

Fast Reactor and FEM William Christopher Dawn

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Conclusions

# Simulation of Fast Reactors with the Finite Element Method and Multiphysics Models

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### Disclaimer

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# Why are we here?

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# Why are we here?

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Model a nuclear reactor.



### Why are we here?

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# Model a nuclear reactor.

- Neutron distribution.
- Thermal hydraulics.
- Thermal expansion.



### **Current Simulation Procedure**

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- Heuristically estimate material temperatures.
- Manually calculate thermally expanded dimensions.
- Manually homogenize assembly number densities.
- Run DIF3D and collect  $k_{eff}$  and power distribution.



### **Current Simulation Procedure**

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- Heuristically estimate material temperatures.
- Manually calculate thermally expanded dimensions.
- Manually homogenize assembly number densities.
- Run DIF3D and collect  $k_{eff}$  and power distribution.

No thermal feedback or multiphysics simulation capability. Modern numerical methods can be implemented.



### Goals

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- Easy user input with intuitive keywords.
  - ► Reactor geometry via VTK mesh.
  - Temperature dependent cross sections either plain-text or ISOTXS format.
  - ► Pin and assembly dimensions.
  - Material compositions.
- Simulate thermal expansion and thermal hydraulics internally.
- Collect k<sub>eff</sub>, reactor power distribution, and average material temperatures.



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# Multigroup Neutron Diffusion Equation

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$$\begin{split} &-\nabla\cdot(D_g(\mathbf{r})\nabla\phi_g(\mathbf{r}))+\Sigma_{r,g}(\mathbf{r})\phi_g(\mathbf{r})=\\ &\frac{\widetilde{\chi_g}(\mathbf{r})}{k_{eff}}\sum_{g'=1}^G \nu\Sigma_{f,g'}(\mathbf{r})\phi_{g'}(\mathbf{r})+\sum_{\substack{g'=1\\g'\neq g}}^G \Sigma_{s,g'\to g}(\mathbf{r})\phi_{g'}(\mathbf{r}) \end{split}$$

$$D_g(\mathbf{r})$$
 = diffusion coefficient for energy group  $g$  [cm],

$$\phi_g(\mathbf{r})$$
 = scalar neutron flux for energy group  $g\left[\frac{1}{\mathrm{cm}^2 \mathrm{s}}\right]$ ,

$$\Sigma_{r,g}(\mathbf{r}) = \text{macroscopic removal cross section for energy group } g \left[\frac{1}{\text{cm}}\right],$$
 $\widetilde{\chi}_g(\mathbf{r}) = \text{effective fission spectrum for energy group } g,$ 

$$k_{eff}$$
 = effective neutron multiplication factor,

$$\nu \tilde{\Sigma}_{f,g}(\mathbf{r})$$
 = number of fission neutrons times macroscopic fission cross section in energy group  $g\left[\frac{1}{cm}\right]$ ,

$$\Sigma_{s,g'\to g}(\mathbf{r}) = \text{macroscopic scatter cross section from energy group } g' \text{ to}$$
 energy group  $g \left[ \frac{1}{cm} \right]$ ,

$$G$$
 = total number of energy groups (typically  $G = 33$ ).

# **Boundary Conditions**

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For problem domain  $\Omega$  and boundary  $\partial \Omega$ .  $\hat{\mathbf{n}}$  is the outward normal direction on the boundary.

Mirror.

$$\nabla \phi_g(\mathbf{r}) \cdot \hat{\mathbf{n}} = 0 \text{ for } \mathbf{r} \in \partial \Omega$$

Albedo.

$$D_g(\mathbf{r})\nabla\phi_g(\mathbf{r})\cdot\hat{\mathbf{n}} + \alpha\phi_g(\mathbf{r}) = 0 \text{ for } \mathbf{r} \in \partial\Omega$$

 $\alpha \in \mathbb{R}$  is a scalar constant specified by the user. For non-reentrant (vacuum) boundary condition,  $\alpha = \frac{1}{2}$ .

