Verification of the Cardinal Multiphysics Solver for 1-D Coupled Heat Transfer and Neutron Transport

L.I. Gross¹, A.J. Novak², P. Shriwise², and P.P.H. Wilson¹

¹University of Wisconsin – Madison

1500 Engineering Drive, Madison, WI 53706

²Argonne National Laboratory 9700 S Cass Avenue, Lemont, IL 60439

ligross@wisc.edu, anovak@anl.gov, pshriwise@anl.gov paul.wilson@wisc.edu

ABSTRACT

Cardinal is a multiphysics software tool that couples OpenMC Monte Carlo transport and NekRS Computational Fluid Dynamics (CFD) to the Multiphysics Object-Oriented Simulation Environment (MOOSE). In this work, we verify Cardinal for coupled heat conduction and neutron transport using a 1-D analytic solution from previous work by the Naval Nuclear Laboratories. This numerical benchmark includes S_2 transport, Doppler-broadened cross-sections, thermal conduction and expansion, and convective boundary conditions. The goals of this work are to verify Cardinal's basic multiphysics modeling capabilities for coupled neutronics and heat conduction. The benchmark provides analytical solutions for the temperature and flux distributions, as well as the k-eigenvalue. Using these solutions, an L_2 error norm was computed for each spatial discretization (finite element heat conduction mesh, Monte Carlo cells). The temperature error showed linear convergence on a log-log plot of mesh element number vs error with a slope of -0.9982. The flux was near linear convergence, with a slope of -0.9496 on a log-log plot of number of cells vs error, but the higher-than-expected error of the last element deviated from linear behavior. The eigenvalue k_{eff} agrees well with the benchmark value for each mesh size. The outcome of this work is verification of coupled Monte Carlo-thermal conduction modeling using Cardinal.

KEYWORDS: Cardinal, MOOSE, OpenMC, multiphysics, verification

1. INTRODUCTION

With recent advancements in methods, software, and computing, high-fidelity multiphysics Modeling and Simulation (M&S) is becoming an important component of the nuclear engineer's "toolbox." These high-fidelity models substitute more conservative safety factors with predictive science. This can reduce uncertainty in analyses, enabling tighter margins to realize improved economics and licensing certainty. However, analytical benchmarks and comparison to experimental data are required to assess the stability, convergence, and predictive capability of these high-fidelity models for reactor design and analysis.

Cardinal [?,1] (https://cardinal.cels.anl.gov/) is an open-source code that couples OpenMC [2] Monte Carlo particle transport and NekRS CFD to MOOSE [3]. This coupling brings high-fidelity multiphysics feedback to the MOOSE "ecosystem." Cardinal couples OpenMC and NekRS to MOOSE simulations by copying data between the internal code data structures (e.g. a vector of tally results in OpenMC) and a MooseMesh, or the unstructured mesh class in MOOSE. MOOSE's mesh-to-mesh interpolation system then communicates between the MooseMesh "mirror" of the external code's solution and an arbitrary coupled MOOSE application in the form of boundary conditions (such as for conjugate heat transfer with NekRS) or source terms (such as for volumetric heating with OpenMC). Convergence is obtained with Picard iteration.

For coupled neutronics-thermal-fluid simulations with OpenMC, each Picard iteration consists of several steps: 1) a MOOSE application (e.g. BISON, Pronghorn, NekRS via Cardinal, ...) solves for temperatures

and densities; 2) Cardinal transfers temperatures and densities to the OpenMC model; 3) OpenMC solves for the nuclear heating; and 4) Cardinal transfers the tally values to the MooseMesh "mirror."

These steps continue until convergence criteria is achieved. In this work, we pursue verification of these multiphysics aspects of Cardinal using a 1-D analytical benchmark from the Naval Nuclear Laboratories [4]. This work does not require CFD, and thus NekRS will be left out of the discussion from this point on.

The remainder of this paper is organized as follows. In Section 2, we summarize the analytical benchmark modeled in this work. Section 3 then describes the Cardinal computational model of the benchmark. Section 4 presents comparisons between Cardinal and the analytical benchmark. Finally, Section 5 presents conclusions and outlines ongoing and future efforts in the verification and validation of Cardinal.

