A 3DHZETRN Code in a Spherical Uniform Sphere with Monte Carlo Verification

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Abstract

The computationally efficient HZETRN code has been used in recent trade studies for lunar and Martian exploration and is currently being used in the engineering development of the next generation of space vehicles, habitats, and extra vehicular activity equipment. A new version (3DHZETRN) capable of transporting High charge (Z) and Energy (HZE) and light ions (including neutrons) under space-like boundary conditions with enhanced neutron and light ion propagation is under development. In the present report, new algorithms for light ion and neutron propagation with well-defined convergence criteria in 3D objects is developed and tested against Monte Carlo simulations to verify the solution methodology. The code will be available through the software system, OLTARIS, for shield design and validation and provides a basis for personal computer software capable of space shield analysis and optimization.

Introduction

High charge (Z) and Energy (HZE) ion transport developed parallel to the experimental studies of Schimmerling et al. [1986] at the Lawrence Berkeley Laboratory and involved solving the Boltzmann transport equation for mono-energetic ion beams in the context of the continuous slowing down approximation (CSDA) [Wilson et al. 1984]. The main purpose was to identify the source of computational limitations. This was found in part to be the inadequacy of available nuclear data through comparing computational results with the experimental ionization data for a broad beam of $^{20}\text{Ne}$ ions [Wilson et al. 1984]. Emphasis was soon overtaken by the need to establish the scope of the galactic cosmic ray (GCR) protection problem, and marching procedures were used to get first order estimates of shielding requirements [Wilson and Lamkin 1975, Lamkin 1974, Wilson and Badavi 1986, Wilson et al. 1991]. Testing the new computational marching model against atmospheric air shower data again pointed to the inadequacy of the available nuclear data [Wilson and Badavi 1986], and development of a semi-empirical nuclear fragmentation model (NUCFRG) followed, leading to improved comparisons with experimental deep penetration data [Wilson et al. 1987a, 1987b]. This new NUCFRG model likewise improved comparisons with Schimmerling’s laboratory studies [Shavers et al. 1993, Wilson et al. 1991]. The next dozen years emphasized spaceflight validation of the marching solution and nuclear fragmentation model improvements [Wilson et al. 1995, 1998; Badhwar et al. 1996; Shinn et al. 1998; Cucinotta et al. 1998; Hugger et al. 2003; Nealy et al. 2006; Wilson et al. 2005, 2006, 2007; Slaba et al. 2011a, 2013; Norman et al. 2012, 2013].


To support the need for keeping records of astronaut exposures in space operations, considerable time has been spent in validating and testing in Shuttle flight operations with an array of instruments [Badhwar et al. 1995, 1996, 2001]. As a result of the need to design and test new systems for future exploration, there has been renewed interest in flight validation using the International Space Station (ISS) as a measurements platform in low Earth orbit (LEO) [Hugger et al. 2003; Wilson et al. 2005, 2006, 2007; Nealy et al. 2006; Slaba et al. 2011a, 2013].

In the present report, the current status of transport code development is reviewed with emphasis on extending these developments into a 3D version of the HZETRN code. These advances will use available Monte Carlo codes Geant4 [Agostinelli et al. 2003, Geant4 Collaboration 2012a, 2012b], FLUKA [Fasso et al. 2005, Battistoni et al. 2007], and PHITS [Sato et al. 2013] to judge the veracity of these developments especially with regard to their 3D aspects. Upon completion, the 3DHZETRN code will be integrated into the web based OLTARIS.
Deterministic Code Development

The relevant transport equations are the time-independent linear Boltzmann equations derived on the basis of conservation principles [Wilson et al. 1991] for the flux density \( \phi_j(x, \Omega, E) \) of type \( j \) particles as

\[
\Omega \cdot \nabla \phi_j(x, \Omega, E) = \sum_k \int_{E}^{\infty} \int_{4\pi} \sigma_{jk}(E, E', \Omega, \Omega') \phi_k(x, \Omega', E') d\Omega' dE' - \sigma_j(E) \phi_j(x, \Omega, E),
\]

and solved subject to a boundary condition over the enclosure of the solution domain as shown in Fig. 1.

![Fig. 1. Geometric relations of quantities useful in solving equation (1). The symbol \( \vec{n} \) is a unit normal vector.](image)

The \( \sigma_j(E) \) and \( \sigma_{jk}(E, E', \Omega, \Omega') \) are the media macroscopic cross sections. The double differential cross section, \( \sigma_{jk}(E, E', \Omega, \Omega') \), represents all those processes by which type \( k \) particles moving in direction \( \Omega' \) with energy \( E' \) produce a type \( j \) particle in direction \( \Omega \) with energy \( E \) (including decay processes). Note that there may be several reactions that produce a particular product, and the appropriate cross sections for equation (1) are the inclusive ones. Exclusive processes are functions of the particle fields and may be included once the particle fields are known. The total cross section \( \sigma_j(E) \) for the medium may be written for each particle type as

\[
\sigma_j(E) = \sigma_j^{(at)}(E) + \sigma_j^{(el)}(E) + \sigma_j^{(nuc)}(E),
\]

where the first term \( (at) \) refers to collision with atomic electrons, the second term \( (el) \) is for Coulomb elastic scattering on the atomic nuclei, and the third term \( (nuc) \) describes nuclear scattering and reactions where the minor nuclear inelastic processes (excitation) have been ignored. The corresponding differential cross sections are similarly ordered. Many atomic collisions (~10^6) occur in a centimeter of ordinary matter, whereas ~10^3 nuclear Coulomb elastic collisions occur per centimeter, while nuclear processes are separated by a fraction to many centimeters depending on energy and particle type. Solution methods for equation (1) use perturbations based on the ordering of the physical cross sections with atomic interactions as the first physical perturbation; special methods are used for neutrons for which atomic cross sections are zero. Coulomb scattering is the second order physical perturbation process. The nuclear processes are treated in detail as the third order physical perturbation.

The atomic/molecular interaction terms of equations (1) and (2) may be written as [Wilson et al. 1991]
\[
\sum_k \int_E^\infty \int_{4\pi} \sigma_{jk}^{(at)}(E, E', \Omega, \Omega') \phi_k(x, \Omega, E') d\Omega' dE' - \sigma_j^{(at)}(E) \phi_j(x, \Omega, E) \\
= \sum_n \sigma_j^{(at)}(E + A_j^{-1} \varepsilon_n) \phi_j(x, \Omega, E + A_j^{-1} \varepsilon_n) - \sigma_j^{(at)}(E) \phi_j(x, \Omega, E) \\
= \sum_n \left\{ \sigma_j^{(at)}(E) \phi_j(x, \Omega, E) + A_j^{-1} \varepsilon_n \frac{\partial}{\partial E} \left[ \sigma_j^{(at)}(E) \phi_j(x, \Omega, E) \right] \right\} - \sigma_j^{(at)}(E) \phi_j(x, \Omega, E) + O(\varepsilon_n^2),
\]

(3)

where \(A_j\) is the mass of particle type \(j\), and the stopping power \(S_j(E)\) is given by

\[
S_j(E) = \sum_n \varepsilon_n \sigma_j^{(at)}(E).
\]

(4)

In equations (3) and (4), \(n\) denotes the electronic excitation levels, and \(\varepsilon_n\) refers to the corresponding excitation energies. Evaluation of the stopping power by equation (4) appears deceptively simple. However, all of the excited states including continuum states of the atomic/molecular system need to be evaluated. Furthermore, the projectile remains a bare ion except at low energies where the projectile ion atomic orbital states begin to resonate with the relative speed of the electrons of the media leading to electron capture and lowering of the ion charge (the stopping power is then related to RMS charge, called effective charge). These issues are further discussed in Wilson et al. [1991, 2005, 2006] and Tai et al. [1997]. The higher order terms of equation (3) are neglected in the CSDA and are further discussed elsewhere in connection to mono-energetic ion beams [Wilson et al. 2005, 2006]. The second physical perturbation, Coulomb scattering, results in angular diffusion about the ion direction leading to small angular corrections that are of little interest in the nearly omnidirectional space environment [Wilson et al. 1974, 2005, 2006] and are therefore ignored. Henceforth, the total cross sections and the double differential cross sections appearing in equation (1) are the nuclear scattering and reaction values, and the superscript \(\text{(nuc)}\) will not be written.

Equation (1) can be written in the CSDA, ignoring multiple Coulomb scattering, as

\[
B[\phi_j(x, \Omega, E)] = \sum_k \int_E^\infty \int_{4\pi} \sigma_{jk}(E, E', \Omega, \Omega') \phi_k(x, \Omega', E') d\Omega' dE',
\]

(5)

where the differential operator on the left hand side is defined as

\[
B[\phi_j(x, \Omega, E)] \equiv \Omega \cdot \nabla \phi_j(x, \Omega, E) - \frac{1}{A_j} \frac{\partial}{\partial E} \left[ S_j(E) \phi_j(x, \Omega, E) \right] + \sigma_j(E) \phi_j(x, \Omega, E).
\]

(6)

The right-hand side of equation (5) excludes the atomic/molecular processes now appearing on the left as the energy shifting operator in addition to the spatial drift and nuclear absorption terms. Neutral particles would have null atomic cross sections (except for negligible magnetic coupling) for which the stopping power term of \(B\) in equation (6) is set to zero. Application of CSDA in both laboratory and space shielding has been wide and the resulting errors discussed elsewhere [Tweed 2006a, 2006b]. Equation (5) can be rewritten as a Fredholm equation [Wilson 1977] given by

\[
\phi_j(x, \Omega, E) = \frac{S_j(E) P_j(E)}{S_j(E) P_j(E) + \sum_k \int_E^\infty \int_{4\pi} \sigma_{jk}(E', \Omega', \Omega, \Omega') \phi_k(x, \Omega', \Omega, E') d\Omega' dE'} \phi_j(x, \Omega, \Omega, E)
\]

(7)

with
\[ E_j = R_j^{-1} [\rho - d + R_j(E)], \quad (8) \]

\[ x_{\Omega} = x + [R_j(E) - R_j(E')]_{\Omega} \quad (9) \]

In equations (7) and (8), \( \Gamma(\Omega, x) \) is the point on the boundary connected to \( x \) along \( -\Omega \), \( \rho \) is the projection of \( x \) onto \( \Omega \) (Fig. 1), \( d \) is the projection of \( \Gamma(\Omega, x) \) onto \( \Omega \), \( R(E) \) is the distance a type \( j \) ion with energy \( E \) will travel before losing all of its energy to excitation/ionization of atomic electrons, and \( P_j(E) \) is the probability that a type \( j \) ion with energy \( E \) will not have a nuclear interaction in coming to rest in the media. The zero charge limit of equation (7) is discussed elsewhere [Wilson et al. 1991]. One obstacle to solving either equations (5) or (7) is the need to evaluate the integral \( d\Omega' \) at arbitrary locations within the media and development of methods to handle this limitation expediently in ways amenable to efficient computational practice.

