## Modeling and Simulation of Online Reprocessing in the Thorium-Fueled Molten Salt Breeder Reactor

Andrei Rykhlevskii<sup>a</sup>, Jin Whan Bae<sup>a</sup>, Kathryn D. Huff<sup>a,\*</sup>

<sup>a</sup>Dept. of Nuclear, Plasma, and Radiological Engineering, University of Illinois at Urbana-Champaign, Urbana, IL 61801

## Abstract

Today climate change pushes humankind in search new ways to generate carbon-free, reliable base load power. Thus, interest in advanced nuclear energy and particularly Molten Salt Reactors (MSRs) has resurged, with multiple new companies pursuing commercialization of MSR designs (e.g. Transatomic, Terrapower, Terrestrial, Moltex Energy, Thorcon). To further develop these MSR concepts, researchers need simulation tools for analyzing liquid fueled molten salt reactor depletion and fuel processing. However, most contemporary nuclear reactor physics software is unable to perform high-fidelity full-core depletion calculations in an online reprocessing regime. This paper introduces a Python package, SaltProc, which couples with the Monte Carlo code, SERPENT2, and simulates MSR online reprocessing by modeling the changing isotopic composition of MSR fuel salt. SaltProc capabilities were demonstrated for full-core highfidelity model of the commercial Molten Salt Breeder Reactor (MSBR) concept and compared well to previous results from lower fidelity analyses.

*Keywords:* molten salt reactor, molten salt breeder reactor, python, depletion, online reprocessing, nuclear fuel cycle, salt treatment

<sup>\*</sup>Corresponding Author

Email address: kdhuff@illinois.edu (Kathryn D. Huff)

## 1. Introduction

The Molten Salt Reactor (MSR) is an advanced nuclear reactor which was developed at Oak Ridge National Laboratory (ORNL) in the 1950s and was operated in the 1960s. More recently, the Generation IV International Forum <sup>5</sup> (GIF) included MSRs among the six advanced reactor concepts promising for further research and development. MSRs offer significant improvements "in the four broad areas of sustainability, economics, safety and reliability, and proliferation resistance and physical protection" [1]. To achieve the goals formulated by the GIF, MSRs attempt to simplify the reactor core and improve inherent safety <sup>10</sup> by using liquid fuel.

In the thermal spectrum MSR, fluorides of fissile and/or fertile materials (i.e.  $UF_4$ ,  $ThF_4$ ,  $PuF_3$ ,  $TRU^1F_3$ ) combine with carrier salts to form a liquid fuel that circulates in a loop-type primary circuit [2]. Immediate advantages over traditional, solid-fueled, reactors include near-atmospheric pressure in the

- <sup>15</sup> primary loop, relatively high coolant temperature, outstanding neutron economy, improved safety parameters, reduced fuel preprocessing, and the ability to continuously remove fission products and add fissile and/or fertile elements [3]. The thorium-fueled Molten Salt Breeder Reactor (MSBR) was developed in the early 1970s by ORNL specifically to explore the promise of the thorium fuel
- <sup>20</sup> cycle, which uses natural thorium instead of enriched uranium. With continuous fuel reprocessing, the MSBR realizes the advantages of the thorium fuel cycle because the <sup>233</sup>U bred from <sup>232</sup>Th is almost instantly <sup>2</sup> recycled back to the core [4]. The chosen fuel salt, LiF-BeF<sub>2</sub>-ThF<sub>4</sub>-UF<sub>4</sub>, has a melting point of 499°C, a low vapor pressure at operating temperatures, and good flow and heat transfer
- <sup>25</sup> properties [5]. In the matter of nuclear fuel cycle, the thorium cycle produces a reduced quantity of plutonium and minor actinides (MAs) compared to the traditional uranium fuel cycle. Finally, the MSR also could be employed as

<sup>&</sup>lt;sup>1</sup> Transuranic elements

 $<sup>^2</sup>$   $^{232}$  Th transmutes into  $^{233}$  Th after capturing a neutron. Next, this isotope decays to  $^{233}$  Pa ( $\tau_{1/2}=21.83$ m), which finally decays to  $^{233}$  U ( $\tau_{1/2}=26.967$ d).

a converter reactor for transmutation of spent fuel from current Light Water Reactors (LWRs).

- Liquid-fueled systems present a challenge to existing neutron transport and depletion tools which are typically designed to simulate solid-fueled reactors. To handle the material flows and potential online removal and feed of liquid-fueled systems, early MSR simulation methods at ORNL integrated neutronics and fuel cycle codes (i.e., Reactor Optimum Design (ROD) [6]) into operational plant
- tools (i.e., Multiregion Processing Plant (MRPP) [7]) for MSR and reprocessing system design. Based on this approach, recent tools from universities and research institutions can approximate online refueling [8]. Recent research efforts in Europe and Asia mainly focus on fast spectrum reactor fuel cycle analysis. These couple external tools to neutron transport while depletion codes take into
  account continuous feeds and removals in MSRs. A summary of recent efforts

are listed in table 1
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Neutronic code	Authors	Spectra
MCNP/REM [9, 10]	Doligez et al., 2014; Heuer et al.,	fast
	2014 [11, 12]	
ERANOS [13]	Fiorina <i>et al.</i> , 2013 [14]	fast
KENO-IV/ORIGEN [15,	Sheu <i>et al.</i> , 2013 [17]	fast
16]		
SERPENT 2 [18]	Aufiero <i>et al.</i> , 2013 [19]	fast
MCODE/ORIGEN2 [20,	Ahmad <i>et al.</i> , 2015 [22]	thermal
21]		
MCNP6/CINDER90 [23]	Park et al., 2015; Jeong et al., 2016	thermal
	[24, 25]	
SCALE/TRITON [26, 27]	Powers et al., 2014; Betzler et al.,	thermal
	2017 [27, 28, 29]	
SERPENT 2	Rykhlevskii et al., 2017 [30]	thermal
MCNP/REM	Nuttin et al. [31]	thermal

Table 1: Tools and methods for fast spectrum system fuel cycle analysis.

Most of these methods are also applicable to thermal spectrum MSRs. Additional tools developed specifically for thermal MSR applications are also listed in table 1.

<sup>45</sup> References [11, 12, 17, 19] simulate some form of reactivity control, and methods [11, 12, 19, 22, 24, 25, 30, 31] use a set of all nuclides in depletion calculations.

Many liquid-fueled MSR designs rely on online fuel processing in which material moves to and from the core continuously or at specific time steps (batch-wise). In the batch-wise approach, the burn-up simulation stops at a given time and restarts with a new liquid fuel composition (after removal of discarded materials and addition of fissile/fertile materials). ORNL researchers have developed ChemTriton, a Python-based script for SCALE/TRITON which

uses the batch-wise approach to simulate a continuous reprocessing. ChemTriton

<sup>55</sup> models salt treatment, separations, discharge, and refill using a unit-cell MSR SCALE/TRITON depletion simulation over small time steps to simulate continuous reprocessing and deplete the fuel salt [27]. Methods listed in references [14, 17, 24, 25, 28, 29, 30] as well as the current work employ a batch-wise approach.

Accounting for continuous removal or addition presents a greater challenge since it requires adding a term to the Bateman equations. In SCALE [26], ORI-GEN [16] solves a set of Bateman equations using spectrum-averaged fluxes and cross sections generated from a deterministic transport calculation. Meanwhile, approaches listed in references [11, 12, 19, 31] model true continuous feeds and removals.

