

Advancing tritium self-sufficiency in fusion power plants: insights from the BABY experiment

Original

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





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Advancing tritium self-sufficiency in fusion power plants: insights from the BABY experiment

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Abstract

In the pursuit of fusion power, achieving tritium self-sufficiency stands as a pivotal challenge. Tritium breeding within molten salts is a critical aspect of next-generation fusion reactors, yet experimental measurements of Tritium Breeding Ratio (TBR) have remained elusive. Here we present the results of the Build A Better Yield blanket experiment, which represents a pioneering effort in tritium research by utilizing high-energy (14 MeV) neutron irradiation of molten salts, a departure from conventional low-energy neutron approaches. Using a small-scale (100 ml) molten salt tritium breeding setup, we not only simulated, but also directly measured a TBR (3.57×10^{-4}). This innovative approach provides crucial experimental validation, offering insights unattainable through simulation alone. Moreover, our findings reveal a surprising outcome: tritium was predominantly collected as HT, contrary to the expected TF. This underscores the complexity of tritium behavior in molten salts, highlighting the need for further investigation. This work lays the foundation for a more sophisticated experimental setup, including increasing the volume of the breeder, enhancing neutron detection, and refining tritium collection systems. Such improvements are crucial for advancing our understanding of fusion reactor feasibility and paving the way for future experiments.

Keywords: tritium breeding, TBR, molten salts

(Some figures may appear in colour only in the online journal)

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Glossary

BABY	Build a better yield blanket
D–T	Deuterium–tritium
FPP	Fusion power plant
LIB	Liquid immersion blanket
LIBRA	Liquid immersion blanket: robust accountancy
LSC	Liquid scintillation counting
TBR	Tritium breeding ratio

1. Introduction

Most proposed FPPs are fueled by D–T fusion reactions [1], but tritium is scarce [2, 3]. It is essential to develop tritium breeding blanket and fuel cycle technologies that can generate and maintain a self-sustaining supply of tritium within the FPP. Tritium self-sufficiency refers to the ability of the FPP to breed and process enough tritium to run the plant without relying on an external source [4–7]. Achievement of tritium self-sufficiency is intrinsically tied to the design and performance of the tritium breeding blanket.

However, tritium self-sufficiency on a large scale has never been demonstrated. The LIBRA project was proposed at the MIT Plasma Science and Fusion Center (PSFC) to address critical research gaps in understanding tritium breeding and tritium chemistry in molten FLiBe exposed to a fusion neutron environment [8]. FLiBe has been proposed as a breeder material for the LIB of ARC-class FPPs [9]. In an ARC-class FPP, the first wall and vacuum vessel structures are coupled and located inside a tank of molten FLiBe salt. The LIB is a self-cooled blanket concept [10], and the FLiBe serves as breeder, neutron shield, and coolant for the vacuum vessel [11]. The beryllium in FLiBe acts as a neutron multiplier, which significantly enhances TBR. Although there are many challenges associated with working with beryllium because of its toxicity, FLiBe remains the most promising breeder choice for the ARC-class FPP due to its cooling capabilities [12], and its ability to achieve high TBR [13]. Recent models have shown that the FLiBe LIB should enable the ARC-class FPP to achieve tritium self-sufficiency [6].

The primary parameter used to describe the efficiency of a breeding blanket is the TBR. The TBR is defined as the ratio of tritium produced in the blanket to tritium consumed in the plasma. To achieve self-sufficiency, the TBR of a commercial fusion reactor must be greater than unity, meaning that more tritium is generated than consumed during the fusion process. The TBR required by a given FPP design depends on the plant's performance requirements, the efficiency of its fuel cycle components, expected tritium losses, and reserve inventory needs. Designing a breeding blanket with a TBR high enough is a critical engineering challenge on the path to commercial fusion power.

The long-term goal of LIBRA is to demonstrate a $TBR \geq 1$ in a large volume (1000 kg \sim 500 l) of FLiBe molten salt using D–T neutron generators. Note that a full-scale LIB in an ARC-class FPP will require $\sim 250\,000$ l of FLiBe, hence the importance of understanding tritium behavior in large salt volumes. LIBRA presents major safety and infrastructure

challenges related to beryllium, high-energy neutrons, tritium, high-temperature fluids (>750 K) and hydrofluoric acid formation. Therefore, the PSFC is undertaking a scaled approach to develop the knowledge, team, and techniques necessary to achieve the goals of LIBRA, beginning with the BABY campaign, which is the focus of this paper.

The ITER and DEMO research programs have adopted a similar scaled approach through the test blanket module program [14], which evaluates fabrication, cooling, and tritium production capabilities using blanket mock-ups for the Water Cooled Lithium Lead (WCLL), Helium Cooled Ceramic Pebble (HCCP), Helium Cooled Ceramic Breeder (HCCB), and Water Cooled Ceramic Breeder (WCCB) concepts [15–17]. Neutronics experiments were conducted on the mock-ups with 14 MeV neutrons at low neutron fluxes ($<5 \times 10^8$ ncm $^{-2}$ s $^{-1}$), aimed at validating numerical tools, developing neutron detection techniques, and quantifying tritium production rates [18, 19]. Similar neutronics experiments were carried out for the WCCB blanket of the China fusion engineering test reactor (CFETR) [20].

