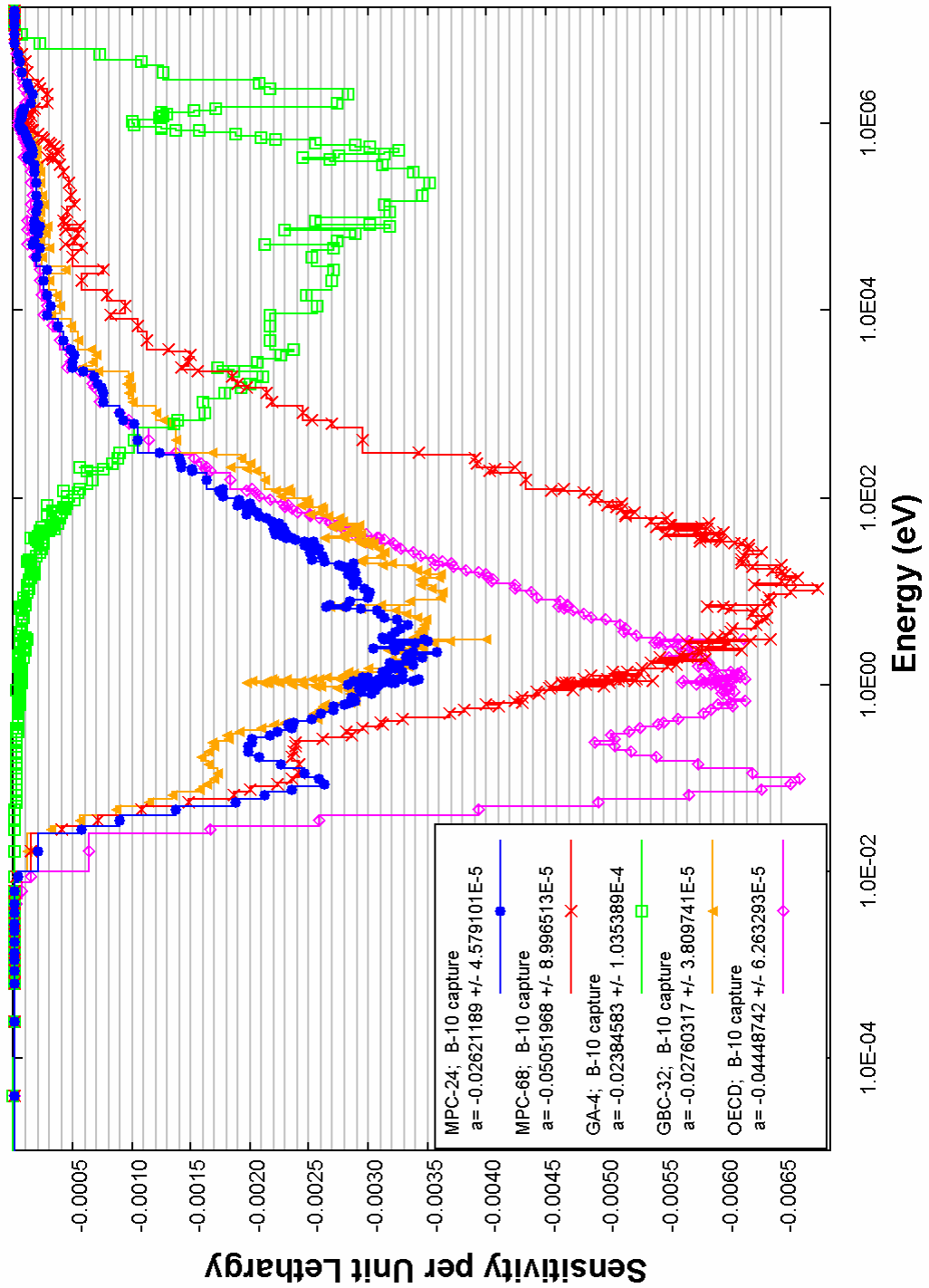


Figure 4 Capture sensitivity profiles of ¹⁰B for fuel

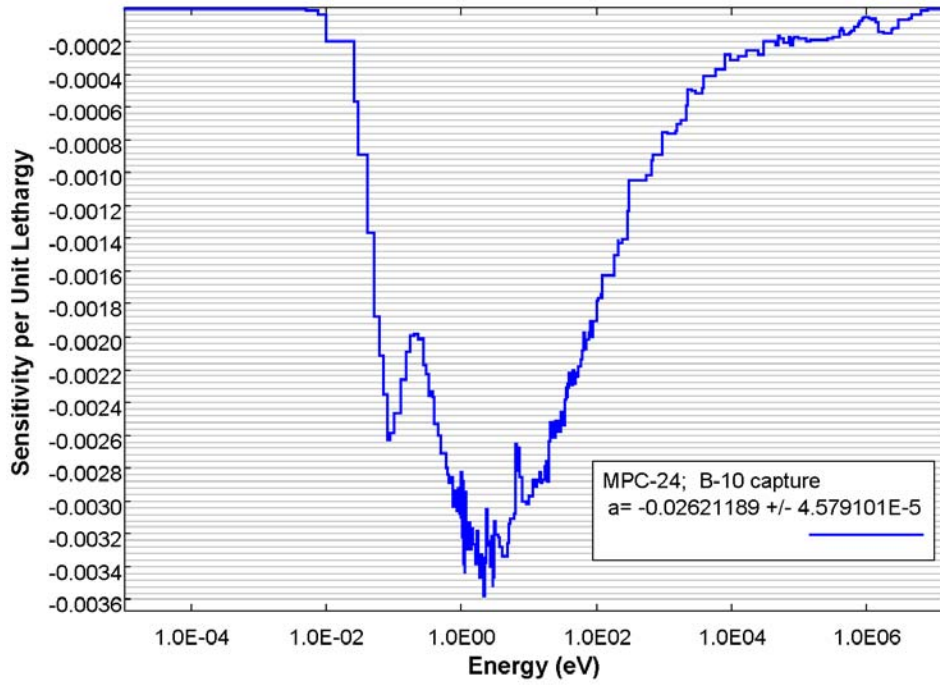


Figure 5 Capture sensitivity profile of ^{10}B for MPC-24 fuel cask

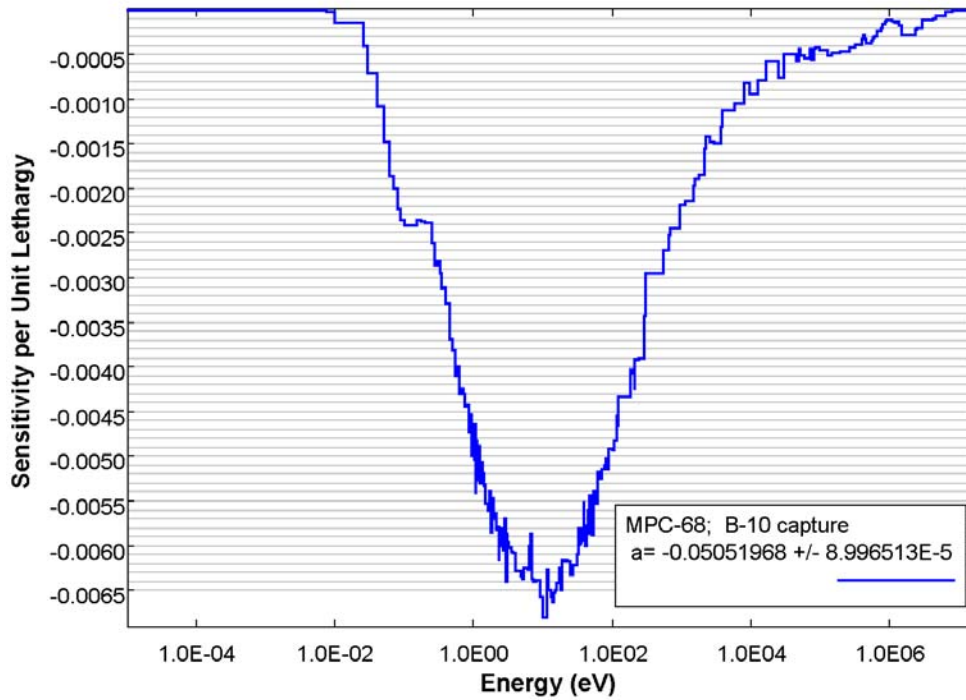


Figure 6 Capture sensitivity profile of ^{10}B for MPC-68 fuel cask

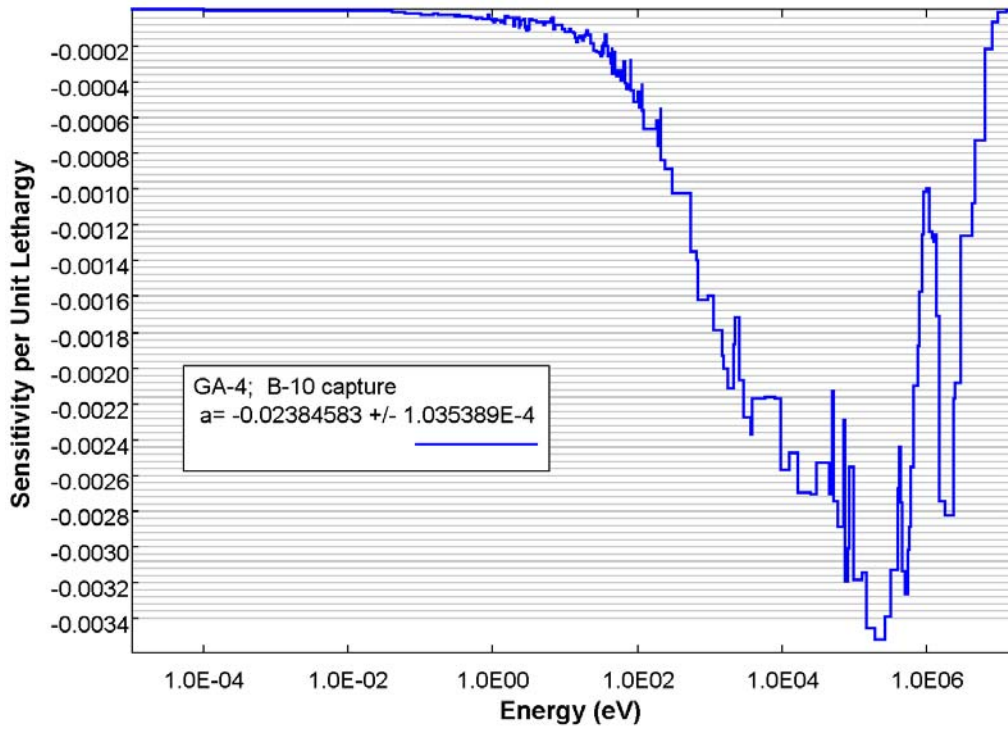


Figure 7 Capture sensitivity profile of ¹⁰B for GA-4 fuel cask

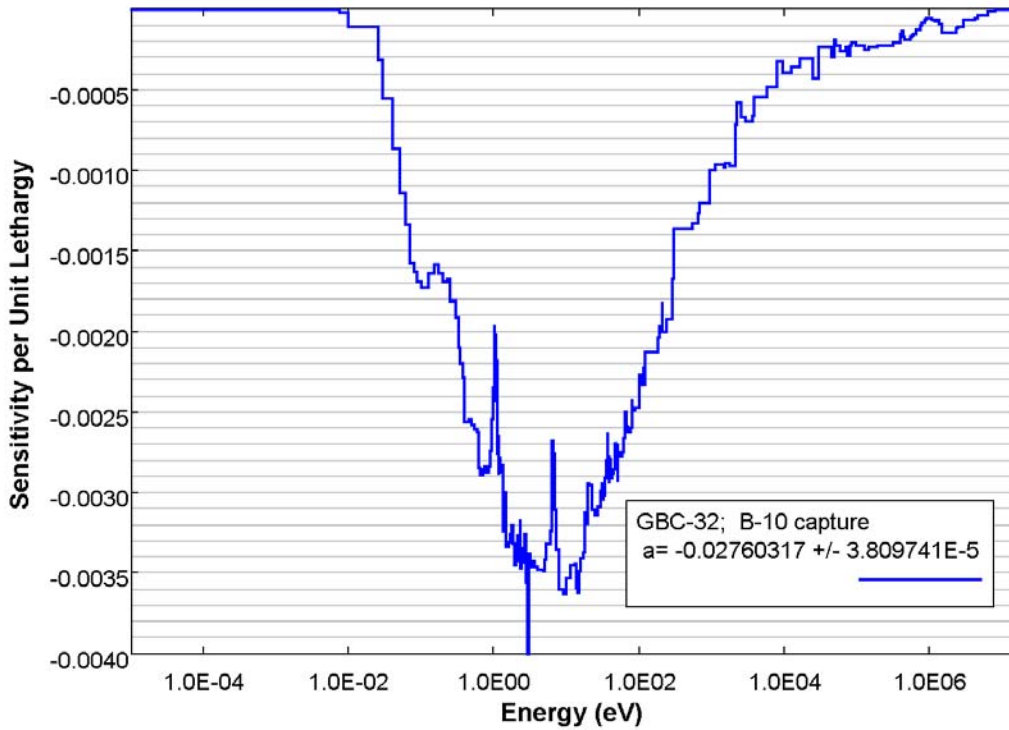


Figure 8 Capture sensitivity profile of ¹⁰B for GBC-32 fuel cask

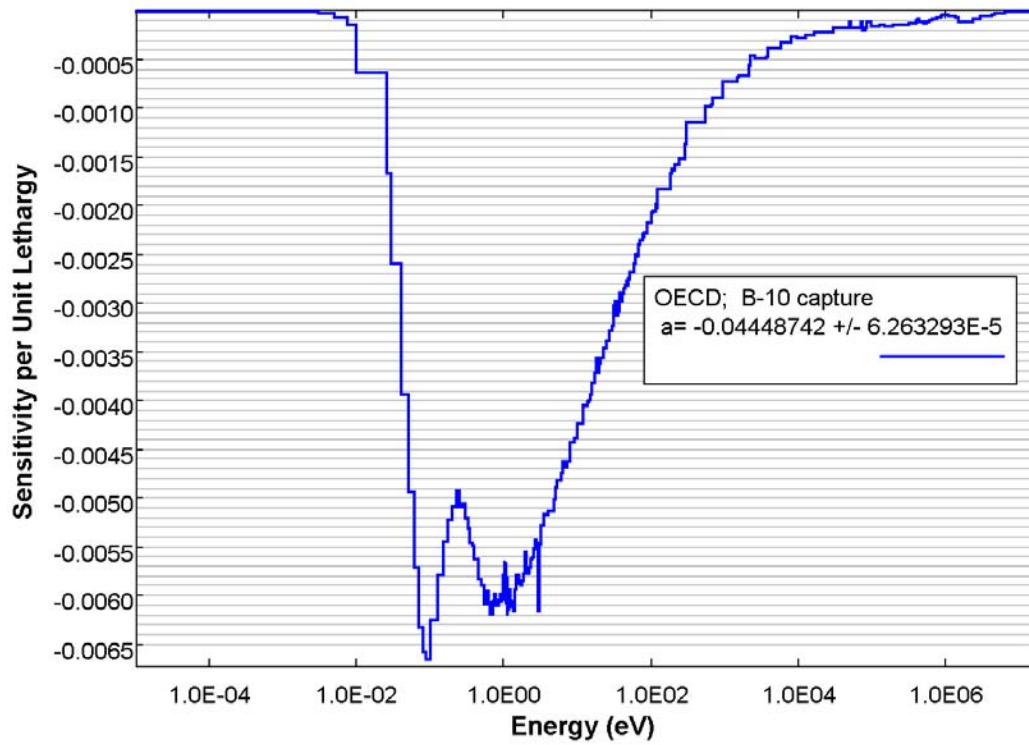


Figure 9 Capture sensitivity profile of ^{10}B for OECD fuel cask

4 ANALYSIS OF NUCLEAR FUEL CASK MODELS

As can be seen from the data in Table A.1 of the Appendix, the available benchmark experiments utilizing boron as a neutron poison between fuel assemblies are generally not very sensitive to changes in the ^{10}B capture reaction. With the notable exception of the experiments in LEU-COMP-THERM-051, the ^{10}B capture sensitivity coefficients for the experiments range from $-6.364\text{E}-04$ to $-2.137\text{E}-02$, meaning that a 1% increase in the ^{10}B capture cross section across all energy groups or in the number density would result in a decrease in k_{eff} ranging from 0.0006 to 0.0214%. Assuming linearity, this would also mean that removal of the ^{10}B from the model (100% decrease in the number density) would result in an increase in k_{eff} ranging from 0.06 to 2.14%. The experiments in LEU-COMP-THERM-051 are distinctive because their ^{10}B capture sensitivity coefficients are much higher than those for the other experiments, ranging from $-5.025\text{E}-02$ to $-1.399\text{E}-01$.

A comparison of the data in Tables 1 and A.1 as illustrated in Figure 3 shows that the cask models span the same approximate range of EALF as the set of benchmark experiments, but the absolute magnitude of their ^{10}B capture sensitivity coefficients is greater, on average, than that of the benchmark set. Only the experiments from LEU-COMP-THERM-051 fall in the same range of sensitivities.

4.1 Evaluation of E_{sum} Integral Parameter

The values of E_{sum} for each benchmark relative to each of the five cask models are given in Table 2. Based on these data, it would appear that most of the benchmarks are very similar to the cask models and are appropriate for code validation, with the exception of the GBC-32 cask. However, the values of E_{sum} are strongly dominated by the ^{235}U fission and ^1H scattering reactions. Therefore, these values are more indicative of the use of water-moderated LEU (low-enriched uranium) fuel in both the benchmarks and the cask models. Also note that E_{sum} values for the GBC-32 cask are significantly lower than the E_{sum} values for the other casks. This is most likely due to ^{235}U fission or ^1H scatter sensitivities not being similar to the benchmarks. The reason becomes clear after detailed analysis of these sensitivities using the g value (see Section 4.3).

Table 2 Values of E_{sum} for each benchmark/cask model pair