Thorium-fueled MSBR-like reactors similar to the one in this work are described in [24, 25, 27, 28, 29, 30, 31]. Nevertheless, most of these efforts considered only simplified unit-cell geometry because depletion computations for a many-year fuel cycle are computationally expensive even for simple models.

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Nuttin *et al.* broke up reactor core geometry into three MCNP cells: one for salt channels, one for the salt plena above and below the core and the last cell for the annulus, consequently, the two-region reactor core was approximated by one region with averaged fuel/moderator ratio [31]. A similar approach was used by Powers *et al.*, Betzler *et al.*, and Jeong *et al.* [27, 28, 4, 29, 32, 25]. This

- <sup>75</sup> approach misrepresents the two-region breeder reactor concept. The unit-cell or one-region models may produce reliable results for homogeneous reactor cores (i.e. Molten Salt Fast Reactor (MSFR), Molten Salt Actinide Recycler and Transmuter (MOSART)) or for one-region single-fluid reactor designs (i.e. Molten Salt Reactor Experiment (MSRE)). However, a two-region MSBR must
- <sup>80</sup> be simulated using a whole-core model to capture different neutron transport characteristics in the inner and outer regions of the core. In particular, most fissions happens in the inner region while breeding occurs in the outer zone.

Aufiero *et al.* extended the Monte Carlo burnup code SERPENT 2 and employed it to study the material isotopic evolution of the MSFR. The developed

- extension directly accounts for the effects of online fuel reprocessing on depletion calculations and features a reactivity control algorithm. The extended version of SERPENT 2 was assessed against a dedicated version of the deterministic ERANOS-based EQL3D procedure [13] and adopted to analyze the MSFR fuel salt isotopic evolution. We employed this extended SERPENT 2 for a simplified unit-cell geometry of thermal spectrum thorium-fueled MSBR and obtained
- results which contradict existing MSBR depletion simulations [25]. The present work introduces the online reprocessing simulation code, Salt-

Proc, which expands the capability of the continuous-energy Monte Carlo Burnup calculation code, SERPENT 2 [18], for simulation liquid-fueled MSR operation

<sup>95</sup> [33]. It also reports the application of the coupled SaltProc-SERPENT 2 system to the MSBR, an extension of the work presented in [34, 30]. In this work, we analyze MSBR neutronics and fuel cycle to find the equilibrium core composition. The additional objective is to compare predicted operational and safety parameters of the MSBR at both the initial and equilibrium states. Finally,
<sup>100</sup> <sup>232</sup>Th feed rate will be determined and MSBR fuel cycle performance will be analyzed.

The complex MSBR geometry is challenging to describe in software input, and, usually, researchers make significant geometric simplifications to model it [24]. This study leverages extensive computational resources to avoid these <sup>105</sup> geometric approximations in order to accurately capture breeding behavior.

#### 2. Methods

The ability of liquid-fueled systems to continuously remove fission products and add fissile and/or fertile elements is the main challenge for depletion simulations. The python package introduced in this work, SaltProc, takes into account online separations and feeds using the SERPENT 2 continuous-energy Monte Carlo neutron transport and depletion code.

#### 2.1. Molten Salt Breeder Reactor design and model description

The MSBR vessel has a diameter of 680 cm and a height of 610 cm. It contains a molten fluoride fuel-salt mixture that generates heat in the active <sup>115</sup> core region and transports that heat to the primary heat exchanger by way of the primary salt pump. In the active core region, the fuel salt flows through channels in moderating and reflecting graphite blocks. Fuel salt at 565°C enters the central manifold at the bottom via four 40.64-cm-diameter nozzles and flows upward through channels in the lower plenum graphite. The fuel salt exits at <sup>120</sup> the top at about 704°C through four equally spaced nozzles which connect to

the salt-suction pipes leading to primary circulation pumps. The fuel salt drain lines connect to the bottom of the reactor vessel inlet manifold.

Figure 1 shows the configuration of the MSBR vessel, including the "fission" (zone I) and "breeding" (zone II) regions inside the vessel. The core has two
radial zones bounded by a solid cylindrical graphite reflector and the vessel wall. The central zone, zone I, in which 13% of the volume is fuel salt and 87% graphite, is composed of 1,320 graphite cells, 2 graphite control rods, and 2 safety<sup>3</sup> rods. The under-moderated zone, zone II, with 37% fuel salt, and radial reflector, surrounds the zone I core region and serves to diminish neutron leakage.

 $<sup>^{3}</sup>$  These rods needed for emergency shutdown only.

<sup>130</sup> Zones I and II are surrounded radially and axially by fuel salt (figure 2). This space for fuel is necessary for injection and flow of molten salt.

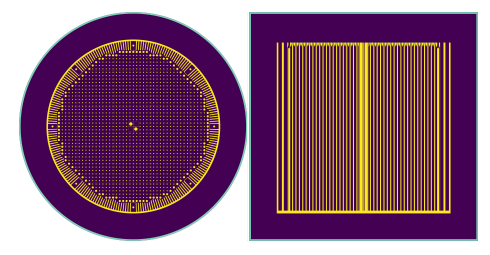


Figure 1: Plan and elevation views of SERPENT 2 MSBR model developed in this work.

Since reactor graphite experiences significant dimensional changes due to neutron irradiation, the reactor core was designed for periodic replacement. Based on the experimental irradiation data from MSRE, the core graphite lifetime is about 4 years and reflector graphite lifetime is 30 years [5].

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There are eight symmetric graphite slabs with a width of 15.24 cm in zone II, one of which is illustrated in Figure 2. The holes in the centers are for the core lifting rods used during the core replacement operations. These holes also allow a portion of the fuel salt to flow to the top of the vessel for cooling the top

- <sup>140</sup> head and axial reflector. Figure 2 also shows :w the 5.08-cm-wide annular space between the removable core graphite in zone II-B and the permanently mounted reflector graphite. This annulus consists entirely of fuel salt, provides space for moving the core assembly, helps compensate for the elliptical dimensions of the reactor vessel, and serves to reduce the damaging flux at the surface of the
- <sup>145</sup> graphite reflector blocks. In this work, all figures of the core were generated using the built-in SERPENT 2 plotter.

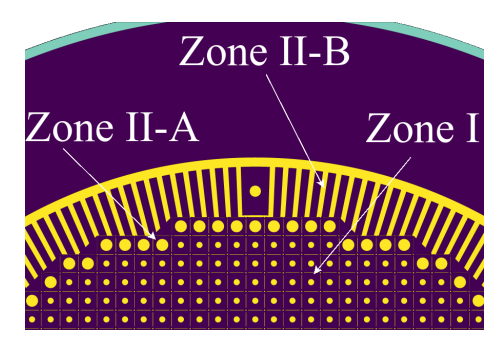


Figure 2: Detailed view of MSBR zone II model. Yellow represents fuel salt, purple represents graphite, and aqua represents the reactor vessel.

#### 2.1.1. Core zone I

The central region of the core, called zone I, is made up of graphite elements, each 10.16cm×10.16cm×396.24cm. Zone I has 4 channels for control rods: two for graphite rods which both regulate and shim during normal operation, and two for backup safety rods consisting of boron carbide clad to assure sufficient negative reactivity for emergency situations.