The range-energy relationship is given (Fig. 2) by

\[ R_j(E) = A_j \int_0^E \frac{dE'}{S_j(E')}, \quad (10) \]

and the nuclear attenuation function is given (Fig. 3) by

\[ P_j(E) = \exp \left[ -A_j \int_0^E \frac{\sigma_j(E')}{S_j(E')} dE' \right], \quad (11) \]

where \( \sigma_j(E') \) includes nuclear elastic and reactive processes.

The nuclear reactive and scattering differential cross sections can be written in the following form

\[ \sigma_{jk}(E, E', \Omega, \Omega') = \frac{1}{4\pi} \sigma_{jk, iso}(E, E') + \sigma_{jk, for}(E, E', \Omega, \Omega'), \quad (12) \]

where the first term (iso) is isotropic and associated with lower energy particles produced, including target fragments, and the second term (for) is highly peaked in the forward direction (\( \Omega \approx \Omega' \)) and is associated mainly with direct quasi-elastic events and projectile fragmentation products [Wilson 1977, Wilson et al. 1988].

![Fig. 2. Range-energy relationship of ions in aluminum. Results for \( Z = 1 \) and \( Z = 2 \) are nearly overlapping.](image-url)
Fig. 3. Probability of nuclear reaction of ions in aluminum. Results for $Z = 1$ and $Z = 6$ are nearly overlapping.

Fig. 4. Isotropic (iso) and forward (for) neutron spectra produced by 500 MeV protons in aluminum.

Surprisingly, even nucleon-induced reactions follow this simple form, and the isotropic term extends to relatively high energies (see Fig. 4). For nucleon-induced reactions, the following form of the differential cross section has been used in versions of FLUKA [Ranft 1980] as follows

$$
\sigma_{jk}(E, E', \Omega, \Omega') = \nu_{jk}(E')\sigma_k(E')f_{jk}(E, E')g_R(\theta, E, A_T),
$$

where $\nu_{jk}(E')$ is average multiplicity, $f_{jk}(E, E')$ is the energy spectrum of produced fragments, and the Ranft angular factor used in early versions of FLUKA is

$$
g_R(\theta, E, A_T) = \begin{cases} 
N_R \exp(-\theta^2 / \lambda_R), & 0 \leq \theta \leq \pi / 2 \\
N_R \exp(-\pi^2 / 4\lambda_R), & \pi / 2 < \theta \leq \pi
\end{cases}
$$

with $\cos \theta = \Omega \cdot \Omega'$ and
\[ \lambda_N = \left( 0.12 + 0.00036A_T \right) / E . \]  

(15)

In equations (14) and (15), \( A_T \) is the mass of the target nucleus, and \( N_E \) is a normalization constant. The Ranft formalism was modified by adding a quasi-elastic component [Wilson et al. 1988] and was instrumental in achieving the correct neutron KERMA [Wilson et al. 1991]. The isotropic component of equation (12) is determined by integrating equation (13) over the back hemisphere (i.e. \( \cos(\theta) < 0 \)) according to

\[ \sigma_{jk,iso}(E',E') = 2 \int_{2\pi B} \sigma_{jk}(E, E', \Omega, \Omega') \, d\Omega', \]  

(16)

where \( 2\pi B \) refers to the back hemisphere. The separation in phase space given by equations (12) - (16) is exploited in obtaining corrections to the straight-ahead approximation for light particles \((Z \leq 2)\). The heavy ion \((Z > 2)\) projectile fragment cross sections are further represented by

\[ \sigma_{jk}(E, E', \Omega, \Omega') = N_t \frac{\sigma_{jk}(E')}{\sqrt{2\pi \varepsilon_{jk}^2}} \exp \left[ -2m \sqrt{E'} (1 - \Omega \cdot \Omega') / \varepsilon_{t,jk} \right] \exp \left[ -(E - E' + \lambda_{jk})^2 / 2 \varepsilon_{jk}^2 \right], \]  

(17)

where \( m \) is the nucleon rest mass, \( \lambda_{jk} \) is related to the momentum downshift, \( \varepsilon_{jk} \) is related to the longitudinal momentum width, \( \varepsilon_{t,jk} \) is related to the transverse momentum width, \( N_t \) is the transverse normalizing factor, and the fragmentation cross sections, \( \sigma_{jk}(E') \) are obtained herein from NUCFRG3. Since the transverse width is small compared to the projectile-fragment energy, the transverse function is highly peaked about the forward direction \((\Omega \cdot \Omega' \approx 1)\).

Atomic interactions limit the contributions of charged particles in the transport process. For example, the protons and alpha particles produced in aluminum below 100 A MeV contribute to the fluence only within a few centimeters of their collisional source and the heavier ions are even more restricted (see Fig. 2). This is an important factor in that the transported secondary charged particle flux tends to be small at low energies and the role of additional nuclear reactions by these low-energy ions are likewise limited (see Fig. 3), an important fact to be implemented later in the current development.

**Straight-ahead solution \((N = 1)\)**

The approach to a practical solution of equation (5) is to develop a progression of solutions from the simple to increasingly complex, allowing early implementation of high-performance computational procedures and establishing a converging sequence of approximations with established accuracy criteria and means of verification/validation. The first step leading to the lowest order solution uses the straight-ahead approximation as guided by the nucleon transport studies of Alsmiller et al. [1965] using Monte Carlo methods in which the differential cross sections were approximated as

\[ \sigma_{jk}(E, E', \Omega, \Omega') = \sigma_{jk}(E, E') \delta(\Omega - \Omega'), \]  

(18)

resulting in dose and dose equivalent per unit fluence to be within the statistical uncertainty of the Monte Carlo result obtained using the fully angle dependent cross sections in slab geometry. The relation of angular dependent cross sections to spacecraft geometry in space applications was further examined by Wilson and Kandelwal [1974] using asymptotic expansions about angular divergence parameters, demonstrating errors in the straight-ahead approximation to be on the order of the square of the ratio of distance of divergence (few centimeters) to radius of curvature of the shield (few to several m), resulting in a small error \( \sim 10^{-4} \) in most human rated space systems [Wilson and Khandelwal 1974]. Introducing approximation (18) into equation (7) reduces the Fredholm equation to a Volterra equation that can be solved using marching procedures [Wilson and Lamkin 1975, Wilson 1977, Wilson and Badavi 1986]. Considering the success of the straight-ahead approximation for nucleons encourages its use in HZE transport where the mass of heavier ions reduces even further the angle of scatter. The success of equation (18) results from the fact that lateral diffusion from a given ray is fully compensated in a flat plate (i.e. slab with infinite lateral dimensions) by lateral diffusion along adjacent rays. The verification and validation processes are described elsewhere [Wilson et al. 2005, 2006].
Initial quantification of the usefulness of the code evaluated dose and dose equivalent (as given by the ICRP 26 [ICRP 1977]) in 30 cm slab of water behind a 20 g/cm$^2$ slab shield of aluminum (and alternately of iron) for the Webber [1966] approximation of the February 23, 1956 solar particle event (SPE) spectrum given as

$$\phi_p(E) = \frac{10^9(E + 938)}{200p(E)} \exp\left\{\frac{239.1 - p(E)}{100}\right\},$$  \hspace{1cm} (19)

with $p(E) = \sqrt{E(E + 1876)}$. Comparisons were made against 3D Monte Carlo results (an early version of HETC) by Scott and Alsmiller [1967]. The results are shown in Fig. 5 for dose and Fig. 6 for dose equivalent. Statistical uncertainty of the Monte Carlo result is on the order of ten to fifteen percent, and the early version of HZETRN certainly lies within these bounds.

![Fig. 5](image1.png)

**Fig. 5.** Dose in water target protected by 20 g/cm$^2$ aluminum shield (left pane) and iron shield (right pane) exposed to Webber SPE spectrum (from Wilson et al. [1991]).

![Fig. 6](image2.png)

**Fig. 6.** Dose equivalent [ICRP 1977] in water target protected by 20 g/cm$^2$ aluminum shield (left pane) and iron shield (right pane) exposed to Webber SPE spectrum (from Wilson et al. [1991]).
The newly developed dynamic LEO environmental models, shield models, and transport algorithm are shown to have an end-to-end uncertainty estimated to be 20 percent by Badhwar et al. [1995, 1996, 2001]. Aside from its accuracy, this version of HZETRN provides a computationally efficient procedure allowing rapid dosimetric evaluations and providing a basis for development of advanced engineering design methods capable of design optimization processes [Wilson et al. 2003a, 2004b]. A sample of results is given in Table 1 and Fig. 7. A sequence of corrections to the straight-ahead approximation is now taken up for light ions and neutrons. The heavy ion nuclear cross sections, represented by equation (17), indicate that the straight-ahead approximation is suitable for heavy ions for most applications, including space, as evidenced by laboratory validation with mono-energetic beams [Shavers et al. 1993, Shinn et al. 1998, Walker et al. 2006].

![Fig. 7. Observed dose rate versus calculated dose rate [Badhwar, Chapter 21 of Wilson et al. 1997].](image)

Table 1. Comparison of straight-ahead HZETRN transport model with Shuttle flight data [Wilson et al. 2003b].