Benchmark	MPC-24	MPC-68	GA-4	GBC-32	OECD
LCT009-05	0.9322	0.9098	0.9370	0.5811	0.9418
LCT009-06	0.9342	0.9086	0.9384	0.5849	0.9449
LCT009-07	0.9344	0.9131	0.9380	0.5829	0.9438
LCT009-08	0.9306	0.9090	0.9335	0.5784	0.9369
LCT009-09	0.9332	0.9130	0.9373	0.5798	0.9434
LCT012-02	0.9803	0.9607	0.9862	0.6210	0.9258
LCT012-03	0.9782	0.9581	0.9857	0.6208	0.9212
LCT012-04	0.9817	0.9617	0.9886	0.6230	0.9277
LCT012-05	0.9814	0.9605	0.9871	0.6223	0.9272
LCT012-06	0.9791	0.9599	0.9851	0.6193	0.9276
LCT012-07	0.9793	0.9615	0.9863	0.6198	0.9251
LCT013-02	0.9768	0.9573	0.9726	0.6005	0.9084
LCT013-03	0.9782	0.9566	0.9743	0.6051	0.9134
LCT013-04	0.9773	0.9541	0.9738	0.6034	0.9064
LCT016-08	0.9530	0.9322	0.9637	0.6070	0.9423
LCT016-09	0.9535	0.9323	0.9647	0.6089	0.9459
LCT016-10	0.9538	0.9306	0.9637	0.6084	0.9472
LCT016-11	0.9510	0.9313	0.9609	0.6041	0.9428
LCT016-12	0.9573	0.9348	0.9664	0.6101	0.9481
LCT016-13	0.9550	0.9335	0.9650	0.6084	0.9457
LCT016-14	0.9499	0.9298	0.9591	0.6040	0.9384
LCT042-02	0.9834	0.9663	0.9902	0.6251	0.9267
LCT042-03	0.9849	0.9707	0.9909	0.6249	0.9271
LCT042-04	0.9843	0.9661	0.9890	0.6274	0.9281
P3314BA	0.9753	0.9607	0.9710	0.5988	0.9040
P3314BC	0.9791	0.9652	0.9735	0.5991	0.9111
P3314BF1	0.9817	0.9645	0.9733	0.6026	0.9096
P3314BF2	0.9796	0.9637	0.9724	0.6015	0.9051
P3314BS3	0.9672	0.9466	0.9617	0.5944	0.8996
P3314BS4	0.9699	0.9450	0.9646	0.5993	0.9101
P3602BS1	0.9832	0.9659	0.9886	0.6277	0.9275
P62FT231	0.9785	0.9587	0.9737	0.6019	0.8933
P71F14F3	0.9839	0.9715	0.9777	0.6055	0.9026
P71F14V3	0.9799	0.9700	0.9746	0.5998	0.8931

Table 2 (Continued)

Benchmark	MPC-24	MPC-68	GA-4	GBC-32	OECD
P71F14V5	0.9852	0.9685	0.9794	0.6076	0.9020
P71F214R	0.9816	0.9674	0.9753	0.6028	0.8967
PAT80L1	0.9793	0.9572	0.9754	0.5961	0.9661
PAT80L2	0.9803	0.9611	0.9783	0.5955	0.9607
PAT80SS1	0.9775	0.9542	0.9758	0.5963	0.9652
PAT80SS2	0.9810	0.9620	0.9784	0.5969	0.9640
LCT051-10	0.9851	0.9736	0.9890	0.6230	0.9331
LCT051-11	0.9849	0.9744	0.9875	0.6186	0.9327
LCT051-12	0.9833	0.9697	0.9856	0.6178	0.9314
LCT051-13	0.9595	0.9487	0.9621	0.5960	0.9132
LCT051-14	0.9654	0.9556	0.9686	0.6063	0.9204
LCT051-15	0.9497	0.9386	0.9528	0.5893	0.9070
LCT051-16	0.9615	0.9477	0.9650	0.6060	0.9194
LCT051-17	0.9358	0.9271	0.9379	0.5759	0.8938
LCT051-18	0.9493	0.9374	0.9518	0.5956	0.9117
LCT051-19	0.9574	0.9456	0.9631	0.6167	0.9268

4.2 Evaluation of c_k Integral Parameter

The values of c_k for each benchmark relative to each of the five cask models are given in Table 3. As with E_{sum} , it would appear that most of the benchmarks are very similar to the cask models and are appropriate for code validation, with the exception of the GBC-32 cask. However, the values of c_k are also strongly dominated by the ^{235}U fission and ^1H scattering reactions. Therefore, they are more indicative of the use of water-moderated LEU fuel in both the benchmarks and the cask models. Also once again note that c_k values for the GBC-32 cask are significantly lower than the c_k values for the other casks. This is also due to ^{235}U fission or ^1H scatter sensitivities not being similar to the benchmarks.

