

A HIGH-ORDER LOW-ORDER ALGORITHM WITH EXPONENTIALLY-CONVERGENT MONTE CARLO FOR THERMAL RADIATIVE TRANSFER

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ABSTRACT

We have demonstrated the potential of a new high-order low-order (HOLO) algorithm for solving thermal radiative transfer problems. The low-order (LO) solver is based on spatial and angular moments of the transport equation and a linear discontinuous finite-element spatial representation. The LO solver is fully implicit in time and resolves the non-linear temperature dependence at each time step. The solution to the LO solver produces a fixed-source, pure absorber transport problem as the high-order (HO) system. The HO solver utilizes exponentially-convergent Monte Carlo (ECMC) to give a globally accurate solution for the angular intensity. This global solution is used to compute consistency terms that require the HO and LO solutions to converge to the same solution. The use of ECMC eliminates instabilities caused by statistical noise introduced to the LO consistency terms. Herein, we discuss the method in detail and compare initial results with an implicit Monte Carlo code for one-dimensional gray test problems, as well as demonstrate the efficiency of ECMC over SMC in this algorithm.

Key Words: List of at most five key words

1. INTRODUCTION

Thermal radiative transfer (TRT) physics are relevant in many high-temperature physical applications, e.g., inertial confinement fusion and supernovae. Such problems feature a strong coupling between the material and radiation energy balance equations, as well as non-linear material and source term temperature dependences. TRT problems often require solutions in a mix of streaming and diffusive regions of the problem due to absorption-reemission physics and opacity temperature dependencies. The diffusive regions are time consuming to solve with higher-rank transport solutions and can be accurately represented by a diffusion solution. However, streaming regions require the accuracy of a full transport treatment.

Moment-based hybrid Monte Carlo (MC) methods have been proven useful for solving non-linear, TRT problems. Recent work has focused on picard iteration high-order low-order (HOLO) approaches [2,3,?]. Such methods utilize a low-order (LO) operator based on angular moments of the transport equation, formulated over a coarse spatial mesh. Physics operators that are time consuming for the HO transport solver to resolve, e.g., the re-emission source and physical scattering, are moved to the LO system. Newton methods allow for non-linearities in the LO equations to be efficiently solved [2]. This allows for non-linear terms at each time step to be accurately resolved, eliminating the need for approximate linearizations and short time steps relevant to other methods.

High-fidelity solutions can be achieved with HOLO methods by using standard MC simulations to solve the HO transport equation. The HO solution is used to construct consistency terms that require the LO solution to be consistent with the HO solution. These consistency terms preserve the accuracy of the MC solution method in the LO operator. Such HOLO algorithms suffer from stability issues caused by statistical noise introduced into the consistency terms that are estimated via MC simulation. In this work, we demonstrate the utility of a unique LO operator in conjunction with an exponentially-convergent Monte Carlo (ECMC) method[?] for the HO solver. The ECMC algorithm allows for statistical noise to be reduced to the same order as the HOLO iteration error with significantly less particles than standard MC. We have derived the LO operator directly from the transport equation, using a linear-discontinuous (LD) finite-element (FE) spatial discretization, such that the HO and LO solutions are consistent upon convergence. The LD spatial representation mitigates issues with energy propagating faster than the speed of light by providing an accurate spatial representation of the Planckian emission term within a cell. We present results for various Marshak wave test problems and reference are results against IMC.

2. High-Order Low-Order Algorithm for Thermal Radiative Transfer

2.1. Governing Equations

We have implemented a high-order low-order (HOLO) algorithm for the case of gray, one spatial dimension TRT problems. For simplicity, we will also consider constant density and specific heats throughout. The governing equations are the radiation and material energy balance equations, i.e.,

$$\frac{1}{c} \frac{\partial I}{\partial t} + \mu \frac{\partial I}{\partial x} + \sigma_t I = \frac{\sigma_s}{2} \phi + \frac{1}{2} \sigma_a a c T^4 \quad (1)$$

$$\rho c_v \frac{\partial T}{\partial t} = \int_{-1}^1 \sigma_a I^{n+1}(x, \mu) d\mu - \sigma_a a c (T^4), \quad (2)$$

where $\phi(x) = \int_{-1}^1 I(x, \mu) d\mu$ is the scalar radiation intensity, related to the energy density $E(x) = \phi(x)/c$. Here, x is the position, μ is the x -direction cosine, and a , c , ρ , c_v , σ_a , σ_s , and σ_t are the radiation constant, speed of light, mass density, specific heat, absorption, scattering, and total opacities (cm^{-1}), respectively. The desired fundamental unknowns are the material temperature $T(x)$ and radiation intensity $I(x, \mu)$. In general, the material properties are a function of T . The equations are coupled through the gray Planckian emission source $\sigma_a a c T^4$ and absorption term $\sigma_a \phi$. The notation of the scalar intensity ϕ is used throughout rather than the typical radiation energy density $E = \phi(x)/c$.

MC solution to the TRT equations is typically achieved by the well-known implicit Monte Carlo (IMC) method, first introduced by Fleck and Cummings [?]. This method linearizes the emission source in the time derivative of the material energy equation to eliminate the material energy equation from the system. The remaining transport equation is solved by MC with an effective scattering cross section representing emission and reemission over the time step. The time derivative in the transport equation is solved continuously. This solution method is computationally expensive in diffusive regions of the problem due to the high effective scattering, and is limited in time step size.

A HOLO approach is a desirable alternative because the LO method can easily resolve the solution in these diffusive regions. For simplicity, our HOLO method will use a backwards Euler discretization in time, as well as constant heat capacity and cell-wise constant opacity. The time discretized equations are

$$\mu \frac{\partial I^{n+1}}{\partial x} + \left(\sigma_t + \frac{1}{c \Delta t} \right) I^{n+1} = \frac{\sigma_s}{2} \phi^{n+1} + \frac{1}{2} (\sigma_a a c T^4)^{n+1} + \frac{I^n}{\Delta t c} \quad (3)$$

$$\rho c_v \frac{T^{n+1} - T^n}{\Delta t} = \int_{-1}^1 \sigma_a I^{n+1}(x, \mu) d\mu - \sigma_a a c (T^4)^{n+1}. \quad (4)$$

where Δt is the time step size and the superscript n is used to indicate the n -th time step. The HOLO method will use moments of the above two equations to resolve the non-linear temperature dependence between the two equations. ECMC will be used to solve the fixed source version of Eq. (3) to provide consistency terms in the LO system to correctly solve the problem.

