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# Reduced order models in reactor kinetics: A comparison between point kinetics and multipoint kinetics



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#### ABSTRACT

Due to the heavy computational burden of full reactor kinetics modeling, reduced-order models are usually employed to simulate transients. Among those, point and multipoint kinetics have small computation time and provide satisfying results for many applications. We implemented point and multipoint kinetics (Avery's and Kobayashi's models) in the APOLLO3 code. Then these models are applied to study two simple transients, one in a coupled fast-thermal configuration, the other in a fast reactor. As at present we focus on the neutronic response only (neutron and precursor populations), the study is limited to step-change transients with no thermal feedback. This work permits to better delineate the potential of these methods and opens interesting perspectives. As an example, multipoint kinetics allows accounting for very fast shape transients that, depending on the coupling of the system, may result in global population changes occurring before the conventional prompt jump and altering significantly its quantitative value.

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# 1. Introduction

In this paper, we investigate the main reduced-order approaches to transient analysis in reactor physics. In particular we focus on point kinetics (hereafter PK) (Henry, 1958) and multipoint kinetics (hereafter MPK) (Dulla and Picca, 2006; Avery, 1958; Kobayashi, 1992). MPK is an advanced zero-dimensional modeling technique that allows to subdivide the reactor in different regions and still consider them as point-like but coupled with each other. In this way we allow for a relative degree of freedom among them and the neutron population is not forced to evolve rigidly. Several versions of MPK exist (Dulla and Picca, 2006) but in this paper, we focus on two of them, the one proposed by Avery (1958) and the one proposed by Kobayashi (Kobayashi, 1992). Aside from the theoretical motivation (Avery, 1958; Kobayashi, 1992), very few applications of MPK can be found in literature (Dulla and Picca, 2006; Tommasi and Palmiotti, 2019) and its interest over PK remains unclear. The purpose of this paper is to stress the differences between those two techniques to better characterize their descriptive capabilities. The novelty of this study is a more precise analysis of behaviors that MPK is able to model but that PK can't reproduce; this is particularly important since some of those behavior can be

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associated with important changes in the total neutron populations. Thanks to our developments in the neutronic platform APOLLO3 (Schneider et al., 2016) we were able to apply these modeling techniques to two transients. In order to focus on neutronics and on the kinetic response of the systems, the two transients are chosen to be step changes without thermal feedback. In Section 2 we briefly describe the various models from the mathematical point of view, trying to focus on their differences; the purpose is to provide an overview of the methods and reference notations. For more complete expositions the reader can consult (Henry, 1958; Avery, 1958; Kobayashi, 1992; Tommasi and Palmiotti, 2019). To stress the differences between the two modeling approaches we designed a 1D model that, despite being geometrically simple and allowing for a fast calculation of needed quantities, has non-homogeneous properties and permits to test the limit of PK. This model is detailed in Section 3. In Section 4 we present some results about an early version of the ASTRID reactor (Varaine et al., 2012) that were already obtained in Tommasi and Palmiotti (2019) using the code ECCO/ERANOS (Ruggieri et al., 2006) but that we recompute using the neutronic platform APOLLO3 (Schneider et al., 2016) and a more accurate  $S_N$  solver (MINARET Moller et al., 2011). Finally, we present our conclusions and perspectives about these methods based on the experience acquired so far.

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#### 2. Mathematical description and notation

The purpose of this section is to fix the notation and give a global overview of the models. Rigorously the neutron flux in a nuclear reactor can be described by the Boltzmann equation:

$$\begin{cases} \frac{1}{v} \frac{\partial \psi}{\partial t} + \mathscr{A}\psi = \mathscr{F}_{p}\psi + \sum_{i} \frac{\chi_{i}}{4\pi} \lambda_{i} C_{i} \\ \frac{\chi_{i}}{4\pi} \frac{dC_{i}}{dt} = \mathscr{F}_{di}\psi - \frac{\chi_{i}}{4\pi} \lambda_{i} C_{i} \end{cases}, \tag{1}$$

where  $\mathscr{F}$  is the fission production operator,  $\mathscr{A}$  is the operator containing the transport, removal and scattering term,  $\chi_i$  is the spectrum and  $\lambda_i$  the decay constant of the i-th delayed neutron family,  $\psi$  is the time dependent neutron flux and  $C_i$  is the concentration of precursors of the i-th family. When dealing with the fission operator the convention adopted in this paper is that  $\mathscr{F}$  without subscript refers to the total fission operator, with subscript p to the prompt one, with subscript di to the i-th delayed family and with subscript d to the total delayed operator.