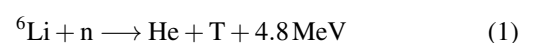
This paper presents the first results of the BABY experiment. BABY is a small-scale (100 ml) molten FLiBe salt tritium breeding experiment that uses 14 MeV D–T neutrons and is the initial experimental step towards the construction and operation of LIBRA. Smaller-scale experiments enable the LIBRA team to gain experience in neutron generation, neutron detection, and tritium accountancy and refine the experimental plans for LIBRA while minimizing safety hazards. Furthermore, it enables the near-term generation of useful new data on tritium breeding in molten salts. For the initial test results presented here, the chemical hazard associated with FLiBe was alleviated by using a surrogate salt: CLiF, a mixture of lithium fluoride (LiF) and lithium chloride (LiCl). By avoiding the presence of beryllium for the initial tests, the team was able to build and troubleshoot the BABY experimental setup before introducing the additional safety protocols and equipment required for beryllium work.

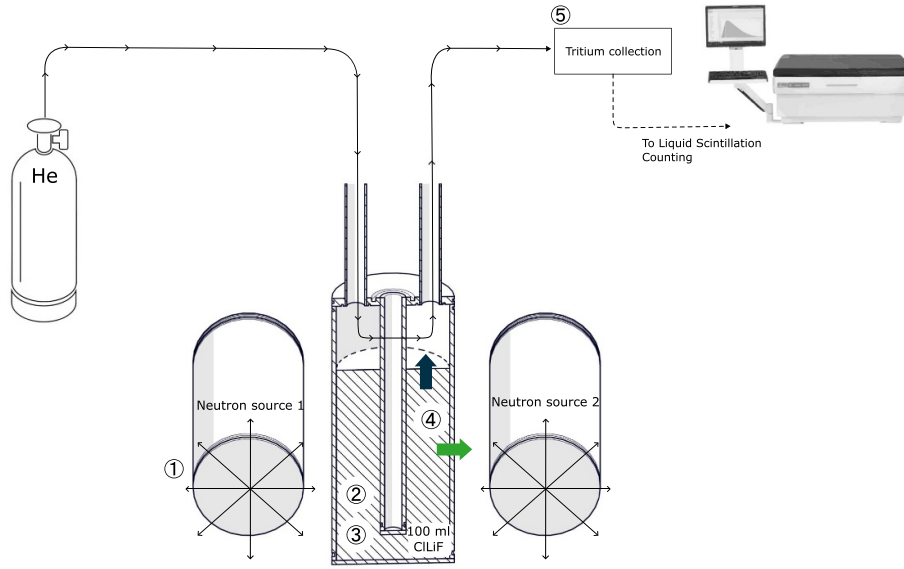
2. Principle

This section describes the general principle of the BABY experiment. Higher level methodology details are available in section 5.

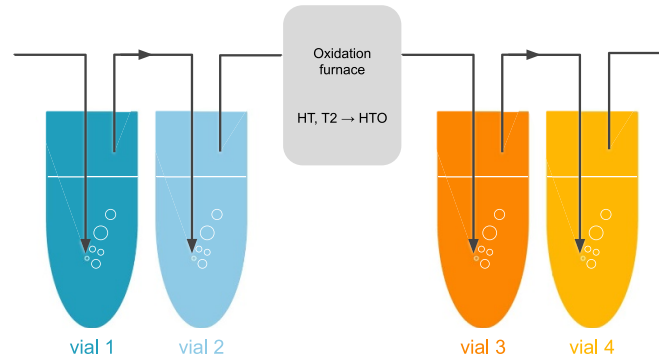
The BABY experiment is composed of a crucible containing 100 ml of molten salt, two sources of 14 MeV neutrons, a gas system, a neutron detection system and a tritium collection system (see figure 1(a)):

1. The system is conditioned with 3.5% H₂ in He for 4.5 h at 30 ccm followed by at least 1 h of pure He at 20 ccm. The salt is then brought to 700 °C. It takes a few hours for the salt temperature to equilibrate.
2. Once the system is at equilibrium, 14 MeV neutrons are generated.
3. Neutrons interact with lithium in the salt to produce tritium according to the following reactions:



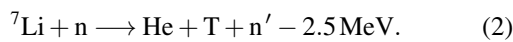


(a) Experimental setup. The thermal insulation around the crucible is not shown here.



(b) Tritium detection principle. The first two vials (in blue) collect tritium in soluble form (HTO, TCl and TF). Insoluble tritium (HT and T₂) is then passed through an oxidation furnace, where it is converted into HTO and collected in the second pair of vials (in orange).

Figure 1. BABY experimental setup.



4. Tritium is transported in the salt through diffusion and/or advection due to natural convection currents in the molten salt [21].
5. Tritium eventually reaches the top surface of the molten salt, which is in contact with the sweep gas, and the tritium is released into the gas stream, or reaches the salt-metal interface at the walls of the crucible, permeates the metal wall, and is released into the room.
6. Tritium in the gas stream is collected and counted.

The ratio of the convective mass transfer to the diffusive mass transfer is expressed by the Sherwood number:

$$\text{Sh} = \frac{k}{D/L} \quad (3)$$

where k is the mass transport coefficient (m s^{-1}), D is the diffusion coefficient ($\text{m}^2 \text{s}^{-1}$), and L is the characteristic length of the system (m). When $\text{Sh} \gg 1$ the dominant mode of mass transfer is advection. When $\text{Sh} \ll 1$, the dominant mode is diffusion. In the BABY experiment, it is expected that $\text{Sh} \gg 1$.

2.1. Molten salt crucible

A mass of 190 g of molten CLiF salt, with stoichiometric composition 30.5% LiF–69.5% LiCl (eutectic), were placed in a 1.6 mm thick Inconel-625 crucible (see figure 1(a)). Natural lithium was used. The internal height of the crucible is 101 mm and its internal diameter is 42 mm. Detailed drawings and CAD models are available at [22]. Helium was swept across the top surface of the CLiF inside the crucible to carry away the released tritium. The molten salt was kept at 700 °C using a re-entrant heater in the center of the crucible. The crucible was wrapped in alumina wool for thermal insulation. The heater

was connected by a temperature controller and a sheathed thermocouple immersed in the salt.