Table 3 Values of c_k for each benchmark/cask model pair

Benchmark	MPC-24	MPC-68	GA-4	GBC-32	OECD
LCT009-05	0.9330	0.8310	0.8890	0.3883	0.9491
LCT009-06	0.9295	0.8263	0.8847	0.3860	0.9477
LCT009-07	0.9344	0.8327	0.8910	0.3902	0.9492
LCT009-08	0.9305	0.8268	0.8855	0.3872	0.9460
LCT009-09	0.9311	0.8277	0.8866	0.3878	0.9468
LCT012-02	0.9794	0.8892	0.9623	0.4653	0.8706
LCT012-03	0.9786	0.8878	0.9613	0.4658	0.8675

Table 3 (Continued)

Benchmark	MPC-24	MPC-68	GA-4	GBC-32	OECD
LCT012-04	0.9813	0.8929	0.9677	0.4705	0.8669
LCT012-05	0.9806	0.8915	0.9662	0.4694	0.8667
LCT012-06	0.9810	0.8922	0.9666	0.4690	0.8683
LCT012-07	0.9810	0.8926	0.9667	0.4699	0.8664
LCT013-02	0.9823	0.8884	0.9439	0.4453	0.8853
LCT013-03	0.9824	0.8882	0.9449	0.4471	0.8830
LCT013-04	0.9824	0.8883	0.9446	0.4468	0.8826
LCT016-08	0.9526	0.8597	0.9284	0.4235	0.9270
LCT016-09	0.9511	0.8569	0.9262	0.4225	0.9262
LCT016-10	0.9520	0.8584	0.9275	0.4234	0.9259
LCT016-11	0.9499	0.8548	0.9239	0.4210	0.9259
LCT016-12	0.9549	0.8630	0.9323	0.4268	0.9264
LCT016-13	0.9513	0.8571	0.9264	0.4226	0.9262
LCT016-14	0.9530	0.8595	0.9291	0.4248	0.9259
LCT042-02	0.9849	0.9027	0.9750	0.4740	0.8692
LCT042-03	0.9855	0.9047	0.9786	0.4782	0.8653
LCT042-04	0.9855	0.9043	0.9781	0.4781	0.8649
P3314BA	0.9867	0.8968	0.9536	0.4526	0.8838
P3314BC	0.9890	0.9003	0.9562	0.4529	0.8880
P3314BF1	0.9889	0.9002	0.9572	0.4542	0.8871
P3314BF2	0.9882	0.8993	0.9567	0.4546	0.8848
P3314BS3	0.9780	0.8811	0.9364	0.4396	0.8857
P3314BS4	0.9782	0.8811	0.9366	0.4401	0.8856
P3602BS1	0.9851	0.9027	0.9752	0.4747	0.8691
P62FT231	0.9827	0.8932	0.9509	0.4563	0.8639
P71F14F3	0.9920	0.9102	0.9678	0.4661	0.8728
P71F14V3	0.9906	0.9061	0.9633	0.4631	0.8719
P71F14V5	0.9897	0.9035	0.9615	0.4625	0.8700
P71F214R	0.9880	0.9014	0.9568	0.4583	0.8715
PAT80L1	0.9807	0.8941	0.9518	0.4361	0.9437
PAT80L2	0.9802	0.8912	0.9494	0.4341	0.9450
PAT80SS1	0.9810	0.8943	0.9521	0.4365	0.9438
PAT80SS2	0.9808	0.8927	0.9506	0.4349	0.9452
LCT051-10	0.9879	0.9129	0.9835	0.4771	0.8741
LCT051-11	0.9877	0.9134	0.9836	0.4765	0.8751
LCT051-12	0.9872	0.9125	0.9824	0.4755	0.8744

Table 3 (Continued)

Benchmark	MPC-24	MPC-68	GA-4	GBC-32	OECD
LCT051-13	0.9860	0.9127	0.9817	0.4739	0.8739
LCT051-14	0.9839	0.9063	0.9742	0.4634	0.8919
LCT051-15	0.9853	0.9120	0.9808	0.4736	0.8725
LCT051-16	0.9830	0.9065	0.9742	0.4631	0.8912
LCT051-17	0.9845	0.9103	0.9787	0.4721	0.8725
LCT051-18	0.9811	0.9066	0.9735	0.4624	0.8903
LCT051-19	0.9780	0.8987	0.9658	0.4552	0.9021

4.3 Evaluation of g Integral Parameter and Computational Penalty

The nuclear fuel cask applications were analyzed using the g parameter method to determine if any of the benchmarks are more sensitive to the ^{10}B capture than the applications. The maximum g value for ^{10}B capture and the benchmark that provides the maximum g value are listed in Table 4 for each cask application. Sensitivities for ^{235}U fission and ^1H scatter along with the maximum g values for these nuclide-reaction pairs are also listed in Table 4. As the g values in the table show, the benchmarks provide poor coverage for ^1H scatter for the GBC-32 cask. This is the reason that the E_{sum} value for this cask is small.

The sensitivity profiles for each application and the corresponding benchmark that provides the greatest g value are shown in Figures 10 through 14. As these g values indicate, none of the benchmarks exhibit ^{10}B capture sensitivities greater than those for the individual applications across the entire energy range. For the energy groups for which the application exhibits greater sensitivity to the ^{10}B capture cross section than the benchmark, the sum of differences in the sensitivities makes up more than 20% (more than 70% for the GA-4 cask) of the application's sensitivity to the ^{10}B capture cross section. Note that $(1 - g)$ is defined as the summed difference between the application and the benchmark for all energy groups where the application's sensitivity is greater than that of the benchmark, normalized with respect to the application's total sensitivity (see Section 2.1.3). Sensitivities for ^{235}U fission and ^1H scatter for the applications are two orders of magnitude higher than the ^{10}B capture sensitivities. The maximum g values for ^{235}U fission and ^1H scatter are much higher than the maximum g values for the ^{10}B capture indicating that the benchmarks provide good coverage for ^{235}U fission and ^1H scatter but that the coverage for ^{10}B capture is poor. Also, the applications are much more sensitive to the ^{235}U fission and ^1H scatter than to ^{10}B capture. Therefore, the E_{sum} and c_k values that were reported in the previous sections indicate good system-wide coverage despite poor coverage for ^{10}B capture.

Table 4 Maximum g values for all applications

Cask	Maximum g value for ^{10}B capture	Benchmark that provides the highest g value		
MPC-24	0.79	LCT051-10		
MPC-68	0.46	LCT051-10		
GA-4	0.29	PAT80SS1		
GBC-32	0.73	PAT80SS1		
OECD	0.75	LCT051-10		

Cask	^{235}U Fission sensitivity	^1H Scatter sensitivity	Maximum g value for ^{235}U fission	Maximum g value for ^1H scatter
MPC-24	0.325	0.253	0.97	0.93
MPC-68	0.344	0.214	0.93	0.84
GA-4	0.367	0.345	0.99	0.85
GBC-32	0.154	0.233	1.00	0.68
OECD	0.376	0.148	0.99	0.86

A comparison of the ^{10}B capture sensitivity profiles for the five casks as shown in Figure 4 gives some clues as to why there are no benchmark experiments that are applicable to the MPC-68 and GA-4 casks relative to the ^{10}B capture reaction. The MPC-68 and GA-4 casks are more sensitive to ^{10}B capture cross section in the higher energy region than are the other three casks. In the GA-4 cask, this difference is very significant. The set of benchmarks does not contain any experiments with a sensitivity profile similar to these two casks.

Further analysis reveals that the five casks have a wide range of surface density values for ^{10}B , meaning that the atoms of ^{10}B per square centimeter (number density times the thickness) of the separating material vary widely (see Figure 2). The GA-4 cask contains more than 100 times the boron surface density of the OECD cask, and the separating material is dense enough to effectively stop almost all of the thermal neutron flux from passing through. As seen in Table 1, the EALF for the GA-4 cask is also significantly higher than that for the other casks. Of the five casks, the MPC-68 has the next highest EALF value and has about 4 times the boron surface density of the OECD casks.

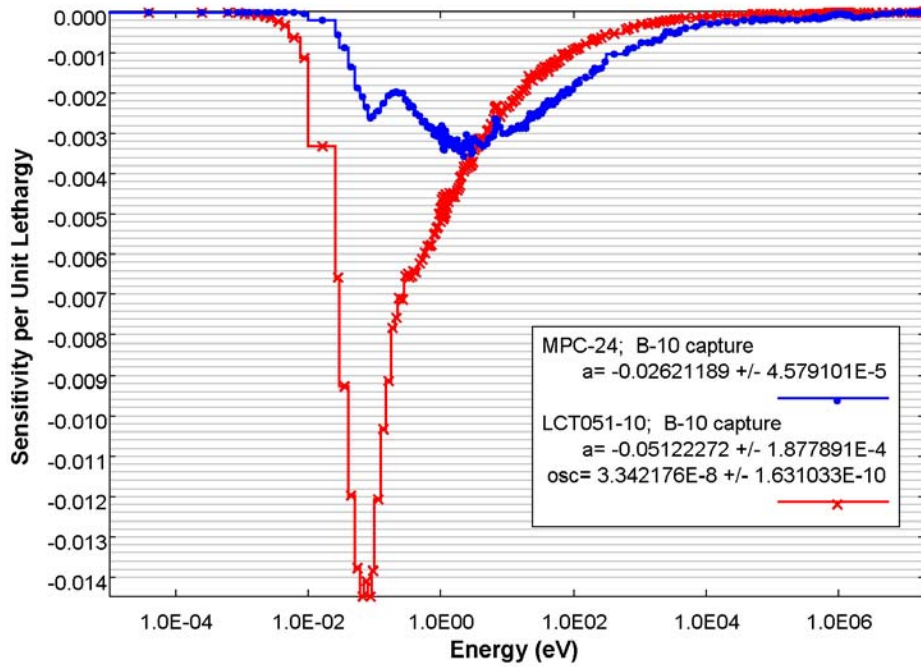


Figure 10 Comparison of ^{10}B capture sensitivity profiles for MPC-24 and LCT051-10

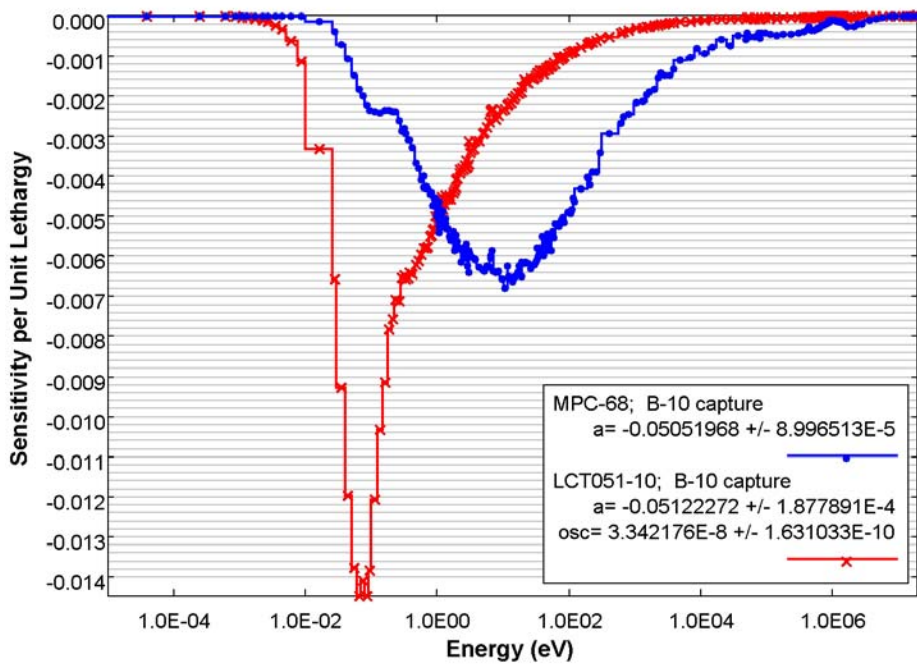


Figure 11 Comparison of ^{10}B capture sensitivity profiles for MPC-68 and LCT051-10