This formulation of the Boltzmann equation, however, is used to study nuclear reactors that operate in time dependent condition. In the case of steady state conditions, the second equation in system (1) can be substituted in the first one and we have an homogeneous problem. To allow for a positive non zero solution we have to introduce an eigenvalue or, as is usually done in reactor physics, its reciprocal that is called k-effective ( $k_{\rm eff}$ ). The resulting equation is:

$$\left(\mathscr{A} - \frac{\mathscr{F}}{k_{\text{off}}}\right)\phi = 0. \tag{2}$$

At this point it is worth introducing also the adjoint formulation of Eq. (2) that is:

$$\left(\mathscr{A}^* - \frac{\mathscr{F}^*}{k_{\rm eff}}\right)\phi^* = 0, \tag{3}$$

where the superscript \* denotes the adjoint operator or flux. This adjoint flux can be interpreted as the importance of the neutrons relatively to their capability to participate in the chain reaction (Usachev et al., 1956).

A huge effort is devoted to the optimization of the steady state (or critical) calculation and the addition of the time variable leads to very expensive calculations for which significant simplifying assumptions are usually made. The most popular model used in reactor kinetics is PK, that is dealing with the reactor as if it were point-like. In this paper, we are going to analyze how this compares with two others reduced-order models that go under the name of MPK. The MPK approach also consists of dealing with a zero-dimensional model but, instead of using just one point as in the previous method, we split the reactor into different zones and we use a point for each one of them. The consequence of this modeling is that the ordinary differential equation describing the evolution of each region is coupled with the other ones. How to describe this coupling depends on which version of the MPK method we are using.

### 2.1. Point kinetics

The PK equations were proposed initially by Fermi by means of physical considerations (Fermi, 1942) but can be obtained rigorously introducing a factorization of the flux in an amplitude (scalar) and a shape (phase-space distribution), multiplying the time dependent Boltzmann equation by a weight function (usually the adjoint flux), integrating on all variables except time and imposing constraints over the time dependence of the flux shape (Henry, 1958). The equations that we obtain at the end are the following:

$$\begin{cases} \frac{dT}{dt} = \frac{\rho - \bar{\beta}}{\Lambda} T(t) + \sum_{k} \lambda_{k} \bar{c}_{k}(t) \\ \frac{d\bar{c}_{k}}{dt} = -\lambda_{k} \bar{c}_{k}(t) + \frac{\bar{\beta}_{k}}{\Lambda} T(t) \end{cases}, \tag{4}$$

where T is the total amplitude of the flux and the other parameters can be found in Table 1.

As we can see, Eq. (4) is a system of ordinary differential equations. From the computational point of view, the solution of these equations is not a critical task and this problem can be tackled with good accuracy and small computation time (Tommasi and Palmiotti, 2019; Ganapol, xxxx).

The main issue left is the computation of the kinetic parameters that can be obtained as in Table 1. In our convention,  $\langle . \rangle$  is the integration over the energy, space, and angular variables and  $\phi^*$  is the static adjoint flux at t = 0. Even though any weight function could be used, the standard practice is to use the adjoint flux due to its physical meaning (Usachev et al., 1956) and mathematical properties. No approximation would have been introduced if the forward flux used in the formulas in Table 1 were the time depended flux  $\psi$ as it appears in Eq. (1). However, since all the coefficients are obtained through a ratio of fluxes and therefore do not depend on the flux amplitude but just on its shape, the standard way to proceed, in order to avoid to compute the direct time dependent flux, is to use always the direct flux at time 0 and therefore to neglect any shape change (PK approximation). In conclusion, to implement this method it is enough to compute the critical forward and adjoint fluxes and use them to compute kinetic parameters. Thanks to the solver MINARET (Moller et al., 2011) we were already able to compute forward and adjoint fluxes but we had to implement a fast method to compute integrals and we had to introduce a delayed nuclear data model to properly define the needed operators (Valocchi et al., 2020). Once all the parameters were obtained the ordinary differential equations were solved thanks to scipy modules (Virtanen et al., xxxx).