2.2. Neutron source

A Thermo-Fisher™ A-325 and a Thermo-Fischer™ P-383, both sealed-tube neutron generators, were used for the BABY experiment. Sealed-tube neutron generators house a deuterated and tritiated metal target that is surrounded by a high voltage electrode. High voltage is applied across the electrode, creating an electric field that accelerates deuterium ions into the target, where they slow down due to Coulomb collisions. A fraction of deuterium ions undergo D–T nuclear fusion reactions, producing a He nucleus and a 14 MeV neutron. The sealed design ensures that the radioactive tritium fuel remains contained, and the generated neutrons can be employed for various applications. It is worth noting that the energy spectrum of these sources is not purely 14 MeV. First, DD reactions also occur producing ≈ 2 MeV neutrons. Moreover, as the neutrons interact with the materials of the source itself, they are attenuated resulting in an energy distribution (see section 3.1.1).

2.3. Neutron detection

The neutron detection system on the BABY experiment is a combination of activation foils and diamond detectors.

Activation foil analysis is a neutron detection technique based on the principle of inducing nuclear reactions in target materials known as activation foils [23]. The populations of radioactive isotopes created in the foil are characterized by using a suitable detection system (e.g. gamma ray spectroscopy), and these can be directly correlated to the energy spectrum and fluence of the incident neutrons. Foils can be strategically placed to characterize the neutron flux at different locations of interest.

Diamond detectors were used to measure the neutron flux near the crucible. Diamond detectors operate on the basis of the neutron-induced displacement damage process in the crystalline lattice of a diamond [24]. The resulting vacancy and interstitial defects act as electrically active centers within the diamond structure and create a measurable electrical signal that can be correlated to the characteristics of the impinging neutron flux. The most common type of diamond detector for neutron detection utilizes synthetic diamond doped with isotopic impurities such as nitrogen-vacancy (NV) centers. The NV centers serve as sensitive charge carriers and enhance the detection efficiency. Diamond detectors exhibit high sensitivity to both thermal and fast neutrons, making them versatile tools for neutron spectroscopy.

2.4. Tritium detection

The tritium detection method is LSC [25]. The gas containing traces of tritium first passes through a volume of water, stripping out all soluble forms of tritium (HTO and TF) (see figure 1(b)). It is then passed through an oxidizing furnace,

which converts insoluble forms of tritium (HT, T₂) into HTO. The gas is then passed through another volume of water to collect the remaining tritium. Therefore, it is possible to discriminate between the soluble and insoluble forms of tritium. The radioactivity of tritiated water is then measured by LSC.

3. Results

All data, analysis and post-processing scripts are available at <https://github.com/LIBRA-project/insights-from-BABY-experiment-paper> [22].

3.1. TBR measurement

3.1.1. Neutron fluence measurement. The diamond detector was used to obtain a real-time measurement of the neutron count rate. The (n, α) peak can be seen between channel 1180 and 1300 of the uncalibrated neutron energy spectrum (see figure 2(a)). Counts within these energy channels corresponding to the (n, α) peak were binned with a bin size of 1000 s (see figure 2(b)). During most of the irradiation, this peak count rate remained steady at about 3.5 to 3.6 counts per second (CPS), but during the first irradiation period, a problem occurred with the P383 neutron generator, resulting in the generator being turned on and off and tuned repeatedly, explaining the dip seen between $t = 16000$ s and $t = 25000$ s in figure 2(b). Additionally, for an unknown reason, the count rate increased to an average rate of 3.9 CPS between $t = 43000$ s and $t = 50000$ s. During the second irradiation (90 000 s to 133 000 s), the peak count rate remained constant at 3.6 CPS.

The results plotted in figure 2 represent the count rate of the detector pulses from the diamond detector, but to convert these results into an actual neutron rate requires a measurement of the intrinsic efficiency of the diamond detector. This measurement is planned for future experiments.

However, by also employing activation foil analysis, the total neutron fluence and therefore the average neutron rate were measurable. Niobium activation foils were placed on top of each generator and aligned with the target plane. The neutron rates of the P383 and A325 generators were calculated to be $2.71 \times 10^8 \text{ ns}^{-1}$ and $1.16 \times 10^8 \text{ ns}^{-1}$, respectively, with a relative error of 8.9%. The total neutron fluence over 24 h of irradiation is therefore $(3.35 \pm 0.30) \times 10^{13} \text{ n}$.

3.1.2. Tritium production measurement. 21 Bq of tritium³ were collected in total (see figure 4). The collection vials were changed every 12 h before day 2 and then regularly until no more tritium was collected. The tritium in the water was then counted with the LSC. Based on the collected amounts, the overall collection efficiency (with two vials) was calculated to be $>99.7\%$.

Because each neutron is the product of one triton being consumed in a D–T reaction in the generators, the TBR of the

³ For convenience, tritium quantities are given in Bq based on the tritium specific activity $3.57 \times 10^{14} \text{ Bq g}^{-1}$.