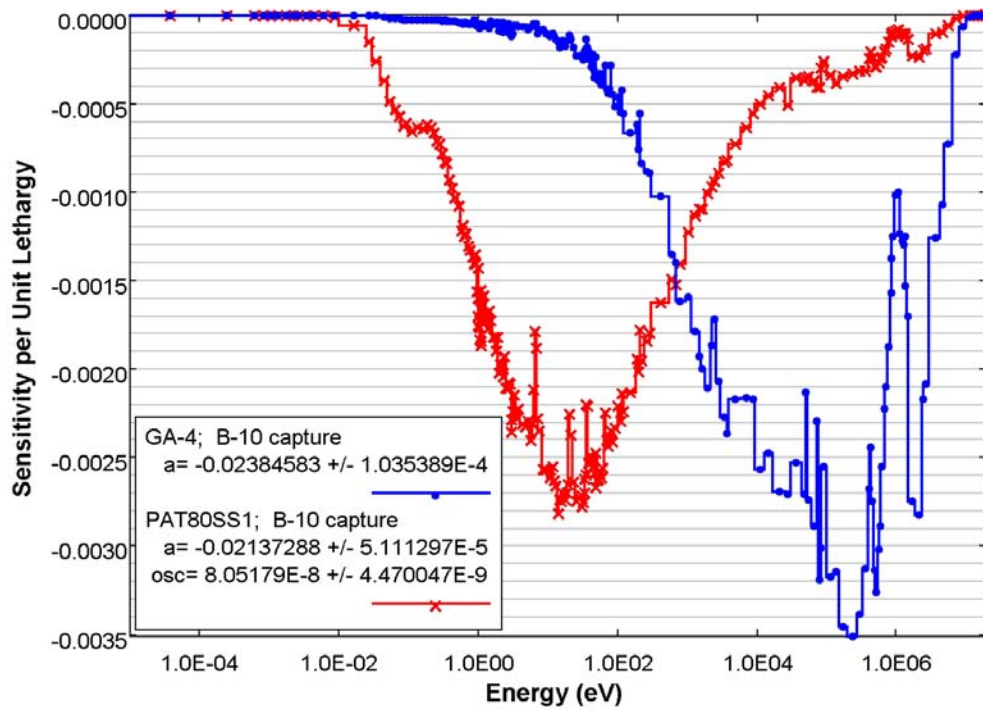


Figure 12 Comparison of ¹⁰B capture sensitivity profiles for GA-4 and PAT80SS1

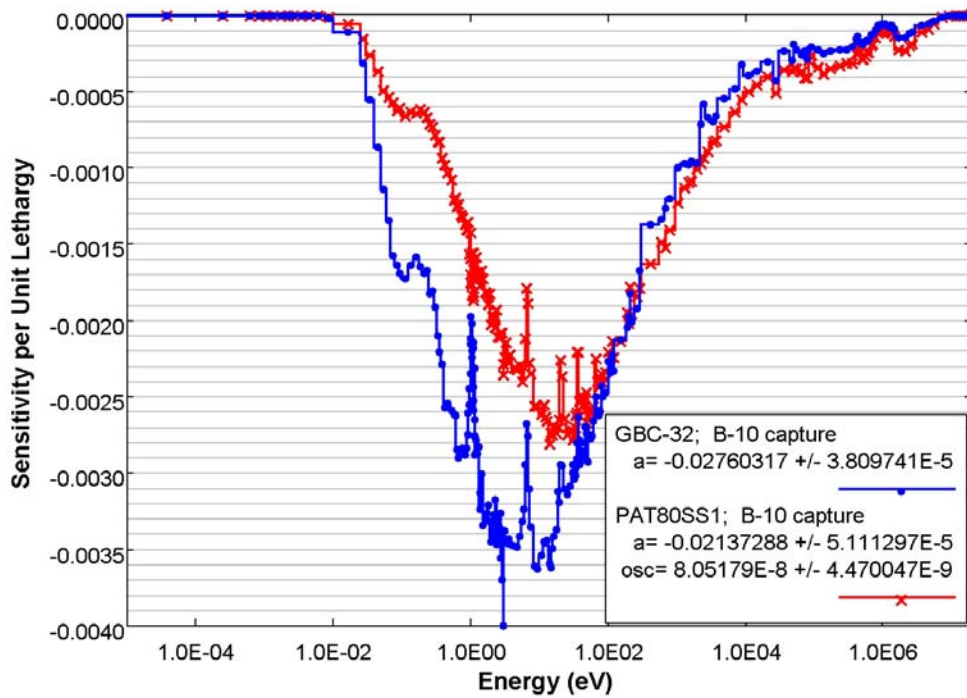


Figure 13 Comparison of ¹⁰B capture sensitivity profiles for GBC-32 and PAT80SS1

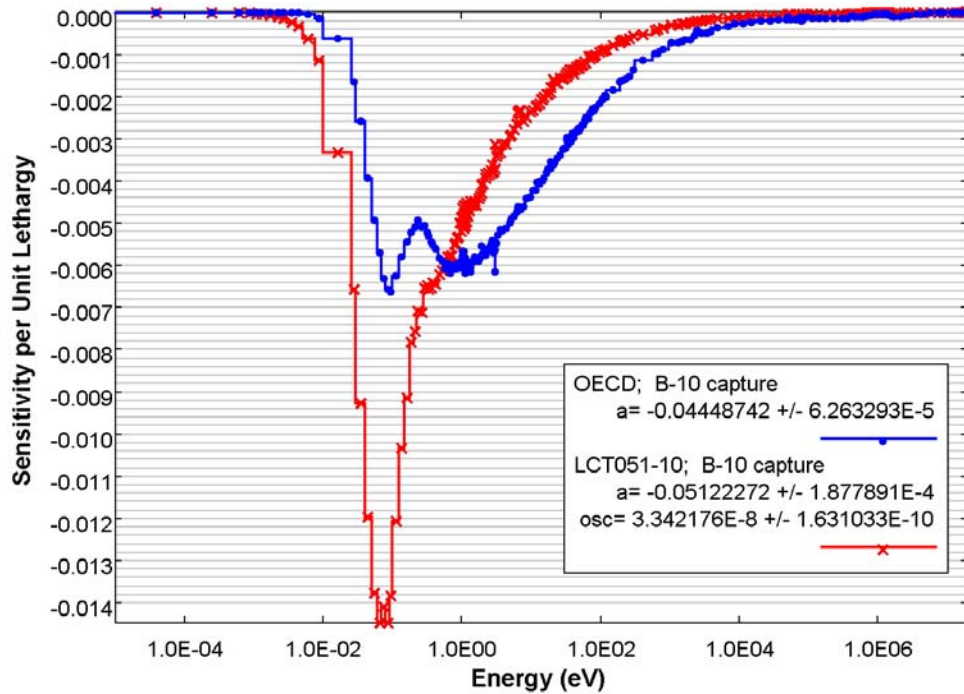


Figure 14 Comparison of ^{10}B capture sensitivity profiles for OECD and LCT051-10

To investigate the effect of the higher boron surface density on the ^{10}B capture sensitivity profiles, the MPC-68 and GA-4 cask models were altered by reducing the boron concentration until the k_{eff} for the cask was approximately one. The TSUNAMI-3D sequence was used to calculate the energy-dependent sensitivity values that were used to plot sensitivity profiles for ^{10}B capture. Figure 15 shows a comparison of the profiles before and after the boron concentration was reduced. Because the sensitivity values at the extremes of the energy range (high and low energy boundaries) are very small compared to the peak values, the differences are most pronounced when using a log-log scale plot. Since negative values cannot be plotted on a logarithmic scale, the negative sensitivities have been inverted in sign before plotting. This procedure is indicated in the legend of the plot by labeling these lines accordingly. Both modified casks show sensitivity profiles that are much more similar to those of the other three cask models shown in Figure 4 (i.e., the energy of the highest sensitivity is around 1 eV). Note that even the modified casks have very small ^{10}B capture sensitivity values, indicating that the casks are not very sensitive to the ^{10}B capture.

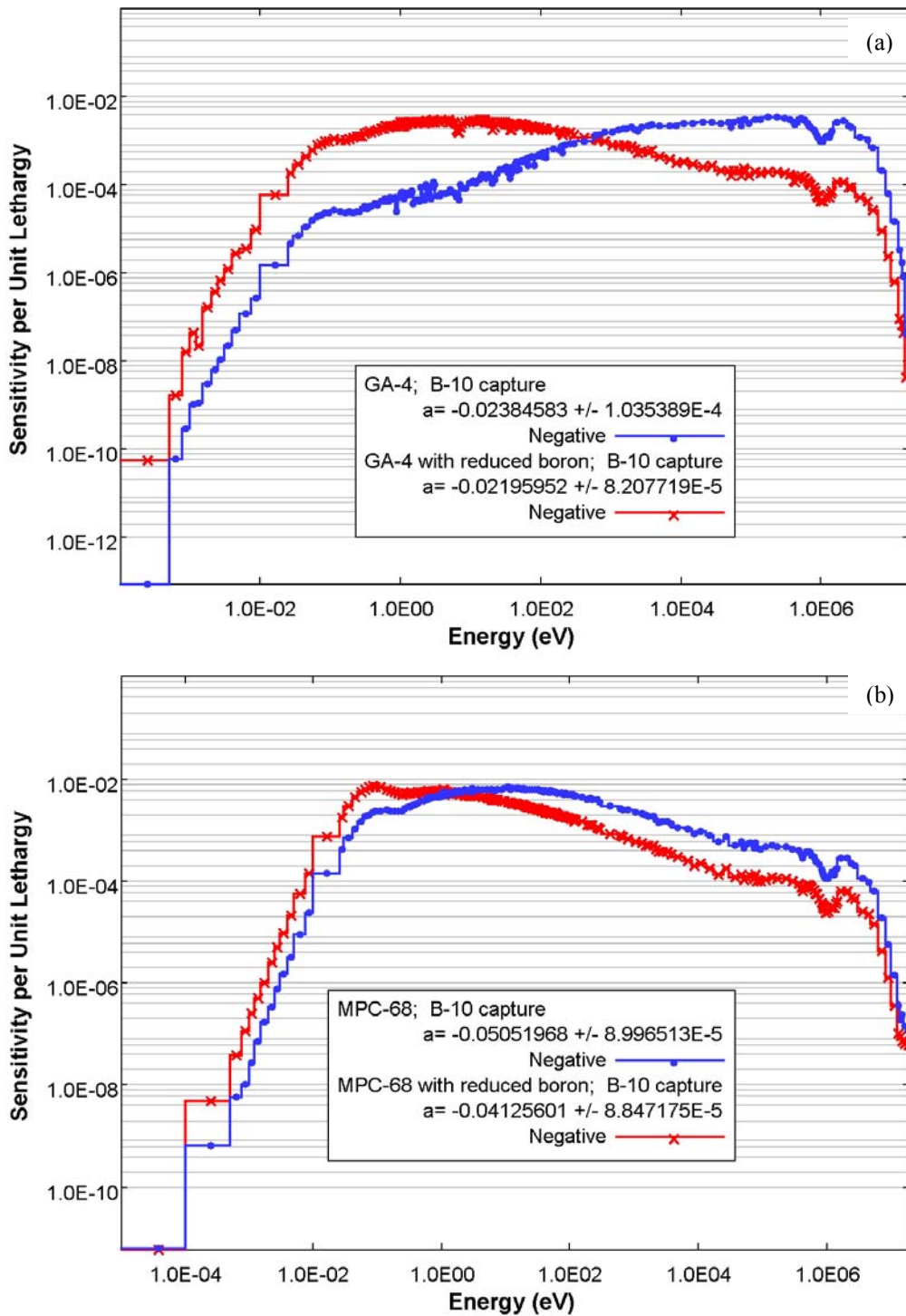


Figure 15 Comparison of ¹⁰B capture sensitivity profiles for casks with (a) normal and (b) reduced boron concentrations

To determine if the set of benchmarks, when used together, provides better coverage than the benchmark that results in the highest g value for the application casks, the ^{10}B capture sensitivity differences were analyzed on a group-wise basis. The number of energy groups (out of 238) that are not covered by any of the benchmarks and the sum of the minimum differences in the group-wise sensitivity coefficients (Z_a) are presented in Table 5. The number of benchmarks that provide coverage for ^{10}B capture for each energy group for each application are plotted in Figures 16 through 22 as a function of neutron energy group. As the results indicate, the greatest sensitivity differences are observed for the MPC-68 and GA-4 casks. However, the highest of these differences is on the order of 10^{-2} , which corresponds to a difference on the order of 10^{-4} in the value of calculated k_{eff} (fourth digit of the calculated k_{eff}) for a 1% change in the ^{10}B capture cross section.