2.2. Overview of the HOLO Algorithm

In the HOLO context, the LO solver handles the physical scattering and resolves the material energy spatial distribution. The LO equations are based on angular integrals and spatial moments formed over a finite element mesh of Eq. (4) and Eq. (3). The LO equations for the radiation energy balance are similar to double- P_0 equations, but with spatially varying consistency parameters that are analogous to a variable Eddington factor. These consistency parameters are lagged in each LO solve, estimated from the previous HO solve. If the angular consistency parameters were estimated exactly, then the LO equations are exact with respect to the chosen spatial discretization. The material energy equation consists of only spatial moments and does not contain any consistency terms, providing energy conservation. The HO solver is not required to conserve energy.

The solution to the LO system is used to construct a spatially LD representation of the scattering and emission sources on the right hand side of Eq. (3). This defines a fixed-source, pure absorber transport problem for the HO operator. This transport problem, which we refer to as the HO problem, is solved using ECMC with adaptive mesh refinement. The HO problem defines a characteristic method that uses MC to invert the continuous streaming plus removal operator with an LD representation of the source terms (including inscattering). The ECMC algorithm allows for the statistical noise in the MC solution to be efficiently reduced to any desired precision within the limits of computational memory. Thus, the HO solve produces a globally-accurate, LD representation of the angular intensity $\tilde{I}(x, \mu)$. Here, we emphasize that the ECMC algorithm computes a projection of the exact solution onto a LDFE space-angle trial space, rather than a standard FE solution. This is in general far more accurate than a standard FE solution.

Once computed, the projected LD angular intensity $\tilde{I}(x, \mu)$ is used to evaluate the LO consistency parameters for the next LO solve. Since there is a global, functional representation of the angular intensity, LO parameters are estimated using quadrature and do not require additional tallies. The HO solver does not produce a new temperature in the thermal radiative transport (TRT) context; it is only used to estimate the angular parameters in the LO solution, which eliminates typical operator splitting stability issues that require linearization of the emission source.

The process of a LO estimate of the transport sources for the HO solver, followed by an ECMC solve to estimate LO parameters, represents one HOLO fixed-point iteration. The consistency terms force the HO and LO solutions for $\phi(x)$ to be consistent to the order of the current HOLO iteration error. The solution for $T(x)$ is only estimated via the LO solution (which preserves the accuracy of the HO solution). The HOLO iterations are performed until the solutions are converged to a desired precision, within each time step. Currently, the HOLO convergence criteria is based on the convergence of the FE representation of $\phi^{n+1}(x)$ between successive HOLO iterations k . The HOLO algorithm, for each time step, is

1. Perform a diffusion-equivalent solve of the LO system to produce $T^0(x)$ and $\phi^{(0)}(x)$ at t_{n+1}
2. Solve the HO system for $\tilde{I}^{k+1/2}(x, \mu)$ using ECMC, based on the current LO estimate of the right hand emission and scattering sources.
3. Compute LO consistency parameters with $\tilde{I}^{k+1/2}$.
4. Solve the LO system using new HO consistency parameters to produce a new estimate of ϕ^{k+1} and T^{k+1} at t_{n+1} .

5. Repeat 2 – 4 until convergence is achieved.
6. Move to the next time step.

3. The Low-Order Solver for the Radiative Transfer Equations

The LO system is formed by taking spatial basis moments and half-range angular integrals of Eq. (3) and Eq. (4). The spatial moments are taken over each spatial cell i : $x \in [x_{i-1/2}, x_{i+1/2}]$, weighted with the standard linear Lagrange interpolatory basis functions. For example, the L moment operator is defined by

$$\langle \cdot \rangle_{L,i} = \frac{2}{h_i} \int_{x_{i-1/2}}^{x_{i+1/2}} b_L(x) (\cdot) dx, \quad (5)$$

where $h_i = x_{i+1/2} - x_{i-1/2}$ is the width of the spatial element and $b_L(x) = (x_{i+1/2} - x)/h_i$ is the FE basis function corresponding to position $x_{i-1/2}$. The right moment $\langle \cdot \rangle_R$ is defined with weight function $b_R(x) = (x - x_{i-1/2})/h_i$. Here, it is emphasized that $\phi_{L,i}$ and $\phi_{R,i}$ represent the values of the scalar intensity at $x_{i-1/2}$ and $x_{i+1/2}$ that preserve the zeroth and first moments over the cell, not to be confused with $\langle \phi \rangle_L$ and $\langle \phi \rangle_R$, which represent a volume-averaged moments. The positive and negative half-range integrals of the angular flux are defined as $\phi^+(x) = \int_0^1 I(x, \mu) d\mu$ and $\phi^-(x) = \int_{-1}^0 I(x, \mu) d\mu$, respectively. Thus, in terms of half-range quantities, $\phi(x) = \phi^-(x) + \phi^+(x)$.

Pairwise application of the L and R basis moments with the $+$ and $-$ half-range integrals to Eq. (3) equation ultimately yields four radiation balance equations, for each cell. For example, the equation result from application of the L moment and positive half-range integral is

$$\begin{aligned} -2\mu_{i-1/2}^{n+1,+} \phi_{i-1/2}^{n+1,+} + \langle \mu \rangle_{L,i}^{n+1,+} \langle \phi \rangle_{L,i}^{n+1,+} + \langle \mu \rangle_{R,i}^{n+1,+} \langle \phi \rangle_{R,i}^{n+1,+} + \left(\sigma_t^{n+1} + \frac{1}{c\Delta t} \right) h_i \langle \phi \rangle_{L,i}^{n+1,+} \\ - \frac{\sigma_s h_i}{2} \left(\langle \phi \rangle_{L,i}^{n+1,+} + \langle \phi \rangle_{L,i}^{n+1,-} \right) = \frac{h_i}{2} \langle \sigma_a^{n+1} ac T^{n+1,4} \rangle_{L,i} + \frac{h_i}{c\Delta t} \langle \phi \rangle_{L,i}^{n,+}, \end{aligned} \quad (6)$$

where the $\langle \phi^\pm \rangle$ terms represent volumetric and angular averaged fluxes, and the $\phi_{i-1/2}$ term represents an angular average over a face. The moments of the source term will be defined later on. The $-$ direction equations are essentially the same with $+$ \rightarrow $-$ (there is also a difference in the upwinding and extrapolated face terms, explained below). The angular consistency terms are defined in terms of half-range averages, e.g.,

$$\langle \mu \rangle_{L,i}^+ = \frac{\frac{2}{h_i} \int_0^1 \int_{x_{i-1/2}}^{x_{i+1/2}} \mu b_L(x) I^{n+1}(x, \mu) dx d\mu}{\frac{2}{h_i} \int_0^1 \int_{x_{i-1/2}}^{x_{i+1/2}} b_L(x) I^{n+1}(x, \mu) dx d\mu}. \quad (7)$$

The consistency terms for the R basis moment and $\mu_{i\pm 1/2}^\pm$ face terms are defined analogously.