2. BENCHMARK PROBLEM DESCRIPTION

The analytical benchmark couples three physics: S_2 neutron transport with Doppler broadening, heat conduction, and thermal expansion. S_2 transport restricts the neutron direction to only the $\pm x$ direction. A summary of the governing Ordinary Differential Equations (ODEs) and boundary conditions in the 1-D slab is shown in Fig. 1.

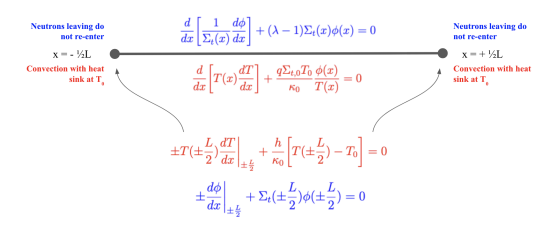


Figure 1: The domain, ODEs, and boundary conditions for the slab.

This benchmark uses a one-group assumption for the neutron cross-sections. As neutrons transport, heat from fission events deposit volumetric power in the slab, causing thermal expansion and affecting the temperature distribution; thermal expansion is restricted to only the x-direction. This slab elongation feeds back into neutronics and heat conduction by influencing the domain length and material density. The slab has convective boundary conditions at the endpoints $x = \pm \frac{L}{2}$ with the heat sink temperature T_0 . The Doppler-broadened total, microscopic cross-section follows an inverse-root temperature relationship,

$$\sigma_t(x) = \sigma_{t,0} \sqrt{\frac{T_0}{T(x)}}. (1)$$

Due to thermal expansion, the slab density varies as

$$\rho(x) = \rho_0 \sqrt{\frac{T_0}{T(x)}}. (2)$$

This gives a Doppler-broadened, macroscopic, total cross-section that accounts for changes in density due to temperature as

$$\Sigma_t(x) = \frac{\rho_0 \sigma_{t,0} N_A}{A} \frac{T_0}{T(x)} = \Sigma_{t,0} \frac{T_0}{T(x)},$$
(3)

where $\sigma_{t,0}$ is the total, microscopic cross-section at T_0 , N_A is Avogadro's number, and A is the mass number of the medium. The conduction equation governs heat flow in the slab and can be described in terms of the

thermal conductivity $\kappa[T(x)]$, the energy released per reaction q, the total, macroscopic cross-section Σ_t , and the neutron flux ϕ .

$$\frac{d}{dx} \left[\kappa [T(x)] \frac{dT(x)}{dx} \right] + q \Sigma_t(x) \phi(x) = 0.$$
(4)

Note that typically q represents the energy per fission, but in this benchmark, it is per total reaction. The main assumption used to craft the analytical solution is that $T(x) = f\phi(x)$. This assumption is guaranteed by manipulating the ODEs, inserting the cross-section temperature dependence, and matching coefficients so that the two ODEs are of the same form. The matching of coefficients imposes two constraints that give equations for the total, microscopic cross-section $\sigma_{t,0}$ and the heat transfer coefficient h, in terms of the slab parameters. For full details on the analytical solution's derivation, see [4].

3. COMPUTATIONAL MODEL

This section describes the OpenMC and MOOSE computational models, followed by the convergence criteria used for coupling and a description of the mapping between OpenMC and MOOSE geometries. At the time at which this benchmark modeling was performed, Cardinal did not support moving geometries in OpenMC. Instead of using thermomechanics for thermal expansion, our simulation simply uses the analytic solution for the equilibrium length L from [4] that accounts for all three physics effects. The change in density due to temperature is accounted for in this simulation by using the total, macroscopic cross-section – which absorbs density to become a $\frac{1}{T}$ dependence – from (3), as was mentioned in Section 2. We note that, in a companion paper in this conference [5], that Cardinal now supports mesh-based geometry (and deforming-mesh problems) through the DAGMC plugin, and hence incorporating the thermomechanics feedback will be included in our future work.