#### 2.2. Multipoint kinetics

# 2.2.1. Avery's model

The first version of MPK has been proposed by Avery (1958) using mainly physical considerations. In this case, we define the partial direct flux and the partial adjoint flux of each region as follows:

$$\mathscr{A}\phi_{j} = \frac{1}{k_{eff}}\mathscr{F}_{j}\phi \qquad \qquad \mathscr{A}^{*}\phi_{j}^{*} = \frac{1}{k_{eff}}\mathscr{F}_{j}^{*}\phi^{*}, \tag{5}$$

where partial fluxes are defined over the whole reactor and the source term is obtained by restricting the multiplication operator to region j and by using the total flux computed previously (Eqs. (2) and (3)). In this case, if we split the reactor in n regions, we have  $n^2$  equations for the evolution of the neutron population and n times the number of precursor families equations for the precursor concentrations. The resulting system is the following

$$\begin{cases} l_{jk} \frac{dS_{jk}}{dt} = k_{jk} (1 - \beta_{jk}) \sum_{m} S_{km} - S_{jk} + \sum_{i} k_{jk,i}^{d} \lambda_{i} C_{ki} \\ \frac{dC_{ki}}{dt} = -\lambda_{i} C_{ki} + \beta_{ki} \sum_{m} S_{km} \end{cases},$$

$$(6)$$

**Table 1**Point kinetics parameters.

$ ho = rac{\langle \phi^*(\mathscr{F}-\mathscr{A})\phi angle}{\langle \phi^*\mathscr{F}\phi angle}$	$ar{\Lambda} = rac{\langle \phi^* rac{1}{p} \phi  angle}{\langle \phi^* \mathscr{F} \phi  angle}$	$\bar{\beta}_i = \frac{\langle \phi^* \mathscr{F}_{di} \phi \rangle}{\langle \phi^* \mathscr{F} \phi \rangle}$
$ar{eta} = \sum_i ar{eta}_i$	$ar{c}_i = rac{\langle \phi^* rac{\chi_d^d}{4\pi} C_i  angle}{\langle \phi^* rac{1}{v} \phi  angle}$	

**Table 2** Avery's MPK parameters.

$$\begin{aligned} k_{jk} &= \frac{\langle \mathcal{F}_{j} \phi \rangle \langle \phi^* \mathcal{F}_{jk} \phi \rangle}{\langle \mathcal{F}_{j} \phi \rangle \langle \phi^* \mathcal{F}_{jk} \phi \rangle} k_{eff} & k_{jk,i}^d &= \frac{\langle \mathcal{F}_{j} \phi \rangle \langle \phi^* \mathcal{F}_{ak} \phi \rangle}{\langle \mathcal{F}_{i} \phi \rangle} k_{eff} \\ l_{jk} &= \frac{\langle \phi^*_{j} \mathcal{F}_{j} \phi \rangle}{\langle \phi^*_{j} \mathcal{F}_{j} \phi \rangle} k_{eff} & S_{jk} &= \frac{\langle \mathcal{F}_{j} \phi \rangle \langle \phi^* \mathcal{F}_{ak} \phi \rangle}{\langle \phi^*_{j} \mathcal{F}_{j} \phi \rangle} \\ \beta_{jk} &= \frac{\langle \phi^*_{j} \mathcal{F}_{ak} \phi \rangle}{\langle \phi^*_{j} \mathcal{F}_{j} \phi \rangle} & \beta_{ki} &= \frac{\langle \mathcal{F}_{ak} \phi \rangle}{\langle \mathcal{F}_{k} \phi \rangle} & C_{ki} &= \frac{\langle \mathcal{F}_{ak}^d \mathcal{C}_{i} \rangle_{k}}{\langle \mathcal{F}_{ak} \phi \rangle} \end{aligned}$$

where each parameter can be expressed as in Table 2, again with the approximation that the time-dependent fluxes are replaced with the static ones of Eqs. (2) and (3).

In this model, the partial neutron populations  $S_{jk}$  represent neutrons produced in region j by neutrons born in region k and we have an equation for each of them.