To derive the LO material energy equations, first $T(x)$ is assumed to be in the LD trial space, i.e., $T(x) \simeq T_{L,i} b_L(x) + T_{R,i} b_R(x)$, $x \in (x_{i-1/2}, x_{i+1/2})$. Similarly, the emission term is represented in the material and radiation equations with the LDFE interpolant $\sigma_{a,i} ac T^4(x) = \sigma_{a,i} ac [T_{L,i}^4 b_L(x) + T_{R,i}^4 b_R(x)]$. The L and R spatial moments are taken of the material energy equation, using the LDFE definition for $T(x)$ and $T^4(x)$. For example, the final LO material energy

equation resulting from application of the L moment is

$$\begin{aligned} \frac{\rho c_v}{\Delta t} \left[\left(\frac{2}{3} T_{L,i} + \frac{1}{3} T_{R,i} \right)^{n+1} - \left(\frac{2}{3} T_{L,i} + \frac{1}{3} T_{R,i} \right)^n \right] + \sigma_a^{n+1,k+1} \left(\langle \phi \rangle_{L,i}^+ + \langle \phi \rangle_{R,i}^- \right)^{n+1} \\ = \sigma_a^{n+1} ac \left(\frac{2}{3} T_{L,i}^4 + \frac{1}{3} T_{R,i}^4 \right)^{n+1}. \quad (8) \end{aligned}$$

3.1. Closing the Non-Linear System

The six degrees of freedom (DOF) over each cell i are the four moments $\langle \phi \rangle_{L,i}^+$, $\langle \phi \rangle_{R,i}^+$, $\langle \phi \rangle_{L,i}^-$, and $\langle \phi \rangle_{R,i}^-$ and the two spatial LDFE edge values $T_{L,i}$ and $T_{R,i}$. The four radiation and two material energy equations define a system of equations for the six DOF without additional discretization. However, the consistency parameters (e.g., Eq. (7)) are not known a priori, and there is no relation between the volume and face averaged quantities. We will use the HO solver to close these equations.

A lagged estimate from the previous HO solve is used to estimate the angular consistency parameters. In the HOLO context, the equations for LO unknowns at iteration $k + 1$ use consistency parameters computed using Eq. (7) with the latest HO solution $\tilde{I}^{n+1,k+1/2}$ as an approximation for I^{n+1} . For the initial LO solve, all average μ parameters are set to $\pm 1/\sqrt{3}$, yielding a solution equivalent to an S_2 solution. For this initial S_2 solve, it is necessary to renormalize radiation boundary conditions to get an accurate solution and preserve conservation, which is standard in S_N methods. To close the LO system spatially, the usual LD upwinding approximation is used. For example, for Eq. (6), the face terms $\mu_{i-1/2}$ and $\phi_{i-1/2}$ terms are upwinded from the previous cell $i - 1$ or from a boundary condition; the terms at $x_{i+1/2}$ are linearly extrapolated, computed using the L and R basis moments. Because the HO solver uses an LD representation, this LO spatial closure is inherently consistent. Because there are no spatial temperature derivatives, there is no evaluation of T at faces and thus no need for an upwinding closure.

3.2. Alternative Closure Near the Wave Front

The LD closure is not strictly positive. In particular, for optically thick cells with a highly varying temperature distribution, the solution is driven negative. In thick regions of TRT problems, reasonably fine spatial cells can still be on the order of millions of mean free paths; negativities with LD are simply unavoidable in these cells and mesh refinement is of minimal use. Typically, for a standard LDFE method, the equations are lumped to produce a strictly positive solution. However, standard FE lumping procedures would make computing the consistency terms from the HO solver difficult. An alternative discontinuous spatial discretization is used that uses a relation between the spatial moments and outflow that produces the same result as the standard FE lumping procedure.

The L and R moments are defined the same as before, preserving the same average within a cell, but the relation between the moments and the outflow is modified to force a strictly positive outflow. For example, for positive μ , the outflow is now defined as

$$\phi_{i+1/2}^+ = \langle \phi \rangle_R^+. \quad (9)$$

This is different than the standard extrapolated outflow for LD given by $\phi_{i+1/2}^+ = 2\langle \phi \rangle_R^+ - \langle \phi \rangle_L^+$. Because the basis function $b_R(x)$ is strictly positive, the above moment and thus outflow is inherently positive. This spatial discretization is second order, as compared to the third order of standard LD. The same upwinding as before is used for this discretization. This closure is only used in cells in which negativities occur.

4. Solving the Non-Linear LO system

We have used Newton's method to solve the global system of coupled LO equations, based on a typical linearization of the Planckian source with opacities evaluated at lagged temperatures. Because we have only considered problems with constant densities and heat capacities, the following derivation is depicted in terms of temperature T rather than material energy for simplicity without loss of consistency.

Application of the first order Taylor expansion in time of the gray emission source $B(T)$, about some temperature T^* at some time near t^{n+1} gives

$$\sigma_a^* acT^{4,n+1} \simeq \sigma_a^* ac [T^{*4} + (T^{n+1} - T^*)4T^{*3}] \quad (10)$$

where the superscript $*$ denotes evaluation at T^* . This expression is substituted for the emission term in the time discretized material energy equation given by Eq. (??). The resulting equation is manipulated further to solve for T^{n+1}

$$T^{n+1} = \frac{\frac{\sigma_a^* \Delta t}{\rho c_v} [\phi^{n+1} - acT^{*4}] + (T^n - T^*)}{1 + \sigma_a^* ac \Delta t \frac{4T^{*3}}{\rho c_v}} + T^*. \quad (11)$$

This provides an explicit expression for T^{n+1} as a function of T^* and the radiation scalar intensity ϕ^{n+1} , i.e.,

$$T^{n+1} = \frac{1}{\rho c_v} f \sigma_a^* \Delta t (\phi^{n+1} - caT^{*4}) + fT^n + (1 - f)T^*. \quad (12)$$

where $f^* = (1 + \sigma_a^* c \Delta t \beta^*)^{-1}$ with $\beta^* = \frac{4aT^{*3}}{\rho c_v}$ and the superscript $*$ indicates evaluation at T^* . The expression for T^{n+1} can be substituted back into Eq. (10) to form an explicit approximation for the emission source at t_{n+1} as

$$\sigma_a acT^{4,n+1} \simeq \sigma_a^* (1 - f^*) \phi^{n+1} + f^* \sigma_a^* acT^{4,n} + \rho c_v \frac{1 - f^*}{\Delta t} (T^n - T^*) \quad (13)$$

The material temperature is updated at the end of the time step using Eq. 12.