3.1. OpenMC Model

The geometry used in OpenMC must be finite, despite the benchmark being infinite in the y and z directions. The slab is divided evenly into N identically shaped rectangular cells along the x-dimension; the cases here are N=[50,100,250,500,1000] cells. In general, reflective boundary conditions can be used to simulate infinite dimensions. Thus, this benchmark can be represented with vacuum boundary conditions at $x=\pm \frac{L}{2}$ and reflective boundary conditions at the y and z boundaries. The cell boundaries have transmissive boundary conditions, which just allows particles to pass without modifying their state. Note that particles only move in the $\pm x$ direction, so no y or z boundaries are crossed, but with non- S_2 scattering, the reflective boundaries would come into play. In addition to being finite for the model, the y and z dimensions need to be set to 1 cm in order for Equation (19) from the benchmark to uphold its one dimensional power integral, i.e. that integration in y and z is implied to contribute a factor of 1 [4]. Fig. 1 shows a diagram of the 1-D geometry, governing equations, and boundary conditions for the different physics.

The benchmark's one-group assumption was satisfied using OpenMC's multigroup mode. Since the benchmark uses a fictitious material with a known function for the temperature dependence, the simulation took advantage of OpenMC's capability for user-defined cross-sections via the XSdata class. The cross-section for each reaction was specified for 50 evenly spaced temperatures between 308 K and 358 K (a range slightly larger than, but including the analytic temperature range from the benchmark) and was exported to a library. Then, when running OpenMC in multigroup mode, that library is loaded to determine the appropriate cross-section in each region as temperature changed via input from Cardinal from iteration to iteration.

OpenMC required slight source code modifications to accommodate S_2 -like transport. In typical OpenMC simulations, the physics of each reaction describes the scattering dynamics. The Monte Carlo algorithm is agnostic to the direction particles move, but it is not typically constrained to a discrete directional distribution. However, when modeling this benchmark, any history with a particle moving perpendicular to the x-direction would attenuate particles in less x-distance than if particles were constrained to either $\pm x$. To address this, we first use OpenMC's PolarAzimuthal distribution to restrict the birth direction of particles to the $\pm x$ direction. We then also modified OpenMC in a patched branch to mimic S_2 transport in two ways. First, when determining the angular cosine of scattering events, μ , particles either continue forward

 $(\mu=1)$ or are back-scattered $(\mu=-1)$ with equal probability, as opposed to sampling the reaction physics for μ . Second, since the simulation uses k-eigenvalue mode, neutrons born in subsequent generations of the simulation also need to have their angular birth distributions restricted to $\pm x$.

Due to the stochastic nature of Monte Carlo, an adequate number of histories must be used to obtain estimators that have reasonable uncertainties. In eigenvalue mode, there are inactive batches followed by active batches. The difference is that inactive batches run the simulation, but do not tally the requested estimators. The only quantity that is tracked is the fission source, which is passed from each inactive batch to the next. The goal is to converge the fission source before tallying, as the initial guess may be inaccurate. Thus, the number of inactive batches is an important parameter. Shannon Entropy computations can help determine fission source convergence [6]. A Shannon Entropy study for this system was conducted on the 1000 cell case, as increasing the cell size improves statistics – at least until there are too few cells to capture spatial dependence. This problem's analytical fission source actually turns out to be independent of position, due to the multiplication of cross-section by flux, where the inverse T and T relationships cancel, so it is safe to assume that if the fission source converges for 1000 cells, it would converge for all the fewer cell cases using the same number of histories. It was found the Shannon Entropy converges in fewer than 5 iterations. This may seem like a low number, but the problem is 1-D and the initial guess provided to OpenMC is a uniform distribution, which is the same as the actual converged solution. In order to be safe, the k-eigenvalue simulation used 50,000 particles per batch, with 50 inactive batches and 50 active batches for every case.