These graphite elements have a mostly rectangular shape with lengthwise ridges at each corner that leave space for salt flow elements. Various element sizes reduce the peak damage flux and power density in the center of the core to prevent local graphite damage. Figure 3 shows the elevation and plan views of graphite elements of zone I [5] and their SERPENT model [34].

## 2.1.2. Core zone II

Zone II which is undermoderated, surrounds zone I. Combined with the bounding radial reflector, zone II serves to diminish neutron leakage. Two kinds

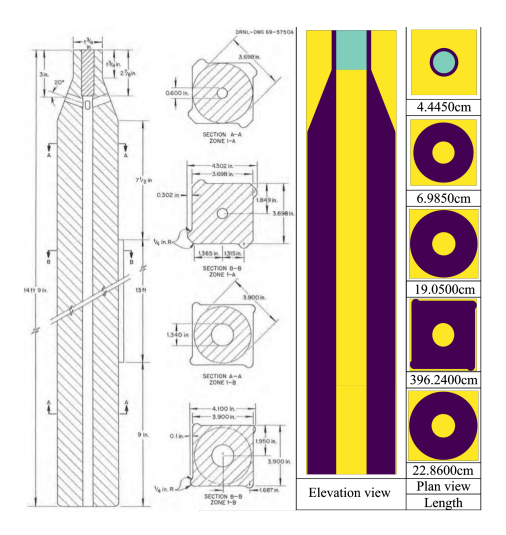


Figure 3: Graphite moderator elements for zone I [5, 34]. Yellow represents fuel salt, purple represents graphite, and aqua represents the reactor vessel.

of elements form this zone: large-diameter fuel channels (zone II-A) and radial graphite slats (zone II-B).

Zone II has 37% fuel salt by volume and each element has a fuel channel diameter of 6.604cm. The graphite elements for zone II-A are prismatic with elliptical dowels running axially between the prisms. These dowels isolate the fuel salt flow in zone I from that in zone II. Figure 4 shows the shapes and dimensions of these graphite elements and their SERPENT model. Zone II-B elements are rectangular slats spaced far enough apart to provide the 0.37 fuel salt volume fraction. The reactor zone II-B graphite 5.08cm-thick slats vary in

the radial dimension (average width is 26.67cm) as shown in figure 2. Zone II serves as a blanket to achieve the best performance: a high breeding ratio and a low fissile inventory. The harder neutron energy spectrum in zone II enhances the rate of thorium resonance capture relative to the fission rate, thus limiting the neutron flux in the outer core zone and reducing the neutron leakage [5].

The sophisticated, irregular shapes of the fuel elements challenge an accurate representation of zone II-B. The suggested design [5] of zone II-B has 8 irregularly-shaped graphite elements as well as dozens of salt channels. These graphite elements were simplified into right-circular cylindrical shapes with central channels. Figure 2 illustrates this core region in the SERPENT model. The

volume of fuel salt in zone II was kept exactly 37%, so that this simplification did not considerably change the core neutronics. Simplyfying the eight edge channels was te only simplification made to the MSBR geometry in this work.

## 2.1.3. Material composition and normalization parameters

The fuel salt, reactor graphite, and modified Hastelloy-N<sup>4</sup> are all materials created at ORNL specifically for the MSBR. The initial fuel salt used the same density (3.35 g/cm<sup>3</sup>) and composition LiF-BeF<sub>2</sub>-ThF<sub>4</sub>-<sup>233</sup>UF<sub>4</sub> (71.75-16-12-0.25 mole %) as the MSBR design [5]. The lithium in the molten salt fuel is fully enriched in <sup>7</sup>Li because <sup>6</sup>Li is a very strong neutron poison and becomes tritium upon neutron capture.

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The JEFF-3.1.2 neutron library provided cross section generation, [35]. The specific temperature was fixed for each material to correctly model the Dopplerbroadening of resonance peaks when SERPENT generates the problem-dependent nuclear data library. The isotopic composition of each material at the initial state was described in detail in the MSBR conceptual design study [5] and has

<sup>&</sup>lt;sup>4</sup> Hastelloy-N is very common in reactors now but have been studied and developed at ORNL in a program that started in 1950s.

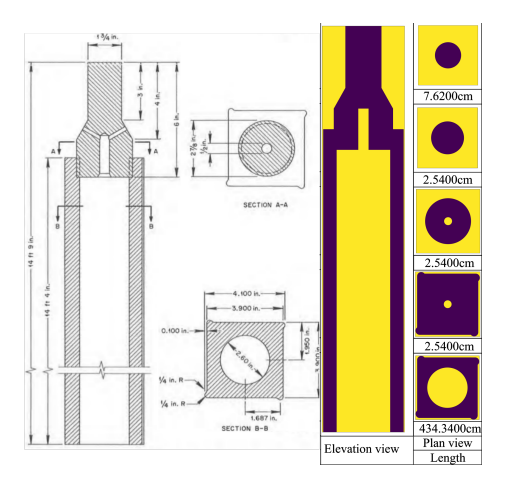


Figure 4: Graphite moderator elements for zone II-A [5, 34]. Yellow represents fuel salt and purple represents graphite.

<sup>195</sup> been applied to the SERPENT model without any modification. Table 2 is a summary of the major MSBR parameters used by this model [5].

## 2.2. Online reprocessing method

Removing specific chemical elements from a molten salt requires intelligent design (e.g., chemical separations equipment design, fuel salt flows to equipment) and has a considerable economic cost. All liquid-fueled MSR designs involve varying levels of online fuel processing. Minimally, volatile gaseous fission products (e.g. Kr, Xe) escape from the fuel salt during routine reactor operation

Table 2: Summary of principal data for MSBR [5].							
Thermal capacity of reactor	2250  MW(t)						
Net electrical output	1000 MW(e)						
Net thermal efficiency	44.4%						
Salt volume fraction in central zone I	0.13						
Salt volume fraction in outer zone II	0.37						
Fuel salt inventory (Zone I)	$8.2 \text{ m}^3$						
Fuel salt inventory (Zone II)	$10.8 \text{ m}^3$						
Fuel salt inventory (annulus)	$3.8 \text{ m}^3$						
Total fuel salt inventory	$48.7 \text{ m}^3$						
Fissile mass in fuel salt	1303.7 kg						
Fuel salt components	$\mathrm{LiF}\text{-}\mathrm{BeF}_2\text{-}\mathrm{ThF}_4\text{-}^{233}\mathrm{UF}_4$						
Fuel salt composition	$71.75\text{-}16\text{-}12\text{-}0.25~\mathrm{mole}\%$						
Fuel salt density	$3.35 \mathrm{~g/cm^3}$						

and must be captured. Additional systems might be used to enhance removal of those elements. Most designs also call for the removal of noble and rare earth metals from the core since these metals act as neutron poisons. Some designs suggest a more complex list of elements to process (figure 5), including the temporary removal of protactinium or other regulation of the actinide inventory [22].