Based on a guess for T^* , the above equation gives an expression for the Planckian emission source in the right hand side of Eq. (6) with an additional effective scattering source. This allows for four linear equations for the four remaining radiation unknowns to be fully defined. The final equation for the left basis moment and positive flow, for constant f^* and σ_a^* over a cell, becomes

$$\begin{aligned} -2\mu_{i-1/2}^{n+1,+} \phi_{i-1/2}^{n+1,+} + \langle \mu \rangle_{L,i}^{n+1,+} \langle \phi \rangle_{L,i}^{n+1,+} + \langle \mu \rangle_{R,i}^{n+1,+} \langle \phi \rangle_{R,i}^{n+1,+} + \left(\sigma_t^* + \frac{1}{c \Delta t} \right) h_i \langle \phi \rangle_{L,i}^{n+1,+} \\ - \frac{h_i}{2} (\sigma_s^* + \sigma_a^* (1 - f^*)) \left(\langle \phi \rangle_{L,i}^{n+1,+} + \langle \phi \rangle_{L,i}^{n+1,-} \right) = \\ \frac{1}{2} h_i \sigma_a^* ac f^* \left(\frac{2}{3} T_{L,i}^{4,n} + \frac{1}{3} T_{R,i}^{4,n} \right) + \frac{h_i}{c \Delta t} \langle \phi \rangle_{L,i}^{n,+} \end{aligned} \quad (14)$$

Once these linear equations have been solved for ϕ^{n+1} , a new estimate of T^{n+1} can be determined using Eq. (12), providing energy conservation. To account for spatial dependence, Eq. (12) can simply be evaluated at the edge values in a cell, e.g., with ϕ_L^{n+1} and T_L^* to get T_L^{n+1} .

Based on these equations, the algorithm for solving the LO system is defined as

1. Guess T_L^* and T_R^* , typically using T^n .

2. Build the LO system based on the effective scattering $\sigma_a^*(1 - f^*)$ and emission terms (i.e., evaluation of Eq. (14)).
3. Solve the linearized LO system to produce an estimate for ϕ^{n+1} .
4. Evaluate a new estimate of the $T_{L,i}$ and $T_{R,i}$ at the end of the time step \tilde{T}^{n+1} using Eq. (??).
5. $T^* \leftarrow \tilde{T}^{n+1}$.
6. Repeat 2-5 until \tilde{T}^{n+1} and ϕ^{n+1} are converged.

5. The ECMC High Order Solver

The HO solver must solve a fixed source, pure absorber transport problem. We will solve this equation using ECMC. A more detailed description of the ECMC method can be found in [?]. In the HOLO context, the sources are estimated by the previous LO solve. The equation with HOLO iteration indices is

$$\mu \frac{\partial I^{n+1,k+1/2}}{\partial x} + \left(\sigma_t^k + \frac{1}{c\Delta t} \right) I^{n+1,k+1/2} = \frac{\sigma_s^k}{2} \phi^{n+1,k} + \frac{1}{2} \left(\sigma_a^k a c T^4 \right)^{n+1,k} + \frac{\tilde{I}^n}{c\Delta t} \quad (15)$$

where k represents the outer HOLO iteration index. Material property indices will be suppressed from now on. Here, $k + 1/2$ denotes the HO solve within HOLO iteration k , whereas k and $k + 1$ represent successive LO solves. In operator notation, the previous equation can be written as

$$\mathbf{L}^k \psi^{k+1/2} = q^k \quad (16)$$

where $\psi^{k+1/2}$ is the solution of the angular intensity I^{n+1} based on the k -th estimate of q^k . The linear operator \mathbf{L}^k is the streaming plus removal operator defined by the left hand side of Eq. (3), and q^k is defined by the LD representation of the right hand side, as estimated via the k -th LO solve.

To define the ECMC algorithm, the HO-LO iteration indices k are dropped, as the LO estimated q^k and \mathbf{L}^k remain constant over the entire HO solve. The i -th approximate solution to Eq. (16) (i represents inner HO batches) is represented as $\tilde{\psi}^{(i)}$. The i -th residual is

$$r^{(i)} = q - \mathbf{L} \tilde{\psi}^{(i)}. \quad (17)$$

Addition of $\mathbf{L} \psi - q = 0$ to the above equation and manipulation of the result yields the error equation

$$\mathbf{L}(I^{n+1} - \tilde{\psi}^{(i)}) = \mathbf{L} \tilde{\epsilon}^{(i)} = r^{(i)} \quad (18)$$

where I^{n+1} is the exact solution and $\tilde{\epsilon}^{(i)}$ is finite element representation of the error in $\tilde{\psi}^{(i)}$. The above equation is inverted yielding the Monte Carlo estimate of the error in $\tilde{\psi}^{(i)}$, i.e.,

$$\tilde{\epsilon}^{(i)} = \mathbf{L}^{-1} r^{(i)} \quad (19)$$

where \mathbf{L}^{-1} is the Monte Carlo inversion of the streaming and removal operator. The solution $\tilde{\psi}^{(i)}$ represents the projection of the exact Monte Carlo solution onto the trial space, rather than a standard finite element solution. For example, the LD trial space preserves the zeroth and first spatial moment over a cell; the zeroth moment is computed using a standard path-length volumetric flux tally, which is equivalent to typical volumetric averages computed in Monte Carlo calculations. The primary truncation error is in the LD representation of the right hand side source terms and residual at each iteration. The only tallies used are volumetric flux tallies over all of the finest space-angle elements; tallies are computed for the average, slope in x , and slope in μ of $I^{n+1}(x, \mu)$.

