Introduction to Moltres: An application for simulation of Molten Salt Reactors

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Abstract

Moltres is a new physics application for modeling coupled physics in fluid-fuelled, molten salt reactors. This paper describes its neutronics model, thermal hydraulics model, and their coupling in the MOOSE framework. Neutron and precursor equations are implemented using an action system that allows use of an arbitrary number of groups with no change in the input card. Results for many-channel configurations in 2D-axisymmetric and 3D coordinates are presented and compared against other coupled models as well as the Molten Salt Reactor Experiment.

1. Introduction

Molten salt reactor concepts garnered considerable interest in the 1950s and 60s with development of the Aircraft Reactor Experiment (ARE) and later the Molten Salt Reactor Experiment (MSRE) at Oak Ridge National Laboratory (ORNL). With the inclusion of the Molten Salt Reactor (MSR) among the Generation-IV reactor designs (GIF, 2008, 2015), this reactor concept has gained renewed research interest in the past decade, with many new nuclear companies proposing both fluid-fuelled and solid-fuelled commercial MSR designs (Roderick and McWhirter, 2015; David, 2015; ThorCon, 2017; Scarlat et al., 2014; Transatomic Power Corporation, 2016). The key advantages of MSRs generally pertain to improved fuel utilization and reactor safety. In contrast to legacy reactors, only moderator fast neutron damage and fuel chemistry evolution limit burnup. A clever configuration of moderator as in Engel et al. (1980) can enable reactor operation without opening the vessel for thirty or more years. Further, several fission products selectively precipitate onto nickel surfaces in fluoride salt, as documented in Engel et al. (1980), thus reducing unwanted neutron absorption. Lastly, the epithermal spectrum of graphite-moderated salt reactors incinerates plutonium more efficiently, thus reducing long-lived transuranic waste production (Engel et al., 1980). The sum of these characteristics implies the MSR uniquely ameliorates spent fuel burden whilst extending nuclear fuel resources. To top all these benefits off, xenon transients become moot in MSRs due to its insolubility in salt, thus narrowing transient analysis focus to thermalhydraulic concerns.

Simulation tools developed by many authors successfully describe steady-state and transient behavior of myriad MSR concepts. Krepel et al. extended the in-house Light Water Reactor (LWR) diffusion code DYN3D to consider drift of delayed neutron precursors alongside the reactor temperature profile, re-casting the extended code as DYN3D-MSR (Kreipel et al., 2007). That work compared DYN3D-MSR against experimental MSRE data and then used it to simulate local fuel channel blockages as well as local temperature perturbations.

In a similar vein, Kophazi et al. used iterative coupling between in-house three-dimensional neutronic and one-dimensional heat conduction models DALTON and THERM to analyze normal MSRE operation as well as channel-blocking-incident transients (Kópházi et al., 2009). The Kophazi model added entrance effects of heat transfer coefficients as well as thermal coupling between fuel channels through moderator heat conduction. More recently, Cammi et al. performed a 2D-axisymmetric single-channel analysis...
of the Molten Salt Breeder Reactor (MSBR) using the commercial finite element package COMSOL Multiphysics (Cammi et al., 2011). That work directly solved the fuel salt velocity field, used heterogeneous group constants in fuel and moderator regions, and employed the COMSOL Multiphysics (COMSOL) software package intrinsically designed for coupled multi-physics simulation. Fiorina, Lathouwers, and their colleagues conducted a benchmarking exercise (Fiorina et al., 2014) in which this Politecnico di Milano approach was expanded to a multi-channel model of the Molten Salt Fast Reactor (MSFR) and compared to code from the University of Delft (De Zwaan et al., 2007; van der Linden, 2012) based on the approach in Kópházi et al. (2009). These models showed good agreement for multiple accident transients. Meanwhile, leveraging lessons learned from these efforts has resulted in a multiscale approach from Zanetti et al. (2015) successfully combines high and low geometric fidelity for graphite-moderated MSRs.

More recently, Auffiero et al. (2014) have begun to approach transient simulations in the MSFR within the finite volume OpenFOAM multiphysics toolkit (Weller et al., 1998). This approach benefits from pre-implemented turbulence models available in the OpenFOAM library and captures the full-core three-dimensional geometry of the reactor primary circuit. OpenFOAM Computational Fluid Dynamics (CFD) has additionally been shown by Laureau et al. (2017) to couple well with Transient Fission Matrix neutronics within the MSFR.