A few tallies were used in order to compare to the analytical solution: a flux cell tally, a kappa-fission rate cell and global tally, and the eigenvalue k_{eff} . Each tally uses a track-length estimator. As particles transport, their path may stay inside a given cell or span more cells. The track-length estimator contributes a tally score when individual particle tracks pass through a region. Though flux and the eigenvalue are the only quantities to compare with the benchmark, these other tallies are needed in order to compute a source strength. OpenMC reports flux, denoted as $\hat{\phi}$, in units of $\frac{n-cm}{sp}$, where sp indicates "source particle." Typical flux, denoted as ϕ , has units of $\frac{n}{cm^2-s}$. In fixed source modes, the source strength is typically known, but in eigenvalue mode, it must be computed, as it depends on the physical source and fission source. The source strength is given by

source strength
$$=\frac{P}{kfV_{cell}}$$
 THUS $\phi=\frac{P}{kfV_{cell}}\hat{\phi},$ (5)

where $P=1e22\frac{ev}{s}$ is the slab integrated power from the benchmark, kf is the tallied kappa-fission rate across the slab, and V_{cell} is the volume the cell containing the tally. The same source strength could be used to convert the kappa-fission tally from OpenMC, which has units of $\frac{ev}{sp}$, to more physically meaningful units of $\frac{ev}{cm^3-s}$.

3.2. MOOSE Heat Conduction Model

MOOSE is used to solve for the temperature distribution within the slab via the Heat Conduction Module. In this MOOSE model, a mesh is used that has identical dimensions to the cells used in the OpenMC model. While this is not required by Cardinal, it allows for 1-to-1 feedback between the temperature computed in MOOSE and the heat source computed in OpenMC. MOOSE solves the conduction equation using the Finite Element Method (FEM):

$$-\nabla \cdot [\kappa_s(\mathbf{r}, T_s) \nabla T_s(\mathbf{r})] = \dot{q}_s, \tag{6}$$

where κ_s is the thermal conductivity in the solid and \dot{q}_s is the heat source, in this case from fission. The boundary conditions used by MOOSE match temperature boundary condition in red shown in Fig (1). MOOSE is coupled to OpenMC by receiving the heat source in each element from the kappa-fission tally. During each constituent iteration, it recomputes the temperature distribution from the heat source and boundary conditions, and then sends the temperatures back to OpenMC for the next transport solve.

3.3. Convergence Criteria

MOOSE uses iterative methods to compute temperature. It uses a combination of linear and non-linear iterations to update the solution. It begins with a non-linear iteration, then linear iterations are run until

convergence criteria are met. These trigger the next non-linear iteration and this process repeats until both of the convergence criteria are met. This study used convergence criteria of 10^{-7} for absolute tolerance and 10^{-9} for relative tolerance. Monte Carlo is a stochastic method, so convergence is achieved by using an appropriate number of batches and histories per batch. The configurations used here were mentioned in Section 3.1.

In terms of converging global iterations across all single physics, each mesh cases used 200 Picard Iterations. To assist with convergence of Monte Carlo quantities, Robbins-Monro relaxation was applied to the flux and kappa-fission tallies. This updates the quantities of interest for the n+1th iteration as an average over the newest solution and the n previous iterations [7]. So the flux at Picard Iteration n would be given by

$$\Phi_n = \frac{1}{n+1} \sum_{i=0}^n \phi_i, \tag{7}$$

where Φ_n is the relaxed flux at step n and ϕ_i is the flux output from the ith Monte Carlo solve. In order to assess the convergence of each physics, plots of error versus mesh element size were used and will be shown in Section 4.

One MOOSE convergence feature that was not used here is steady-state detection. By setting a steady-state tolerance, MOOSE will iterate until the solution norm changes by less than this tolerance. In this benchmark, a high number of Picard Iterations was used in order to ensure convergence, but other modelers may not want to incur the computational cost of running 200 Picard Iterations. They can use steady-state detection instead.