The HO solver requires an LD representation of the Planckian emission sources. For cell-wise constant cross section, the representation takes the form $B(T, x) = \sigma_a a c (T_L^4 b_L(x) + T_R^4 b_R(x))$. For reference, for Eq. (3) the residual at iteration i in the HO solve is

$$r^{(i),k+1/2} = \frac{\sigma_s}{2} \phi_{LD}^{n+1,k} + \frac{1}{2} (\sigma_a^* a T^4)_{LD}^{n+1,k} + \frac{\tilde{I}^n}{\Delta t c} - \left(\frac{\partial \tilde{I}^{n+1,k+1/2}}{\partial x} + \left(\sigma_t + \frac{1}{c \Delta t} \right) \tilde{I}^{n+1,k+1/2} \right)^{(i)} \quad (20)$$

where the k terms are LD in space on the coarsest mesh and are not recalculated at any point during the HO solve. The functional form of \tilde{I}^n is defined over the finest mesh from the last HOLO iteration of the previous time step. For simplicity, \tilde{I}^n is projected onto the current mesh in the HO solve.

An initial guess for the angular intensity is computed based on the previous solution for I^n , or for a later HOLO iteration from the last batch estimate from the previous HO solve. This is an important step, and significantly reduces the required number of particles per time step because I does not change drastically between time steps in highly diffusive problems. The initializations are computed as a projection of the solution onto the initial coarsest mesh.

The ECMC algorithm is

1. Initialize the guess for $\tilde{I}^{n+1,(0)}$ to \tilde{I}^n or the projection of \tilde{I}^{n+1} from the latest HO solve
2. Compute $r^{(i)}$.
3. Solve $\tilde{\epsilon}^{(i)} = \mathbf{L}^{-1} r^{(i)}$
4. Compute a new estimate of the intensity $\tilde{I}^{n+1,(i+1)} = \tilde{I}^{n+1,(i)} + \tilde{\epsilon}^{(i)}$
5. Repeat steps 2 – 4 until desired convergence criteria is achieved.

Currently, the convergence criteria is based on the $\|\tilde{\epsilon}^{(i)}\|_2$, i.e.,

$$\|\tilde{\epsilon}^{(i)}\|_2 = \int_{\mathcal{D}} \int_{-1}^1 \tilde{\epsilon}^{(i)}(x, \mu) d\mu dx \leq \text{TOL}. \quad (21)$$

Exponential convergence is obtained because with each inversion of \mathbf{L} a better estimate of the solution is being used to compute the new residual, decreasing the magnitude of the Monte Carlo source each iteration i . Each Monte Carlo estimate for the error has a sample mean that contains a statistical distribution that decreases by the standard $1/N$ reduction in variance, however the angular intensity can obtain a higher convergence rate because of the new information about the solution introduced each iteration. If the statistical estimate of $\tilde{\epsilon}$ is not sufficiently accurate, then the iterations would diverge. Currently, the error is monitored and ensured to not be too large, relative to ϵ .

Because the exact angular intensity does not in general lie within the LD trial space, the iterative estimate of the error will eventually stagnate once the error cannot be sufficiently represented by a given LD FE mesh. An adaptive h -refinement algorithm is used to allow the system to continue converging towards the exact solution. Once the error has stagnated, refinement is performed on some percentage of the cells based on the jump errors over each face of a space-angle cell. Each space-angle cell to be refined is divided into four equal-sized cells. The previous estimate of the angular intensity is projected onto the new cells (resulting in a continuous solution over each of the cells); this is an inaccurate solution over the new mesh, and thus the first iteration after refinement will result in an increase in the error. The error is also increased due to the new, more accurate projection of \tilde{I}^n onto the new mesh. However, typically within one more iteration the error will have decreased past the error prior to refinement.

It is also noted that the trial space representation of the HO angular intensity contains most of the information necessary to estimate the exact Monte Carlo projection of the LO parameters (the average and first moments of the angular intensity in angle and space), except for the cross moment ($\langle x\mu \rangle$). However, because the LD angular intensity representation is accurate over the refined HO mesh, that functional form can be used to estimate the cross moment on the LO mesh with a similar degree of accuracy.

5.1. Continuous Energy Deposition Tracking

As in [?], we will exponentially attenuate particles to reduce statistical variance in the tallies. Because the ECMC problem solves a pure-absorber problem, particles can be allowed to stream, with their weight deterministically modified along the path. Because the particles are exponentially attenuated, the weight is adjusted as $w(x) = w(x_0) \exp(-|(x - x_0)/\mu|)$. The tallies are modified to account for this weight exponential attenuation by evaluating the integral along the path length.

5.2. Source Bias

A modified stratified sampling method has been used to effectively distribute particle histories throughout the problem. The residual gives a good indication of where particles are most likely to contribute to the error, particularly in diffuse problems. In higher dimensions a more accurate weighting that accounts for the probability of particles in one region contributing to particles in another region may be necessary.

In the modified sampling algorithm the number of particle histories sampled in each cell is proportional to the magnitude of the residual in each cell, unless the number of particles in a cell is below some minimum. In this case, a minimum number of histories, specified by the user, is placed in cells below the minimum. This is to prevent bad statistics in certain cells, particularly in the case of adaptively refined meshes. There is also a relative probability cutoff such that cells with an insignificant residual will have no histories sampled there. In these regions the problem is remaining in steady state and the solution is known exactly. The cutoff is set relative to equal probability of being born in each cell and is adjusted to account for the fact that smaller cells have lower probability. By ensuring sampling of each significant region of phase space, energy is locally conserved more accurately.

The implementation of the above equation is modified such that the total number of histories is as close to original requested number of histories as possible. This is done by first filling the cells with N_{min} histories and distributing the remaining number of histories proportional to p_j .

5.3. Fixup Near the Wave-Front

In cells near the wave front, the LD trial space results in negativities as with the LO solver. Currently, we just check the consistency terms at the end of the HO solve and detect if they lie in the appropriate half-space. If the terms are non-physical, then they are set to diffusion-equivalent terms. This may lead to artificially fast wave-speeds in certain cells. A better alternative would be to switch to another trial space in these cells that is strictly positive.

6. Results

To demonstrate this method we will compare the results of our method with IMC with source tilting, which should be effectively solving the same problem. It is not easy to compare the two results with out an analytical solution for a Marshak wave. These results are presented to verify that with sufficient particle histories to converge the problem our code is getting accurate solutions and to demonstrate issues to be resolved in future work. Future publications will focus on iteration counts and using a limited number of particles. The radiation energy distribution is plotted as an equivalent temperature given by

$T_r = \sqrt[4]{\phi/(ac)}$. Although Thompson scattering can be handled by the LO solver in this method, we have only considered pure absorber problems here. We have demonstrated this method with scattering included in the Eigenvalue solves (??).