The present work introduces the open source simulation tool, Moltres (Lindsay, 2017), for simulating MSRs. By implementing deterministic neutronics and thermal hydraulics in the context of the Multiphysics Object-Oriented Simulation Environment (MOOSE) finite element modeling framework, Moltres solves arbitrary-group neutron diffusion, temperature, and precursor governing equations on a single mesh in anywhere from one to three dimensions and can be deployed on an arbitrary number of processing units.

Moltres is an open source code licensed under Lesser GNU Public License (LGPL) terms so the MSR community can freely use, interrogate, and improve it. Its openness is a defining characteristic of open literature, is tailored to molten salt reactor physics. Accordingly, Moltres joins a veritable parade of open academic nuclear engineering software such as OpenMC (Romano et al., 2015), OpenMOC (Boyd et al., 2014), and PyNE (Bates et al., 2014; Biondo et al., 2014).

Moreover, Moltres depends on the MOOSE framework, Gaston et al. (2015) another LGPL code that itself leans on LibMesh (Kirby et al., 2006), a LGPL finite element library, and PetSc (Balay et al., 2015), a Berkeley Software Distribution (BSD)-licensed toolkit for solving nonlinear equations yielded by discretizing PDEs. MOOSE and LibMesh translate weak PDE forms defined by applications (e.g. Moltres) into residual and Jacobian functions. These functions are the inputs into PetSc Newton-Raphson solution routines. All codes use MPI for parallel communication and are easily deployed on massively-parallel cluster-computing platforms. MOOSE applications by default use monolithic and implicit methods ideal for closely-coupled and multi-scale physics, such as the model problem described in this work. However, Moltres can also use explicit time-stepping routines as well as segregated solution methods, making it extensible to myriad future modeling challenges.

2. Methods

Moltres (Lindsay, 2017) is implemented as an application for use atop the MOOSE (Gaston et al., 2015) framework. Accordingly, Moltres includes physics kernels and boundary conditions for solving for neutron fluxes, temperature, and precursor concentrations. In MOOSE jargon, kernels are C++ classes that contain methods for computing residual and Jacobian contributions corresponding to individual pieces of governing equations. Developing the code-base in this way allows modular construction of equation systems; e.g. the kernel used to represent heat conduction can also represent generic chemical diffusion. Moltres also features neutron and precursor “actions.” These actions automatically construct the systems of equations for an arbitrary number of neutron and precursor groups. Therefore, as long as group constants are provided in an appropriate tabular form, a user only has to modify a couple of lines in a Moltres input file to change from say two to forty-four neutron groups.

In Moltres, neutrons are described with time-dependent multi-group diffusion theory as shown in Eq. (1):

$$\frac{1}{v_g} \frac{\partial \phi_g}{\partial t} - \nabla \cdot D_g \nabla \phi_g + \Sigma_T \phi_g = \sum_{g' \neq g} \Sigma_{g' \to g} \phi_{g'} + \chi_g \sum_{i=1}^G (1 - \beta) \nu \Sigma_f \phi_g + \sum_{i=1}^G \lambda_i C_i$$

(1)