3.4. Data Mapping

One feature of Cardinal is that it does not require the geometry models used in each single-physics subapplication to exactly match the others. This flexibility is a great asset, but it requires the modeler to choose an appropriate geometry model so that the coupling of solution fields can resolve the physics feedback effects. At the beginning of every Cardinal simulation, a mapping between MOOSE's mesh and other geometry representations is established. For a MOOSE-OpenMC coupling, the centroid of every mesh element is mapped to a corresponding OpenMC region, which could either be a cell or a mesh element; here cells are used. Cardinal then creates a tally for every requested type in the input file in each of the regions of OpenMC that a MOOSE centroid falls in. This mapping is used for transferring relevant quantities back and forth between iterations. If the modeler is not careful, this mapping may not be meaningful, as important OpenMC regions may not get a tally to participate in feedback, or too many MOOSE regions could be linked to the same OpenMC region. For this simulation, the MOOSE geometry and OpenMC geometries were intentionally created to have the same dimensions so that the mapping is 1-to-1. Each region in the MooseMesh computes a temperature and sends it to a geometrically identical OpenMC region. This same region in an OpenMC solve computes a heat source via kappa-fission tally and sends it back to MOOSE for the next conduction solve. A table below shows the configurations of each case in terms of MOOSE mesh elements and OpenMC cells.

Table 1: Simulation geometry configurations.

Number of MOOSE Mesh Elements	Number of OpenMC Cells
50	50
100	100
250	250
500	500
1000	1000

4. RESULTS

The results to compare with the benchmark are the temperature distribution, flux distribution, and k-eigenvalue. The benchmark provides analytical solutions for all of these quantities. An example temperature distribution is shown in the figure below for the 50 mesh element case.

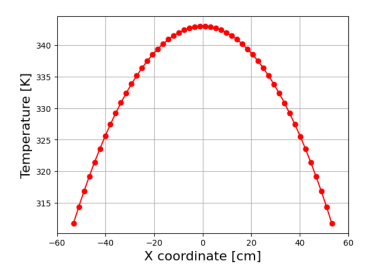


Figure 2: The computed temperature for 50 mesh elements.

In order to assess the accuracy of the temperature solution and verify that refining the mesh increases the accuracy, the analytical solution was used to compute an L_2 error norm for temperature. The error computed, ε_T , is given by

$$\varepsilon_T = \frac{||T_a - T_x||_2}{||T_a||_2},\tag{8}$$

where T_a is the analytical solution evaluated at the x-centroid of each mesh element and T_x is the temperature computed for that voxel from the multiphysics simulation. The figure below shows how the temperature error norm relates to the number of mesh elements

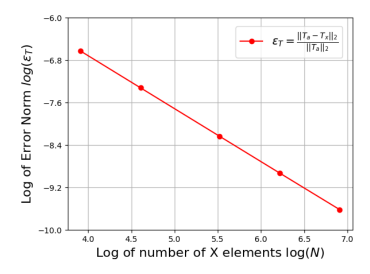


Figure 3: The number of mesh elements vs the log of the L_2 error norm for the computed vs analytical temperature.

Linear convergence is achieved, as the slope of the line on the $\log(N)$ vs $\log(\varepsilon_T)$ plot is -0.9981. The same convergence test was used on the flux solution as well. The flux error norm, ε_{ϕ} , is defined as

$$\varepsilon_{\phi} = \frac{||\phi_a - \phi_x||_2}{||\phi_a||_2}.\tag{9}$$

The figure below shows how the flux error norm relates to the number of mesh elements

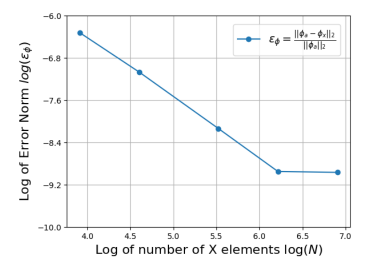


Figure 4: The number of mesh elements vs the log of the L_2 error norm for the computed vs analytical flux.

The slope of the line of best fit for $\log(N)$ vs $\log(\varepsilon_\phi)$ plot is -0.9496. This deviation from first order is clearly being caused by the error in the finest case (N=1000). The error for the N=1000 cases does decrease compared to N=500, but not be enough to fall in line with linear convergence. This is the case where the volume of each cell/element is the smallest. In Monte Carlo, as the volume of a region holding a tally gets smaller, fewer events will happen in that region per history, and thus more histories are required to get the same uncertainty. The N=1000 case appears to be experiencing issues related to the element size. One possible source of error could be the refinement in the cross-section temperature library. All cases used a library with 50 temperatures, so it is possible that this case is reaching a limit where more data refinement would be required to achieve the expected convergence that matches the other cases.