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<th>DLOC</th>
<th>TLD† (µGy/d)</th>
<th>Calc. (µGy/d)</th>
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</table>

* Deep River Neutron Monitor count rate, † GCR corrected TLD-100 data

**Bi-directional neutron transport (N = 2)**

It is clear that the straight-ahead approximation provides accurate dosimetric results and a good approximation to the particle fluence in many circumstances (see Badhwar et al. [1995]). One limitation of the straight-ahead approximation is near the front boundary on a flat plate where the calculated neutron fluence vanishes (unless they are part of the external radiation environment). This results from the straight-ahead approximation itself and can be improved by connecting to 3D corrections containing backward leakage to the front boundary [Marchuk and Lebedev 1986]. This limitation is resolved by treating the leakage that is an important process of neutron transport in space systems but of less importance to the charged particle transport that is limited by range/energy
relations. This bi-directional approximation provides the first correction to the straight-ahead approximation and provides major improvements to the neutron fluence spectra.

In the early transport studies of Wilson and Lamkin [1975] and Lamkin [1974], it was demonstrated that once neutrons are produced by a proton beam, the recoupling of neutrons back to the proton field is limited, but the neutron fields tend to build up with increasing penetration depth. This mainly occurs since many protons produced by neutron induced reactions are of lower energy than the forward components of equation (12) and are quickly removed by atomic interactions (especially these isotropically produced protons). The probability of even a 100 MeV proton to undergo a nuclear reaction is only several percent (Fig. 3). The low energy neutrons on the other hand have no effective atomic interactions and propagate until a nuclear interaction occurs or they escape from the media through a boundary. Yet, these low energy protons and other light ions produced in tissue through neutron collisions contribute significantly to biological injury and must be included. This was the basis of studies starting with Clowdsley et al. [2000, 2001] and Heinbockel et al. [2003], with recent improvements provided by Slaba et al. [2010b]. Herein, we will concentrate on its extension to 3D transport methodologies. Future work will be a direct evaluation of equation (7) quantifying the propagated error of the 3D neutron code.

A bi-directional neutron transport code was first investigated by Clowdsley et al. [2000, 2001] using multigroup methods and recoupling to the light ions using an analytic solution as a quadrature over the light ion collision source term. These types of approximations were further studied and improved by Slaba et al. [2010b] and later compared to Monte Carlo simulations in slab geometry [Heinbockel et al. 2011a, 2011b] and on the lunar surface [Slaba et al. 2011b]. The model has been implemented into OLTARIS and is the basis of the ray-by-ray transport methodology [Slaba et al. 2011b, Singleterry et al. 2011]. In that transport model, the charged particle components were first solved in the straight-ahead approximation with the neutron coupling represented only by the forward component of the differential production cross section. The isotropically produced neutrons are then solved using a coupled bi-direction (forward/backward) transport to complete the neutron solution. These isotropically produced neutrons further collide with the media, giving rise to an isotropic source of ions that are transported assuming only atomic interactions as an approximation to the complete solution. Neglected in this solution is the minor coupling between low energy charged particles and any produced fragments. This assumption is generally accurate since the range of the low energy isotropic ions is much smaller than their nuclear mean free path lengths (see Figs. 2 and 3).

The governing equation in this approach is first given by equation (5). The solution methodology requires particle fluxes to be separated into forward \( \phi_{j,\text{for}} \) and isotropic components \( \phi_{j,\text{iso}} \) according to

\[
\phi_j(x, \Omega, E) = \phi_{j,\text{for}}(x, \Omega, E) + \phi_{j,\text{iso}}(x, \Omega, E).
\]

The differential cross section for neutron fragments \( j = n \) is also separated into forward and isotropic components according to equations (12) and (16). The differential cross section for charged fragments is treated in the straight-ahead approximation by equation (18). Upon substitution of the separated differential cross sections and particle fluxes into equation (5), the forward component of the charged particle \( j \neq n \) fluxes is obtained by solving the straight-ahead equation

\[
\mathbf{B}[\phi_{j,\text{for}}(x, \Omega, E)] = \sum_k \int_{E}^{\infty} \sigma_{jk}(E, E') \phi_{k,\text{for}}(x, \Omega, E') dE',
\]

and the forward neutron component is obtained by solving

\[
\mathbf{B}[\phi_{n,\text{for}}(x, \Omega, E)] = \sum_k \int_{E}^{\infty} \sigma_{nk,\text{for}}(E, E') \phi_{k,\text{for}}(x, \Omega, E') dE',
\]

for which the stopping power term in the operator \( \mathbf{B} \) is zero for neutrons. Note that the full cross section of equation (18) appears in the ion transport of equation (21), but only the forward component of the cross sections appears in the transport of neutrons in equation (22). Equations (21) and (22) are solved simultaneously using methods previously developed [Wilson et al. 2006, Slaba et al. 2010a].

The forward fluxes give rise to an isotropic source of low energy neutrons which are treated by solving the equation
\[ B[\phi_{n,iso}(\mathbf{x}, \Omega, E)] = \int_E^\infty \int_{4\pi} \sigma_{nn}(E, E', \Omega, \Omega') \phi_{n,iso}(\mathbf{x}, \Omega', E') d\Omega' dE' + \sum_k \int_E^\infty \int_{4\pi} \sigma_{nk,iso}(E, E', \Omega, \Omega') \phi_{k,for}(\mathbf{x}, \Omega', E') d\Omega' dE'. \]  

(23)

Whereas equations (21) and (22) take on value only if an external source is present at the boundary, equation (23) has no external sources, and the resulting field develops from the internal isotropic neutron source induced by the forward solution. Development of an algorithm for solving equation (23) is expedited by separating the isotropic neutron fluence into forward and backward propagating components according to

\[ \phi_{n,iso}(\mathbf{x}, \Omega, E) = \phi_{n,iso(f)}(\mathbf{x}, \Omega, E) + \phi_{n,iso(b)}(\mathbf{x}, -\Omega, E), \]  

(24)

and approximating the double differential cross sections as

\[ \sigma_{nn}(E, E', \Omega, \Omega') = \sigma_{nn,f}(E, E') \delta(\Omega - \Omega') + \sigma_{nn,b}(E, E') \delta(\Omega + \Omega'), \]  

(25)

\[ \sigma_{nn,f}(E, E') = \int_{2\pi F} \sigma_{nn}(E, E', \Omega, \Omega') d\Omega' = \sigma_{nn,for}(E, E') + \frac{2\pi}{4\pi} \sigma_{nn,iso}(E, E') / 4\pi, \]  

(26)

\[ \sigma_{nn,b}(E, E') = \int_{2\pi B} \sigma_{nn}(E, E', \Omega, \Omega') d\Omega' = 2\pi \sigma_{nn,iso}(E, E') / 4\pi, \]  

(27)

\[ \sigma_{nk,iso(f)}(E, E') = \int_{2\pi F} \sigma_{nk,iso}(E, E', \Omega, \Omega') d\Omega' = 2\pi \sigma_{nk,iso}(E, E') / 4\pi, \]  

(28)

\[ \sigma_{nk,iso(b)}(E, E') = \int_{2\pi B} \sigma_{nk,iso}(E, E', \Omega, \Omega') d\Omega' = 2\pi \sigma_{nk,iso}(E, E') / 4\pi. \]  

(29)

In equations (26) - (29), \(2\pi F\) and \(2\pi B\) represent integration over the forward and backward hemispheres with respect to \(\Omega'\), respectively. The subscript (f) and (b) in equation (24) refers to the forward hemisphere and backward hemisphere components of the isotropic neutron flux, respectively. Using equations (25) - (29) in equation (23) results in

\[ [\Omega \cdot \nabla + \sigma_n(E) \phi_{n,iso(f)}(\mathbf{x}, \Omega, E)] = \int_E^\infty \sigma_{nn,f}(E, E') \phi_{n,iso(f)}(\mathbf{x}, \Omega, E') dE' \]

\[ + \int_E^\infty \sigma_{nn,b}(E, E') \phi_{n,iso(b)}(\mathbf{x}, -\Omega, E') dE' \]

\[ + \sum_k \int_E^\infty \sigma_{nk,iso(f)}(E, E') \phi_{k,for}(\mathbf{x}, \Omega, E') dE' \]

\[ + \sum_k \int_E^\infty \sigma_{nk,iso(b)}(E, E') \phi_{k,for}(\mathbf{x}, -\Omega, E') dE'. \]  

(30)

for the forward component of the isotropic neutron flux. The backward moving isotropic neutron field satisfies a similar equation

\[ [-\Omega \cdot \nabla + \sigma_n(E) \phi_{n,iso(b)}(\mathbf{x}, \Omega, E)] = \int_E^\infty \sigma_{nn,f}(E, E') \phi_{n,iso(b)}(\mathbf{x}, \Omega, E') dE' \]

\[ + \int_E^\infty \sigma_{nn,b}(E, E') \phi_{n,iso(f)}(\mathbf{x}, -\Omega, E') dE' \]

\[ + \sum_k \int_E^\infty \sigma_{nk,iso(f)}(E, E') \phi_{k,for}(\mathbf{x}, -\Omega, E') dE' \]

\[ + \sum_k \int_E^\infty \sigma_{nk,iso(b)}(E, E') \phi_{k,for}(\mathbf{x}, \Omega, E') dE'. \]  

(31)
Note that $\phi_{k,dir}(x, -\Omega, E') = 0$ in equations (30) and (31). Equations (30) and (31) represent a coupled set of differential equations that are solved efficiently using a Neumann series approach as shown by Slaba et al. [2010b]. This series is expected to converge quickly as $\sigma_{in,f}(E, E') \gg \sigma_{in,b}(E, E')$.

The isotropically produced neutrons initiate isotropic light ion production in collisions with the media resulting in

$$B[\phi_{j,iso}(x, \Omega, E)] = \int_{E}^{\infty} \sigma_{jn}(E, E') \phi_{n,iso}(x, \Omega, E') dE',$$

(32)

for which a semi-analytic solution can be efficiently implemented [Slaba et al. 2010b]. Equations (30) and (31) are solved simultaneously for no external fluence incident on the media directed along $\Omega$ (see Fig. 1). Note that the implicit assumption of equations (30) and (31) is that lateral diffusion from the ray $\Omega$ is fully compensated by diffusion from adjacent rays. This is strictly true for a flat plate slab with infinite lateral dimensions as will now be demonstrated and represents the use of periodic boundary conditions as an importance sampling method common to Monte Carlo simulation practice.