#### 2.2.2. Kobayashi's model

Kobayashi's version of MPK has been derived more recently focusing more on a mathematical rigorous approach to the problem (Kobayashi, 1992). The resulting importances used as weight functions are closer to the Green function definition and are defined as follows:

$$\mathscr{A}^*G_i = \nu \Sigma_{fi},\tag{7}$$

where the partial importance here is computed by using as a source the total production cross section restricted to region j, and without using the adjoint flux.

The resulting system of equations is:

$$\begin{cases} I_j \frac{dS_j}{dt} = \sum_k k_{jk} (1 - \beta_{jk}) S_k - S_j + \sum_i \lambda_i \sum_k k_{ijk}^d C_{ki} \\ \frac{dC_{ki}}{dt} = -\lambda_i C_{ki} + \beta_{ik} S_k \end{cases},$$
(8)

where the unknown neutron populations  $S_j$  are the populations of each region independently from where the neutrons which generate them were coming. Therefore the resulting system has fewer equations for the population evolution (as many as the number of regions) but the same amount for the precursors concentrations. Parameters can be computed as in Table 3.

#### 2.3. Comments

The main differences among the methods described so far result from the weight functions selected. PK uses the adjoint flux that is a measure of the importance of neutrons relative to their capability to participate in the chain reaction (Usachev et al., 1956). Avery's MPK uses a partial version of forward and adjoint fluxes (Eq. (5)) that add to total the respective fluxes used in PK. Kobayashi's importances instead can be seen as the capability of the neutron to reproduce neutrons, but at the next generation only, not over the total chain reaction (Tommasi and Palmiotti, 2019). The calculation of these importance functions has been implemented in APOLLO3

We would like also to add that all the methods described so far can be expressed in the compact form:

$$\frac{dS}{dt} = \mathscr{M}S,\tag{9}$$

where the size of vector S and of matrix  $\mathcal{M}$  depends on the method we are using, in particular, if S is vector of length  $n, \mathcal{M}$  is a  $n \times n$ 

**Table 3** Kobayashi's MPK parameters.

$k_{jk}=rac{\langle G_j\mathscr{F}_k\phi angle}{\langle\mathscr{F}_k\phi angle}$	$k_{ijk}^d = rac{\langle G_j \mathscr{F}_{dik} \phi  angle}{\langle \mathscr{F}_{dik} \phi  angle}$	$l_j = rac{\langle G_j rac{1}{ u} \phi  angle}{\langle \mathscr{F}_j \phi  angle}$	$\beta_{jk} = \frac{\langle G_j \mathscr{F}_{dk} \phi \rangle}{\langle G_j \mathscr{F}_k \phi \rangle}$
$eta_{ij} = rac{\langle \mathscr{F}_{dij} \phi  angle}{\langle \mathscr{F}_{j} \phi  angle}$	$S_j = \langle \mathscr{F}_j \phi  angle$	$C_{ki} = \langle \frac{\chi_i^d}{4\pi} C_i \rangle_k$	

square matrix. In our case, since we are using the JEFF 3.1.1 nuclear data library, we use 8 precursor families (Rudstem et al., xxxx). Therefore for PK the S vector has 9 elements (8 for the precursor families and one for the neutron population). For Kobayashi's MPK, if the number of regions in which we are subdividing our reactor is *n*, the *S* vector has 9*n* elements, *n* for the neutron populations, and 8n for the precursors in each region. For Avery's MPK, S has (8+n)n elements,  $n^2$  for the neutron populations, and 8n for the precursors in each region. The study of the eigenvalues of *M* can provide useful information about the transient, as it is discussed in the next sections. Finally we would like to define the prompt sub-matrix as the square matrix that we obtain from the total one (M) by eliminating lines and columns concerning precursors. This matrix will be the one associated to a model in which precursors do not exist. To clarify this definition we make the example for the associated model in the Kobayashi's case:

$$l_j \frac{dS_j}{dt} = \sum_k k_{jk} (1 - \beta_{jk}) S_k - S_j. \tag{10}$$

The comparison between the eigenvalues of the prompt submatrix and of the total one can help us to identify the source of the different behaviors.