where
\( n_g \) = speed of neutrons in group \( g \)  
\( \phi_g \) = flux of neutrons in group \( g \)  
\( t \) = time  
\( D_k \) = Diffusion coefficient for neutrons in group \( g \)  
\( \Sigma'_g \) = macroscopic cross-section for removal of neutrons from group \( g \)  
\( \Sigma'_g \) = macroscopic cross-section of scattering from group \( g \) to group \( g' \)  
\( \chi'_g \) = prompt fission spectrum neutrons in group \( g \)  
\( G \) = number of discrete groups. \( g \)  
\( v \) = number of neutrons produced per fission  
\( \Sigma'_g \) = macroscopic cross section for fission due to neutrons in group \( g \)  
\( \chi'_g \) = delayed fission spectrum, neutrons in group \( g \)  
\( \beta \) = delayed neutron fraction  
\( \lambda_i \) = average decay constant of delayed neutron precursors in precursor group \( i \)  
\( C_i \) = concentration of delayed neutron precursors in precursor group \( i \)  
\( \beta C_i = \sum_{g=1}^{G} \beta_g \Sigma'_g \phi_g - \lambda_i C_i - \frac{\partial}{\partial z} u C_i \)  
\( \rho_i c_{pf} \frac{\partial T_f}{\partial t} + \nabla \cdot \left( \rho_i c_{pf} \vec{u} \cdot \nabla T_f - k_f \nabla T_f \right) = Q_f \)  
\( \rho_f \) = density of fuel salt  
\( c_{pf} \) = specific heat capacity of fuel salt  
\( T_f \) = temperature of fuel salt  
\( \vec{u} \) = velocity of fuel salt  
\( k_f \) = thermal conductivity of fuel salt  
\( Q_f \) = source term  
\( Q_f = \sum_{g=1}^{G} \epsilon_{fg} \Sigma'_f \phi_g \)  
\( Q_f = \gamma V_{core}^1 \int_{core} Q_f dV \), where \( \gamma \) is a factor representing heat dissipation by gamma and neutron irradiation in the moderator, called the graphite to fuel power density ratio. Robertson’s (1971) original MSBR analysis included a calculation of \( \gamma \). Krepel et al. (2007), Zhang et al. (2009), Cammi et al., 2011 all calculated gamma heating through such a factor. We follow Cammi et al. (2011) and set \( \gamma = 0.0144 \). Notably, the Moltres physics kernel for radiation heating can be calculate the volume average fission heat in a variety of means: whole-core and local averages are possible. Whole-core averages are employed for simplicity and in accordance with prior literature. Knowledge of fission heat rates, however, require knowledge of neutron fluxes, implying a need for group constants.

Group constants are generated by the modeler with either Serpent (Leppänen et al., 2015) or SCALE (DeHart and Bowman, 2011). Moltres interpolates group constant temperature dependence from prepared tables, which must be constructed separately for fuel and moderator regions. For this report, we generated group constants with SCALE with an infinite square pitch lattice of cylindrical fuel channels surrounded by graphite. This model maintained a fuel fraction of 0.225 to be consistent with the MSRE. Subsequently, a critical buckling calculation was applied. The SCALE input files used for generating the group constants appear in the io/msre_conc_cuboid_lattice directory of the github.com/arfc/scale_io repository.

2.1. Performance

Building on the massively parallelizable MOOSE framework allows Moltres to run on super-computing platforms like the Blue Waters supercomputer at the National Center for Supercomputing Applications (NCSA). For some three-dimensional simulations, the number of elements in the mesh and total number of degrees of freedom exceed one million and ten million respectively. To handle problems of this size, we ran Moltres on up to 608 cores. However, reducing the problem dimension from three to two and using a structured mesh, which can be much more coarse in the axial direction, allows the problem to be solved on a single core in under five minutes.

3. Model description

The model molten salt reactor closely emulates the MSRE. When developing new physics or investigating different types of transients, one can reduce the model problem to a two dimensional axisymmetric one for rapid proof of concept. To approximately simulate the lattice structure of the MSRE under 2D conditions, a geometry is constructed with 14 repeating fuel/moderator regions, as shown in Fig. 1. The fuel and moderator radii are chosen such that the resulting area/volume fraction of fuel is 0.225 as for the MSRE. The base 2D mesh has a characteristic size of 10 cm in the axial direction and 4 cm in the radial direction to capture the variation from moving between fuel and moderator subdomains. To determine whether results were converged, a mesh convergence study was conducted with up to three levels of isotropic refinement (e.g. each element was in half in both axial and radial directions, resulting in four new elements). The metric used to assess convergence was the integrated fast group flux. The result of the mesh convergence study is shown in Fig. 2. From refinement level 2 to 3 the metric changes by only 1%. We regard the results as sufficiently converged at level 2 for the purposes of our study; consequently, the results reported in the following section correspond to a radial element dimension of .1 cm and an axial element dimension of 2.5 cm.

The model fuel composition is the Beginning-of-Life (BOL) enriched uranium composition in the MSRE and is given in Table 1 (Robertson, 1965).

Other simulation inputs are outlined in Table 2. We chose a reactor simulation height of 151.75 cm to produce an approximately critical reactor configuration corresponding to BOL MSRE.