A composite profile is one that is created by comparing, on a group-wise basis, the sensitivity value of an application with that of all benchmarks and selecting the sensitivity value of the benchmark that is closest to the application's value. If some benchmarks have larger sensitivity values than that of the application, then the sensitivity value of the composite profile is set equal to the application's sensitivity value. Mathematically the composite profile can be expressed as $S_{composite}^{j,x,i} = S_a^{j,x,i} - Z_a^{j,x,i}$, where $Z_a^{j,x,i}$ is the minimum difference in the group-wise sensitivity coefficient for nuclide j , reaction x , and neutron energy group i and where $S_a^{j,x,i}$ is the application's sensitivity. Note that the benchmark that provides the closest sensitivity may be different for each group. Thus, the composite profile shows the energy groups where none of the benchmarks has a sensitivity as great as the application and how the sensitivity of the most sensitive benchmark for this energy group compares with that of the application.

The ^{10}B capture sensitivity profiles for the application casks have been plotted against the composite profiles in Figures 23 through 29. As evidenced in these figures, the MPC-68 and GA-4 casks have the largest noncoverage.

Table 5 Coverage by group

Cask	Number of groups without coverage	Sum of minimum sensitivity differences in ^{10}B capture (Z_a)
MPC-24	28	-4.76E-04
MPC-68	179	-1.91E-02
GA-4	69	-1.69E-02
GBC-32	72	-1.65E-03
OECD	107	-5.73E-03
MPC-68m*	83	-3.16E-03
GA-4m*	14	-1.81E-04

* Modified by reducing boron concentration.

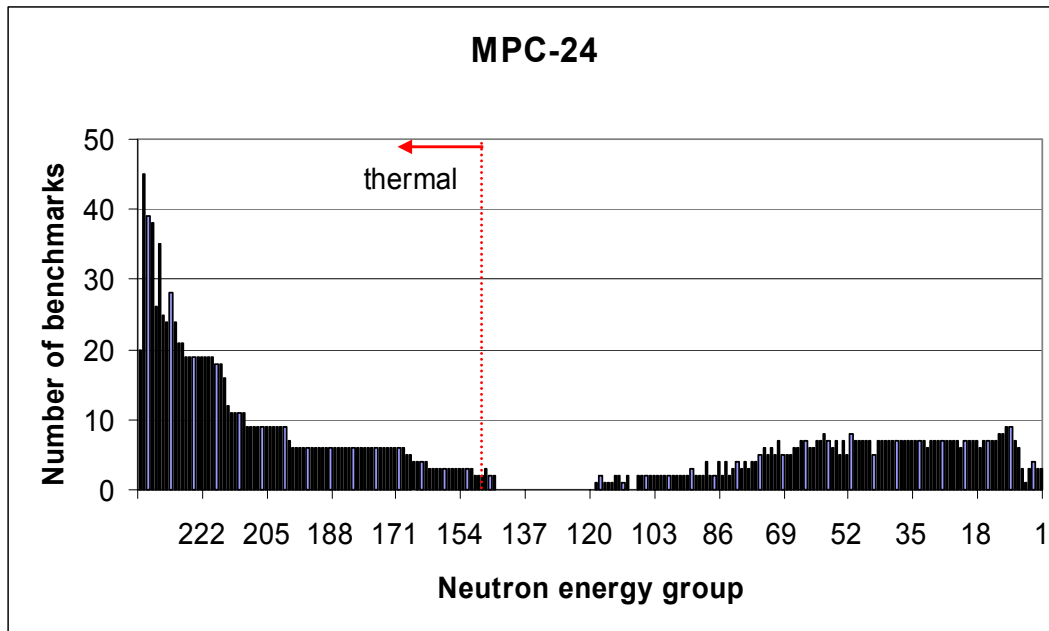


Figure 16 Number of benchmarks that cover ^{10}B capture for each energy group for MPC-24

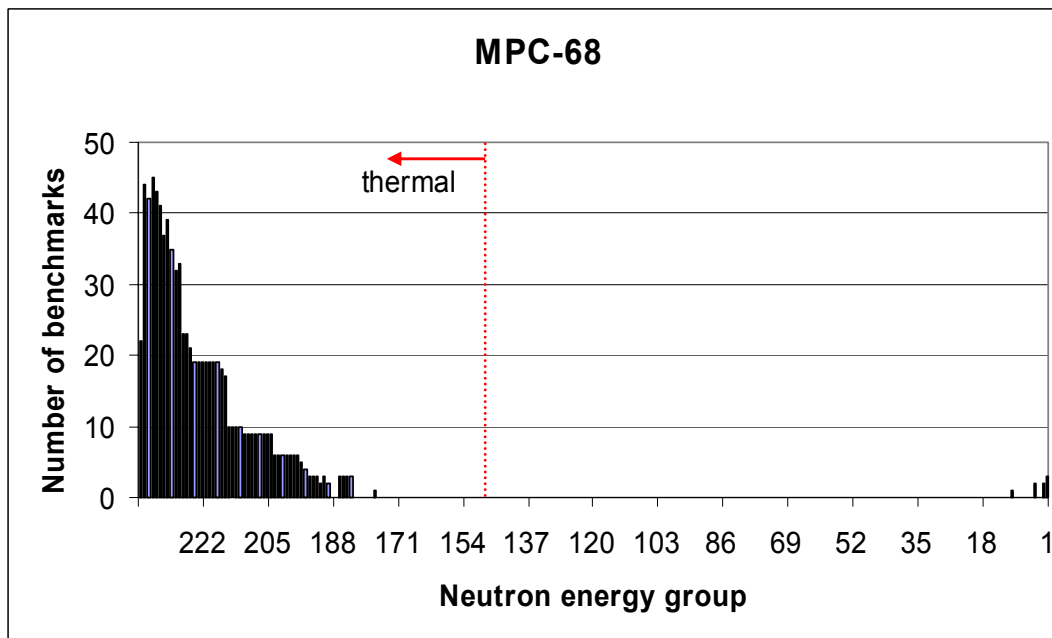


Figure 17 Number of benchmarks that cover ^{10}B capture for each energy group for MPC-68.

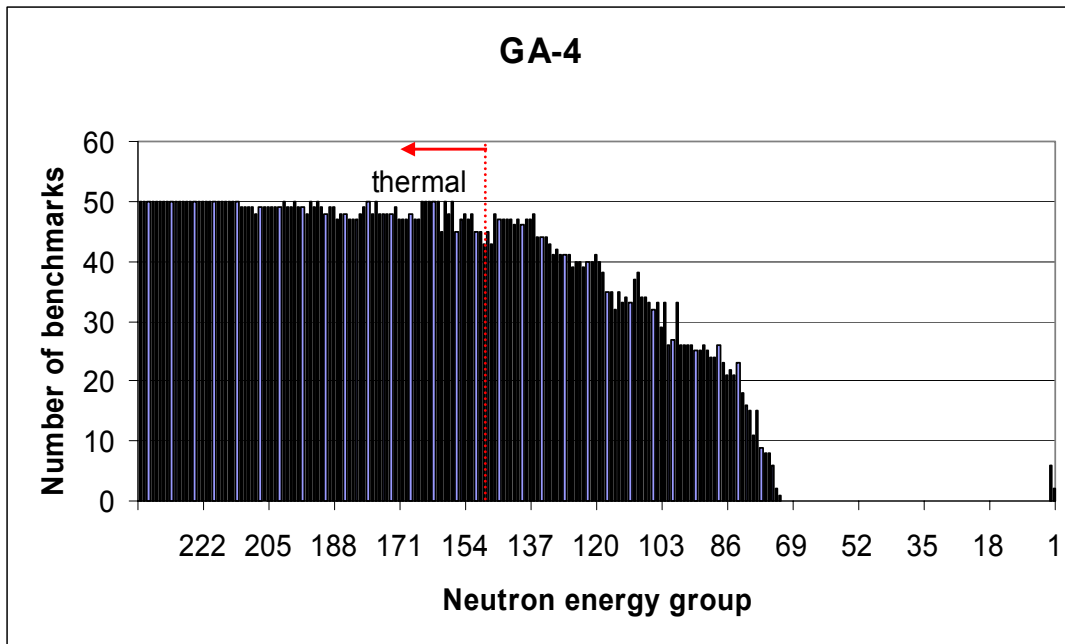


Figure 18 Number of benchmarks that cover ^{10}B capture for each energy group for GA-4

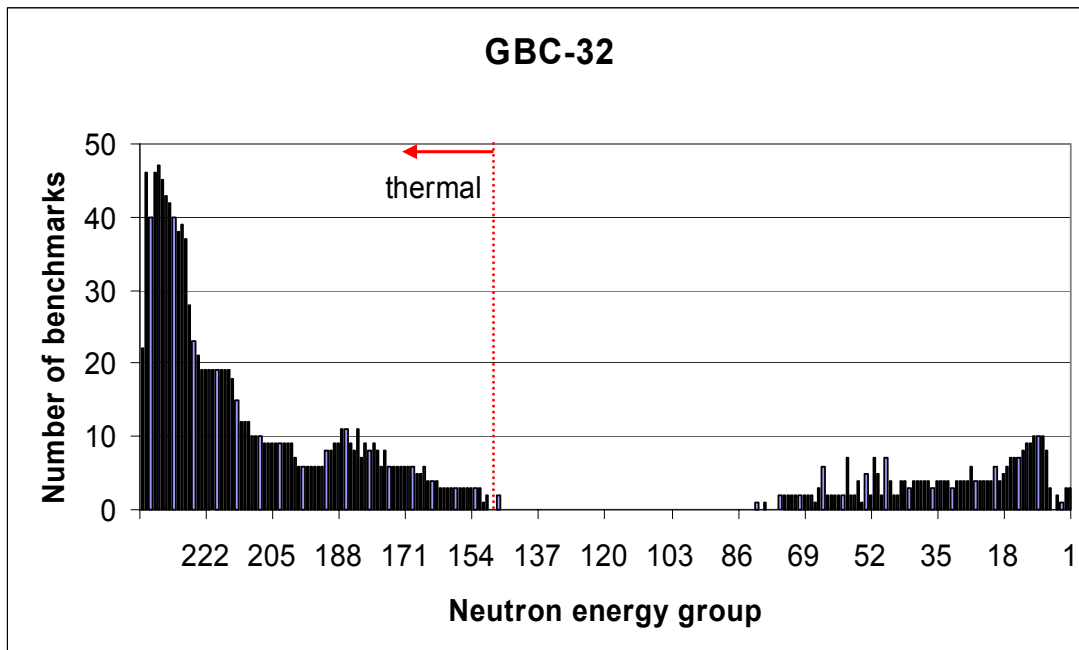


Figure 19 Number of benchmarks that cover ^{10}B capture for each energy group for GBC-32