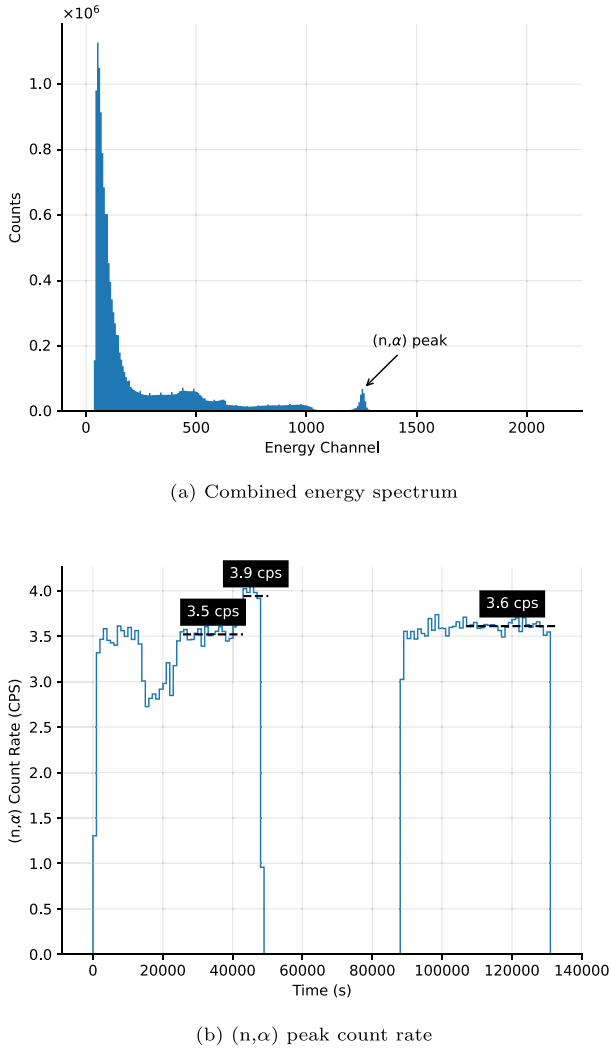


Figure 2. Diamond neutron detector results during BABY irradiation experiment.

experiment can be calculated from the ratio of total tritium production in BABY to measured neutron fluence:

$$\text{TBR} = \frac{1.17 \times 10^{10} \text{ T}}{3.35 \times 10^{13} \text{ n}} = 3.57 \times 10^{-4}. \quad (4)$$

3.1.3. Modeling TBR. The experiment was modeled using OpenMC [26]. The model geometry included the neutron sources, crucible, thermal insulation, salt, and different gases (see figure 3(a)). Modeling the crucible and the two sources was deemed sufficient for a robust TBR evaluation. A parametric study was carried out to maximize the TBR achievable in BABY. This study considered parameters such as the position of the source relative to the crucible, as well as the implementation of moderators and reflectors. The study showed that minimizing the source distance from the crucible was the most effective parameter for increasing the TBR, as this increases the total fluence of neutrons in the salt. The modeled TBR was sufficiently high that the inclusion of reflectors and moderators were deemed unnecessary. During the experiment, the

two neutron sources were positioned as close to the crucible as was possible without exceeding their maximum operating temperatures. The same model was applied to test and compare the TBR of different breeders at a 100 ml scale. The OpenMC study included FLiBe, which is the tritium breeder of choice for the ARC reactor concept [9, 11], PbLi—which is considered for some DEMO reactor breeding concepts [10], as well as FLiNaK and CLiF, which are lithium-based molten salts that do not contain beryllium. They are easier to handle than FLiBe and can breed tritium, although their breeding potential is lower than FLiBe, since beryllium acts as a neutron multiplier. All breeders studied have natural abundance of Li-6 ($\sim 7.6\%$).

At this scale, TBR is about four orders of magnitude lower than the value required by commercial fusion reactors (see figure 3(b)). However, with the irradiation schedule and tritium detection systems foreseen for this experiment, a TBR on the order of 10^{-4} was considered just sufficient to detect tritium. The TBR is low because only 2% of the neutron generator flux is incident on the salt (corresponding to a solid angle ≈ 0.3 sr). Future experiments in the LIBRA campaign will achieve higher TBR (~ 1) by placing the neutron generator inside the salt volume, achieving higher solid angle coverage.

At a volume of 100 ml, the TBR values for FLiBe and CLiF are comparable. Due to the purely fast neutron flux in the crucible, Li-7 is the dominant tritium breeding isotope, with a tritium yield at least one order of magnitude higher than Li-6. At these energies Fluorine has a nonnegligible contribution to the TBR of all molten salts, as well as beryllium in the case of FLiBe.

In our study, we observed that the measured TBR fell significantly short of the predictions made by the OpenMC simulation (see figure 3(c)). This discrepancy suggests that a substantial portion of the tritium produced during the experiment was not effectively collected. Since the bubbler collection efficiency was found to be close to 100%, we hypothesize that the loss of tritium through permeation across the container walls could explain this observed discrepancy. This phenomenon is further elucidated in the following section, where we discuss the transient tritium release model in detail.

3.2. Transient tritium release

Following the modeling strategy proposed by Kumagai *et al* [27], the temporal evolution of the salt tritium concentration c_{salt} is described by:

$$V \frac{dc_{\text{salt}}}{dt} = S - Q_{\text{wall}} - Q_{\text{top}} \quad (5)$$

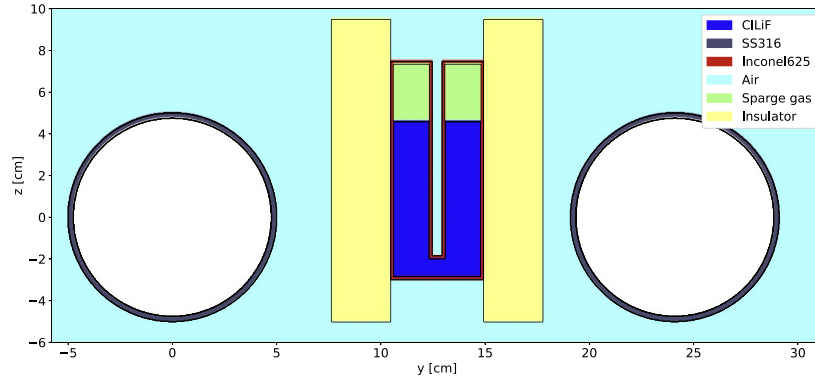
where V is the salt volume and S is the source of tritium in T s^{-1} expressed by:

$$S = \text{TBR} \cdot \Gamma_n \quad (6)$$

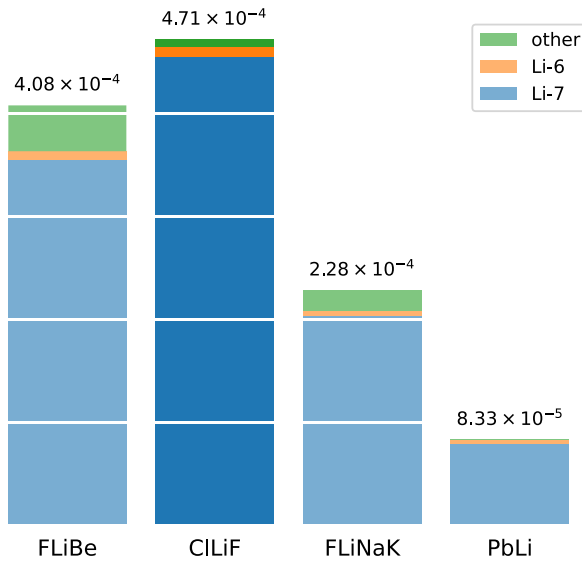
where Γ_n is the neutron rate in neutron s^{-1} .

The tritium release rates Q_i in T s^{-1} are expressed as:

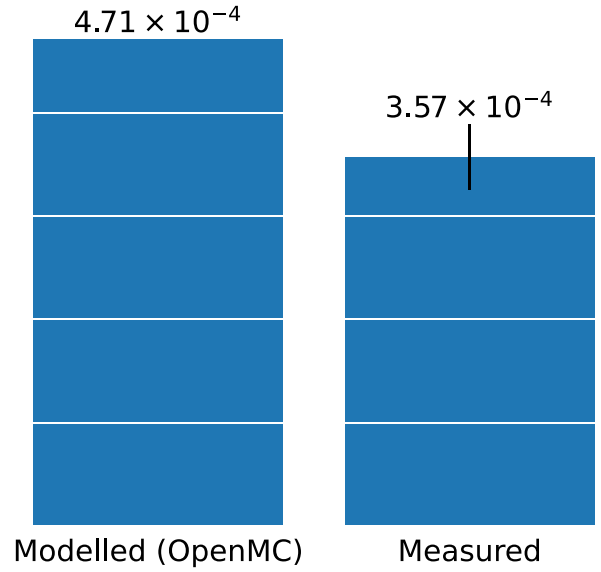
$$Q_i = A_i k_i (c_{\text{salt}} - c_{\text{external}}) \quad (7)$$



(a) Geometry used to model the BABY experiment in OpenMC.



(b) TBR values for FLiBe, CILiF, FLiNaK, and PbLi from OpenMC simulations of the BABY experiment. The contribution of lithium isotopes to tritium production is shown.



(c) Comparison between the experimentally measured TBR for BABY using CILiF salt and the TBR calculated in the corresponding OpenMC simulation.

Figure 3. Results of the OpenMC model. The reported values represent the statistical mean derived from Monte Carlo evaluations, with a relative standard deviation below 0.1%.

$$\approx A_i k_i c_{\text{salt}} \quad (8)$$

where A_i is the surface area of each release pathway in m^2 , c_{external} is the tritium concentration in the gas phase (assumed negligible), and k_i is the mass transport coefficient in m s^{-1} .

Equation (5) is solved numerically and the temporal evolution of c_{salt} is obtained. The release rates Q_i can be obtained from c_{salt} and the cumulative release can be expressed as $\int_0^{t_f} Q_i dt$. This is the quantity measured by the tritium detection system. The mass transport coefficients k_i are then varied in order to fit the experimental data.

It should be noted that the total production of tritium $\int_0^{t_f} S dt$ in the salt can be calculated independently of the mass transport coefficients. Indeed, by integrating equation (5), one can obtain:

$$V[c_{\text{salt}}(t = t_f) - c_{\text{salt}}(t = 0)] = \int_0^{t_f} S dt - \int_0^{t_f} \sum_i Q_i dt \quad (9)$$

where t_f is the final time.

If $c_{\text{salt}}(t = t_f) = c_{\text{salt}}(t = 0) = 0$, which is equivalent to $Q_i(t = t_f) = Q_i(t = 0) = 0$, then:

$$\int_0^{t_f} S dt = \int_0^{t_f} \sum_i Q_i dt \quad (10)$$

Assuming S is constant during irradiation periods:

$$S \Delta t_{\text{irradiation}} = \int_0^{t_f} \sum_i Q_i dt \quad (11)$$

$$S = \Delta t_{\text{irradiation}}^{-1} \int_0^{t_f} \sum_i Q_i dt \quad (12)$$

where $\Delta t_{\text{irradiation}} = 24 \text{ h}$ is the total irradiation time.

In other words, the source term of tritium S can be determined by measuring the cumulative tritium release over the span of the experiment as long as the initial and final concentrations of tritium in the salt are zero. Furthermore, the TBR can

Table 1. Parameters used in the tritium release model.

Parameter	Value
V	100 cm^3
A_{top}	13.80 cm^2
A_{wall}	116.18 cm^2
k_{top}	$9.11 \times 10^{-7} \text{ m s}^{-1}$
k_{wall}	$3.65 \times 10^{-8} \text{ m s}^{-1}$
S	$1.83 \times 10^5 \text{ T s}^{-1}$

be experimentally measured by a ratio of the total amount of tritium produced to the total amount of neutrons produced.