\( Q_g = \gamma V_{core}^1 \int_{core} Q_f dV \), where \( \gamma \) is a factor representing heat dissipation by gamma and neutron irradiation in the moderator, called the graphite to fuel power density ratio. Robertson’s (1971) original MSBR analysis included a calculation of \( \gamma \). Krepel et al. (2007), Zhang et al. (2009), Cammi et al., 2011 all calculated gamma heating through such a factor. We follow Cammi et al. (2011) and set \( \gamma = 0.0144 \). Notably, the Moltres physics kernel for radiation heating can be calculate the volume average fission heat in a variety of means: whole-core and local averages are possible. Whole-core averages are employed for simplicity and in accordance with prior literature. Knowledge of fission heat rates, however, require knowledge of neutron fluxes, implying a need for group constants.

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4. Results & discussion

Group fluxes are shown in Figs. 3 and 4. The cosinusoidal shapes in radial and axial directions are caused by the vacuum boundary conditions. Both the fast and thermal fluxes are striated, with the composition. This differs from the actual MSRE height, which was 162.56 cm.

### Table 1

<table>
<thead>
<tr>
<th>Component</th>
<th>Mass fraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li-7</td>
<td>.1090</td>
</tr>
<tr>
<td>Li-6</td>
<td>5 × 10⁻⁶</td>
</tr>
<tr>
<td>F-19</td>
<td>.6680</td>
</tr>
<tr>
<td>Be-9</td>
<td>.0627</td>
</tr>
<tr>
<td>U-235</td>
<td>.0167</td>
</tr>
<tr>
<td>U-238</td>
<td>.0344</td>
</tr>
</tbody>
</table>

Fuel salt composition is the BOL enriched uranium composition in the MSRE design (Robertson, 1965).
fast group preferring the fuel and the thermal group preferring the moderator.

In Fig. 5 the temperature rises along the reactor height because of fuel advection. The temperature gradient is negative in the radial direction, as expected.

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Table 2
Simulation input parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Units</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inlet temp.</td>
<td>922</td>
<td>K</td>
<td>MSRE nominal (Robertson, 1965)</td>
</tr>
<tr>
<td>Wall temp.</td>
<td>922</td>
<td>K</td>
<td>MSRE nominal (Robertson, 1965)</td>
</tr>
<tr>
<td>Neutron groups</td>
<td>2</td>
<td>I</td>
<td>User</td>
</tr>
<tr>
<td>Precursor groups</td>
<td>6</td>
<td>I</td>
<td>User</td>
</tr>
<tr>
<td>Reactor radius</td>
<td>72.5</td>
<td>cm</td>
<td>MSRE radius (70.2 cm) (Robertson, 1965)</td>
</tr>
<tr>
<td>Reactor height</td>
<td>151.75</td>
<td>cm</td>
<td>User</td>
</tr>
<tr>
<td>$k_f$</td>
<td>0.0553</td>
<td>W cm$^{-1}$ K$^{-1}$</td>
<td>Robertson (1965)</td>
</tr>
<tr>
<td>$c_p$</td>
<td>1967</td>
<td>kg cm$^{-3}$ K$^{-1}$</td>
<td>Robertson (1965)</td>
</tr>
<tr>
<td>$\rho_f$</td>
<td>2.146 $10^{-3}$ e$^{-(r_f - 0.022)}$</td>
<td>J kg$^{-1}$</td>
<td>Robertson (1965)</td>
</tr>
<tr>
<td>$\sigma_f$</td>
<td>2.12 $10^{-4}$</td>
<td>kg cm$^{-3}$</td>
<td>Haubenreich and Engel (1970)</td>
</tr>
<tr>
<td>$k_x$</td>
<td>0.312</td>
<td>J K$^{-1}$</td>
<td>Cammi et al. (2011)</td>
</tr>
<tr>
<td>$c_{p,x}$</td>
<td>1760</td>
<td>W cm$^{-1}$ K$^{-1}$</td>
<td>Cammi et al. (2011)</td>
</tr>
<tr>
<td>$\rho_x$</td>
<td>1.86 $10^{-3}$ e$^{-(r_x - 0.022)}$</td>
<td>kg m$^{-3}$</td>
<td>Robertson (1965)</td>
</tr>
<tr>
<td>$\sigma_x$</td>
<td>1.8 $10^{-5}$</td>
<td>K$^{-1}$</td>
<td>Haubenreich and Engel (1970)</td>
</tr>
</tbody>
</table>

---

Fig. 3. The group 1 flux in this 2-D cylindrical axisymmetric model has the anticipated magnitude and canonical cosine shape ($r = 0$ is center of core).