For all problems, a relatively high HOLO iteration relative error tolerance of 1.E-02 between radiation energy distributions was used, leading to only one HO solve being performed per time step to limit simulation times. A tolerance of 1.E-06 was used for the LO solver and 5.E-03 for the HO solver. To ensure convergence of the HO solver, a relatively large number of histories per batch > 150000 was used, unless otherwise noted. This is a sufficiently high number of histories that for the majority of time steps HO solver is converged within 2 batches; in reality more efficient error reduction could be achieved by using more batches with less particles per batch. It is important to note that because there is no scattering, each history can be solved more efficiently than in IMC. Adaptive mesh refinement and more batches were used as necessary (typically in optically thin regions) by the solver, testing this aspect of the code. All meshes begin with 200 spatial cells and two angular cells in each half-range.

6.1. Constant Solution Infinite Medium

To check the basic functionality of the HOLO algorithm a constant solution representing an infinite medium was performed, with results shown in Fig 1. Even though the LO solver can get the solution exactly right, we forced the residual to sample 2500 particles per time step. Because of the initial guess of \tilde{I}^n for the radiation, the residual source is very small in this case and the statistical noise is on the order of roundoff for the ECMC solver. In general, regions that are in steady state will not sample residual particles. For the IMC results, statistical noise is relevant even for a constant solution for the unless significantly more than 2500 particles per time step is used.

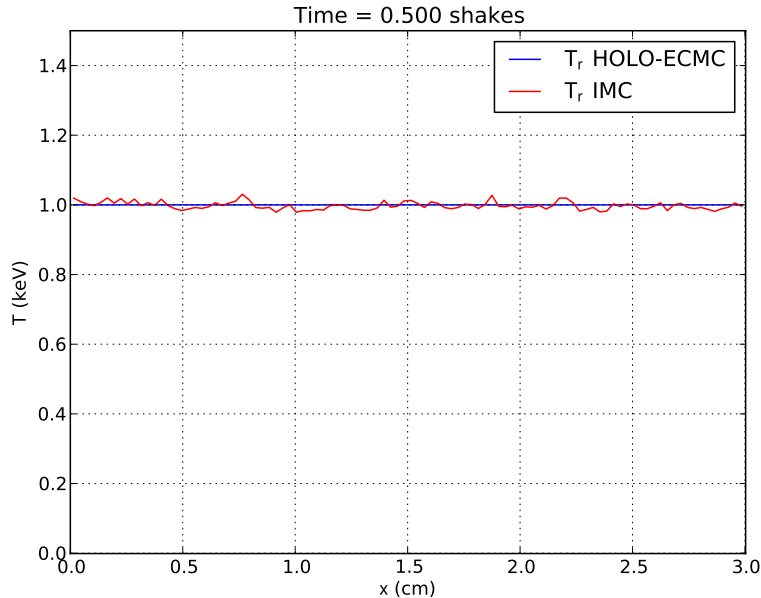


Figure 1. Comparison of IMC and HOLO-ECMC results for an equilibrium solution.

6.2. Two Material Problem

This problem consists of an optically thin and an optically thick material region, with constant cross sections. This problem demonstrates the utility of the hybrid method in both the streaming (left) and

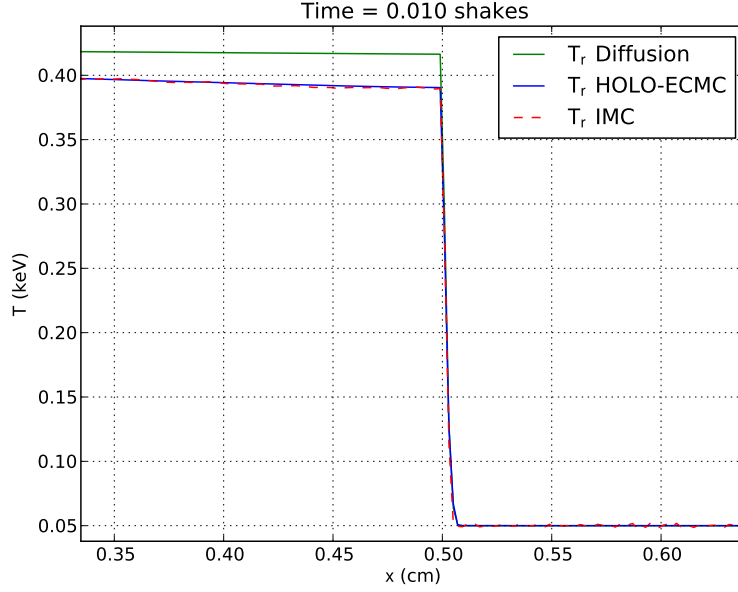


Figure 2. Comparison of radiation temperatures for HOLO-ECMC, IMC, and diffusion for two material problem after 10 time steps.

diffuse (right) regions of the domain. The material properties are given in Table I. Initially the radiation and material energies are in equilibrium at a temperature of 0.05 keV. An isotropic incident intensity of 0.500 keV is applied at $x = 0$ at $t = 0$; the incident intensity on the right boundary is 0.05 keV. The simulation was ran for 5 shakes with a time step size of 0.001 shakes.

Fig. 2 compares the HOLO, IMC, and diffusion computed radiation temperatures at $t = 0.010$ sh. Both curves plot the cell-wise averages rather than the LD representation. At this point in the simulation, the radiation is in the optically thin region of the problem, so the diffusion solution has propagated too quickly. The IMC and HOLO methods agree well. Fig 3 compares the IMC and HOLO radiation temperature profiles at the end of the simulation. The wave fronts show good agreement. These results were computed using the lumping-equivalent discretization for the LO solver, noting that this is not consistent with the HO solver. A consistent discretization in the HO solver would likely produce a more accurate solution.