The parameters in the tritium release model are presented in table 1. The source S was assumed constant during the irradiation periods (as indicated by the neutron measurements in section 3.1). The TBR calculated by OpenMC was used as well as the measured neutron rate to determine the tritium source term. The values of the mass transport coefficient for release of tritium through the top surface of the salt, k_{top} , and the mass transport coefficient for tritium moving through the crucible walls, k_{wall} , were optimized to minimize the mean square error with the cumulative release measurements. The model agrees very well with the experimental data (see figures 4(a) and (b)). k_{top} was determined to be almost twice the value measured by Kumagai *et al* [27] ($9.11 \times 10^{-7} \text{ m s}^{-1}$ versus $4.9 \times 10^{-7} \text{ m s}^{-1}$, respectively). k_{wall} is also almost twice the value measured by Kumagai ($1.9 \times 10^{-8} \text{ m s}^{-1}$). The discrepancies are attributable to the different salt (FLiNaK instead of CILiF), different geometries, and lower temperature (500°C versus 700°C here) used in [27].

The Sherwood number can be computed (see equation (3)) from the measured mass transfer coefficient. Using the height of the crucible as characteristic length L and the diffusivities of FLiNaK and FLiBe measured in the right temperature range (HTM database v0.14 [28]), it ranges from 0.8 to 21.6 with an average value of 10.3. This suggests that convective mass transfer dominates tritium transport in the crucible. No suitable mass transport correlations were found in the literature for this problem [29]. In future work, computational studies will be performed in order to estimate the Sherwood number from a numerical model using the tritium transport code FESTIM [30].

A sensitivity study was performed and the value of k_{top} was varied by $\pm 30\%$ from its optimal value (k_{wall} was varied by the same factor). As explained above, the value of the total cumulative release is independent of the mass transport coefficient (see figure 4(c)). As the value of the mass transport coefficient increases, the time required to release all tritium from the salt decreases (i.e. the characteristic time of the system is shorter).

3.3. Chemical forms of tritium

The entirety of the collected tritium was in non-soluble forms (HT or T_2) during the described runs, which used pure He as a sweep gas. This result differs from previous observations [27, 31, 32] in which tritium was found as soluble TF or

HTO in FLiNaK and FLiBe when swept with pure He. Several explanations are possible. First, we are using a different salt which affects the chemistry of tritium speciation: more oxidizing salts preferentially produce soluble forms of tritium whereas reducing salts preferentially produce insoluble forms of tritium. To confirm this, we are planning an experiment in which europium fluoride is added to the salt to make it more oxidizing and expect tritium to be released mainly as TF/TCl. Another hypothesis was that the potential presence of H_2 (residual from the initial conditioning step with 3% H_2 (see section 2)) led to the formation of HT by isotopic exchange. We tested this hypothesis by removing this conditioning step, but observed no change in speciation.