## 2.2.1. Fuel material flows

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# The <sup>232</sup>Th in the fuel absorbs thermal neutrons and produces <sup>233</sup>Pa which then decays into the fissile <sup>233</sup>U. Furthermore, the MSBR design requires online reprocessing to remove all poisons (e.g. <sup>135</sup>Xe), noble metals, and gases (e.g. <sup>75</sup>Se, <sup>85</sup>Kr) every 20 seconds. Protactinium presents a challenge, since it has a large absorption cross section in the thermal energy spectrum. Accordingly, <sup>233</sup>Pa is continuously removed from the fuel salt into a protactinium decay tank to allow <sup>233</sup>Pa to decay to <sup>233</sup>U without poisoning the reactor. The reactor

reprocessing system must separate  $^{233}$ Pa from the molten-salt fuel over 3 days,

HIDROGEN H LITHERH SOCIUM Na 201745344	<ul> <li>Elements that escape from fuel salt</li> <li>Elements that can be removed without processing</li> <li>Elements that can be removed only by chemical processing of fuel salt</li> </ul>																
K	Ca	SCAROUR	Ti		Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Germannum	AREENIC	Se	Br	Kr
Rb	STRONTFUIM	YTTHUM Y	Zr	NDELUM	Мо	Тс	Ruthensium	Rh	Pattabilit	Ag	CADMUM	In	Sn	Sb	Те		Xe
CAESUM	Ва		HAFNEH	Та	TUNESTER W	Re	OSMUM Os	lr Ø	PLATINUM	Au	Hg		Pb	Bi	Ро	At	Rn
Fr <sub>P</sub>	Ra		Rf	Db	Sg	Bh	HASSOUN	Mt	Ds	Rg	Cn	Uut	Fi	Uup	Lv	Uus	Uuo
		Lanthumut	Се	Pr	Neccontration Nd	Prometinium Pm	Sumation Sm	Eu	Gd	Tb Tb	DYSMOSUUM Dy es	НОСИЛИМ	Er	Tm	YTTERRUM Yb 72	Lu	
		ACTINUM	Th			Np	Putonium	Am	Cm	Bk	Cf	Es	Fm	Md	No	LANNELNELUU Lr	

Figure 5: Processing options for MSR fuels. Reproduced from [22] where it was adapted from a chart courtesy of Nicolas Raymond, www.freestock.ca.

hold it while <sup>233</sup>Pa decays into <sup>233</sup>U, and return it back to the primary loop. This feature allows the reactor to avoid neutron losses to protactinium, lowers <sup>220</sup> in-core fission product inventory, and increases the efficiency of <sup>233</sup>U breeding. Table 3 summarizes full list of nuclides and the cycle times used for modeling salt treatment and separations [5].

Th removal rates vary among nuclides in this reactor concept which dictate the necessary resolution of depletion calculations. If the depletion time intervals are very short, an enormous number of depletion steps are required to obtain the equilibrium composition. On the other hand, if the depletion calculation time interval is too long, the impact of short-lived fission products is not captured. To compromise, the time interval for depletion calculations in this model was selected as 3 days to correlate with the removal interval of <sup>233</sup>Pa and <sup>232</sup>Th was continuously added to maintain the initial mass fraction of <sup>232</sup>Th.

#### 2.2.2. The SaltProc modeling and simulation code

The SaltProc tool [33] is designed to expand SERPENT 2 depletion capabilities for modeling liquid-fueled MSR for continuous reprocessing. The Python

Processing group	Nuclides	Cycle time (at full
		power)
Rare earths	Y, La, Ce, Pr, Nd, Pm, Sm, Gd	50 days
	Eu	500  days
Noble metals	Se, Nb, Mo, Tc, Ru, Rh, Pd, Ag,	20 sec
	Sb, Te	
Seminoble metals	Zr, Cd, In, Sn	200 days
Gases	Kr, Xe	20 sec
Volatile fluorides	Br, I	60 days
Discard	Rb, Sr, Cs, Ba	$3435 \mathrm{~days}$
Salt discard	Th, Li, Be, F	$3435 \mathrm{~days}$
Protactinium	<sup>233</sup> Pa	3 days
Higher nuclides	<sup>237</sup> Np, <sup>242</sup> Pu	16 years

Table 3: The effective cycle times for protactinium and fission products removal (reproduced from [5]).

package uses HDF5 [36] to store data, and the PyNE Nuclear Engineering
Toolkit [37] for SEPRENT output file parsing and nuclide naming. SaltProc is an open-source tool that uses a semi-continuous approach to simulate continuous feeds and removals in MSRs.

The tool structure and capabilities of SaltProc are similar to the ChemTriton tool developed in ORNL for SCALE [27]. SaltProc is coupled with the Monte <sup>240</sup> Carlo SERPENT 2 software to simulate online reprocessing for irregular full-core geometry with high fidelity. The primary function of SaltProc is to manage material streams while SERPENT 2 performs the computationally heavy work, namely neutron transport and depletion calculations. Saltproc is defined as a python class, where each material stream is defined as a isotopic atomic density

<sup>245</sup> vector variable. This allows tracking of time-sensitive material streams such as the <sup>233</sup>Pa tank in the MSBR. The user can define the reprocessing parameters, such as the reprocessing interval and removal efficiency. In addition, SaltProc provides a set of functions for each stream: read and write isotopic data in/from database, separate out specific isotopes from stream with defined efficiency, feed

- in specific isotopes to stream, and maintain constant number density of specific nuclide in the core. These attributes and functions are crucial to simulating the operation of a complex, multi-zone, multi-fluid MSR and sufficiently general to represent myriad reactor systems.
- SaltProc, currently in active development in Github (https://github.com/ arfc/saltproc), leverages unit tests and continuous integration for sustainable development. There is also documentation generated through Sphinx document generator for ease of use. In future releases, we plan to implement support for entirely user-customized reprocessing strategies, two-region MSR modeling capabilities, and decay modeling.
- Figure 6 illustrates the online reprocessing simulation algorithm coupling SaltProc and SERPENT 2. To perform a depletion step, SaltProc reads a user-defined SERPENT 2 template file. This file contains input cards with parameters such as geometry, material, isotopic composition, neutron population, criticality cycles, total heating power, and boundary conditions. After the depletion
  <sup>265</sup> calculation, SaltProc reads the depleted fuel composition file ( .bumat1) and stores the depleted composition isotopic vector in an HDF5 database.

SaltProc only stores and edits the isotopic composition of the fuel stream, which makes SaltProc a flexible tool to model any geometry: an infinite medium, a unit cell, a multi-zone simplified assembly, or a full core. This flexibiliity allows the user to perform simulations of varying fidelity and computational intensity.

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SaltProc can manage as many material streams as desired. It also may work with multiple depletion materials. At the end of a each depletion step, SaltProc reads the depleted compositions and tracks each material stream individually. Following this, it applies chemical separation functions to fuel stream vectors.

<sup>275</sup> These vectors then form a matrix (isotopics x timesteps) which SaltProc stores in an HDF5 database and prints into the SERPENT 2 composition file for the next depletion calculation.

Saltproc datasets are timseries, meaning that every value is recorded every

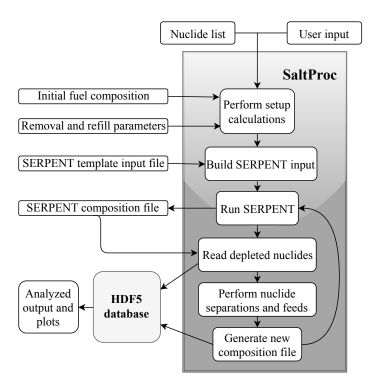


Figure 6: Flow chart for the Saltproc python package.

timestep. The datasets SaltProc produces are listed below, where the values inside the parenthesis are the dataset sizes:

- core adensity before reproc (number of isotopes x timesteps)
- core adensity after reproc (number of isotopes x timesteps)
- Keff\_BOC (1 x timesteps)
- Keff\_EOC (1 x timesteps)
- Th tank adensity (number of isotopes x timesteps)
  - iso codes (number of isotopes x 1)

In addition, SaltProc is able to define time-dependent material feed and removal rates to investigate their impacts. These rates need not be constant in SaltProc. They can be defined as piecewise functions or set to respond to

conditions in the core. For instance, SaltProc might increase the fissile material feeding rate if the effective multiplication factor,  $k_{eff}$ , falls below a specific limit (e.g., 1.002). These capabilities allow SaltProc to analyze fuel cycle of a generic liquid-fueled MSR. In summary, the development approach of SaltProc focused on producing a generic, flexible and expandable tool to give the SERPENT 2

<sup>295</sup> Monte Carlo code the ability to conduct advanced in-reactor fuel cycle analysis as well as simulate a myriad of online refueling and fuel reprocessing systems.