# 3. 1D Model

The first system we choose to model is a simplified 1D model of a coupled thermal-fast reactor. Although being an extremely simplified configuration, this model aims at having short computation time while retaining some heterogeneity that can highlight the differences among the modeling techniques presented so far. The description of this system can be found in (Ros, 2017) and a synthetic description of the materials and its properties can be found in (Ros et al., 2016). Originally it was described by an RZ geometry but we preferred to adopt a 1D geometry since it is simpler and can be handled by our solver (MINARET (Moller et al., 2011). The core is divided into 6 regions as can be seen in Fig. 1 where reflection condition is applied to the left boundary and vacuum condition to the right one; from left to right there is a central fast zone (Fast, F), a filter to absorb thermal neutron (Absorber, A), a natural uranium transition zone (Buffer, B), an enriched uranium plate (Converter, C), a thermal zone (Thermal, T) and a water reflector

The material used for the fast zone is the cell used in the ERMINE V program of the MINERVE experimental reactor, in the center of the ZONA-1 core (Martin-Deidier, 1979). This cell is composed of 6 MOx rodlets, 2 natural UOx rodlets, and 8 sodium rodlets, combined in a 2 inch square. The filter to absorb thermal neutron is made up of natural  $B_4C$  rodlets and the natural uranium in the transition zone is in the form of  $UO_2$  rodlets. The  $UO_2$  rodlets in region C are enriched at 30% in  $^{235}U$  and the  $UO_2$  in the thermal zone has 3.7% enrichment. Water is present only in the thermal zone and in the reflector. All the materials have been homogenized by ECCO/ERANOS (Ruggieri et al., 2006) and provided to APOLLO3® using a 33 groups energy mesh (more detail can be found in Ros (2017) and Ros et al. (2016)).

The basic idea behind this configuration is that the absorber, the buffer and the converter help to decouple the fast zone from the thermal one, being these the two zones where most of the fissions take place.

We can then introduce a perturbation, removing the absorber and the converter. In this way, we change the kinetic parameters of the reactor keeping the reactivity insertion positive but below prompt criticality ( $\beta$  = 586 pcm and  $\rho$  = 284 pcm). In order to do so we artificially replace the material in the converter zone

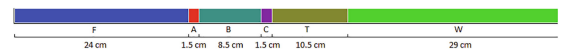


Fig. 1. 1D model of a coupled fast-termal reactor.

(enriched uranium) with the one in the buffer zone (natural uranium) and remove 92% of the <sup>10</sup>B in the absorber zone.

In order to keep the transient simple and to focus on the neutronics, we simulate an instantaneous change without considering thermal feedback. The system is supposed to be at t=0 in steady state in the nominal configuration and to shift instantaneously to the perturbed configuration. In order to have a steady state system at t=0, we divide all the coupling matrices and generation times by the k-effective of the nominal configuration, that is 1.11706.

#### 3.1. Results

In this section, we present five sets of results. One set is obtained by applying PK, and four are obtained by applying MPK. For MPK we have 2 sets using four regions, one for each homogeneous fissile zone, that are F, B, C and T in Fig. 1, both with Kobayashi's and with Avery's models, and two sets using 6 regions, where we split in half the 2 main fissile regions that are F and T.

As an example, we present the main kinetic parameters and coupling matrices we obtained for the nominal case. The  $k_{eff}$  and  $k_{ii}$  are listed in Table 4, mean generation time and relative MPK versions in Table 5 and beta effective and  $\beta_{ii}$  in Table 6. A first observation can immediately be made on the data presented in Table 4, the first of the four eigenvalues of both  $k_{ii}$  matrices is equal to  $k_{eff}$ , as it should be by construction. This is a first verification of our implementation and shows how, in MPK,  $k_{ij}$  has a role similar to the one that  $k_{eff}$  has in PK. Regarding the data in Table 5 they represent the mean neutron generation life time. These data are mainly sensitive to the neutron spectrum and show how the spectrum gets softer moving towards the thermal zone (last line). We can also observe that, looking at the Avery's  $l_{ii}$ , discriminating the various regions where neutrons are generated (different columns represent different departing regions) has limited effects with respect to the importance that has the region where the neutrons end (different lines represent different ending regions). Finally looking at data in Table 6 we can see how the delayed neutron fraction has a very heterogeneous distribution, mainly depending on where neutrons are produced (therefore presenting similar values along each column). We can also notice, in the second line, how the neutrons ending in the natural uranium zone (zone B in Fig. 1) have considerably smaller effective delayed fraction with respect to the neutrons born from the same region but ending in a different one. This can be explained thanks to the <sup>238</sup>U fission threshold that prevents most of the delayed neutrons from any region (that have a softer spectrum compared to the prompt ones) to participate in the chain reaction by inducing fission in the buffer region containing mainly <sup>238</sup>U as fissile nuclide.