These two methods (straight-ahead ($N = 1$) and bi-directional ($N = 2$)) are now compared to values simulated by the Monte Carlo codes Geant4, FLUKA, and PHITS for the Webber SPE spectrum normally incident on a 40 g/cm$^2$ slab of aluminum with infinite lateral dimensions. All Monte Carlo fluence results presented herein are plotted as mean values with estimated statistical uncertainty shown as vertical error bars placed at the energy bin midpoint. The points are connected with lines to improve plot clarity. If not visible, vertical error bars are smaller than the symbols on the plot. Details of the Monte Carlo simulation setups are given in Appendix A. Results are shown in Fig. 8. The straight-ahead solution gives good agreement for charged components at most depths. Compare the left and right panes of Fig. 8. The neutron fluence reaches a broad maximum across 20 to 35 g/cm$^2$ wherein the neutron fluence is nearly isotropic at low energies and the straight-ahead ($N = 1$) fluence approaches the bi-directional ($N = 2$) fluence from below before the maximum and over estimates beyond the maximum. On the back face, leakage is an important process leaving the straight-ahead fluence as an over estimate at low neutron energies where the neutron fluence is reduced by about half of its interior value due to leakage through the back surface (compare left and right panes of Fig. 8). The resonance structure seen in the Monte Carlo neutron fluences below 1 MeV is a result of elastic cross section databases that have not been included in 3DHZETRN.

![Fig. 8. Comparison of fluence spectra at 35 g/cm$^2$ (left pane) and 40 g/cm$^2$ (right pane) in a 40 g/cm$^2$ aluminum slab (infinite in lateral dimensions) exposed to Webber SPE spectrum.](image)
It is clear that the bi-directional solution for the semi-infinite slab where lateral leakage is negligible is a reasonable approximation. The success of the straight-ahead approximation in validation in human rated systems (Shuttle, ISS) clearly meets the requirements for a useful solution of the transport problem. Attention is now given to transport in finite objects in which lateral leakage plays an important role.

3D Marching Procedures

A sequence of approximations is now sought that will span the formalism from the simple bi-directional approximation to more complex propagation algorithms. With the demonstrated success of the bi-directional approximation, the solution is divided into forward and isotropic components, as before. The governing equation in the CSDA ignoring Coulomb scattering is given by equation (5). The particle flux is separated into forward and isotropic components, as in equation (20). Similarly, the double differential cross sections for neutron production are separated into forward and isotropic components given by equations (12) and (16). The charged particle production cross sections are assumed to take the form of equation (18). Upon substitution of the separated fluxes and production cross sections into equation (5), the forward components are obtained by solving

$$B[\phi_{j,\text{for}}(x,\Omega, E)] = \sum_k \int_{E'}^\infty \sigma_{jk}(E, E')\phi_{k,\text{for}}(x, \Omega, E') dE'$$  \hspace{1cm} (33)

for charged particles ($j \neq n$), and for neutrons ($j = n$),

$$[\Omega \cdot \nabla + \sigma_n(E)]\phi_{n,\text{for}}(x, \Omega, E) = \sum_k \int_{E'}^\infty \sigma_{nk,\text{for}}(x, \Omega, E')\phi_{k,\text{for}}(x, \Omega, E') dE'$$  \hspace{1cm} (34)

The space environment (external source) is nearly uniform in position and varies slowly in angle (if not isotropic), and for the forward flux, each directional component of the boundary condition may be propagated according to equations (33) and (34). Note that $\Omega$ enters equations (33) and (34) as a parameter and can be evaluated according to the boundary condition we have set as the incident flux from direction $\Omega_0$ (see Fig 9). Once the forward flux is evaluated for direction $\Omega_0$ over the domain of $x$ and $E$, one must yet solve for the induced isotropic neutron flux within the shield configuration. Coupling to the isotropically produced neutron field is taken as

$$[\Omega \cdot \nabla + \sigma_n(E)]\phi_{n,\text{iso}}(x, \Omega, E) = \int_E^\infty \int_{4\pi} \sigma_{mn}(E, E', \Omega, \Omega')\phi_{n,\text{iso}}(x, \Omega', E') d\Omega' dE' + \xi_{n,\text{iso}}(x, \Omega, \Omega_0, E), \hspace{1cm} (35)$$

where the isotropic neutron source is given by

$$\xi_{n,\text{iso}}(x, \Omega, \Omega_0, E) = \sum_k \int_{E'}^\infty \sigma_{nk,\text{iso}}(E, E', \Omega, \Omega_0)\phi_{k,\text{for}}(x, \Omega_0, E') dE'$$  \hspace{1cm} (36)

Fig. 9. Geometry of 3D marching procedure.
Note that the isotropic neutron source is generated by collisions between the forward flux and target media, for which the forward flux is a function of only the penetration depth \(z\) along the direction \(\Omega_0\). This is denoted by writing the isotropic neutron source as

\[
\chi_{n,\text{iso}}[z(x), \Omega, \Omega_0; E] = \xi_{n,\text{iso}}(x, \Omega, \Omega_0; E),
\]

where \(z(x)\) is the penetration depth along \(\Omega_0\) to the point \(x\). Equation (35) is now rewritten as

\[
[\Omega \cdot \nabla + \sigma_n(E)]\phi_{n,\text{iso}}(x, \Omega, E) = \int_E^{\infty} \int_{\Omega} \sigma_{nn}(E, E', \Omega, \Omega')\phi_{n,\text{iso}}(x, \Omega', E') d\Omega' dE' + \chi_{n,\text{iso}}[z(x), \Omega, \Omega_0; E]
\]

Equation (38) also provides a basis for introducing anisotropic source terms as well. Note that while the forward propagating solution has its source fixed by the boundary condition, the isotropic solution (having no inbound fluence) is driven by internal sources generated by the collisions of the inbound fluence (as modified by the forward propagator in penetrating to \(x\) along direction \(\Omega_0\)).

To solve equation (38), the neutron fluence \(\phi_{n,\text{iso}}(x, \Omega', E')\) must be known at every \(x\) from all directions \(\Omega'\) approaching \(x\) from adjacent locations as was the case of equation (23). Although the use of approximation (25), based on the assumption that losses along the ray \(\Omega\) are compensated by diffusion from adjacent rays, is a demonstrated useful assumption in a semi-infinite slab, it is an approximate overestimate in a sphere. Yet, using equation (25) in evaluation of equation (38) seems a reasonable approximation in most human rated vehicles leading to an extension of transport along an arbitrary ray \(\Omega\) as a solution to the bi-directional equations. The forward component of the isotropic neutron field satisfies

\[
[\Omega \cdot \nabla + \sigma_n(E)]\phi_{n,\text{iso}(f)}(x, \Omega, E) = \int_E^{\infty} \sigma_{nn,f}(E, E')\phi_{n,\text{iso}(f)}(x, \Omega, E') dE' + \int_E^{\infty} \sigma_{nn,b}(E, E')\phi_{n,\text{iso}(b)}(x, -\Omega, E') dE' + \chi_{n,\text{iso}}[z(x), \Omega, \Omega_0; E],
\]

and the backward moving field satisfies

\[
[-\Omega \cdot \nabla + \sigma_n(E)]\phi_{n,\text{iso}(b)}(x, \Omega, E) = \int_E^{\infty} \sigma_{nn,b}(E, E')\phi_{n,\text{iso}(b)}(x, -\Omega, E') dE' + \chi_{n,\text{iso}}[z(x), -\Omega, \Omega_0; E].
\]

It is clear that \(\Omega\) enters equations (39) and (40) as a parameter and can be solved along an arbitrary number of rays used to construct a numerical representation of \(\phi_{n,\text{iso}}(x, \Omega, E)\) whose \(\Omega\) dependence comes from distance to the boundary \(t(\Omega)\) and the penetration depth \(z\) along \(\Omega_0\). The isotropic neutron source of equation (38) effectively decouples the incident direction \(\Omega_0\) from the neutron direction of propagation \(\Omega\).

These isotropically produced neutrons produce isotropic light ions in collisions with the media resulting in a light ion fluence given by

\[
B[\phi_{j,\text{iso}}(x, \Omega, E)] = \int_E^{\infty} \sigma_{j}(E, E')\phi_{n,\text{iso}}(x, \Omega, E') dE'.
\]

Equations (39) and (40) are solved simultaneously for no external fluence incident on the media directed along \(\Omega\) (see Fig. 9). Nonetheless, efficient numerical methods already developed for the bi-directional model may be utilized. Note also that the implicit assumption of equations (39) and (40) is that lateral diffusion from the ray \(\Omega\) is fully compensated by diffusion from adjacent rays. This has been demonstrated for slab geometry with results in
Fig. 8. The first order lateral leakage in this case is found by solving equations (39) and (40) along each \( \Omega_i \) over the ray limits \( \{-t(-\Omega_i), t(\Omega_i)\} \) for a given set of \( N \) directions \( \Omega_i \). The next sections address convergence in the isotropic neutron and ion fields as a function of the number of directions \( N \) chosen to represent the fields. Numerical solutions for equations (39) - (41) are the same as in the bi-directional solution and described by Slaba et al. [2010b]. The only difference here is that the isotropic source term in each case is computed as a function of penetration depth with respect to the incident source direction, \( \Omega_0 \).

The present transport formalism is related, but quite different from \( S_N \) transport theory [Marchuk and Lebedev 1986]. The main difference between the two formalisms is that solutions along a set of directions \( \Omega_i \) are fully coupled to all other discrete directions in the \( S_N \) treatment. While in the present formalism, coupling is only included through the isotropic source term and for transport along anti-parallel rays (i.e., \( \Omega_i \) and \(-\Omega_i\)). It is demonstrated below that the present formalism compares well to Monte Carlo simulations where such coupling is implicitly included.