#### 3. Results

The SaltProc online reprocessing simulation package is demonstrated in four applications: (1) analyzing MSBR neutronics and fuel cycle to find the equilibrium core composition and core depletion, (2) studying operational and safety parameters evolution during MSBR operation, (3) demonstrating that in a single-fluid two-region MSBR conceptual design the undermoderated outer core zone II works as a virtual "blanket", reduces neutron leakage and improves breeding ratio due to neutron energy spectral shift, and (4) determining the effect of fission product removal on the core neutronics.

The neutron population per cycle and the number of active/inactive cycles were chosen to obtain balance between reasonable uncertainty for a transport problem (≤ 15 pcm<sup>5</sup> for effective multiplication factor) and computational time. The MSBR depletion and safety parameter computations were performed on 64 <sup>310</sup> Blue Waters XK7 nodes (two AMD 6276 Interlagos CPU per node, 16 floatingpoint Bulldozer core units per node or 32 "integer" cores per node, nominal clock speed is 2.45 GHz). The total computational time for calculating the equilibrium composition was approximately 9,900 node-hours (18 core-years.)

 $<sup>^{5}</sup>$  1 pcm =  $10^{-5}\Delta k_{eff}/k_{eff}$ 

3.1. Effective multiplication factor

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Figure 7 shows the effective multiplication factors obtained using SaltProc and SERPENT 2. The effective multiplication factors are calculated after removing fission products listed in Table 3 and adding the fertile material at the end of cycle time<sup>6</sup> which was fixed at 3 days for this work. The effective multiplication factor fluctuates significantly as a result of the batch-wise nature of this online



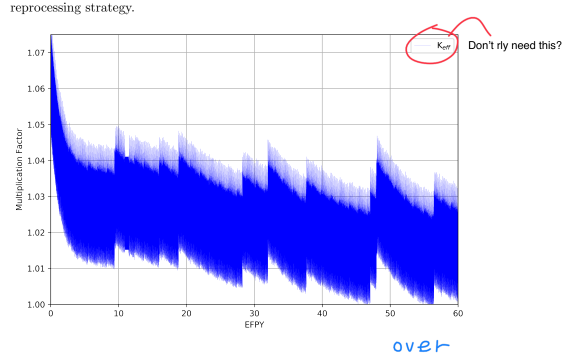


Figure 7: Effective multiplication factor dynamics for full-core MSBR model for a 60-year reactor operation.

First, SERPENT calculates the effective multiplication factor for the beginning of cycle time (fresh fuel composition for the first step). Next, it computes the new fuel salt composition for the end of a 3-day depletion step. The corresponding effective multiplication factor is much smaller than the previous one. Finally, SERPENT calculates  $k_{eff}$  for the depleted composition after applying feeds and

Not rly sure what you mean by this?

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 $<sup>^6</sup>$  The MSBR program defined a "cycle time" as the amount of time required to remove 100% of a target nuclide from a fuel salt.

removals.  $K_{eff}$  this increases accordingly since major reactor poisons (e.g. Xe, Kr) are removed, while fresh fissile material (<sup>233</sup>U) from the protactinium decay tank is added.

Additionally, the presence of rubidium, strontium, cesium, and barium in the core are disadvantageous to reactor physics. In fact, removal of these elements every 3435 days causes the multiplication factor to jump by approximately 450 pcm, and limits using the batch approach for online reprocessing simulation. Overall, the effective multiplication factor gradually decreases from 1.075 to  $k_{eff} \approx 1.02$  at equilibrium after approximately 6 years of irradiation.

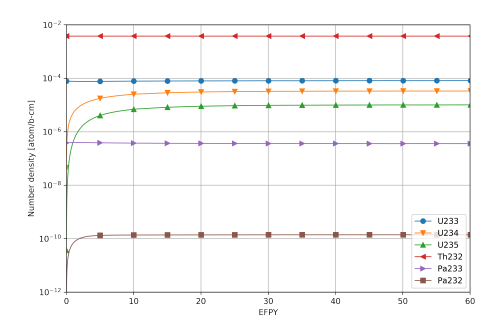
## 335 3.2. Fuel salt composition dynamics

The analysis of the fuel salt composition evolution provides more comprehensive information about the equilibrium state. Figure 8 shows number density of major nuclides which have a strong influence on the reactor core physics. The concentration of <sup>233</sup>U, <sup>232</sup>Th, <sup>233</sup>Pa, and <sup>232</sup>Pa in the fuel salt change insignificantly after approximately 2500 days of operation. Particularly, the <sup>233</sup>U number density fluctuates less than 0.8% in the time interval from 16 to 20 years of operation. Hence, a quasi-equilibrium state was achieved after 16 years of reactor operation. In contrast, a wide variety of nuclides, including fissile isotopes (e.g. <sup>235</sup>U) and non-fissile strong absorbers (e.g. <sup>234</sup>U), keep accumulating in the

- core. Figure 9 demonstrates production of fissile isotopes in the core. In the end of the considered operational time, the core contains significant <sup>235</sup>U ( $\approx 10^{-5}$  atom/b-cm), <sup>239</sup>Pu ( $\approx 5 \times 10^{-7}$  atom/b-cm), and <sup>241</sup>Pu ( $\approx 5 \times 10^{-7}$  atom/b-cm). Meanwhile, the equilibrium number density of the target fissile isotope <sup>233</sup>U was approximately  $7.97 \times 10^{-5}$  atom/b-cm. Thus, production of new fissile materials in the core as well as <sup>233</sup>U breeding make it possible to compensate for negative
  - effects of strong absorber accumulation and keep the reactor critical.

#### 3.3. Neutron spectrum

Figure 10 shows the normalized neutron flux spectrum for the full-core MSBR model in the energy range from  $10^{-8}$  to 10 MeV. The neutron energy spectrum



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Figure 8: Number density of major nuclides during 60 years of reactor operation.

at equilibrium is harder than at startup due to <sup>238</sup>Pu, <sup>239</sup>Pu, <sup>240</sup>Pu, <sup>241</sup>Pu, and <sup>242</sup>Pu accumulation in the core during reactor operation.

Figure 11 shows that zone I produced more thermal neutrons than zone II, corresponding to a majority of fissions occuring in the central part of the core. In the undermoderated zone II, the neutron energy spectrum is harder which leads to more capture of neutrons by <sup>232</sup>Th and helps a achieve relatively high breeding ratio. Moreover, the  $(n,\gamma)$  resonance energy range in <sup>232</sup>Th is from  $10^{-4}$  to  $10^{-2}$  MeV. Therefore, the moderator-to-fuel ratio for zone II was chosen to shift the neutron energy spectrum in this range. Furthermore, in the central core region (zone I), the neutron energy spectrum shifts to a harder spectrum over 20 years of reactor operation. Meanwhile, in the outer core region (zone II) a similar spectral shift takes place at a reduced scale. This results is in a good agreement with original ORNL report [5] and most recent whole-core steady-state study [24].