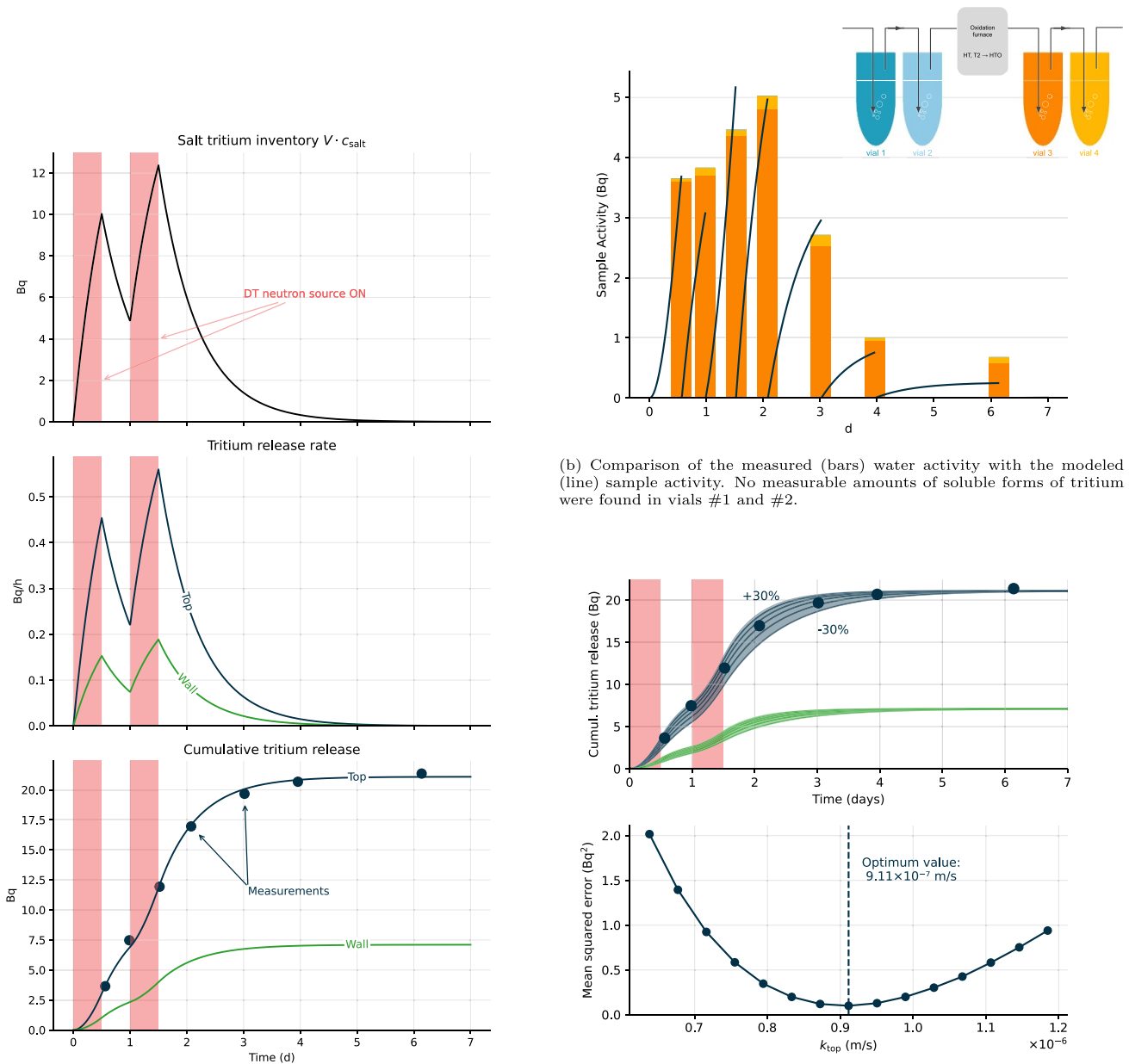
3.4. Reproducibility

These results have been reproduced in other runs (see figure 5). For all runs, the plateau value is similar with a relative standard deviation of 12.28%. Runs #1 and #2 were initial troubleshooting runs during which we did not detect tritium. Moreover, the neutron rate measurements showed similar results (see table 2). The measured TBR values for runs #4, #5, and #6 are also similar, with a standard deviation of 4.67%. The dynamics of the release are also very similar with the exception of run #4 where little tritium was detected before the end of the second irradiation. While the reasons of this behavior are still unclear, one hypothesis is that tritium was being held up somewhere in the apparatus and got released between day 2 and day 3. This could explain the large difference between the fourth and fifth data point that is not observable in runs #3 #5 and #6. The experimental setup was sometimes subject to pipe clogging due to the salt migrating in the tubing and solidifying. Other reasons could be a change in the convective flow of the salt leading to a reduced mass transport coefficient.

4. Discussion

Although these results are extremely encouraging and demonstrate capability in measuring a TBR in molten salts, several points will be improved in future experiments.

First, we found that the measured TBR was 2.3 times less than the expected TBR calculated using OpenMC. Future iterations of the BABY setup surround the crucible with a second vessel. Sweep gas through the interstitial space will capture the release of tritium that permeates through the crucible walls. This will test the hypothesis formulated in section 3.2 that there are nonnegligible permeation losses through the crucible walls. Assuming that the parameterization of the model detailed in this section is correct, the mass transfer coefficient for the wall release is an order of magnitude higher than that calculated by Kumagai *et al* [27]. In one of the experiments, the authors were unable to measure the permeated tritium, as it was below the LSC detection limit. At the high temperatures employed in BABY, the Inconel crucible wall is not expected to act as a tritium permeation barrier. However, it is possible that the components wrapped around the crucible for



(a) Modeling of the temporal evolution of the tritium salt inventory $V \cdot c_{salt}$ (top), tritium release rates Q_i (middle), and cumulative tritium release $\int_0^t Q_i dt$ with comparison to measurements (bottom).

(c) Temporal evolution of the cumulative tritium release (top) and evolution of the mean squared error (bottom) for different values of k_{top} . k_{wall} was varied accordingly.

Figure 4. Comparison of the transient tritium release model, which predicts tritium release through the crucible walls and from the top surface of the salt inside the crucible, with the tritium release measurements (here, only tritium release from the top surface of the salt was measured experimentally). The red-shaded regions identify the irradiation periods.

heating and insulation reduced the permeability of the system. The iteration of BABY reported in this work uses alumina as thermal insulation, which is very porous and is not expected to inhibit permeation. However, future configurations of BABY will attempt to experimentally clarify how much tritium permeates through the walls.

Second, a better characterization of the neutron sources is planned in order to better understand the spatial distribution

of the neutron flux and energy spectrum. This will greatly increase the fidelity of the OpenMC model.

The tritium speciation observed (see section 3.3) does not agree with what was observed in FLiBe or FLiNaK with pure He as a sweep gas [27, 31, 32]. In these experiments, very little to no insoluble tritium (HT, T2) compared to soluble forms of tritium (TF, HTO) was found. The different redox potentials of the salts are likely to be the reason for this discrepancy

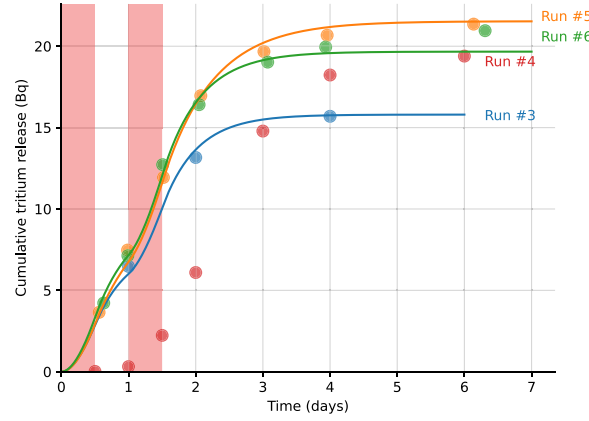


Figure 5. Comparison of the cumulative release measured in runs #3, #4, #5 and #6. The red-shaded regions identify the irradiation periods. Runs #1 and #2 were initial troubleshooting runs during which we did not detect tritium.

Table 2. Neutron rates measurements from activation foil analysis and TBR measurements. A-325 and P-383 refer to the sealed-tube neutron generators described in section 2.2.