	$x \in [0, 0.5) \text{ cm}$	$x \in [0.5, 1.0] \text{ cm}$
$\sigma_a \text{ (cm}^{-1}\text{)}$	0.2	2000
$\rho \text{ (g cm}^{-3}\text{)}$	0.03	10.0
$c_v \text{ (jks/keV-g)}$	0.1	0.1

Table I. Two material problem properties

6.3. Marshak Wave

For this problem, initially the radiation and material energies are in equilibrium at 2.5E-05 keV. An isotropic incident intensity of 0.150 keV is applied at $x = 0$ at $t = 0$; the incident intensity on the right boundary is 2.5E-05 keV. The material properties are $\rho = 1 \text{ g cm}^{-3}$ and $c_v = 0.013784 \text{ jks/keV-g}$. The absorption cross section varies as

$$\sigma(T) = \frac{0.001\rho}{T^3}, \quad (22)$$

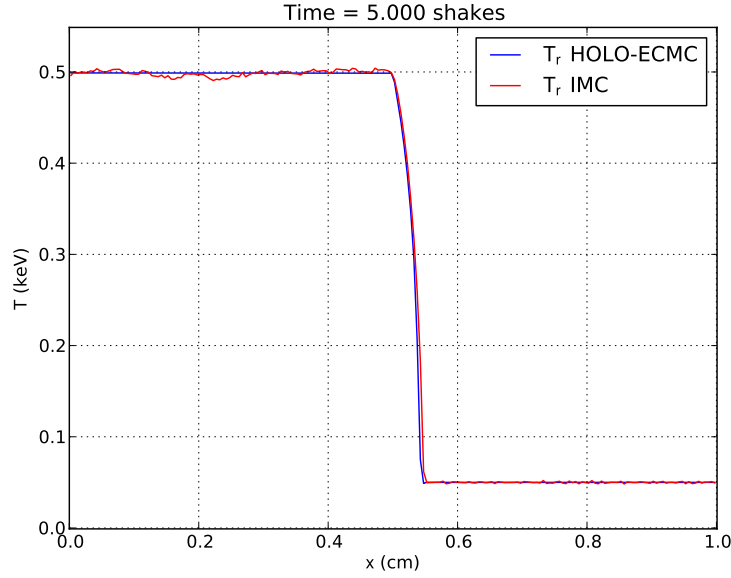


Figure 3. Radiation temperature profiles for two material problems after 5 shakes.

which introduces a strong non-linearity into the problem. The simulation was ran for 5 shakes with a time step size of 0.001 shakes.

Fig. 4 compares the cell average radiation temperatures for the IMC and HOLO methods. The radiation wave front is significantly farther ahead in the HOLO method. Simulations with a refined mesh and time step did not remove this discrepancy, and the IMC solution did not move much. This suggests that the fact that IMC lags opacities is not the cause of the difference. There are a couple notable issues that could be the cause of the discrepancy. Although the averages are positive, the LD representation of the intensity leads to negativities near the steep wave front. The negativities lead to poor consistency terms. Additionally, histories with large negative weights in the wave-front region enter the equilibrium cells to the right and cause poor statistics due to the thick cross section. This leads to consistency terms that are unphysical ($\langle \mu \rangle \notin [-1, 1]$) in cells near the wave front. In these cells diffusion equivalent parameters are used for the parameters that are bad. The diffusion approximation near the wave front ultimately leads to a faster wave speed. There are a couple ideas discussed in future work for correcting the bad statistics on the right side of the wave front.

Fig. 5 plots the LD representation of the material and radiation energies (in this figure the negativities in the radiation are plotted as zero, but do exist at the wave front). The steep, unphysical gradients in the material temperature distribution are a result of using cell-wise constant cross sections. More accurate, spatially varying cross sections will be implemented in future work.

To correct issues introduced into the LO solver by negativities in the radiation equation, the lumped-equivalent spatial discretization discussed previously was implemented for the LO solver. The HO solver was left as LD in space in angle. Thus, the LO discretization preserves the L and R moments of the HO solution, but does not use a consistent outflow. Results with this alternative LO discretization are displayed in Fig. 6. The wave speeds are much closer, but there is still a noticeable discrepancy. The first and easiest method for resolving this issue would be to implement a similar strictly positive discretization in the HO solver to ensure consistency. This will likely resolve some of the issues with noise resulting in bad consistency terms.

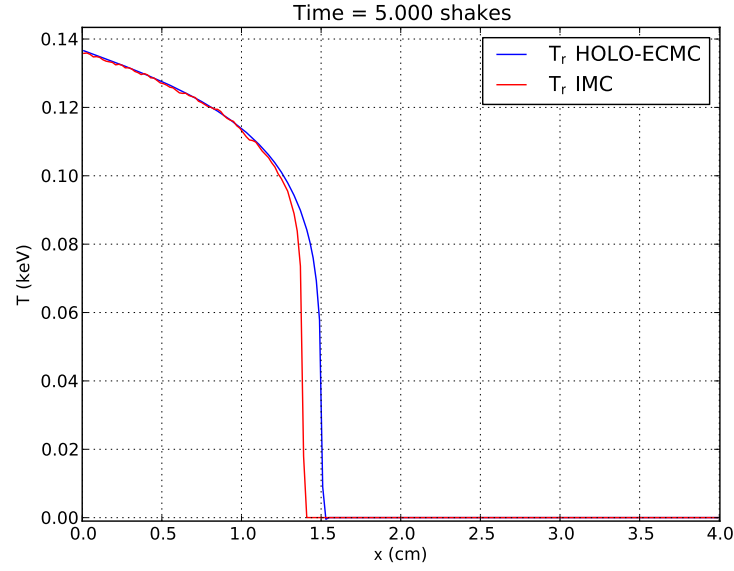


Figure 4. Comparison of cell-averaged radiation temperatures for Marshak wave problem.

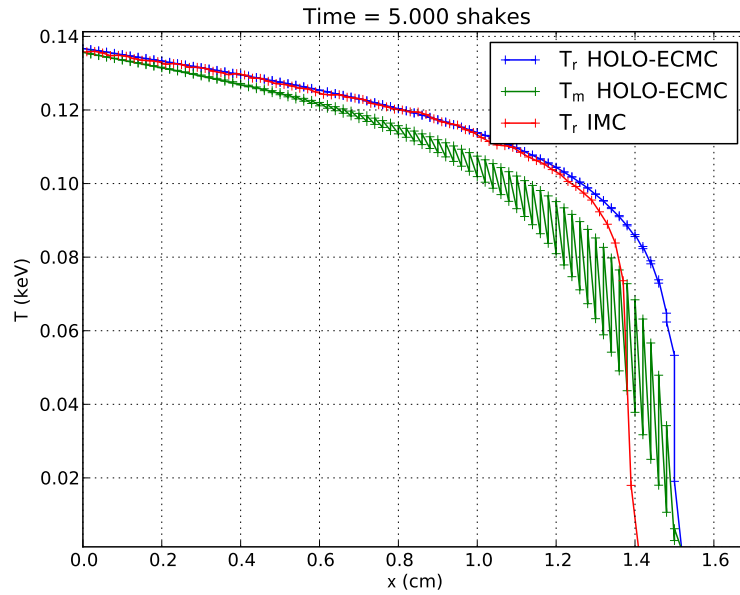


Figure 5. LD representation of material and radiation energy temperatures for HOLO, as compared to cell averages for IMC, for Marshak wave problem.