Fig. 4. The group 2 flux in this 2-D cylindrical axisymmetric model has the anticipated magnitude and canonical cosine shape ($r = 0$ is center of core).

Fig. 6 shows the concentration of the longest lived precursor in the reactor. Not surprisingly, the channel concentrations are higher in fuel channels with higher neutron fluxes and corresponding fission events. Because of the small decay constant of the precursor, the maximum concentration in any given channel occurs at the core outlet due to advection.
With its much larger decay constant, the sixth and last precursor has its maximum concentration around the center-plane of the reactor as shown in Fig. 7. As for all other precursors, its concentration decreases with increasing radius and decreasing neutron flux.

4.1. Three dimensional simulation capability

Figs. 8 and 9, show Moltres physics applied to a three dimensional geometry. The fast group flux (Fig. 8) is in good qualitative agreement with the two dimensional axisymmetric case shown in Fig. 3. Fig. 9 is in similarly good agreement with Fig. 5. Fig. 10 shows the temperature profile at the outlet of the reactor \((z = H)\). This three dimensional case contained 1,155,045 degrees of freedom and took only 2.5 h to solve on 160 Blue Waters cores.

4.2. Comparison with MSRE

Fig. 11 shows a comparison between Moltres predicted temperature profiles with cosinusoidal gamma heating and MSRE design calculations (Briggs, 1964) in the hottest channel and adjacent graphite. The ORNL MSRE design calculations, conducted in 1963–1964, were 32-group calculations using legacy computing tools (GAM-I, MODRIC, and EQUIPOSE, and THERMOS). Those calculations were conducted in two-dimensional R-Z geometry (a cylinder with angular symmetry), with 20 spatial regions. Notably, one limitation of the ORNL model was the control rod thimbles that, due to angular symmetry, were effectively a cylindrical shell of metal.

The profile shapes are in decent qualitative agreement with both Moltres and MSRE calculations showing a peak in graphite temperature before the reactor outlet. Fuel temperature increases monotonically in both Moltres and MSRE models. In the MSRE design, the moderator temperature at the reactor inlet is about 11 K larger than the fuel temperature, whereas the temperatures are about the same in the Moltres model. This difference is likely because the MSRE design model neglected axial heat conduction (Briggs, 1964, p. 99).

Fig. 12 compares the fast and thermal neutron fluxes at the reactor mid-plane \((z = H/2)\) for Moltres and MSRE design models. Local thermal flux growth and fast flux decay in moderator regions and visa versa in fuel regions are apparent in the Moltres calculation. The Moltres flux magnitudes are in good agreement with the magnitudes from the MSRE design calculations (Briggs, 1964, p. 92). The peak fast to thermal flux ratio is approximately 3.5 in the MSRE design calculation as opposed to a ratio of 3 for the Moltres calculation. Control rod thimbles and an extra volume of surrounding fuel not included in the Moltres calculations cause the depression in the thermal flux in the MSRE profile.

Fig. 13 compares the axial flux profiles calculated by Moltres and the ORNL MSRE design model. The radii for the plots are chosen to correspond to the peak of the thermal flux in both cases; for
the ORNL calculations is 21.336 cm (8.4 inches) from the core center-line because of the effect of the control rod thimbles and extra fuel along the center-line. Once again, the plots are in decent agreement. The ORNL calculations include the lower and upper plena which are not included in this report’s Moltres model. Consequently, the MSRE lines extend to lower and higher z-values than the Moltres lines. Additionally, absorption in the plena cause deviation of the thermal flux from a sinusoidal shape in the MSRE design case. The peak power density from the MSRE calculation is 31 kW/L; the corresponding value for Moltres is 29 kW/L.