Run	A-325	P-383	Neutron fluence	T production	TBR
1–3	Not measured	—	—	—	—
4	$1.02 \times 10^8 \text{ ns}^{-1}$	$2.64 \times 10^8 \text{ ns}^{-1}$	$3.17 \times 10^{13} \text{ n}$	$1.08 \times 10^{10} \text{ T}$	3.42×10^{-4}
5	$1.16 \times 10^8 \text{ ns}^{-1}$	$2.71 \times 10^8 \text{ ns}^{-1}$	$3.35 \times 10^{13} \text{ n}$	$1.19 \times 10^{10} \text{ T}$	3.57×10^{-4}
6	$1.06 \times 10^8 \text{ ns}^{-1}$	$2.48 \times 10^8 \text{ ns}^{-1}$	$3.06 \times 10^{13} \text{ n}$	$1.17 \times 10^{10} \text{ T}$	3.83×10^{-4}

and experiments are planned to confirm it. The presence of H2 impurities could also potentially lead to the formation of HT by isotopic exchange.

5. Methods

5.1. Tritium collection and LSC counting

Each vial contains 10 ml of deionized water. When sampling, the tritiated water of each vial is mixed with 10 ml of liquid scintillation cocktail (Ultima Gold™ LLT) and transferred to the LSC equipment (model PerkinElmer Tri-Carb 5110 TR) and the total tritium activity in the sample is measured with a counting time of 2 h. The file for the LSC method is also available as supplementary data.

The collection efficiency of the bubbling system η (see figure 1(b)) is calculated by:

$$\eta = 1 - (1 - \eta_{\text{vial}})^{N_{\text{vials}}} \quad (13)$$

where N_{vials} is the number of vials in series (here there are two vials on each side of the conversion furnace so $N_{\text{vials}} = 2$), and η_{vial} is the collection efficiency of one vial. The latter is calculated by the ratio:

$$\eta_{\text{vial}} = \frac{A_{\text{vial } 3}}{A_{\text{vial } 3} + A_{\text{vial } 4}} \quad (14)$$

where $A_{\text{vial } i}$ is the activity measured by vial i . We measured these efficiency for each sampling time and average them: $\eta_{\text{vial}} \approx 90\%$ and $\eta > 99\%$.

5.2. Activation foil analysis

Niobium activation foils from Shieldwrx were placed in the same plane as shown in figure 3(a) at the top of each neutron generator for runs 4, 5, and 6. After each run, the foils were placed between two $100 \times 100 \times 100$ mm cubic sodium iodide (NaI) scintillation detectors equipped with photo-multiplier tubes (PMTs) and connected to CAEN pulse digitization hardware. The pulses from the counting setup were collected using the ADAQAcquisition data acquisition software. During exposure to the flux of neutrons from the generators, activation occurs through the $^{93}\text{Nb}(n, 2n)^{92\text{m}}\text{Nb}$ reaction. The de-excitation of $^{92\text{m}}\text{Nb}$ results in the emission of gamma rays with energy of 934.4 keV with a half-life of 10.25 days [33]. The efficiency of detecting 934.4 keV photons was inferred from the measurement of the intrinsic efficiency of the detectors with 661.7 keV photons from ^{137}Cs decay and 511 keV and 1274.5 keV photons from ^{22}Na decay.

The time history of the activation foil from irradiation through the final count of gammas emitted from $^{92\text{m}}\text{Nb}$ are used to calculate the average flux of neutrons the foil experienced during the two irradiation periods. Once the neutron flux in $\text{n s}^{-1} \text{ cm}^{-2}$ is calculated, we assume an isotropic distribution to obtain the neutron rate in n s^{-1} based on the position of the foil. The error on the neutron fluence measurement from activation foils is computed from the error on the foil mass measurement, the counts error, the error on the intrinsic efficiency as well as the error on the geometric efficiency of the detector.

Complete description of the post-process of the activation foil analysis is available as supplementary data [22].

5.3. Diamond detector

The Cividec diamond detector (Cx-L Spectroscopic Amplifier) was placed approximately 35 cm away from the center of the crucible and perpendicular to the target plane of both neutron generators. Additionally, it was also placed approximately level with the radial center of the A-325 neutron generator. The detector held a bias voltage of 400 V, as recommended by the manufacturer.

Detector pulses arising from incident neutrons reacting with the diamond lattice were counted using the CAEN CoMPASS software, thus providing a time-dependent count rate, proportional to the sum of the neutron rates from the two generators.

5.4. Neutronics OpenMC model

The (n,Xt) reactions were tallied in the salt region. The neutron generators were modeled as point sources placed on the tritium-based target position. The energy and angular distribution were modeled based on a characterization carried out on the A-325 generator, assuming that the P-383 had similar emission characteristics. The Fusion Evaluated Nuclear Data Library (FENDL-3.1d) library was used for cross sections data. OpenMC v0.14.0 was used. The complete materials composition in the model can be found as supplementary data. Particular attention has been given to the modeling of the neutron sources. The angular and energy distributions of the neutron source A-325 were previously characterized using a diamond detector. We model it as a point source with energy and azimuthal angular distributions based on measurements, assuming axial symmetry in the polar direction. The same distributions are assumed for the P-383 source.

5.5. Geometry

The detailed geometry is available as supplementary data.

6. Conclusion

The BABY experiment represents a significant step forward in the quest for tritium self-sufficiency in FPPs. Despite its preliminary nature, the findings offer valuable insights into the intricate dynamics of tritium breeding within molten salts. A TBR of 3.57×10^{-4} was measured: 21 Bq of tritium produced and a neutron fluence of 3.35×10^{13} n over 24 h of irradiation. In addition to being a stepping stone towards demonstrating high TBR, this provides experimental data to validate numerical models (neutronics, tritium transport, fluid models,...) and better inform future FPP designs.

Future endeavors will be dedicated to reconciling disparities between observed and modeled TBRs, refining neutron source characterization, and better understanding of the tritium speciation observations. Moreover, an effort will be made to develop and validate higher-fidelity tritium transport models. This project will progress towards larger-scale investigations, such as the planned 11 volume experiment with FLiBe salt,

and will continue to explore alternative beryllium-free molten salt breeders.


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