It is important to obtain the epithermal and thermal spectra to produce

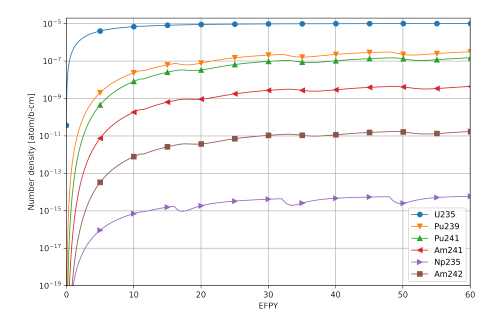


Figure 9: Number density of fissile in epithermal spectrum nuclides accumulation during the reactor operation.

 $^{233}$ U from  $^{232}$ Th because the radiative capture cross section of thorium decreases monotonically from  $10^{-10}$  MeV to  $10^{-5}$  MeV. Hardening the spectrum tends to significantly increase resonance absorption in thorium and decrease the absorptions in fissile and construction materials.

#### 3.4. Neutron flux

Figure 12 shows the radial distribution of fast and thermal neutron flux for both initial and equilibrium composition. The neutron flux has the same shape for both compositions but the equilibrium case has a harder spectrum. A significant spectral shift was observed for the central region of the core (zone I) when for the outer region (zone II) it is negligible for fast but notable for thermal

neutrons. This neutron flux radial distribution is in a good agreement with original ORNL report [5]. Overall, spectrum hardening during MSBR operation should be carefully studied for designing the reactivity control system.

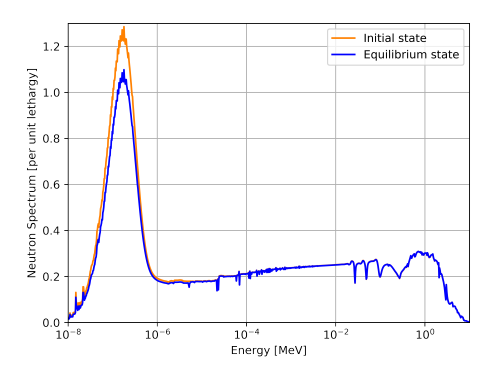


Figure 10: Neutron flux energy spectrum normalized by unit lethargy for initial and equilibrium fuel salt composition.

## 3.5. Power and breeding distribution

Table 4 shows the power fraction in each zone for initial and equilibrium fuel composition. Figure 13 reflects the normalized power distribution of the MSBR quarter core which is the same at both the initial and equilibrium states. For both the initial and equilibrium compositions, fission primarily occurs in the center of the core, namely zone I. The spectral shift during reactor operation results in different power fractions at startup and equilibrium, but most of the power is still generated in zone I at equilibrium. Figure 14 shows the neutron capture reaction rate distribution for <sup>232</sup>Th normalized by the total neutron flux for initial and equilibrium states. The distribution reflects the spatial distribution of <sup>233</sup>Th production in the core. The thorium-232 then  $\beta$ -decays to <sup>233</sup>Pa which is the precursor for <sup>233</sup>U production. Accordingly, this characteristic represents the

<sup>395</sup> breeding distribution in the MSBR core. Spectral shift does not cause significant

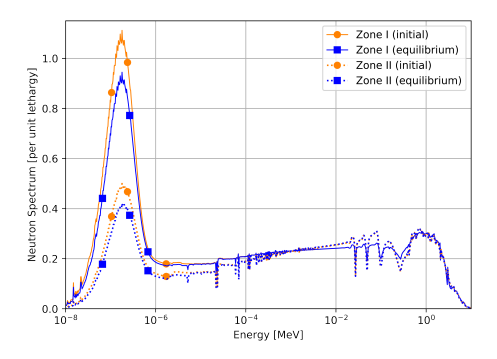


Figure 11: Neutron flux energy spectrum in different core regions normalized by unit lethargy for the initial and equilibrium fuel salt composition.

Core region	Initial	Equilibrium
Zone I	97.91%	98.12%
Zone II	2.09%	1.88%

changes in power nor in breeding distribution. Even after 20 years of operation, most of the power still is generated in zone I though the majority of <sup>233</sup>Th is produced in zone II.

## 3.6. Temperature coefficient of reactivity

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Table 5 summarizes temperature effects on reactivity calculated in this work for both initial and equilibrium fuel composition, and compared with original ORNL report data [5]. Uncertainty for each temperature coefficient also appears in Table 5. The main physical principle underlying the reactor

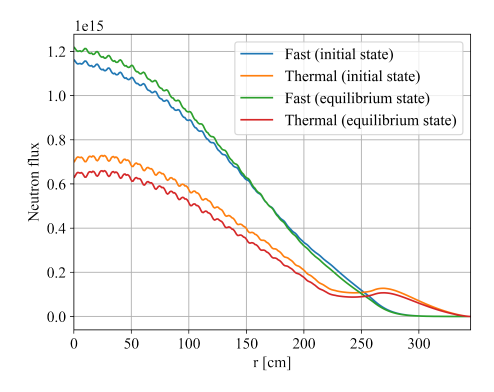


Figure 12: Radial neutron flux distribution for initial and equilibrium fuel salt composition.

- temperature feedback is an expansion of material that is heated. When the
  fuel salt temperature increases, the density of the salt decreases, but at the same time, the total volume of fuel salt in the core remains constant because it is bounded by the graphite. When the graphite temperature increases, the density of graphite decreases creating additional space for fuel salt. To determine temperature coefficients, the cross section temperatures for fuel and moderator
  were changed from 900K to 1000K. Three different cases were considered:
  - 1. Temperature of fuel salt rising from 900K to 1000K.
  - 2. Temperature of graphite rising from 900K to 1000K.
  - 3. Whole reactor temperature rising from 900K to 1000K.

In the first case, changes in the fuel temperature only impact fuel density. In <sup>415</sup> this case, the geometry is unchanged because the fuel is a liquid. However,



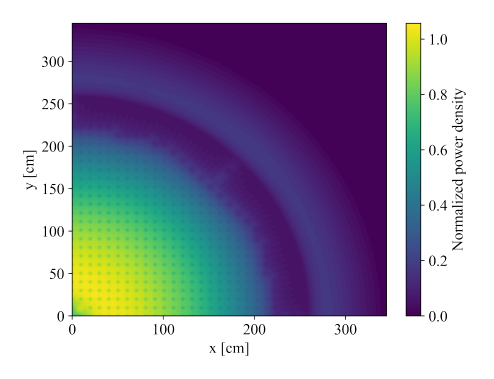


Figure 13: Normalized power density for both initial and equilibrium fuel salt composition.