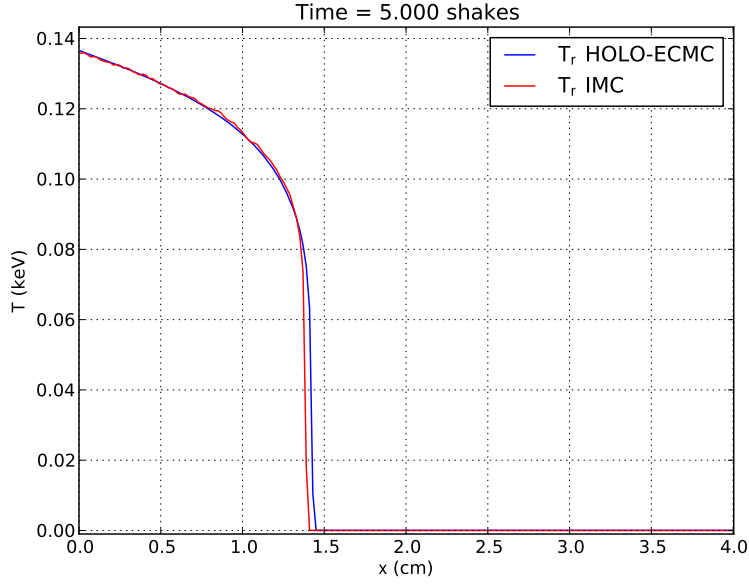


Figure 6. Marshak wave problem with lumped-equivalentLD spatial representation for LO solver.

7. Conclusions

We have been able to reproduce the IMC solutions for certain problems using a new HOLO method. The primary difficulty is in the speed of wave fronts for the Marshak wave problem. The LO solver resolves the non-linearities in the equations resulting in a fully implicit time discretization. The ECMC approach, with initial guesses based on the previous radiation intensity, results in efficient reduction of statistical error and allows for particles to be distributed to largely varying regions of the problem. Analytic transport benchmark results are needed for a direct comparison of the accuracy between the HOLO and IMC methods. However, most of the transport benchmarks utilize reflective boundary conditions. Reflective boundary conditions have been successfully implemented in the LO solver, but have not yet been successfully implemented in the HO solver. An explanation of the difficulties with reflective boundary conditions is detailed in the Appendix. Once the reflective boundary conditions are correctly implemented, a comparison of the accuracy of the two methods can be made.

Other future work will include continuous energy deposition tallies as a form of variance reduction. Also, a strictly positive spatial discretization in the HO system will be implemented. If this does not resolve issues with statistical noise, a form of cell-wise error filtering will be implemented. With the exception of the first batch, it is not expected that the relative error in a cell should be less than 1. If this is the case, then it is likely due to a bad MC estimate. These cases could be rejected and the previous estimate of the intensity in that cell used for the next calculation, or a damped version of the change could be used. Long term work will include adding the ability to handle opacities that vary as ILD with temperature in space. A multigroup formulation will be explored, before extending to multiple spatial dimensions.

A. SECOND OR SUBSEQUENT MAJOR HEADING

This is Section A. It is followed by a Subsection, that is, A.1. For this class file you must use the `\Section{}` to denote sections (note the capital “S”). The style for subsection titles and all text in this template is defined in the *mc2013.cls* file. Make sure to avoid widow/orphan lines.

A.1. Subsection Title: First Character of Each Non-trivial Word is Uppercase

Double-space before and after secondary titles is automatic. Figures and tables should appear as close as possible to where they are first cited, e.g., Fig. 7, in the text. Figures are numbered in Arabic numerals, with the caption centered below the figure, in **boldface**.

Triple-space before the figure, and after the figure caption.

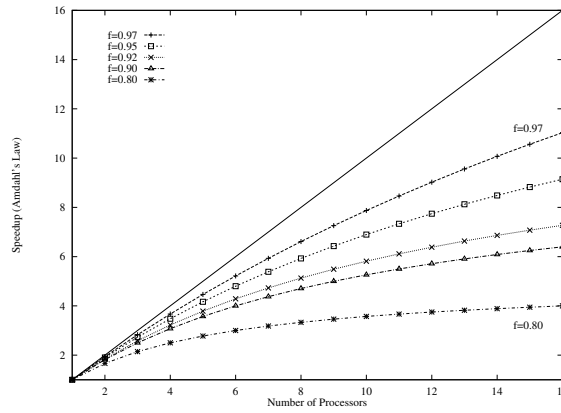


Figure 7. Speedup vs Number of Processors for Varying Parallelizable Fractions in Order to See a Multiple Line Caption

When importing figures or any graphical image please verify two things:

- Any number, text or symbol is in Times font and is not smaller than 10-point after reduction to the actual window in your paper
- That is can be translated into PDF

Equations, such as Eq. (23), should be centered and sequentially numbered to the flush right of the formula.

$$Speedup = \frac{1}{\frac{f}{p} + (1 - f)} \quad (23)$$

The continuation of a paragraph after an equation should not be indented. All paragraphs, as well as section or subsection headings, are separated by just one single empty line.

A.1.1. Sub-section level and lower: only first character uppercase

See Table II for a sample table. The `tbls` package is recommended for improved row and column spacing. Notice the caption appears above the table by setting the `\caption` command immediately after the `\begin{table}`. Tables are numbered in Roman numerals, with the caption centered above the table, in **boldface**. Triple-space before and after the table.

Table II. Parallel Performance for the Sample Problem

Number of Processors	Wall-Clock Time ^a (min)	Speedup (T_s/T_p)	Efficiency (%)
1	100.0	—	—
2	52.6	1.9	95.0

B. CONCLUSIONS

Present your summary and conclusions here.

ACKNOWLEDGEMENTS

This research was performed, in part, using funding received from the DOE Office of Nuclear Energy's Nuclear Energy University Programs.

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