Although the qualitative agreement is decent, there are discrepancies between the MSRE and Moltres calculations. As outlined above there is an 11 K offset between MSRE and Moltres calculations for the fuel temperature in the hottest channel. The peak graphite temperature is around 14 K larger in the Moltres calculation. Fast fluxes are larger in the MSRE calculation by roughly 20%. These are not insignificant differences. However, given the differing nature of the two models—no axial conduction in the MSRE model, 2-group vs 32-group neutronics, exclusion of the control rod thimbles in the Moltres calculation—we believe this variation in quantitative behavior is acceptable for the purpose of this work, which is to introduce Moltres as a simulation tool. One additional feature that could influence Moltres results is inclusion of precursor decay heat; this is under active development. Trustworthy verification of Moltres results, such as spatial flux and temperature profiles, will
require more detailed experimental measurements than those available in the ORNL technical reports, which contain only macroscopic data such as steady-state power and heat-exchanger temperature drops. Additionally, detailed comparison with other modern modeling efforts such as that conducted in Aufiero et al. (2014), Laureau et al. (2017) is desirable. In that vein, a collaboration is under way with the primary author of Aufiero et al. (2014) to try and reproduce calculation results from OpenFOAM. Further, Collins from Turner et al. (2016) has expressed interest in comparing Moltres results against MSR-VERA in order to quantify errors resulting from usage of multigroup diffusion in comparison to 2D-1D method of characteristics fine group solutions. We hope to publish the results of these comparisons in a future work.

4.3. Scaling performance

Parallelization in Moltres is implemented via LibMesh, which includes a set of utilities for massively parallel finite element based computations, including mesh input/output, a finite element library, and an interface for connections with solver packages. Employing LibMesh provides Moltres with significant flexibility including the ability to swap out solver libraries such as PetSc, which includes an expanding suite of parallel linear and nonlinear solvers. Problem domain decomposition relies on LibMesh mesh adaptation capabilities for running on a specific number of processors and can either be performed manually before the start of the simulation or automatically at the parameters of computation.
We conducted strong and weak scaling studies to characterize parallel performance in Moltres. In case of strong scaling, the problem size remains fixed but the number of processors is increased. Strong scaling studies seek to identify an optimal ratio between the number of processors and elements for the most rapid and power-efficient computation for a given problem. We measured Moltres strong scaling with a simple 2D axisymmetric case for various problem sizes separately for intra-node (2,820; 5,640; 11,280 and 28,200 elements) and extra-node (86,655; 173,310; 317,735; 664,355 elements) setup on Blue Waters' XK7 nodes (two AMD 6276 Interlagos CPU per node, 16 floating-point bulldozer core units per node or 32 “integer” cores per node, nominal clock speed is 2.45 GHz).

Fig. 14 shows the simulation speed in seconds per element vs. the number of cores on 1 node (maximum 32 cores). Up to 8 cores, larger problems required considerably more time per element because of cache overhead. However, beyond 8 cores, scaling demonstrates asymptotic dependence on the number of processors due to increasing communication costs. The best parallel efficiency for the intra-node study is approximately 89% and has been achieved for the largest problem (28,200 elements).

Fig. 15 shows Moltres strong scaling up to 768 processors. This takes into account communication costs between nodes. Similar to the intra-node study, when fewer than 128 cores were used, cache overhead causes performance slow down for larger problems. However, beyond 256 cores, the simulation time per element remains almost constant for small cases (86,655 and 173,310 elements) and slightly decreases for the two larger problems. For extra-node scaling, parallel efficiency also grows with the problem size and reaches an optimal value of 73% for 664,355 elements.

For the weak scaling study, the part of the problem (workload) assigned to each processor stays constant and additional elements are used to solve a bigger problem which would not fit in memory on a single node. Thus, the weak scaling measurement is justification for memory-bound application such as multiphysics code. Linear weak scaling is achieved when the execution time stays constant while the workload increasing in direct proportion to the number of cores. We performed Moltres weak-scaling tests on Blue Waters, keeping the workload constant at 581, 985, 1970 and 3940 elements per core. Fig. 16 shows Moltres weak scaling performance measured for \( n_{\text{cores}} \in [1, 32] \) within one Blue Waters node and Fig. 17 demonstrates performance for \( n_{\text{cores}} \in [32, 128] \).