**A Simple Geometry and Source**

Simple geometry and source orientation are chosen to allow efficient Monte Carlo simulation for comparison with the calculation of quantities with the above formalism. In this way, the solution methodology is verified and experience is gained in the formalism in preparation for connecting to the general geometric descriptions generated by the engineering design process for which Monte Carlo simulation is impractical. The geometry herein is taken as a simple sphere of aluminum of 40 g/cm\(^2\) diameter for comparison with the appropriate solutions of the current formalism. Target points (detectors) are placed at depths along the \( z \)-axis (\( x = 0, y = 0 \)). The external radiation environment (external source boundary condition) is assumed to be anti-parallel with the \( z \)-axis, uniform in the \( x-y \) plane, positioned above the sphere, and directed down onto the top of the sphere as shown in Fig. 10.

![Fig. 10. Spherical geometry and external source orientation for benchmark comparisons (not all detectors shown).](image)

The object geometry is specified at a point \( x \) where field values are to be evaluated, about which the thickness of material \( t \) is known over an array of directions \( \Omega_i \) (i.e., \( t_i = t(\Omega_i) \)). Note that each \( \Omega_i \) for \( i \) odd is backed by \( \Omega_{i+1} = -\Omega_i \). The remainder of the solution methodology lies in selection of the arbitrary directions of \( \Omega_i \) of propagation within the target material and evaluation of the penetration depth \( z \) for any transport depth \( x_T \) along each \( \Omega_i \) for the given \( \Omega_0 \) at the boundary. The \( N \) directions are taken as the central ray of an equal division of the unit sphere such that \( \Delta\Omega = 4\pi / N \). In this, the interest is in evaluating the omni-directional field function at an arbitrary location \( x \) about which the surrounding material is known as a thickness distribution of the various materials (in the present only homogenous aluminum). The evaluation will be made for a single direction of incidence \( \Omega_0 \) to highlight 3D transport effects, but this could include an array of incident directions for which the solutions would be integrated over all \( \Omega_0 \) values.
Fig. 11. Isotropic neutron source ($E > 1$ MeV) distribution induced by the Webber SPE spectrum in the spherical benchmark geometry.

In that low-energy neutron transport is dominated by diffusive processes, it is anticipated that the solution will rapidly converge for low values of $N$ (i.e. less than 20). The source of neutrons induced within the sphere by the Webber SPE is shown in Fig. 11. The resulting neutron fluence depends not only on the distance $t(\Omega)$ to the boundary, but the intensity of the source along the ray in reaching the boundary. Clearly for SPEs, the most intense regions must be adequately represented in the domain of the solution. The next three sub-sections describe how the penetration depth, $z$, is determined for $N = 1, 2$ and the general case ($N > 2$).

**N = 1, Straight-ahead approximation**

For each $\Omega_0$, only one value of $\Omega_i$ is allowed and is taken to be that of the incident direction $\Omega_0$. The transport distance $x$ along $\Omega_1$ and penetration depth $z$ are the same, reducing complication in the formulation for $N = 1$. In this case, leakage from the media is only through the distal face. The structure of this code is quite different than the original HZETRN code, although computational results are identical, and the original HZETRN code provides a verification test-case of the new 3DHZETRN code. For $N = 1$, the comparisons in the prior sections describing the verification and validation of HZETRN is fully applicable. If one restricts interest to $N = 1$, then the HZETRN code is preferred in that computational efficiency is higher but the added overhead of 3DHZETRN for $N = 1$ is minimal. In the following sections, comparisons between 3DHZETRN and Monte Carlo simulations will also include results from the HZETRN code ($N = 1$) to highlight shortcomings of the straight-ahead approximation and improvements of the 3D corrections presented herein.

**N = 2, Bi-directional approximation**

The first step towards a full 3D code is the treatment of backward propagating particles. This was first accomplished by Clowdsley et al. [2000, 2001] and later improved by Slaba et al. [2010b] with demonstrated improved results [Slaba et al. 2010b, 2011b; Heinbockel et al. 2011a, 2011b]. In this case, propagation of the isotropic neutrons is represented in two directions by $\Omega_1 = \Omega_0$ and $\Omega_2 = -\Omega_0$. Therefore, the penetration depth $z$ is equal to $x$ along $\Omega_1$ and equal to $t(\Omega_2) + t(\Omega_1) - x$ along $\Omega_2$. Note that lateral leakage is not represented in this approximation although leakage at the forward and backward boundaries is so represented. The neutron transport in semi-infinite slab geometry has extensive Monte Carlo results for verification for which this two-stream approximation, ignoring lateral leakage, is most appropriate as demonstrated in Fig. 8. This approximation is appropriate for slab geometry and works well in applications in which the shield geometry is a collection of thin plates. The next step for treatment of lateral leakage is now discussed.
For general $N$, the transport formalism requires the penetration depth, $z_p$, to be known as a function of transport depth, $x_T$, along the propagation direction $\Omega$ for each $\Omega$. In order to simplify algebraic manipulation and expressions, the center of the sphere is assumed to be positioned at the origin with the positive $z$-axis anti-parallel to the source direction, as shown in Fig. 10. Due to the simplified geometry and source orientation, an analytic expression for the penetration depth may be determined. The point $x_0$ may be written as

$$x_0 = x - \Omega(t - x_T),$$

(42)

where $t$ is the known thickness along direction $-\Omega$ from $x$ to the boundary. The point $x_T$ may then be written in terms of $x_0$ as

$$x_T = x_0 - \Omega z_p.$$  

(43)

The expression for $z_p$ is obtained by writing equation (43) in component form and substituting into the equation of a sphere with radius $R = 20 \text{ g/cm}^2$. This yields

$$z_p = x_0 \cdot \Omega_0 + \sqrt{R^2 - |x_0|^2 + (x_0 \cdot \Omega_0)^2}.$$  

(44)

For preliminary testing and benchmark comparisons against Monte Carlo codes, the solutions for $N = 6, 10, 14, 18,$ and $22$ are considered separately, in addition to the simple $N = 1$ and $N = 2$ cases discussed above. The distributions for all $N$ are shown in Fig 12. Note that the $N = 6, 14,$ and $22$ cases include rays perpendicular to the $z$-axis (i.e. tangent to the sphere surface at the top and bottom detector locations). On the surface of the sphere, these rays correspond to zero shield thickness. This is corrected in the $N = 10$ and $N = 18$ distributions. The directional elements for each $N$ are given in the Appendix B, along with a brief description of how the elements were determined. For each $N$, the omni-directional flux is obtained by integrating the fluxes along each ray direction after the transport procedure.

Fig. 12. Directional components for $N = 1, 2, 6, 10, 14, 18,$ and $22$.

Preliminary testing

A first quantity of interest is the average path length at each detector location where the solution is evaluated to the sphere boundary. For a given $N$, the average path length is computed over the material thicknesses traversed from the target point to the boundary along all rays. This quantity is expected to converge rapidly at most locations with increasing $N$. As shown in Table 2, convergence is reasonably rapid for all locations except on the sphere surface. It is clear that the results are reasonably converged at $N = 18$, especially in comparison to $N = 10$, as will be further demonstrated for transport quantities.
Table 2. Average thickness traversed at different detector locations along z-axis of sphere. The left-most column is the distance from the top of the sphere to the detector.

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<td>19.6</td>
<td>19.6</td>
<td>19.6</td>
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</tr>
<tr>
<td>20</td>
<td>20.0</td>
<td>20.0</td>
<td>20.0</td>
<td>20.0</td>
<td>20.0</td>
<td>20.0</td>
<td>20.0</td>
</tr>
<tr>
<td>25</td>
<td>25.0</td>
<td>20.0</td>
<td>19.6</td>
<td>19.6</td>
<td>19.6</td>
<td>19.6</td>
<td>19.6</td>
</tr>
<tr>
<td>30</td>
<td>30.0</td>
<td>20.0</td>
<td>18.2</td>
<td>18.2</td>
<td>18.2</td>
<td>18.2</td>
<td>18.2</td>
</tr>
<tr>
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<td>35.0</td>
<td>20.0</td>
<td>15.5</td>
<td>15.6</td>
<td>15.6</td>
<td>15.7</td>
<td>15.7</td>
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<tr>
<td>40</td>
<td>40.0</td>
<td>20.0</td>
<td>6.7</td>
<td>10.4</td>
<td>9.4</td>
<td>9.4</td>
<td>9.4</td>
</tr>
</tbody>
</table>

The sequence of solutions for N = 1 to 22 is used as a test on the convergence of the solution method. In addition, a Monte Carlo benchmark is defined for each of the indicated methods. Benchmarks for N = 1 and N = 2 are given in slab geometry in Fig. 8 to be later compared with N = 18 benchmarks in spherical geometry.

Table 3. Dose (cGy/event) at different detector locations along z-axis of sphere for Webber SPE. The left-most column is the distance from the top of the sphere to the detector.

<table>
<thead>
<tr>
<th>d (g/cm²)</th>
<th>N = 1</th>
<th>N = 2</th>
<th>N = 6</th>
<th>N = 10</th>
<th>N = 14</th>
<th>N = 18</th>
<th>N = 22</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>43703.70</td>
<td>43703.77</td>
<td>43703.73</td>
<td>43703.84</td>
<td>43703.79</td>
<td>43703.82</td>
<td>43703.80</td>
</tr>
<tr>
<td>0.3</td>
<td>1216.22</td>
<td>1216.28</td>
<td>1216.33</td>
<td>1216.35</td>
<td>1216.34</td>
<td>1216.34</td>
<td>1216.34</td>
</tr>
<tr>
<td>5</td>
<td>69.95</td>
<td>69.98</td>
<td>70.03</td>
<td>70.02</td>
<td>70.02</td>
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<tr>
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<td>24.81</td>
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<tr>
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<td>4.34</td>
<td>4.34</td>
<td>4.34</td>
<td>4.34</td>
<td>4.34</td>
</tr>
<tr>
<td>30</td>
<td>2.90</td>
<td>2.88</td>
<td>2.85</td>
<td>2.85</td>
<td>2.85</td>
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<tr>
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<td>1.41</td>
<td>1.38</td>
<td>1.38</td>
<td>1.38</td>
<td>1.38</td>
<td>1.38</td>
</tr>
</tbody>
</table>
Table 4. Dose equivalent (cSv/event) at different detector locations along z-axis of sphere for Webber SPE. The leftmost column is the distance from the top of the sphere to the detector.