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_	i i i i i i i i i i i i i i i i i i i	ure coefficients of	reacting	y for mithar and equilibri	um state.			
	Reactivity coeffi-	Initial		Equilibrium	Reference			
_	cient $[pcm/K]$				[5]			
	Fuel salt	$-3.22 \pm 0.044$		$-1.53 \pm 0.046$	-3.22			
•	Moderator	$+1.61 \pm 0.044$		$+0.97 \pm 0.046$	+2.35			
	Total	$-3.1\pm0.04$		$-0.97 \pm 0.046$	-0.87			
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Table 5: Temperature coefficients of reactivity for initial and equilibrium state.

when the moderator heats up, both the density and the geometry change due to thermal expansion of the solid graphite blocks and reflector. Accordingly, the new graphite density was calculated using a linear temperature expansion coefficient of  $1.3 \times 10^{-6} 1/K$  [5]. A new geometry input was created based on this information.

The fuel temperature coefficient (FTC) is negative for both initial and equilibrium fuel composition due to thermal Doppler broadening of the resonance

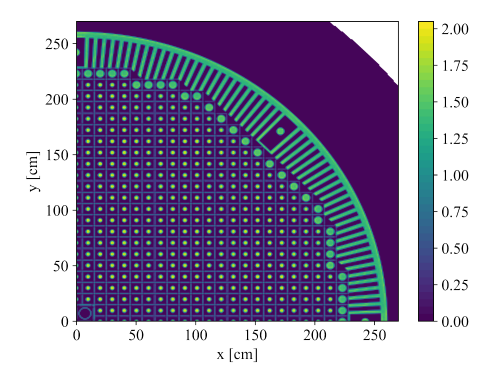


Figure 14: <sup>232</sup>Th neutron capture reaction rate normalized by total flux for both initial and equilibrium fuel salt composition.

capture cross sections in the thorium and is in a good agreement with earlier research [5, 24]. The moderator temperature coefficient (MTC) is positive for
startup composition and decreases during reactor operation because of spectrum hardening with fuel depletion. Finally, the total temperature coefficient of reactivity is negative for both cases, but decreases during reactor operation due to spectral shift. In summary, even after 20 years of operation the total temperature coefficient of reactivity is relatively large and negative during reactor operation, despite positive MTC, and affords excellent reactor stability and control.

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#### 3.7. Reactivity control system rod worth

Table 6 summarizes the reactivity control system worth. During normal operation the control (graphite) rods are fully inserted, and the safety  $(B_4C)$ 

rods are fully withdrawn. To insert negative reactivity into the core, the graphite rods are gradually withdrawn from the core. In an accident, the safety rods would fall down into the core. The integral rod worths were calculated for various positions to separately estimate control graphite moderator  $\mathrm{rods}^7$ , safety (B<sub>4</sub>C) rod, and the whole reactivity control system worth. Control rod integral worth

<sup>440</sup> is approximately 28 cents and stays almost constant during reactor operation. The safety rod integral worth decreases by 16.2% during 20 years of operation because of neutron spectrum hardening and absorber accumulation in proximity to reactivity control system rods. This 16% decline in control system worth should be taken into account in MSBR accident analysis and safety justification.

Table 6: Control system rod worth for initial and equilibrium fuel composition.

Reactivity parameter [cents]	Initial	Equilibrium
Control (graphite) rod integral worth	$28.2\pm0.8$	$29.0\pm0.8$
Safety $(B_4C)$ rod integral worth	$251.8\pm0.8$	$211.0\pm0.8$
Total reactivity control system worth	$505.8\pm0.7$	$424.9\pm0.8$
	I	

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#### 3.8. Six Factor Analysis

The effective multiplication factor could be expressed using formula:

$$k_{eff} = k_{inf} P_f P_t = \eta \epsilon p f P_f P_t$$

Table 7 summarizes the six factors for both initial and equilibrium fuel salt composition. The non-leakage probability for both fast and thermal neutrons does not change during reactor operation because these values are not largely affected by the neutron spectrum shift. In contrast, neutron reproduction factor  $(\eta)$ , resonance escape probability (p), and fast fission factor ( $\epsilon$ ) are considerably different between startup and equilibrium. As indicated in Figure 10 the neutron spectrum is softer at the beginning of reactor life. Neutron spectrum hardening

<sup>&</sup>lt;sup>7</sup>In [5], the graphite rods are referred to as "control" rods.

causes the fast fission to increase through the core lifetime. The opposite is true for the resonance escape probability. Finally, the neutron reproduction factor decreases during reactor operation due to accumulation of fissile plutonium

Factor	Initial	Equilibrium
Neutron reproduction factor $(\eta)$	$1.3960\pm .000052$	$1.3778 \pm .00005$
Thermal utilization factor (f)	$0.9670\pm .000011$	$0.9706 \pm .00001$
Resonance escape probability (p)	$0.6044 \pm .000039$	$0.5761 \pm .00004$
Fast fission factor $(\epsilon)$	$1.3421 \pm .000040$	$1.3609 \pm .00004$
Fast non-leakage probability $(\mathbf{P}_f)$	$0.9999 \pm .000004$	$0.9999 \pm .000004$
Thermal non-leakage probability	$0.9894 \pm .000005$	$0.9912 \pm .00005$
$(\mathbf{P}_t)$		

Table 7: Six factors for the full-core MSBR model for initial and equilibrium fuel composition.

## 3.9. Thorium refill rate

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isotopes.

In MSBR reprocessing scheme the only external feed material flow is <sup>232</sup>Th. Figure 15 shows the <sup>232</sup>Th feed rate calculated for 60 years of reactor operation. The <sup>232</sup>Th feed rate fluctuates significantly as a result of the batch-wise nature of this online reprocessing approach. For example, the large spikes up to 36 kg/day in a thorium consumption occurs every 3435 days. This is required due to batch-wise removal of strong absorbers (Rb, Sr, Cs, Ba). The corresponding effective multiplication factor increase (Figure 7) and breeding intensification leads to additional <sup>232</sup>Th consumption.

The average thorium feed rate increases during the first 500 days of operation and than steadily decreases due to spectrum hardening and accumulation of absorbers in the core. As a result, the average  $^{232}$ Th feed rate over 60 years of

<sup>470</sup> operation is about 2.40 kg/day. This results are in a good agreement with a recent online reprocessing study by ORNL [29]. Specify ORNL results?

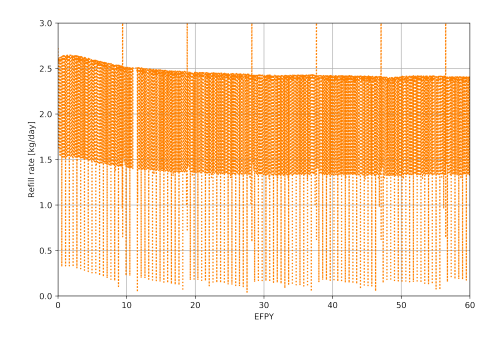


Figure 15:  $^{232}\mathrm{Th}$  feed rate over 60 years of MSBR operation.

## 3.10. The effect of removing fission product from fuel salt

Loading initial fuel salt composition into the MSBR core leads to supercritical configuration (Figure 16). After reactor startup the effective multiplication factor for the case with volatile gases and noble metals removal is approximately 7500 pcm higher than for case with no fission products removal. This significant impact on the reactor core achieved due to immediate removal (20 sec cycle time) and high absorption cross section of Xe, Kr, Mo, and other noble metals removed. The effect of rare earth element removal considerable after few month from startup and achieves approximately 5500 pcm after 10 years of operation. The rare earth elements are removed with slower rate (50 days cycle time). Moreover, Figure 16 demonstrates that batch-wise removal every 3-day step even strong absorbers did not necessarily leads to fluctuation in results but rare earth elements removal every 50 days causes approximately 600 pcm jump in reactivity.