3 Zero Flux.

$$\phi_g(\mathbf{r}) = 0 \text{ for } \mathbf{r} \in \partial \Omega$$



### Finite Element Method (FEM) Discretization

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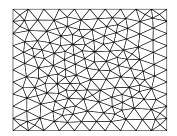
Conclusions

Divide the domain  $\Omega$  into a set of unstructured, non-overlapping, finite elements (e.g. Delaunay triangulation).

$$\Omega = \Omega_1 \cup \Omega_2 \cup \Omega_3 \cup \ldots \cup \Omega_{N_E}$$

$$\Omega = \{\Omega_e\} \text{ for } e = 1, 2, \ldots, N_E$$

$$\Omega_i \cap \Omega_j = \emptyset \text{ for } i \neq j$$



Example Rectangular Mesh.

# **Combining Neutron Source**

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• Neutron sources are combined into a single term.

$$- \, \nabla \cdot (D_g(\mathbf{r}) \nabla \phi_g(\mathbf{r})) + \Sigma_{r,g}(\mathbf{r}) \phi_g(\mathbf{r}) = q_g(\mathbf{r})$$

$$q_g(\mathbf{r}) = q_{g,e} \ \forall \ \mathbf{r} \in \Omega_e$$

$$\overline{\phi}_{g,e} = \frac{1}{N_p} \sum_{i \in \Omega_e}^{N_p} \phi_{i,g}$$

- Neutron source  $q_{g,e}$  is constant over an element  $\Omega_e$ .
- Cross sections are constant within an element.

### Finite Element Method (FEM)

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Multiply the multigroup neutron diffusion equation by a testing function  $v(\mathbf{r}) \in H_1(\Omega)$  and integrate over the problem domain.  $H_1(\Omega)$  is a Sobolev space.

This yields the **Weak Form** of the problem.

$$-\int_{\Omega} \nabla \cdot (D_g(\mathbf{r}) \nabla \phi_g(\mathbf{r})) v(\mathbf{r}) \ d\mathbf{r} + \int_{\Omega} \Sigma_{r,g}(\mathbf{r}) \phi_g(\mathbf{r}) v(\mathbf{r}) \ d\mathbf{r} = \int_{\Omega} q_g(\mathbf{r}) v(\mathbf{r}) \ d\mathbf{r}$$

Partition the integral into a summation of integrals over elements.

$$-\sum_{e=1}^{N_E} D_{g,e} \int_{\Omega_e} \nabla \cdot \nabla \phi_g(\mathbf{r}) v(\mathbf{r}) \ d\mathbf{r} + \sum_{e=1}^{N_E} \Sigma_{r,g,e} \int_{\Omega_e} \phi_g(\mathbf{r}) v(\mathbf{r}) \ d\mathbf{r} = \sum_{e=1}^{N_E} q_{g,e} \int_{\Omega_e} v(\mathbf{r}) \ d\mathbf{r}$$

### Second Green's Theorem

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Use the Second Green's Theorem to rewrite the first integral [Li18].

$$\begin{split} -\sum_{e=1}^{N_E} D_{g,e} \int_{\partial \Omega_e} v(\mathbf{r}) \nabla \phi_g(\mathbf{r}) \cdot \hat{\mathbf{n}} \ ds + \sum_{e=1}^{N_E} D_{g,e} \int_{\Omega_e} \nabla \phi_g(\mathbf{r}) \cdot \nabla v(\mathbf{r}) \ d\mathbf{r} + \\ \sum_{e=1}^{N_E} \Sigma_{r,g,e} \int_{\Omega_e} \phi_g(\mathbf{r}) v(\mathbf{r}) \ d\mathbf{r} = \sum_{e=1}^{N_E} q_{g,e} \int_{\Omega_e} v(\mathbf{r}) \ d\mathbf{r} \end{split}$$

### Galerkin Finite Element Method (FEM)

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Galerkin FEM assumes the solution  $\phi_g(\mathbf{r})$  is a linear combination of chosen basis functions  $\{N_i\}$ .

$$\phi_g(\mathbf{r}) = \sum_{i=1}^{DOF} v_{g,i} N_i(\mathbf{r})$$

 $v(\mathbf{r}) \in H_1(\Omega)$  is arbitrary and is chosen to be a linear combination of the basis functions with unit magnitude.

$$v(\mathbf{r}) = \sum_{j=1}^{DOF} N_j(\mathbf{r})$$

Typically,  $N(\mathbf{r})$  is a polynomial of a chosen order (e.g. linear, quadratic, cubic).