<table>
<thead>
<tr>
<th>d (g/cm²)</th>
<th>(N = 1)</th>
<th>(N = 2)</th>
<th>(N = 6)</th>
<th>(N = 10)</th>
<th>(N = 14)</th>
<th>(N = 18)</th>
<th>(N = 22)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>737745.84</td>
<td>737746.95</td>
<td>737746.21</td>
<td>737748.11</td>
<td>737747.20</td>
<td>737747.89</td>
<td>737747.45</td>
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<tr>
<td>0.3</td>
<td>2871.17</td>
<td>2872.15</td>
<td>2873.07</td>
<td>2873.31</td>
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<td>2873.17</td>
<td>2873.12</td>
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<tr>
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<td>110.33</td>
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<td>111.05</td>
<td>111.02</td>
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<td>19.01</td>
<td>19.04</td>
<td>18.97</td>
<td>19.00</td>
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<td>2.26</td>
<td>2.21</td>
<td>2.19</td>
<td>2.19</td>
<td>2.21</td>
</tr>
</tbody>
</table>

Fig. 13. Particle spectra at detector locations 35 g/cm² (left pane) and 40 g/cm² from top of aluminum sphere (diameter 40 g/cm²) exposed to Webber SPE spectrum for different \(N\) in 3D transport solution.

It is clear in the prior discussion that the main difference between \(N = 2\) and \(N > 2\) is the degree of lateral diffusion and the resulting leakage near boundaries. The ratio

\[
r_N(E) = 1 - \frac{\phi_n(E, N > 2)}{\phi_n(E, N = 2)}
\]

is directly related to the representation of lateral losses for neutrons. The quantity \(r_N(E)\) is fixed by the shape of the object, the scattering angular distribution, and the fidelity of the transport solution represented by \(N\). The quantity \(r_N(E)\) is shown in Fig. 14 for \(N = 6, 10, 14, 18, \) and 22. The value of zero at the highest energy is a result of the straight-ahead approximation for the forward component for which there is no lateral leakage. The largest relative difference between the \(N = 10\) and \(N = 22\) ratios is less than 3% at both depths, and the \(N = 18\) solution appears converged based on the results in Tables 2-4 and Figs. 13 and 14.
19

Fig. 14. The neutron leakage ratio at detector locations 35 g/cm² (left pane) and 40 g/cm² (right pane) from top of aluminum sphere (diameter 40 g/cm²) exposed to Webber SPE spectrum for different $N$ in 3D transport solution. The largest relative difference between the $N = 10$ and $N = 22$ ratios is less than 3% at both depths.

Monte Carlo Benchmarks

In 2005, a systematic benchmark exercise was administered as a means of verifying various computational procedures resulting from the first phase of code development activity (1996-2004). Code verification uses established benchmarks as consisting of standard environments: the Webber SPE spectrum given in equation (19) and the 1977 solar minimum GCR environment.

In earlier SPE benchmarking, the configuration was for a 30 cm semi-infinite slab of water shielded by 20 g/cm² of either aluminum or iron utilizing only dose and dose equivalent (ICRP 26 quality factor [ICRP 1977]) as given in Figs. 5 and 6. For the moment, focus is mainly given to the results for the neutron spectra at the aluminum/water interface as shown in Fig. 15. It is seen from the figure that those codes with roots in the HETC code (HETC-HEDS and MCNPX) form a cluster of results including HZETRN at low energies ($E < 40$ MeV). In distinction, Geant4 and FLUKA form a cluster below HZETRN. All codes show improved agreement above 100 MeV.

Fig. 15. Secondary neutron fluence at the interface between a 20 g/cm² aluminum shield and 30 g/cm² water target exposed to the Webber SPE (from Slaba et al. [2010b]).
It was generally concluded that HZETRN agrees with the Monte Carlo codes as well as the various Monte Carlo codes agreed among themselves. There are currently no compelling reasons to attempt to use Monte Carlo methods in design and operational applications. However, it is further demonstrated below, the Monte Carlo codes provide an indispensable guide to development of the 3DHZETRN code (that does provide a useful basis for space radiation protection). Focus is now shifted towards benchmarking 3DHZETRN, in which the added complications of finite geometry are addressed. Details associated with Monte Carlo simulations presented in Figs. 16 - 21 are given in Appendix A. One needs to be mindful of the intercode comparisons of Fig. 15 when viewing succeeding benchmarks.

**Webber SPE Benchmark**

The benchmark SPE spectrum is the Webber [1966] approximation of the February 23, 1956 SPE (not an accurate representation but a historically useful test spectrum, see Foelsche et al. [1974]) as given in equation (19). The induced neutron and proton fluence spectra were evaluated in a 40 g/cm² diameter aluminum sphere by Monte Carlo codes (Geant4, FLUKA, and PHITS) and 3DHZETRN with the results at 35 and 40 g/cm² depths shown in Fig 16. The proton fluences among the three codes are in reasonable agreement. The larger discrepancy lies with the neutron spectra. One interesting note is that 3DHZETRN agrees with Geant4 below 10 MeV and agrees with FLUKA above 10 MeV. FLUKA and 3DHZETRN exhibit a similar inflection in the neutron spectrum near 10 to 60 MeV that may be related to the use of an early FLUKA approximation to the secondary neutron spectrum used in HZETRN and carried over into 3DHZETRN. Some of the differences in proton spectra on the back surface may be associated with the simplified assumptions of the straight-ahead approximation in low energy isotropic light ion transport. A principle difference in the three codes is the computational efficiency, as discussed in more detail later in this section. It is also worth noting that the 3DHZETRN (N = 18) solution is a substantial improvement over the HZETRN (N = 1), indicating that lateral neutron leakage is more accurately described by the updated transport model.

![Fig. 16. Particle spectra at detector locations 35 g/cm² (left pane) and 40 g/cm² from top of aluminum sphere (diameter 40 g/cm²) exposed to Webber SPE spectrum.](image-url)

The definition of the leakage factor given in equation (45) is specific to the 3DHZETRN formalism presented herein; however, an analogous quantity may also be computed for Monte Carlo results by comparing the ratio of the fluences in a slab (Fig. 8) to those obtained in a sphere (Fig. 16) provided the slab thickness and sphere diameters are identical. The leakage factor evaluated by three Monte Carlo codes (PHITS, FLUKA, and Geant4) are shown in comparison with 3DHZETRN (N = 18) at 35 g/cm² and 40 g/cm² in Fig. 17. The leakage is related to the angular dependence of the cross section. For example, the near zero leakage in the 3DHZETRN results above 200 MeV is a result of the straight-ahead approximation. It is clear that the Monte Carlo results show significant
differences in leakage as determined mainly by differences in angular scattering cross sections. The 3DHZETRN results appear at a slight disadvantage with respect to Monte Carlo evaluations. It should also be clear that the improvement of the 3DHZETRN cross sections could result in more trustworthy results. This would not only improve leakage estimates but also give better representation of the resonance structure seen in the neutron fluences at lower energies.

![Graph showing neutron leakage ratio factor](image)

Fig. 17. The neutron leakage ratio factor at detector locations 35 g/cm² (left pane) and 40 g/cm² (right pane) from top of aluminum sphere (diameter 40 g/cm²) exposed to the Webber SPE spectrum.

The time required by the three codes to run these benchmarks is given in CPU seconds used and listed in the first column of Table 5. The 5 CPU seconds of 3DHZETRN allows for the development of affordable means of design optimization software on a commercial laptop while Geant4, FLUKA, and PHITS software requires a multiprocessing cluster environment to do a single design evaluation and would still be impractical for design optimization. The main limitation of the Monte Carlo methods is the number of particle histories required for reasonable convergence. It should be noted that source biasing, physics biasing, or other optimization strategies could significantly reduce the Monte Carlo runtimes shown in Table 5. However, the fully optimized run times would still be orders of magnitude larger than the 3DHZETRN run times and still require a multi-processor cluster computing environment.

### Table 5. Total CPU seconds required for benchmark calculations. Total number of histories ran in Monte Carlo simulations are shown in parentheses.

<table>
<thead>
<tr>
<th>Model</th>
<th>Webber SPE</th>
<th>GCR hydrogen</th>
<th>GCR helium</th>
<th>GCR carbon</th>
<th>GCR iron</th>
</tr>
</thead>
<tbody>
<tr>
<td>3DHZETRN (N = 1)</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>3DHZETRN (N = 18)</td>
<td>5</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Geant4</td>
<td>9×10⁷ (2×10¹¹)</td>
<td>1×10¹⁸ (2×10¹¹)</td>
<td>3×10⁸ (5×10⁹)</td>
<td>2×10⁸ (2×10⁹)</td>
<td>4×10⁸ (5×10⁹)</td>
</tr>
<tr>
<td>FLUKA</td>
<td>2×10⁸ (3×10¹¹)</td>
<td>2×10⁸ (3×10¹¹)</td>
<td>1×10⁸ (1×10¹⁰)</td>
<td>9×10⁷ (7×10⁹)</td>
<td>2×10⁷ (5×10⁹)</td>
</tr>
<tr>
<td>PHITS</td>
<td>5×10¹⁰ (6×10²⁰)</td>
<td>7×10⁹ (1×10⁰)</td>
<td>1×10⁹ (3×10⁷)</td>
<td>2×10⁸ (1×10⁰)</td>
<td>1×10⁸ (3×10⁷)</td>
</tr>
</tbody>
</table>

**Solar Minimum GCR benchmark.**

The present GCR benchmarks utilized the Badhwar-O’Neill 2010 [O’Neill 2010] model to represent the 1977 solar minimum conditions (solar modulation parameter set to 475 MV). The hydrogen, helium, carbon, and iron spectra were considered separately as external source boundary conditions. The results for the neutron and proton fluence spectra within a 40 g/cm² sphere of aluminum for an incident solar minimum GCR hydrogen spectrum are shown in Fig. 18 at the two depths of 35 and 40 g/cm². The GCR hydrogen results are qualitatively similar to results
obtained for the Webber SPE, only now the 3DHZETRN values favor Geant4 over the entire energy range except for the proton fluence below 100 MeV at the back surface (40 g/cm²). The computation time used for transporting GCR hydrogen is slightly longer than for the Webber SPE transport in the same object. Still, the 3DHZETRN time is a very manageable 8 CPU seconds whereas the Monte Carlo codes take several orders of magnitude longer.