As expected, the largest drop in performance occurs when the number of cores increases from one to \( \approx 8 \), which corresponds to switching from no communication to a 2-D domain decomposition. The further reduction in performance of only about 50% over a range of 32 cores is likely caused by increased communication latency appearing from collective Message Passing Interface (MPI) calls. In the extra-node case, the performance drops by a factor of three, which is most likely due to poor node selection by the Blue Waters job scheduler and significantly increased latency and bandwidth costs.

Moltres scalability study results clearly indicate that parallelization using LibMesh’s automatic domain decomposition is good, but not perfectly efficient. This scaling performance is satisfactory for MSR simulations approached thus far and improved parallel performance would require further optimization within LibMesh. Moreover, Moltres is memory-bound and therefore very sensitive to host memory and memory bandwidth. Consequently, if improved performance is needed, one could consider a transition from CPUs computing to GPU-accelerated computing because GPUs operate on the fly with global memory, avoiding CPU cache storage issues. Another way to improve parallel performance is to force the solver to use “older” information from previous iterations. However, this has been shown to slow convergence in terms of iterations and increased workload (Balay et al., 2015).
4.4. Monolithic vs. segregated solution

Relative performance of monolithic vs. segregated solution methods for large systems of equations is an active area of research. Monolithic solvers are generally regarded to be more robust than their segregated counterparts but are often perceived as being too expensive for large-scale problems. Work conducted by Heil (2008) demonstrates that monolithic solvers are competitive with segregated techniques for weakly coupled physics; moreover, for strongly coupled problems the segregated methods struggle to achieve convergence. It is noted by the authors that use of monolithic solvers on large 3D problems requires efficient preconditioning to be effective.

Monolithic and segregated solution techniques were compared in Moltres using the 2D-axisymmetric problem but with delayed neutron precursors removed. In this case starting from arbitrary initial conditions the neutron multiplication exhibits damped oscillations around the critical state. Feedback between the power deposition by neutrons and temperature modification of neutron macroscopic group constants make this a very tightly coupled problem. Using adaptive time stepping based on the number of non-linear iterations at each time step, steady-state neutron fluxes and temperature profiles are achieved in 358 time steps and in 4 min of compute time on a single core using full-coupling (e.g. neutron fluxes and temperature in the same matrix). Alternatively, a solution was also attempted splitting neutron and temperature solves and performing Picard iterations until convergence. Ultimately, a steady-state solution could not be achieved using the segregated method. After 670 time steps, the Picard solve hits the maximum number of allowed iterations (100) after only halving the neutronics residual. Before reaching the fail-point, the segregated solution required numerous Picard iterations to converge (68, 37, 22 for the previous three time-steps for example), resulting in a total compute time of 37 min before failure. These results suggest that for solving tight coupling between neutronics and temperature, a monolithic solve approach is more effective.

5. Conclusion

This work introduces the open source MSR simulation code Moltres. Moltres solves arbitrary-group neutron diffusion, temperature, and precursor governing equations in any dimensions and can be deployed on an arbitrary number of processing units. The 2D-axisymmetric and 3D models presented here employ heterogeneous group constants for fuel and moderator regions generated with SCALE. Fuel volume fraction and fuel salt composition are based on the MSRE. Neutron fluxes show expected cosinoidal shapes in radial and axial directions with visible striations between fuel and moderator regions. The fast group flux is enhanced in fuel regions while the thermal group flux in enhanced in moderator regions. Due to advection the temperature profile in the fuel increases monotonically in the direction of salt flow, while the moderator temperature exhibits a maximum between the mid-plane and outlet. The role of advection is also seen in precursor concentrations. Long lived precursors exhibit maximum concentrations at the core outlet. As the decay constant increases across precursor groups the maximum concentrations moves towards the reactor center where the precursor production rate is maximum. Results from 2D-axisymmetric and 3D models show good qualitative agreement. Moreover, Moltres results compare favorably with the actual design calculations of the MSRE. Moltres demonstrated strong parallel scaling on a typical model problem. Future Moltres publications will highlight transient simulation cases investigating control rod ejection, single channel blockage, loss of flow, and loss of secondary cooling.

Acknowledgments

All figures in this paper were generated using the python package yat (Turk et al., 2011). The package engauge-digitizer (Mitchell et al., 2017) was used to convert rasterized MSRE line plots into point data for plotting alongside Moltres data.

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