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The effective multiplication factor of the core reduces gradually over operation

time because the fissile material  $\binom{233}{4}$  continuously depletes from the fuel salt due to fission and, at the same time, fission products accumulates in the fuel salt. Eventually, without fission products removal, the reactivity decreases to the subcritical state after approximately 500 and 1300 days of operation for cases with no removal and volatile gases & noble metals removal, respectively. The time when the simulated core reaches subcriticality ( $k_{eff} < 1.0$ ) for full-core model) called the core lifetime. Therefore, removing fission products provides with significant neutronic benefit and enables a longer core lifetime.

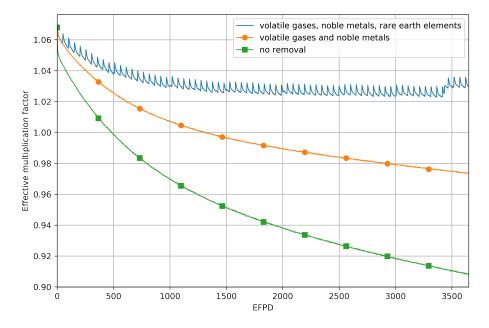


Figure 16: Calculated effective multiplication factor for full-core MSBR model with removal of various fission product groups over 10 years of operation.

## 495 **4. Discussion and conclusions**

This work introduces the open source MSR simulation package SaltProc. SaltProc expands the capability of SERPENT 2, the continuous-energy Monte Carlo code to include online reprocessing in liquid-fueled MSR operation [33]. Benefits of SaltProc include generic geometry modeling, multi-flow capabilities,

- time-dependent feed and removal rates, and the ability to specify removal efficiency. The main goal of this work has been to demonstrate the SaltProc capability to find the equilibrium fuel salt composition (where equilibrium is defined as when the number densities of major isotopes vary less than 1% over several years). A secondary goal has been to compare predicted operational
- and safety parameters (e.g., neutron energy spectrum, power and breeding distribution, temperature coefficients of reactivity) of the MSBR at startup and equilibrium state. A tertiary goal has been to demonstrate benefits of continuous fission products removal for thermal MSR design.
- Toward these goals, a full-core high-fidelity benchmark model of the MSBR <sup>510</sup> was implemented in SERPENT 2. The purpose of the full-core model instead of the simplified single-cell model [29, 30, 38] was to precisely describe the two-region MSBR concept design sufficiently to accurately represent breeding in the outer core zone. When running depletion calculations, the most important fission products and <sup>233</sup>Pa are removed and fertile fissile materials are added to
- the fuel salt every 3 days. Meanwhile, the removal interval for the rare earths, volatile fluorides, and seminoble metals was more than month which causes of effective multiplication factor fluctuation.

#### 4.1. Equilibrium state search

The results of this study indicate that the effective multiplication factor slowly decreases from 1.075 and reaches 1.02 at equilibrium after approximately 6 years of operation. At the same time, the concentration of <sup>233</sup>U, <sup>232</sup>Th, <sup>233</sup>Pa, <sup>232</sup>Pa stabilizes after approximately 2500 days of operation. Particularly, <sup>233</sup>U number density equilibrates<sup>8</sup> after 16 years of operation. Consequently, the core reaches the quasi-equilibrium state after 16 years of operation. On the other <sup>525</sup> hand, a wide diversity of nuclides, including fissile isotopes (e.g. <sup>233</sup>U, <sup>239</sup>Pu) and non-fissile strong absorbers (e.g. <sup>234</sup>U), continue accumulating in the core. The current work/results show that a true equilibrium composition cannot exist

 $<sup>^8 {\</sup>rm fluctuates}$  less than 0.8%

but the balance between strong absorber accumulation and new fissile material production can be achieved to keep the reactor critical.

530 4.2. Spectral shift

We also found that the neutron energy spectrum grows harder as equilibrium approaches because significant heavy fission products accumulate in the MSBR core. Moreover, the neutron energy spectrum in the central core region is much softer than in the outer core region due to lower moderator-to-fuel ratio in the outer zone, and this distribution remains stable during reactor operation. Finally, the epithermal or thermal spectrum is needed to effectively breed <sup>233</sup>U from <sup>232</sup>Th because radiative capture cross section of thorium-232 monotonically decreases from 10<sup>-10</sup> MeV to 10<sup>-5</sup> MeV. A harder spectrum in the outer core region tends to significantly increase resonance absorption in thorium and decrease the absorptions in fissile and structural materials.

The spatial power distribution in the MSBR shows that 98% of the fission power is generated in central zone I, and neutron energy spectral shift did not cause any notable changes in a power distribution. The spatial distribution of neutron capture reaction rate for fertile <sup>232</sup>Th, corresponding to breeding in the <sup>545</sup> core, confirms that most of the breeding occurs in an outer, undermoderated, region of the MSBR core. Finally, the average <sup>232</sup>Th refill rate throughout 60 years of operation is approximately 2.40 kg/day or 100 g/GWh<sub>e</sub>.

We compared the safety parameters for the initial fuel loading and equilibrium compositions using the SERPENT 2 Monte Carlo code. The total temperature coefficient is large and negative at startup and equilibrium but the magnitude decreases throughout reactor operation from -3.10 to -0.94 pcm/K as the spectrum hardens. The moderator temperature coefficient is positive and also decreases during fuel depletion. From reactivity control system efficiency analysis, showed the safety rod integral worth decreases by approximately 16.2% over 16 years of operation, while graphite rod integral worth remains constant. Summing up, neutron energy spectrum hardening during fuel salt depletion has an undesirable impact on MSBR stability and controllability, and should be taken into consideration in further analysis of accident transient scenarios.

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## 4.3. Benefits of fission product removal

For thermal spectrum MSBR removal of volatile gases, noble metals, and rare earths from a fuel salt benefits core performance. Moreover, immediate removal of volatile gases (e.g., xenon) and noble metals increased reactivity by approximately 7500 pcm over a 10-year timeframe. In contrast, the effect of relatively slower removal of rare earth elements (every 50 days cycle instead of 3 days) has less impact (5500 pcm) on the core reactivity after 10 years of operation. In sime, additional study is needed to establish neutronic and economic tradeoffs of removing each element.

4.4. Future work

Continued SaltProc-SERPENT coupled simulation efforts could progress in a number of different directions. First optimization of reprocessing parameters (e.g. time step, feeding rate, protactinium removal rate) could establish the best fuel utilization, breeding ratio, or safety characteristics for various designs. This might be performed with a parameter sweeping outer loop which would change an input parameter by a small increment, run the simulation and analyze output to determine optimal configuration. Alternatively, the existing RAVEN optimization framework [39] might be employed for such optimization studies.

Only the batch-wise online reprocessing approach has been treated in this work. However, the SERPENT 2 Monte Carlo code was recently extended for continuous online fuel reprocessing simulation [19]. This extension must be
verified against existing SaltProc/SERPENT or ChemTriton/SCALE packages, and could be employed for immediate removal of fission product gases (e.g., Xe, Kr) which have a strong negative impact on core lifetime and breeding efficiency. Finally, using the built-in SERPENT 2 Monte Carlo code online reprocessing & refueling material burnup routine would significantly speed up
computer-intensive full-core depletion simulations.

#### 5. Acknowledgments

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