While this case appears to not perform as well from the L_2 error norm, it performs will in another measure of comparison – the Computed to Expected Ratio (C/E). The C/E is the ratio between the computed quantity, C, and the expected answer, E. In this case, C would either be the temperature or flux output from Cardinal, and E would be the corresponding analytical solution. Thus, the C/E ratio r is given by

$$r = \frac{C}{E}. (10)$$

The ideal would be exact equality, or a ratio of r=1. The temperature and flux C/Es for each mesh element and cell, respectively, are shown in the figure below.

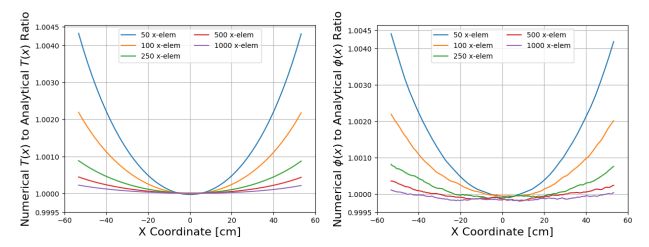


Figure 5: The left shows the C/E for temperature on every mesh and the right shows the C/E for flux in each cell.

Despite the disagreement from linear convergence for the flux, the C/E for the N=1000 case appears to be the closest to the line y=1. This builds confidence that Cardinal is outputting a correct flux distribution for this case. The eigenvalue for each mesh is reported in the table below.

Table 2: Eigenvalue with uncertainty for each mesh size, and the pcm difference from the analytical.

analytical $k_{eff} = 0.29557$		
numerical k_{eff}	pcm (analytical - numerical)	
0.29555 ± 0.00009	-2	
0.29551 ± 0.00011	-6	
0.29560 ± 0.00012	-3	
0.29575 ± 0.00013	18	
0.29551 ± 0.00013	-6	
	numerical k_{eff} 0.29555 ± 0.00009 0.29551 ± 0.00011 0.29560 ± 0.00012 0.29575 ± 0.00013	

The eigenvalue is the metric that agrees with the analytical solution the quickest. This is due to it being a system-wide parameter, which gains contributions from every fission event, no matter which cell they occur in. It converges much quicker than the flux, which only gains contributions when an event occurs in a given cell. All the standard deviations of the computed eigenvalues are below 13 pcm, which is a tightly converged answer. The analytical eigenvalue is within one standard deviation of the computed eigenvalue for all cases except the N=500; this case is off by 18 pcm, but the analytical eigenvalue is only 5 pcm above the upper bound of the 13 pcm uncertainty associated with that tally.

5. CONCLUSIONS

Overall, the numerical results show agreement with the analytical solutions. The convergence is first order for the temperature, and the flux shows near first order convergence, baring the $N=1000\,\mathrm{case}$. Since zeroth order shape functions are used for temperature in the FEM, the first order temperature spatial convergence is to be expected [8]. The C/E approach closer and closer to $y=1\,\mathrm{as}\ N$ increases in each case. The eigenvalue agrees very well for all cases.

While code to code benchmarking is common in for nuclear M&S, agreement with analytical benchmarks increases the confidence in Cardinal's multiphysics coupling. Though typical industry-grade simulations

would not run S_2 transport, this modification allows Cardinal to show it can match against theoretical problems as well. Another study with Cardinal [5], coupled OpenMC Monte Carlo transport to NekRS heat conduction. In principle, this benchmark could also use that coupling, and it will be future work to validate this OpenMC-MOOSE coupling to a study matching the same physics in an OpenMC-NekRs coupling. In addition to the various, ongoing verification efforts, validation against physical experiments would be another next step to increase confidence in Cardinal's modeling capabilities.

NOMENCLATURE

Nomenclature

C/E Computed to Expected Ratio

CFD Computational Fluid Dynamics

FEM Finite Element Method

M&S Modeling and Simulation

MOOSE Multiphysics Object-Oriented Simulation Environment

ODE Ordinary Differential Equation

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