**Table 4**  $k_{eff}$  and  $k_{ij}$  of the nominal configuration (before division by  $k_{eff}$ ).

PK (k <sub>eff</sub> )	1.11706
Kobayashi $(k_{ij})$	[0.88923     0.29334     0.15291     0.05134       [0.03203     0.12125     0.07372     0.02288       [0.12134     0.28680     0.38487     0.19834       [0.16048     0.41637     0.56683     0.77045
Avery $(k_{ij})$	0.89357     0.28779     0.15006     0.05032       0.03211     0.12127     0.07364     0.02286       0.12139     0.28695     0.38502     0.19824       0.16864     0.43211     0.58580     0.75710

**Table 5**  $\Lambda$ ,  $l_i$  and  $l_{ii}$  of the nominal configuration (before division by  $k_{eff}$ ).

PK (Λ) [ <i>μs</i> ]	9.9915					
Kobayashi ( $l_j$ ) [ $\mu$ s]	0.3125 0.4241 3.7227 28.232					
Avery $(l_{ij})$ $[\mu s]$	$\begin{bmatrix} 0.2465 & 0.3491 & 0.3935 & 0.4495 \\ 0.2938 & 0.1370 & 0.2646 & 0.6829 \\ 2.8120 & 2.5776 & 2.4007 & 4.2755 \\ 17.8861 & 18.8878 & 19.1796 & 25.4637 \end{bmatrix}$					

**Table 6**  $\beta_{\text{eff}}$  and  $\beta_{ii}$  of the nominal configuration.

PK (β <sub>eff</sub> ) [pcm]	586.9			
Kobayashi $(eta_{ij})$ [pcm]		1097.1 289.9 1486.3 1371.8	170.0 776.3	
Avery $(\beta_{ij})$ [pcm]	332.7 119.2 339.8 307.7		496.7 170.0 776.2 738.7	298.4 194.3 714.7 809.4

The first quantity on which we would like to focus on is the total neutron population evolution during the transient that can be seen in Fig. 2.

On a short time scale, as we can notice, there is a huge difference between the results obtained with PK and those obtained with MPK, while all the MPK cases provide similar results. In particular, all the MPK curves have a bump before  $10^{-5}$  s that is completely absent in the PK one. After that, there is the classical prompt jump that, however, in the MPK cases has a bigger amplitude. Looking at the differences among the various MPK, subdividing the main regions has virtually no effect on the Avery modeling and leads to a slight increase of the total neutron population modeled by Kobayashi MPK. Compared to Avery, Kobayashi has a slightly bigger population before the prompt jump and slightly

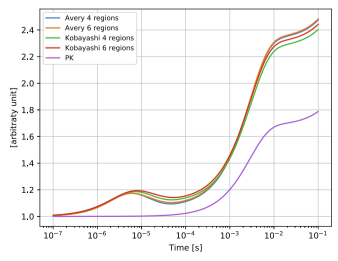


Fig. 2. Total neutron population evolution - 1D model.

smaller afterwards, although all these are minor differences and can be due to the various approximations made during the modeling.

Looking at Fig. 3 we can see the relative distribution of the neutron population in the various regions. We can notice how the redistribution of the neutron population takes place at the same time as the first bump in the total population evolution and is almost complete before the prompt jump. We can also notice that no major difference appears between Kobayashi's and Avery's models.

Concerning the long time scale (Fig. 4), all models present a coherent exponential behavior with Kobayashi's model having a slightly different time constant with respect to the other results and that changes slightly with the further subdivision of the main fissile zones. This effect, even though of second order, could be due to the approximations made (e.g. the use of the k-modes instead of the alpha-modes) and should be further investigated.

To conclude we can look at the eigenvalues of the matrix  $\mathcal{M}$  (Eq. (9)) in the different models, plotted in Fig. 5 where on the horizontal axis we have the real part of the eigenvalues on a symmetric log scale. We have to remark that all the eigenvalues are real, except for Avery's MPK with 6 regions that has a few complex eigenvalues; in all the cases however the real part is much larger than the imaginary part meaning that the damping effect is dominant with respect to the oscillating behavior and therefore we focus only on real parts.