Fig. 18. Particle spectra at detector locations 35 g/cm² (left pane) and 40 g/cm² from top of aluminum sphere (diameter 40 g/cm²) exposed to the 1977 solar minimum GCR hydrogen spectrum.

Fig. 19. Particle spectra at detector locations 35 g/cm² (left pane) and 40 g/cm² from top of aluminum sphere (diameter 40 g/cm²) exposed to the 1977 solar minimum GCR helium spectrum. The proton and alpha results have been scaled by 0.1 and 0.01 to improve plot clarity.

The neutron, proton, and alpha spectra generated by GCR helium ions are shown in Fig. 19. A large contribution to the alpha fluence spectrum is from the incident helium ions. There is generally better agreement with Geant4 results than those generated by FLUKA. The computer resources used in these calculations are given in...
Table 5. Similar results are shown for GCR carbon and iron ions in Figs. 20 and 21 with the corresponding resources used in Table 5. The light ion and neutron production nuclear database is more limited in HZETRN (and 3DHZETRN) when the incident ion has \( Z > 2 \). First, the light ion and neutron fragments from the projectile are assumed to have the same velocity as the fragmenting ion itself. Second, the light ions and neutrons produced as target fragments have been ignored. Furthermore, the light ions are mostly treated within the straight-ahead approximation that mainly affects the light ion solution below 100 MeV/n. The spectral results will improve as these additional nuclear databases and corresponding transport procedures are added to the 3DHZETRN code.

Fig. 20. Particle spectra at detector locations 35 g/cm\(^2\) (left pane) and 40 g/cm\(^2\) from top of aluminum sphere (diameter 40 g/cm\(^2\)) exposed to the 1977 solar minimum GCR carbon spectrum. The proton and alpha results have been scaled by 0.1 and 0.01 to improve plot clarity.

Fig. 21. Particle spectra at detector locations 35 g/cm\(^2\) (left pane) and 40 g/cm\(^2\) from top of aluminum sphere (diameter 40 g/cm\(^2\)) exposed to the 1977 solar minimum GCR iron spectrum. The proton and alpha results have been scaled by 0.1 and 0.01 to improve plot clarity.
Although it is difficult to favor one code over another on the basis of the veracity of the solutions obtained, there are vast differences in computational time required to evaluate fluence spectra even in this simplest of geometries (homogeneous sphere of a single elemental material). Still, Monte Carlo methods provide an important test point for developing codes capable of meeting operational and especially design requirements. Computational speed has proven to be important in spaceflight validation in LEO where the time structure of the environment can be used to test various environmental components as was done using ISS and the Liulin instrument [Wilson et al. 2007; Slaba et al. 2011a, 2013]. The main limitation on such studies remains the uncertainty in the environmental models (especially for the trapped environment) and uncertainty in nuclear cross sections.

Conclusions

The present version of 3DHZETRN shows improved agreement with Monte Carlo results in finite geometries where lateral leakage is non-negligible. Despite the improvements, the enhanced formalism and code has inherent limitations (aside from the simplified geometry of a homogeneous material) that carry over from the available bi-directional methods [Slaba et al. 2010b] in which the main share of light ions are still treated in the straight-ahead approximation as coupled to the forward propagating neutrons. Only the isotropically produced neutrons are first transported in 3D without the coupling to the isotropically produced light ion components. Once the isotropically produced neutrons are evaluated, the final coupling to the light ions is implemented (without further nuclear reactions) in a full 3D manner. Hence, only a fraction of the light ions have a 3D treatment, resulting in transport effects of unquantified magnitude. Future work will require extension to inhomogeneous media, complex geometries, and treatment of the cross-coupling of 3D neutron/light ion transport. Even so, the present expanded treatment of 3D effects shows great promise in the path to a fully 3D transport algorithm.

The present benchmarks provide useful exercises for evaluation of computational procedures and atomic/nuclear database. It is now time for improving the physical description of the transport process that is still hampered by a lack of basic nuclear physics database developments. The codes for space design are still incomplete in both the physical description and corresponding implementation into computational procedures. Even so, the current development appears as state-of-the-art in computational procedures for spacecraft design.

Acknowledgements

This work was supported by the Human Research Program under the Human Exploration and Operations Mission Directorate of NASA and by NASA Grant number NNX09AR20A.

Appendix A: Monte Carlo simulations

In this section, a brief description is given of the Monte Carlo codes Geant4, FLUKA, and PHITS used to simulate particle spectra shown in Figs. 8 and 16 - 21. A discussion of the source, geometry, and simulation setup used for the codes is also given.

Geant4 version 9.4.6 [Agostinelli et al. 2003] was used in the present calculations with the physics list QGSP_BERT_HP. This physics list utilizes the quark gluon string model (QGS) for high energy interactions with nucleons, pions, and nuclei. Post-interaction nuclear de-excitation is handled by the precompound (P) model. The Bertini cascade (BERT) model is used for interactions below 10 GeV. A high-precision (HP) tracking model is used for neutrons below 20 MeV. Nucleus-nucleus collisions are represented by the native Quantum Molecular Dynamics (QMD) model [Koi 2008]. More information about the chosen physics list can be found at the Geant4 website [Geant4 2012a]. The QGSP_BERT_HP physics list has been recommended by the Geant4 collaboration for high energy physics applications [Geant4 2012b] and has been used, with some slight variation, in other space radiation related studies [Bernabeu and Casanova 2007, Hayatsu et al. 2008, Martinez and Kingston 2012, Slaba et al. 2013].

FLUKA is a Monte Carlo program that contains fully integrated physics and performs the transport of elementary particles and ions through materials. It has the ability to transport and calculate interactions with all elementary hadrons, light and heavy ions, and electrons and photons over an energy range which extends up to $10^4$ TeV for all particles, and down to thermal energies for neutrons [Battistoni et al. 2007]. The nuclear models are integrated into the software for all particles and energies, with the exception of neutrons below 20 MeV, where standard international evaluated data files are used. This code has been extensively benchmarked against available accelerator and cosmic ray experimental data, at beam energies as low as a few MeV and as large as cosmic ray energies [Aarnio et al. 1993; Andersen et al. 2004].
PHITS has been used extensively in space radiation transport. It simulates ionization through transport processes and uses the mean free path to determine when collisions occur [Niita et al. 2006, Sihver et al. 2007]. Ionization transport can include angle and energy straggling, showing good agreement for Bragg peaks and fragmentation tails [Niita et al. 2006]. Tabulated nuclear cross section data are used for neutron interactions below 20 MeV, and evaluated data are used for gamma and electron transport below 1 GeV [Niita et al. 2006]. High energy neutron and heavy charged particle interactions are simulated through the Jet AA Microscopic transport model (JAM), which addresses hadron-hadron interactions, and the Japan Atomic Energy Research Institute (JAERI) Quantum Molecular Dynamics model (JQMD), which addresses hadron-nucleus and nucleus-nucleus interactions, up to energies of 200 GeV/n [Niita et al. 2006, Sato et al. 2006, Sihver et al. 2007]. Both JAM and JQMD are used for the dynamic portion of the nuclear reaction, and once the dynamics are completed, the General Evaporation Model is used to address nuclear de-excitation through particle emission in a statistical manner [Niita et al. 2006, Sihver et al. 2007]. The total nucleus-nucleus cross sections used in PHITS are the same as those that have been used in versions of HZETRN [Niita et al. 2006].

In Fig. 8, the geometry setup was for a 40 g/cm² aluminum slab with infinite lateral dimensions and the external source was applied normally to the front face of the slab. Particle spectra were tallied at a thin (0.027 g/cm²) aluminum detector layer placed 35 g/cm² and 40 g/cm² from the front of the slab. In Figs. 16 - 21, the geometry setup was for a sphere with diameter 40 g/cm², and the external source was applied uniformly onto the top of the sphere (covering the entire sphere) parallel with the z-axis as shown in Fig. 10. Particle spectra were tallied at spherical aluminum detectors (radius 0.675 g/cm²) placed along the z-axis down through the sphere. Convergence tests on the detector radius were performed to arrive at the final choice. In general, the chosen detector radius is large enough to allow reasonable tally statistics, but small enough to effectively prevent fragmentation events from occurring within the detector.

For all of the GCR simulations, the source energy spectrum was sampled directly from the input spectrum (i.e. no source energy biasing). For the SPE simulations, the source energy spectrum was biased so that source sampling was performed from a uniform distribution covering 0.01 MeV to 2500 MeV. This biasing technique places greater emphasis on the higher energy portion of the SPE spectrum improving tally statistics for secondary neutrons.