# Linear System of Equations

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Including albedo form of boundary condition and assumption of linear combination of basis functions.

$$\sum_{i=1}^{DOF} v_{i,g} \sum_{j=1}^{DOF} \left( \sum_{e=1}^{N_E} \alpha \int_{\partial \Omega_e} N_i(\mathbf{r}) N_j(\mathbf{r}) ds + \sum_{e=1}^{N_E} D_{g,e} \int_{\Omega_e} \nabla N_i(\mathbf{r}) \cdot \nabla N_j(\mathbf{r}) d\mathbf{r} \right)$$
$$+ \sum_{e=1}^{N_E} \sum_{r,g,e} \int_{\Omega_e} N_i(\mathbf{r}) N_j(\mathbf{r}) d\mathbf{r} = \sum_{i=1}^{DOF} \left( \sum_{e=1}^{N_E} q_{g,e} \int_{\Omega_e} N_i(\mathbf{r}) d\mathbf{r} \right)$$

Rewriting in the form common to the FEM.

$$a_g(N_i, N_j) = f_g(N_i)$$

In the form common to linear systems.

$$\mathbf{A}_g \mathbf{u}_g = \mathbf{f}_g$$
$$\mathbf{u}_g = \{ \nu_{i,g} \}$$



# Properties of $\mathbf{A}_g \mathbf{u}_g = \mathbf{f}_g$

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- Properties of the linear system include:
  - ► Sparse.
  - ▶ Matrix,  $A_g$ , is Symmetric Positive Definite (SPD) [Hug87].
  - Solution,  $\mathbf{u}_g$ , is unique and bounded by Lax-Milgram Lemma [Li18].
- Solution via Conjugate Gradient (CG) method [Kel95].



# Integration

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### Integrals of interest:

$$\int_{\Omega_e} \nabla N_i(\mathbf{r}) \cdot \nabla N_j(\mathbf{r}) d\mathbf{r}$$

$$\int_{\Omega_e} N_i(\mathbf{r}) N_j(\mathbf{r}) d\mathbf{r}$$

$$\int_{\Omega_e} N_i(\mathbf{r}) d\mathbf{r}$$

$$\int_{\partial\Omega_e} N_i(\mathbf{r}) N_j(\mathbf{r}) ds$$

### Options for integration:

- Analytic.
- Numeric (quadrature).
  - Linear (Gaussian).
  - Triangular.



### **Triangular Elements**

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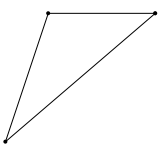
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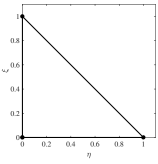
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General Triangle Element.



Reference Triangle.



# Wedge Elements

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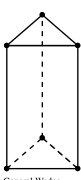
Finite Element Neutron Diffusion

Neutron Diffusion Results

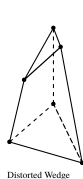
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Thermal Expansion

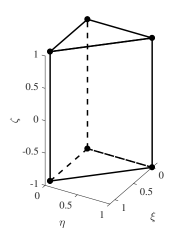
Coupled Results



General Wedge Element.



Distorted Wedge Element.



Description of Reference Wedge.



# RCM Matrix Ordering

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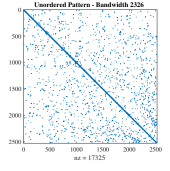
Results

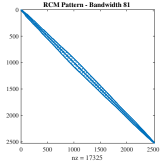
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- Matrix is reordered to increase computational efficiency and compute the same solution.
- Reverse Cuthill-McKee (RCM) order is chosen [Cut69].