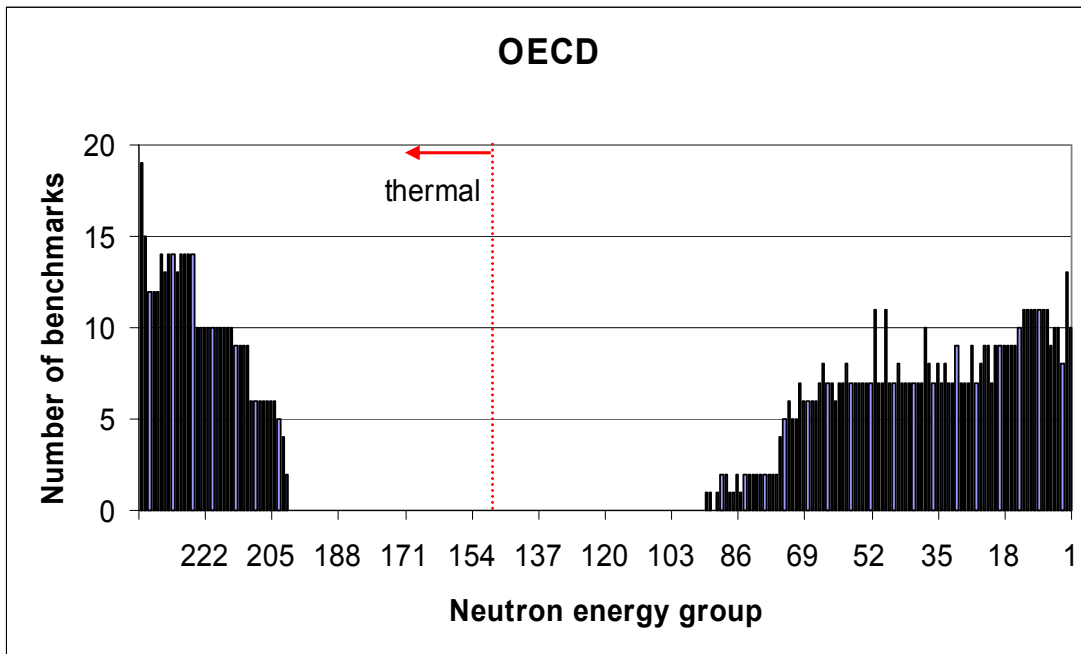


Figure 20 Number of benchmarks that cover ^{10}B capture for each energy group for OECD

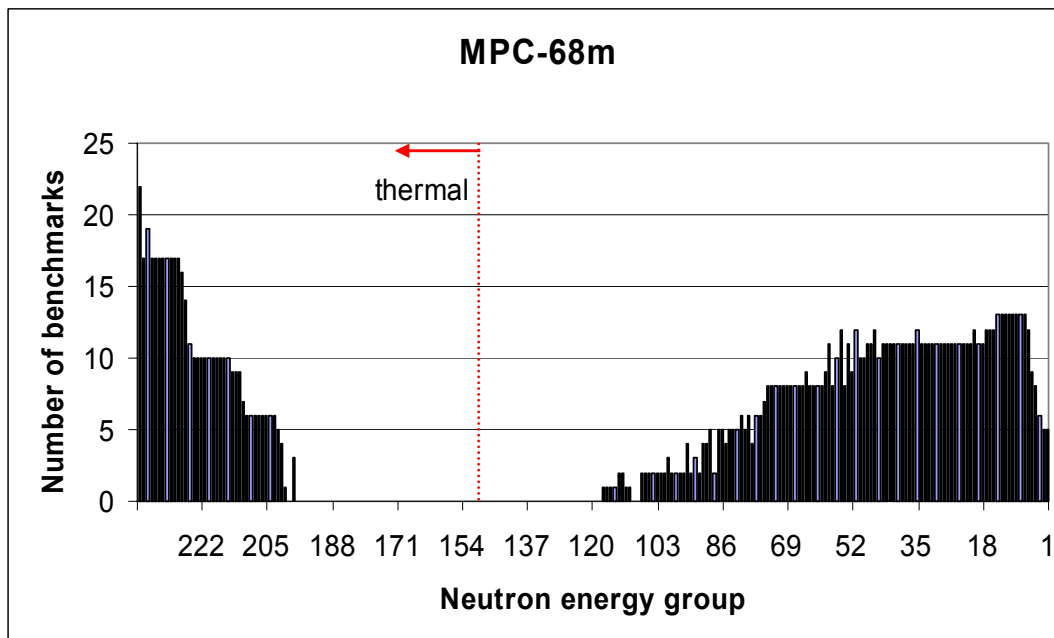


Figure 21 Number of benchmarks that cover ^{10}B capture for each energy group for modified MPC-68

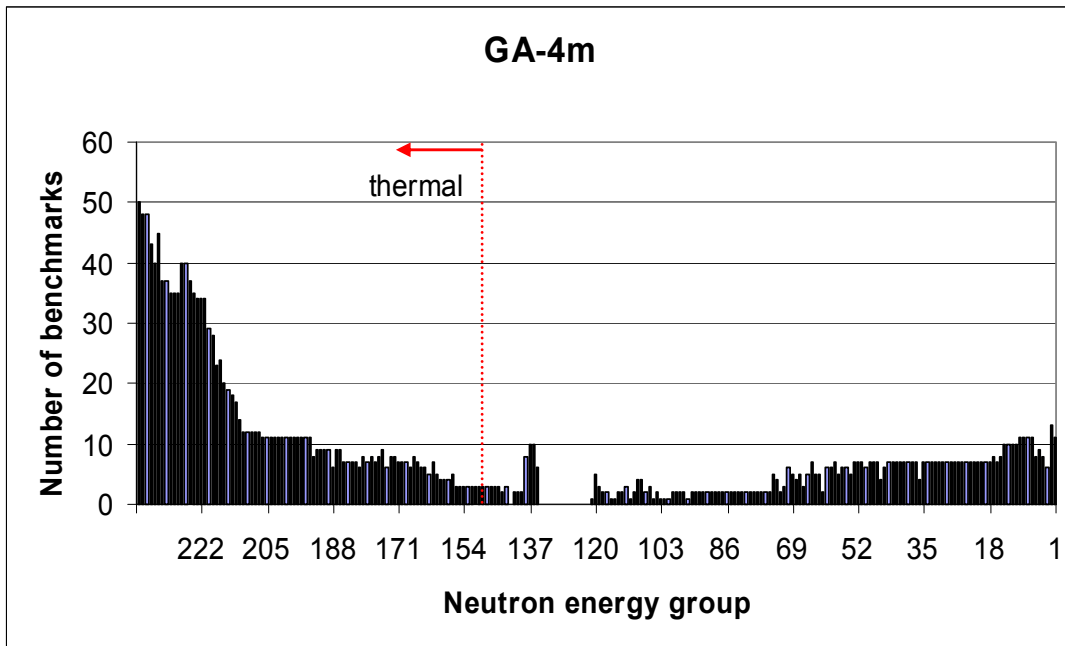


Figure 22 Number of benchmarks that cover ^{10}B capture for each energy group for modified GA-4

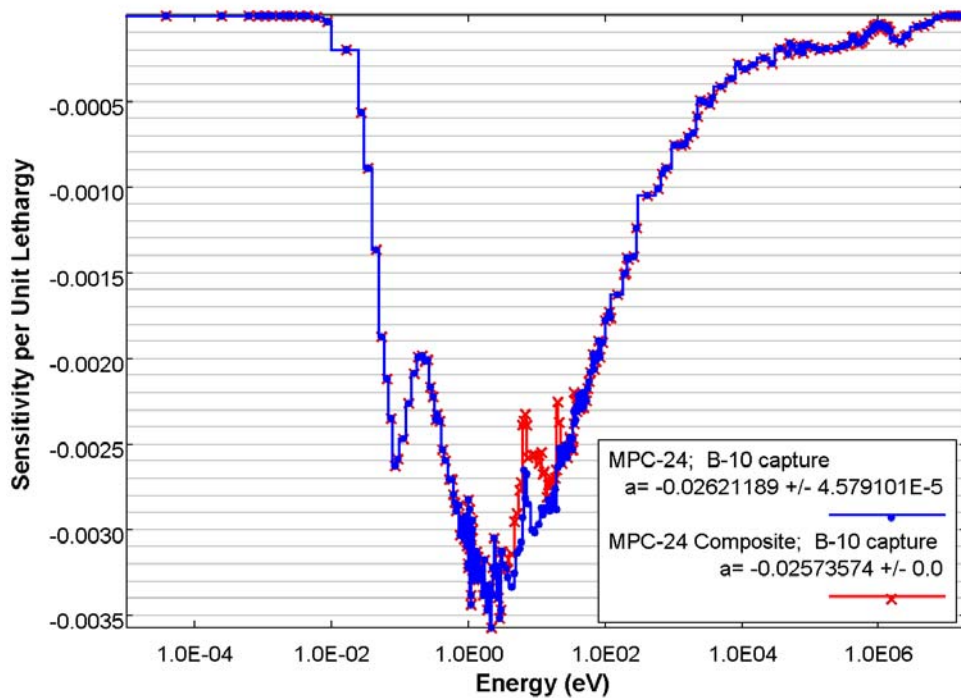


Figure 23 Comparison of MPC-24 ^{10}B capture sensitivity profile and composite profile for MPC-24

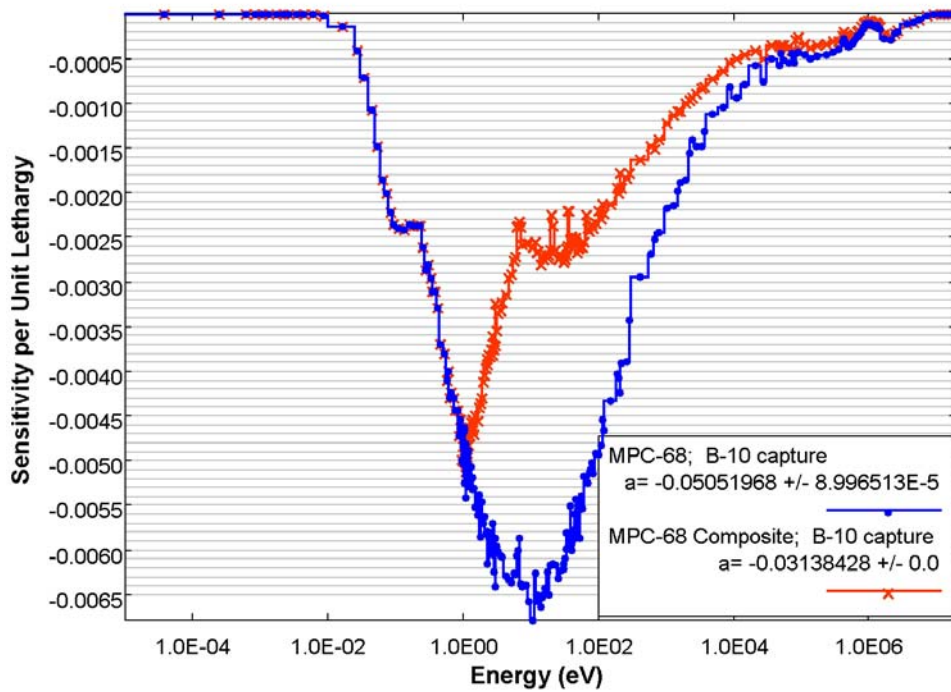


Figure 24 Comparison of MPC-68 ¹⁰B capture sensitivity profile and composite profile for MPC-68