Appendix B: Evaluation of \( \Omega \) for \( N = 6, 10, 14, 18, 22 \)

The \( \Omega \) are herein evaluated with \( \Omega_0 \) as the top hemispheric pole so that \( \Omega_1 = \Omega_0 \), while the bottom pole is at \( \Omega_2 = -\Omega_0 \) and are the central rays of the two polar regions. The remaining directions denoted by \( \Omega_i \) for \( i = 3 \ldots N \) are symmetrically distributed about the mid-latitudes. Generally in application, the \( \Omega_i \) ‘s must be rotated to the body coordinates of the transport media. This is accomplished by a rotation matrix \( R(\theta_0, \phi_0) \) determined from the polar vector \( \Omega_0 \) relative to the body coordinate system (note, \( \theta \) denotes co-latitude with values on 0 to \( \pi \), and \( \phi \) denotes longitude running from 0 to 2\( \pi \)). The regional boundaries are chosen such that

\[
\int_{\Delta \Omega_i} d\Omega_i = 4\pi / N, \tag{46}
\]

where \( \Delta \Omega_i \) denotes the \( i^{th} \) region, \( N \) is the number of regions and is related to the number of \( N_\theta \) segments of \( \Delta \cos \theta \) and the number of \( N_\phi \) segments \( \Delta \phi \) plus the two polar regions such that

\[
N = N_\phi N_\theta + 2, \tag{47}
\]

and defined such that

\[
\sum_{i=1}^{N} \Delta \Omega_i = \sum_{i=1}^{N} \Delta \phi_i \Delta \cos \theta_i. \tag{48}
\]

The polar caps are bound by

\[
\cos \theta_1 = 1 - 2 / N, \tag{49}
\]
\[ \cos \theta_2 = -1 + 2 / N , \]  
and the corresponding central rays \( \Omega_1 \) and \( \Omega_2 \) lie along the \( \pm z \) axis. The middle latitudes are divided into \( N_\phi N_\theta \) regions extending from the polar cap to the equator in each hemisphere. The corresponding longitudinal boundaries are set by the \( N_\phi \) such that the

\[ \Delta \phi_i = 2\pi / N_\phi . \]  

Note that \( N_\phi \) is generally greater than or equal to 4. The remaining boundaries in latitude are given as

\[ \Delta \cos \theta_i = (\cos \theta_1 - \cos \theta_2) / N_\theta . \]  

For \( N = 6 \), there is only one midpoint in latitude taken as

\[ \cos \theta_i = (\cos \theta_1 + \cos \theta_2) / 2 = 0 , \]  

for \( i \geq 2 \). The four longitudinal points are given as

\[ \phi = \left\{ \frac{0, \pi}{2}, \pi, \frac{3\pi}{2} \right\} . \]  

The directional elements for each solid angle region are given by

\[ \Omega_1 = \left\{ \cos \phi_i \sin \theta_i, \sin \phi_i \sin \theta_i, \cos \theta_i \right\} , \]  

and are given for \( N = 6 \) in Table B.1. Similar results for other \( N \) used in this paper are given in Tables B.2 - B.5.

**Table B.1 Directional elements of the \( \Omega \) for \( N = 6 \).**

<table>
<thead>
<tr>
<th>( i )</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>( e_x )</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>-1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>( e_y )</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>-1</td>
</tr>
<tr>
<td>( e_z )</td>
<td>1</td>
<td>-1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

**Table B.2 Directional elements of the \( \Omega \) for \( N = 10 \).**

<table>
<thead>
<tr>
<th>( i )</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td>( e_x )</td>
<td>0</td>
<td>0</td>
<td>0.9165</td>
<td>-0.9165</td>
<td>0</td>
<td>0</td>
<td>-0.9165</td>
<td>0.9165</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>( e_y )</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0.9165</td>
<td>-0.9165</td>
<td>0</td>
<td>0</td>
<td>-0.9165</td>
<td>0.9165</td>
</tr>
<tr>
<td>( e_z )</td>
<td>1</td>
<td>-1</td>
<td>0.4</td>
<td>-0.4</td>
<td>0.4</td>
<td>-0.4</td>
<td>0.4</td>
<td>-0.4</td>
<td>0.4</td>
<td>-0.4</td>
</tr>
</tbody>
</table>

**Table B.3 Directional elements of the \( \Omega \) for \( N = 14 \).**

<table>
<thead>
<tr>
<th>( i )</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>7</th>
<th>9</th>
<th>11</th>
<th>13</th>
</tr>
</thead>
<tbody>
<tr>
<td>( e_x )</td>
<td>0</td>
<td>0.8207</td>
<td>0</td>
<td>-0.8207</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>( e_y )</td>
<td>0</td>
<td>0</td>
<td>0.8207</td>
<td>0</td>
<td>-0.8207</td>
<td>0</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>( e_z )</td>
<td>1</td>
<td>0.5714</td>
<td>0.5714</td>
<td>0.5714</td>
<td>0.5714</td>
<td>0</td>
<td>0</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>( i )</th>
<th>2</th>
<th>4</th>
<th>6</th>
<th>8</th>
<th>10</th>
<th>12</th>
<th>14</th>
</tr>
</thead>
<tbody>
<tr>
<td>( e_x )</td>
<td>0</td>
<td>-0.8207</td>
<td>0</td>
<td>0.8207</td>
<td>0</td>
<td>-1</td>
<td>0</td>
</tr>
<tr>
<td>( e_y )</td>
<td>0</td>
<td>0</td>
<td>-0.8207</td>
<td>0</td>
<td>0.8207</td>
<td>0</td>
<td>-1</td>
</tr>
<tr>
<td>( e_z )</td>
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<td>-0.5714</td>
<td>-0.5714</td>
<td>-0.5714</td>
<td>-0.5714</td>
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</tr>
</tbody>
</table>
Table B.4 Directional elements of the $\Omega$ for $N = 18$.

<table>
<thead>
<tr>
<th>$i$</th>
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<th>5</th>
<th>7</th>
<th>9</th>
<th>11</th>
<th>13</th>
<th>15</th>
<th>17</th>
</tr>
</thead>
<tbody>
<tr>
<td>$e_x$</td>
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<td>0.7454</td>
<td>0</td>
<td>-0.7454</td>
<td>0</td>
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<td>0</td>
<td>-0.9750</td>
<td>0</td>
</tr>
<tr>
<td>$e_y$</td>
<td>0</td>
<td>0</td>
<td>0.7454</td>
<td>0</td>
<td>-0.7454</td>
<td>0</td>
<td>0.9750</td>
<td>0</td>
<td>-0.9750</td>
</tr>
<tr>
<td>$e_z$</td>
<td>1</td>
<td>0.6667</td>
<td>0.6667</td>
<td>0.6667</td>
<td>0.6667</td>
<td>0.2222</td>
<td>0.2222</td>
<td>0.2222</td>
<td>0.2222</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>$i$</th>
<th>2</th>
<th>4</th>
<th>6</th>
<th>8</th>
<th>10</th>
<th>12</th>
<th>14</th>
<th>16</th>
<th>18</th>
</tr>
</thead>
<tbody>
<tr>
<td>$e_x$</td>
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<td>-0.7454</td>
<td>0</td>
<td>0.7454</td>
<td>0</td>
<td>-0.9750</td>
<td>0</td>
<td>0.9750</td>
<td>0</td>
</tr>
<tr>
<td>$e_y$</td>
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<td>0</td>
<td>-0.7454</td>
<td>0</td>
<td>0.7454</td>
<td>0</td>
<td>-0.9750</td>
<td>0</td>
<td>0.9750</td>
</tr>
<tr>
<td>$e_z$</td>
<td>-1</td>
<td>-0.6667</td>
<td>-0.6667</td>
<td>-0.6667</td>
<td>-0.6667</td>
<td>-0.2222</td>
<td>-0.2222</td>
<td>-0.2222</td>
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</tr>
</tbody>
</table>

Table B.5 Directional elements of the $\Omega$ for $N = 22$.

<table>
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<tr>
<th>$i$</th>
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<th>3</th>
<th>5</th>
<th>7</th>
<th>9</th>
<th>11</th>
<th>13</th>
<th>15</th>
<th>17</th>
<th>19</th>
<th>21</th>
</tr>
</thead>
<tbody>
<tr>
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<td>0.6863</td>
<td>0</td>
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<td>0</td>
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<td>0</td>
<td>-0.9315</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>$e_y$</td>
<td>0</td>
<td>0</td>
<td>0.6863</td>
<td>0</td>
<td>-0.6863</td>
<td>0</td>
<td>0.9315</td>
<td>0</td>
<td>-0.9315</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>$e_z$</td>
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<td>0.7273</td>
<td>0.7273</td>
<td>0.7273</td>
<td>0.7273</td>
<td>0.3636</td>
<td>0.3636</td>
<td>0.3636</td>
<td>0.3636</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>$i$</th>
<th>2</th>
<th>4</th>
<th>6</th>
<th>8</th>
<th>10</th>
<th>12</th>
<th>14</th>
<th>16</th>
<th>18</th>
<th>20</th>
<th>22</th>
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</thead>
<tbody>
<tr>
<td>$e_x$</td>
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<td>0.6863</td>
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<td>0.9315</td>
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<td>0</td>
</tr>
<tr>
<td>$e_y$</td>
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<td>0</td>
<td>-0.6863</td>
<td>0</td>
<td>0.6863</td>
<td>0</td>
<td>-0.9315</td>
<td>0</td>
<td>0.9315</td>
<td>0</td>
<td>-1</td>
</tr>
<tr>
<td>$e_z$</td>
<td>-1</td>
<td>-0.7273</td>
<td>-0.7273</td>
<td>-0.7273</td>
<td>-0.7273</td>
<td>-0.3636</td>
<td>-0.3636</td>
<td>-0.3636</td>
<td>-0.3636</td>
<td>0</td>
<td>0</td>
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References


Scott, W.W., Alsmiller, R.G., Comparisons of results obtained with several proton penetration codes. ORNL-RSIC-17. 1967.


A 3DHZETRN Code in a Spherical Uniform Sphere with Monte Carlo Verification

Wilson, John W.; Slaba, Tony C.; Badavi, Francis F.; Reddell, Brandon; Bahadori, Amir A.

The computationally efficient HZETRN code has been used in recent trade studies for lunar and Martian exploration and is currently being used in the engineering development of the next generation of space vehicles, habitats, and extra vehicular activity equipment. A new version (3DHZETRN) capable of transporting High charge (Z) and Energy (HZE) and light ions (including neutrons) under space-like boundary conditions with enhanced neutron and light ion propagation is under development. In the present report, new algorithms for light ion and neutron propagation with well-defined convergence criteria in 3D objects is developed and tested against Monte Carlo simulations to verify the solution methodology. The code will be available through the software system, OLTARIS, for shield design and validation and provides a basis for personal computer software capable of space shield analysis and optimization.