In the first line we plotted the decay constant of each precursor family, so to show how some of the various eigenvalues cluster around the eight  $(-\lambda_i)$  with a multiplicity equal to the region number (as already discussed by Henry (Henry, 1964). As expected, in all the cases, we have one positive eigenvalue with similar value for all the methods (with the biggest discrepancy being between Avery's and Kobayashi's models of about 7%) Then, for MPK, we have the appearance of several eigenvalues all on a very short timescale. For all the MPK models, if we compare these eigenvalues (those  $<-10^2~s^{-1}$ ) with the ones of the relative prompt submatrix, we notice that we obtain the same eigenvalues (maximum relative error below  $10^{-8}$ ), except for the one relative to the prompt jump (around  $-5 \cdot 10^2~s^{-1}$ ) for which we have, in all the models, a relative error of 0.3%. This shows how the bump noticed in Fig. (2) is mainly due to the prompt behavior of the system.

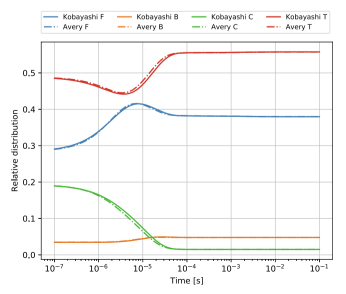


Fig. 3. Relative neutron population evolution - 1D model.

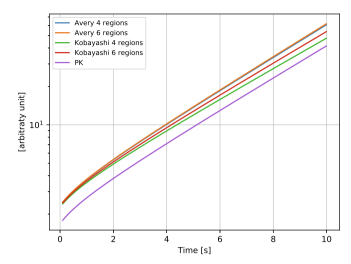


Fig. 4. Asymptotic neutron population behavior – 1D model.

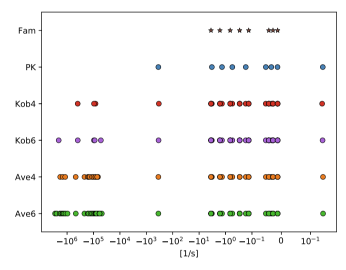


Fig. 5. Eigenvalues of the 1D model.

#### 4. ASTRID

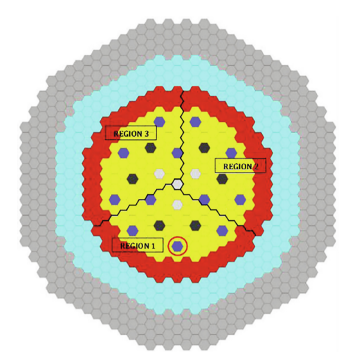
The second system we modeled is an early sodium-cooled fast reactor ASTRID CFV core design (Varaine et al., 2012) shown in Fig. 6. The core has a  $2\pi/3$  rotational symmetry and for the application of the MPK we chose symmetric regions as shown in Fig. 6. The transient we analyzed is a step change where the red-circled rod in region 1 is instantaneously removed from the core. The same transient has been analyzed in Tommasi and Palmiotti (2019) with the ECCO/ERANOS code (Ruggieri et al., 2006).

Also here we are below prompt criticality ( $\beta$  = 369 pcm and  $\rho$  = 349 pcm) but with much smaller margin.

#### 4.1. Results

In this case, we present three sets of results, one for the PK and two for the MPK, one for Avery's model and one for Kobayashi's model. The total population evolution is plotted in Fig. 7. Here, as opposed to what happens in the previous case, there is no major difference between the various modeling techniques for what concerns the total population.

However, if we look at the population repartition in Fig. 8 we can see in the MPK models quick changes in the neutron population distribution taking place in the same timescale as the previous



**Fig. 6.** ASTRID CFV-V1 – Color code: Yellow: inner fuel – Red: outer fuel – Light blue: reflector – Grey: shielding – Dark blue and black: control rods – White: inert sub-assemblies. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

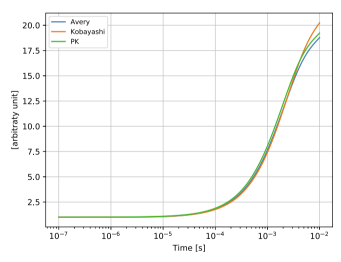


Fig. 7. Total neutron population evolution – ASTRID model.

case and before the prompt jump. The main difference is that, in this case, the redistribution of the neutron population is not associated with a change in the total amplitude.