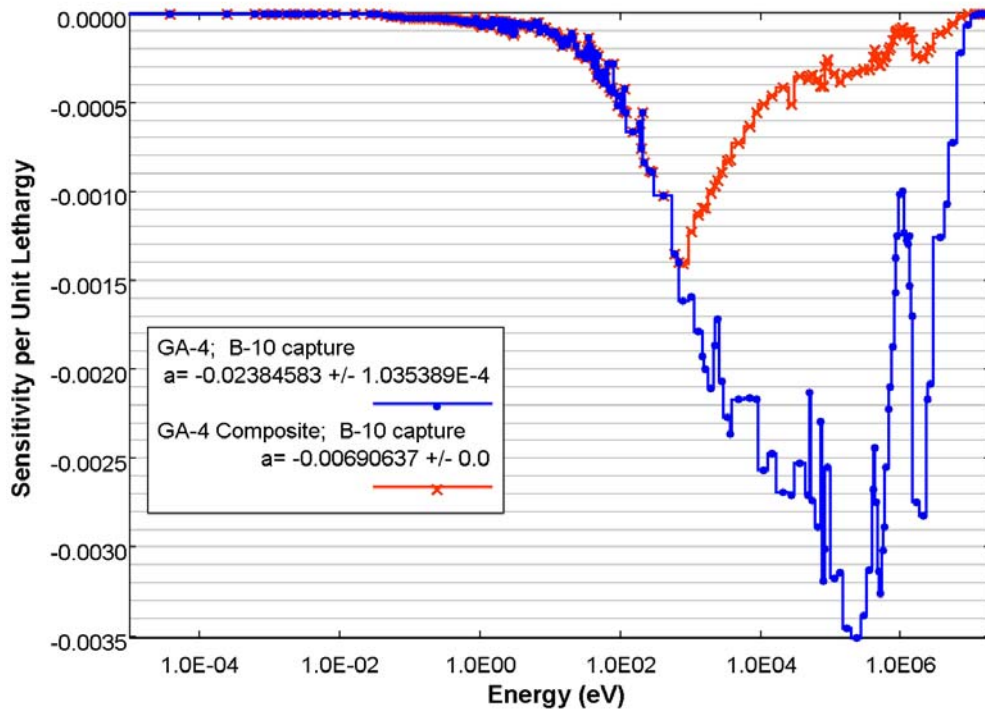


Figure 25 Comparison of GA-4 ¹⁰B capture sensitivity profile and composite profile for GA-4.

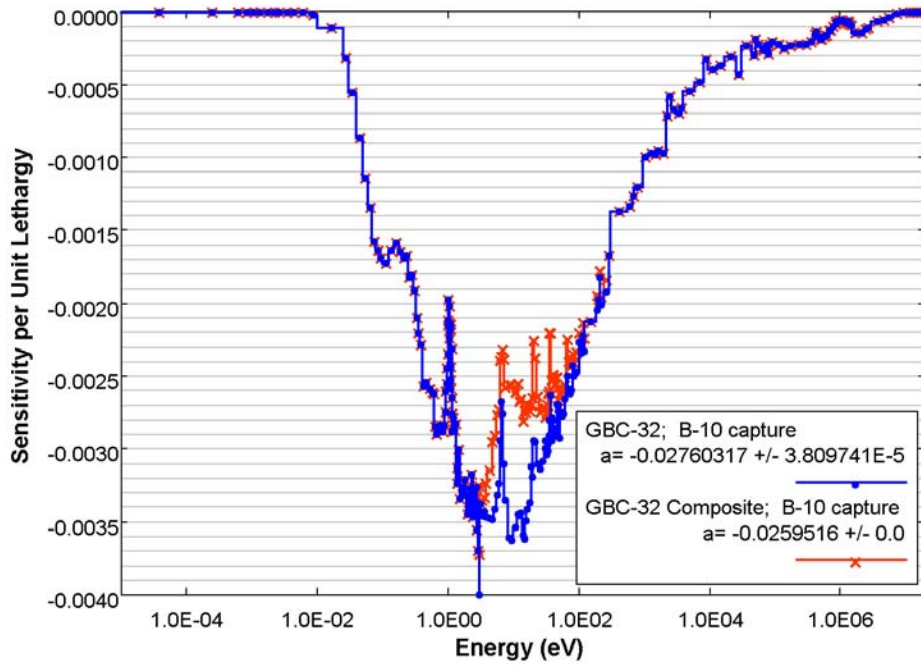


Figure 26 Comparison of GBC-32 ¹⁰B capture sensitivity profile and composite profile for GBC-32

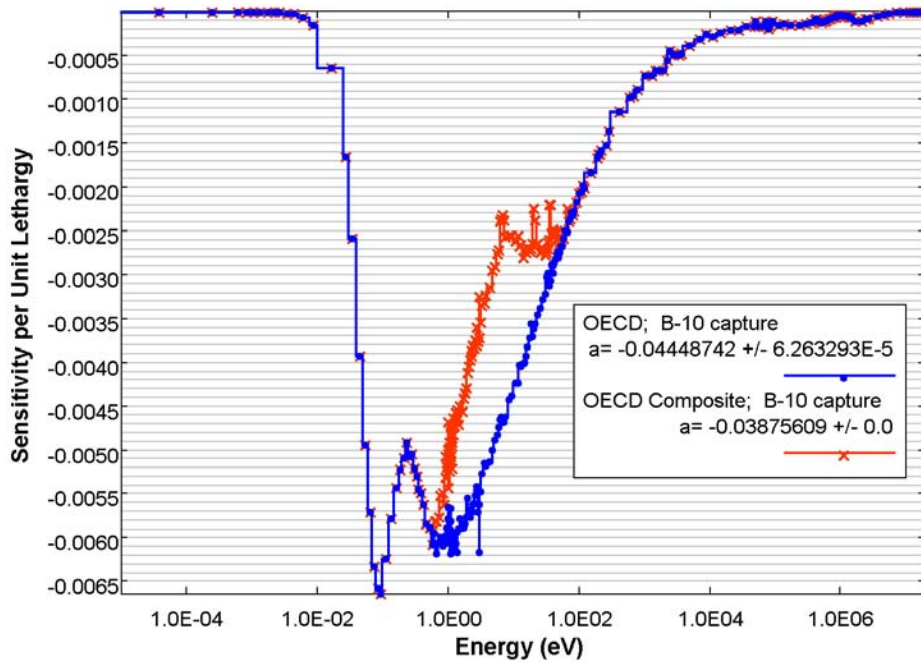


Figure 27 Comparison of OECD ¹⁰B capture sensitivity profile and composite profile for OECD

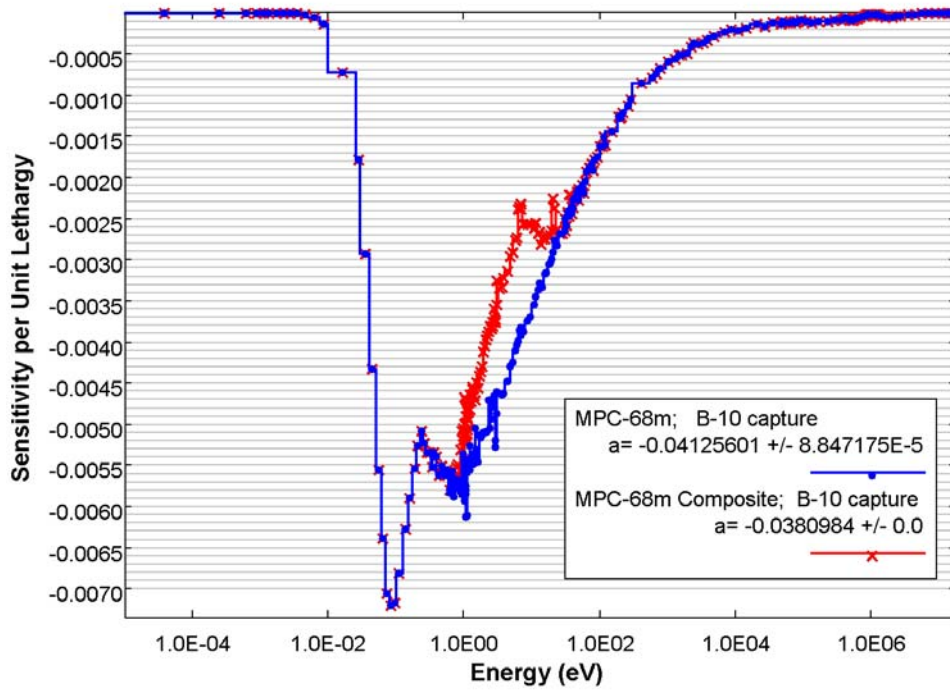


Figure 28 Comparison of modified MPC-68 ¹⁰B capture sensitivity profile and composite profile for modified MPC-68

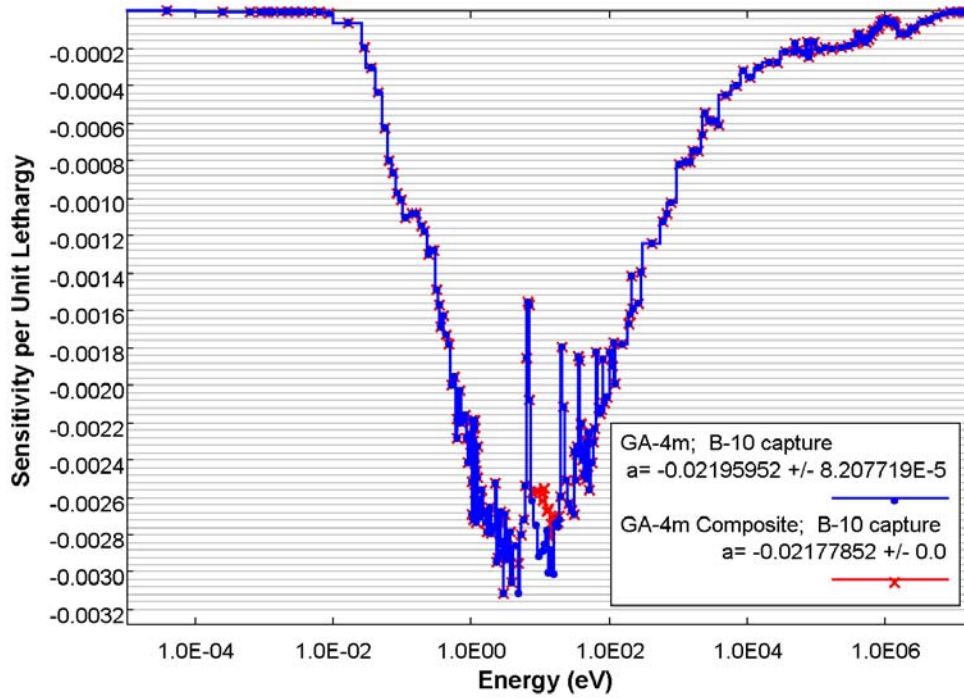


Figure 29 Comparison of modified GA-4 ¹⁰B capture sensitivity profile and composite profile for modified GA-4

Finally, the calculated k_{eff} values for the application casks have been adjusted for the extent of benchmark coverage by applying the methodology from Section 2.2. The ^{10}B (n, α) covariance data have been used in the penalty assessment in lieu of capture covariance data, because the (n, α) reaction sensitivity is four orders of magnitude greater than all other reaction sensitivities that make up the capture reaction. In calculating the adjusted k_{eff} , there was no need to create a hypothetical matrix of covariance data for ^{10}B capture since the cross-section library contained the ^{10}B capture [(n, α)] cross-section uncertainty data. For all nuclide-reaction pairs in the problem for which no covariance data exist in the library, a default value of 0.2 (standard deviation value of 20%) has been used for establishing similarity using c_k values for penalty assessment. When a default standard deviation value for the nuclide-reaction pairs without covariance data is used, the calculated c_k values differ from the ones with no default standard deviation values. Therefore, the c_k values that were used to determine the similarity, and thereby the penalty, were somewhat different from the c_k values listed in Section 4.2.