For what concerns the asymptotic behavior (Fig. 9) no major differences exist among the various models, with existing discrepancies that, also in this case, we think can be explained by the different approximations we did.

Looking at the eigenvalues of the matrix  $\mathcal{M}$  (Eq. (9)) in Fig. 10 we can see how, also in this case, some eigenvalues cluster around the opposites of the decay constants of precursor families. We notice also the single positive eigenvalue with similar value for all the models (with the biggest discrepancy being between PK and Kobayashi's MPK of about 8%) and that MPK presents a set of smaller eigenvalues enabling shape changes on a very short time-scale. Also here, for all the MPK models, if we compare these eigen-

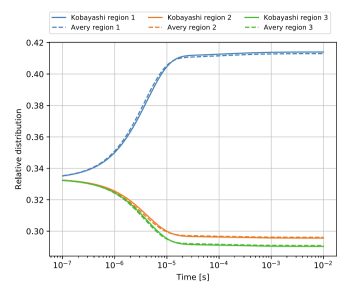


Fig. 8. Relative neutron population evolution – ASTRID model.

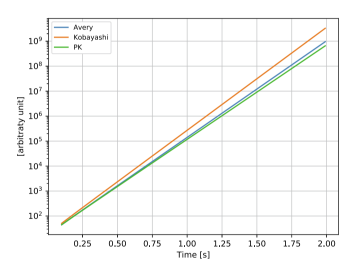


Fig. 9. Asymptotic neutron population behavior – ASTRID model.

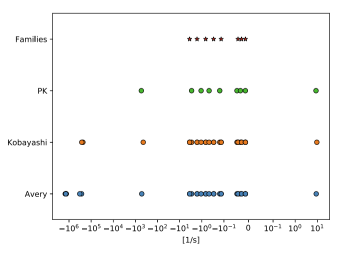


Fig. 10. Eigenvalues of the ASTRID model.

values (those  $<-10^2~s^{-1}$ ) with the ones from the one of the prompt sub-matrix we notice that we get the same eigenvalues (maximum relative error below  $10^{-8}$ ), except for the eigenvalue around  $-5\cdot 10^2~s^{-1}$  for which we have a relative error of 2%. This seems to show that, as the reactor approach prompt criticality, the leading prompt eigenvalue is more influenced by the total system and we have higher deviation from the one of the prompt submatrix alone.

#### 5. Conclusions and perspectives

The discussion of the results presented shows that in the cases considered, there are minor differences between Avery's and Kobayashi's formalisms. We also show how a further subdivision of homogeneous zones, in the cases considered, has a small impact on the overall description of the system. However, in the heterogeneous case, MPK is able to model total flux changes that PK is not able to reproduce and therefore MPK constitutes an important improvement in transient modeling. These results seem to show that, concerning the region subdivision in MPK, it is important to separate regions with different kinetic proprieties but further subdivision of homogeneous regions has a minor impact.

We also noticed that, after a step perturbation, the repartition of the neutron distribution anticipates the prompt jump and the following exponential behavior of the total amplitude.

While in this work we focused on the characterization of PK and MPK, limiting our analysis on the capabilities of the two methods, further work should be done to validate the results presented so far; this can be done by comparison against direct kinetic calculation, either deterministic or Monte Carlo, or against experimental data.

As perspectives, we think it is interesting to further investigate the evolution of the eigenvalues with respect of different criticality situations, including more accurate analyses around prompt criticality. We also think it is interesting to reproduce these results using fluxes different from the ones obtained using the k-modes from Eqs. (2) and (3) [e.g. alpha modes (temporal eigenvectors) or even dynamic fluxes if MPK is included in a quasi-static space–time kinetic scheme] that could be more representative of the analyzed cases. Finally we would like to combine these results with the work done Caron et al. (2017) to predict the flux shape evolution and to optimize shape calculation time step during transients.

#### **CRediT authorship contribution statement**

**G. Valocchi:** Investigation, Software, Validation, Writing - original draft. **J. Tommasi:** Conceptualization, Supervision, Writing - review & editing. **P. Ravetto:** Conceptualization, Supervision, Writing - review & editing.

# **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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