In addition, although the c_k values (with default standard deviation value of 0.2) for the GBC-32 and OECD casks are all below the c_k similarity criteria of 0.9, corresponding computational penalties were still calculated by changing the similarity criteria to 0.0 (i.e., all benchmarks are considered similar to the application). The calculated computational penalties for these casks were deemed appropriate because the sums of the minimum sensitivity differences for ^{10}B capture for both casks are very small indicating that the ^{10}B capture sensitivities in these casks are very similar to those in the experiments. Calculated values for the application casks, along with the adjusted k_{eff} values, are listed in Table 6. Because the penalties in the calculated k_{eff} values due to noncoverage of ^{10}B capture cross section are less than or equal to 10^{-4} (10^{-2} %), the adjusted k_{eff} is identical to the calculated k_{eff} with 4-digit precision. Therefore, the penalty due to noncoverage of ^{10}B capture cross sections is determined to be zero for all practical purposes.

Table 6 Final results

Cask	Calculated k_{eff}	Maximum g for ^{10}B capture	Sum of minimum sensitivity differences in ^{10}B capture ($Z_a^{j,x}$)	Penalty ($\% \Delta k_{eff}/k_{eff}$) due to ^{10}B capture only	Adjusted k_{eff}
MPC-24	0.9458	0.79	-4.76E-04	0.0001	0.9458
MPC-68	0.9349	0.46	-1.91E-02	0.0044	0.9349
GA-4	0.9221	0.29	-1.69E-02	0.0077	0.9221
GBC-32	0.8941	0.73	-1.65E-03	0.0009	0.8941
OECD	1.1303	0.75	-5.73E-03	0.0013	1.1303
MPC-68m*	1.0012	0.83	-3.16E-03	0.0010	1.0012
GA-4m*	0.9959	0.86	-1.81E-04	0.0000	0.9959

* Modified by reducing boron concentration.

5 SUMMARY

The amount of ^{10}B in the nuclear fuel cask applications that were analyzed in this study causes the applications to be more sensitive in many energy groups to the ^{10}B capture cross section than most of the currently available experimental benchmarks that were used in this study. Analyses of these applications with the S/U techniques indicate that, with the exception of MPC-68 cask, the sensitivity for the majority of the energy groups is covered by the composite sensitivity profile of benchmarks that are more sensitive to the ^{10}B capture cross sections than the applications. The differences between the application and composite benchmark sensitivities are very low compared with the subcritical margin that may typically be imposed on these applications. Specifically, the differences amount to changes in k_{eff} on the order of 0.01% (for a 1% change in the cross sections) whereas the subcritical margin is a few percent.

Based on the E_{sum} and c_k values, it appears that all applications, except for the GBC-32 cask, can be validated with the set of benchmarks that were used in this study. However, the E_{sum} and c_k values are dominated by ^{235}U fission and ^1H scattering, as the sensitivities of the applications to these reactions are much higher than their sensitivities to the ^{10}B capture reaction. The g values indicate marginal coverage for the MPC-24, GBC-32, and OECD cask applications.

In determining the coverages and thereby the penalty (if any), as many similar (to the application) benchmarks as possible should be included in the validation benchmark set. The method will take advantage of the similar benchmarks and limit the extent of the noncoverage. If benchmarks that are very similar to the application are omitted or are not available, then the method will yield a higher penalty to reflect that effect. The final penalty value will still depend on the sensitivity to the nuclide-reaction pair of interest and on the uncertainty associated with that pair.

6 CONCLUSIONS

Although the sensitivities of the nuclear fuel cask applications that have been analyzed are not completely covered by the set of benchmarks that were used in this study with regard to the ^{10}B capture cross section, analysis of the coverage using the g parameter method indicates that the effect of lack of coverage on the k_{eff} is minimal. Thus, no additional bias due to boron loading differences should be imposed, and the experimental biases and uncertainties are appropriate for the cask applications.

The above conclusion can readily be extended to other cask applications with similar boron loadings that have similar ^{10}B capture sensitivities. The method will yield small penalty values for the applications that are covered well by the benchmarks. If the noncoverage for an application is large, the method will penalize with a magnitude proportional to the sensitivity to the nuclide-reaction pair and to the uncertainty associated with that pair. The penalty may be small if the system is not sensitive to the nuclide-reaction pair of interest or if the cross sections for that pair are very well known.

7 REFERENCES

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APPENDIX A

A.1 Benchmarks

The benchmarks from International Handbook of Evaluated Criticality Safety Benchmark Experiments (IHECSBE) and NUREG/CR-6361 are listed in the following table. The critical experiments from the benchmark designated by LEU-COMP-THERM-034 in IHECSBE were also investigated for inclusion in the list of benchmarks for the analyses. However, no KENO V.a input models were readily available. Based on the information the evaluators provided in the benchmark, the maximum sensitivity of the experiments to ^{10}B (total) was calculated as -0.036. This value is within the range of sensitivity values from the experiments listed in Table A.1. (The calculated total and capture sensitivities are within 1% of each for all benchmarks in the table.) Therefore, it was decided that the inclusion of LEU-COMP-THERM-034 series experiments would not improve the results considerably and that the conclusions would not change.

Table A.1 List of benchmarks

Case numbers from Figure 1	IHECSBE identifier	NUREG/CR-6361 identifier	Separating material	EALF (eV)	^{10}B Capture sensitivity coefficient
1	LCT009-05		B-Steel	1.156E-01	-2.127E-03
2	LCT009-06		B-Steel	1.150E-01	-1.135E-03
3	LCT009-07		B-Steel	1.160E-01	-2.452E-03
4	LCT009-08		B-Steel	1.151E-01	-1.043E-03
5	LCT009-09	P2615BA	Boral	1.157E-01	-1.345E-03
6	LCT012-02	P3314BS1	B-Steel	1.814E-01	-9.761E-03
7	LCT012-03	P3314BS2	B-Steel	1.823E-01	-9.587E-03
8	LCT012-04		Boral	1.900E-01	-9.351E-03
9	LCT012-05		Boral	1.896E-01	-8.933E-03
10	LCT012-06		Boroflex	1.891E-01	-9.496E-03
11	LCT012-07		Boroflex	1.897E-01	-9.458E-03
12	LCT013-02	P3602BS2	B-Steel	3.073E-01	-6.139E-03
13	LCT013-03	P3602BB	Boral	3.104E-01	-4.248E-03
14	LCT013-04		Boroflex	3.095E-01	-4.627E-03
15	LCT016-08		B-Steel	9.878E-02	-2.076E-03
16	LCT016-09		B-Steel	9.832E-02	-8.885E-04
17	LCT016-10		B-Steel	9.901E-02	-2.114E-03
18	LCT016-11		B-Steel	9.825E-02	-6.364E-04
19	LCT016-12		Boral	9.966E-02	-2.459E-03
20	LCT016-13		Boral	9.859E-02	-9.955E-04
21	LCT016-14	P2438BA	Boral	9.946E-02	-2.058E-03
22	LCT042-02		B-Steel	1.821E-01	-8.942E-03

Appendix A

Table A.1 (continued)

Case numbers from Figure 1	IHECSBE identifier	NUREG/CR-6361 identifier	Separating material	EALF (eV)	¹⁰ B Capture sensitivity coefficient
23	LCT042-03		Boral	1.887E-01	-8.724E-03
24	LCT042-04		Boroflex	1.871E-01	-8.263E-03
25		P3314BA	Boral-A	3.279E-01	-1.686E-02
26		P3314BC	Boral-C	3.221E-01	-2.013E-02
27		P3314BF1	Boroflex	3.195E-01	-2.058E-02
28		P3314BF2	Boroflex	3.231E-01	-1.691E-02
29		P3314BS3	B-Steel	2.980E-01	-9.252E-03
30		P3314BS4	B-Steel	2.996E-01	-8.821E-03
31		P3602BS1	B-Steel	1.819E-01	-9.280E-03
32		P62FT231	Boral	3.671E-01	-7.235E-03
33		P71F14F3	Boral	3.825E-01	-1.452E-02
34		P71F14V3	Boral	3.742E-01	-1.111E-02
35		P71F14V5	Boral	3.739E-01	-8.673E-03
36		P71F214R	Boral	3.737E-01	-7.837E-03
37		PAT80L1	Boral	1.518E-01	-2.095E-02
38		PAT80L2	Boral	1.458E-01	-1.932E-02
39		PAT80SS1	Boral	1.521E-01	-2.137E-02
40		PAT80SS2	Boral	1.469E-01	-1.952E-02
41	LCT051-10	BW1484A1	Boral	2.007E-01	-5.122E-02
42	LCT051-11		Boral	2.007E-01	-5.025E-02
43	LCT051-12		Boral	2.026E-01	-6.342E-02
44	LCT051-13		Boral	2.088E-01	-1.128E-01
45	LCT051-14		Boral	1.754E-01	-7.037E-02
46	LCT051-15		Boral	2.085E-01	-1.255E-01
47	LCT051-16		Boral	1.748E-01	-8.641E-02
48	LCT051-17		Boral	2.096E-01	-1.399E-01
49	LCT051-18		Boral	1.756E-01	-1.070E-01
50	LCT051-19	BW1484A2	Boral	1.559E-01	-5.438E-02

Fifty benchmark experiments were modeled for comparison with the cask models described above. Thirty-four experiments were taken from the IHECSBE, published by the OECD.^{A.1} Sixteen additional experiments were taken from the set used in NUREG/CR-6361.^{A.2} Experiments from the IHECSBE are named using a shortened version of their evaluation identifier. For example, LEU-COMP-THERM-009, experiment 5, is named LCT009-05.

The first 36 experiments in Table A.1 were chosen from various sets of experiments performed by S. R. Bierman and others at the Critical Mass Laboratory at Pacific Northwest Laboratories (PNL). These experiments used various arrangements of fuel pin arrays and neutron poison plates made with Boral, borated stainless steel, or Boroflex and have been the set of experiments normally used in validations for fuel casks. Twenty-four of these experiments have been included in evaluations in the IHECSBE, and the remaining 12 were modeled based on information from Ref. A.2. The experiments were performed with fuel pins arranged in an array that simulated a PWR fuel assembly, and with the assemblies arranged in a 3×1 or 2×2 array separated by the neutron poison plates. The uranium in the fuel pins was enriched to 2.35 or 4.31 wt % ²³⁵U.

The next four experiments in Table A.1 (PAT80x) were performed in France, and used 2×2 arrays of simulated fuel assemblies separated by Boral neutron poison plates.^{A.3} The uranium fuel is enriched to 4.742 wt % ²³⁵U.

The remaining 10 configurations were taken from LEU-COMP-THERM-051. These experiments were performed at the Babcock and Wilcox Lynchburg Research Center and used a 3×3 matrix of simulated fuel assemblies, separated by four neutron poison plates.^{A.4} Each assembly of fuel pins is 14×14 . The uranium enrichment is 2.5 wt % ²³⁵U, and the moderator and reflector are borated water.

A.2 References

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