### Power Iteration Method

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- Solution for largest eigenvalue  $k_{eff}$  and associated eigenvector  $\Phi$ .
- Rewrite the multigroup neutron diffusion equation.

$$\mathbf{B}(\Phi, k_{eff})\,\Phi = \frac{1}{k_{eff}}\mathbf{M}\,\Phi$$

The solution can be written.

$$\Phi = \frac{1}{k_{eff}} \mathbf{R} \, \Phi \quad \text{where} \quad \mathbf{R} = \mathbf{B}^{-1} \, \mathbf{M}$$

- Note: the FEM is used to calculate  $\Phi$ , not **R**.
- The power iteration method proceeds.

$$\Phi^{(s+1)} = \frac{1}{k_{eff}^{(s)}} \mathbf{R} \Phi^{(s)}$$

$$k_{eff}^{(s+1)} = k_{eff}^{(s)} \frac{\langle \mathbf{w}, \Phi^{(s+1)} \rangle}{\langle \mathbf{w}, \Phi^{(s)} \rangle} \qquad s = 1, 2, \dots, \infty$$



# Power Iteration Algorithm

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#### Algorithm 1 General Iteration Scheme

- 1: Read mesh from VTK.
- 2: Initialize  $\overline{\phi}_{\alpha}^{(0)}$ .
- 3: Order the nodes of the mesh into RCM order.
- 4: Calculate  $\Sigma_{s,g'\to g}$ ,  $\Sigma_{r,g}$ , and  $\nu\Sigma_{f,g}$  for each element.
- 5: Calculate finite element matrix  $A_g$  for each group. Store this.
  - 6: while Power Iteration do
    - Update the iteration counter. s = s + 1
  - 8: Update  $q_{fiss,g}$  and  $q_{up,g}$  for all groups from previous data  $\overline{\phi}^{(s-1)}$ .
  - Update  $\widetilde{\chi_g}$  in each element using previous data.
- 10: **for** g = 1, G **do**
- Update  $q_{down,g}$  from current data  $\overline{\phi}_g^{(s)}$
- 12: Calculate total source in each element.
- Update finite element Vector  $\mathbf{f}_g$  with new source.
- Solve  $\mathbf{A}_g \mathbf{u}_g = \mathbf{f}_g$  using an iterative technique (CG).
- Parse  $\mathbf{u}_g$  for  $\phi_g$  solution on nodes.
- 16: Calculate element-average  $\overline{\phi}_g$ .
- Update  $k_{eff}$ .
- 18: Check convergence.
- 19: Perform non-linear update if necessary and update  $\mathbf{A}_g$ .

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### Verification and Validation

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Coupled Results

- "Code Verification"
  - Compare computational results to exact analytic or manufactured results.
  - ▶ Demonstrate the code is solving equations correctly as designed.
  - Quantified numerical errors.
- "Solution Verification"
  - Compare computational results to benchmark results for the intended application of the solver.
  - Computational results from a different method or experimental data.
  - ► Typically verified by others previously.

# Error Analysis

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FEM with linear elements is second-order convergent in space [Li18].

$$\mathbf{e} = \phi(\mathbf{r}) - \phi_{FEM}$$
$$\|\mathbf{e}\|_{\infty} \le ch^2 \|\nabla^2 \phi(\mathbf{r})\|_{\infty}$$

Define Root-Mean-Squared (RMS), maximum, and  $k_{eff}$  errors.

$$\begin{aligned} \text{RMS}(\mathbf{e}) &= \sqrt{\frac{1}{N} \sum_{i=1}^{N} e_i^2} \\ \|\mathbf{e}\|_{\infty} &= \max_{i=1,2,\dots,N} |e_i| \\ k_{eff} \text{ error [pcm]} &= (k_{ref} - k_{eff}) \times 10^5 \end{aligned}$$

The method is second-order spatially convergent.

$$4 = \frac{e^{(i-1)}}{e^{(i)}}$$



### **Analytic Solutions**

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- 6 analytic multigroup neutron diffusion problems.
- Varied number of spatial dimensions, energy groups, and number of materials.

Case	Dimensions	Groups	Criticality	Materials
1	1	1		1
2	1	1	$\checkmark$	1
3	2	1	$\checkmark$	1
4	1	2	$\checkmark$	1
5	1	1	$\checkmark$	2
6	3	1	$\checkmark$	1



# Two-Dimension, One-Group, Criticality

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Refine	$k_{eff}$	k <sub>eff</sub> error [pcm]	$k_{e\!f\!f}$ ratio	RMS	RMS ratio	$\ \mathbf{e}\ _{\infty}$	$\ e\ _{\infty}$ ratio
0	1.983243	1281.65	4.03	1.90E-02	1.66	6.63E-02	1.49
1	1.992884	317.64	3.96	1.15E-02	2.65	4.45E-02	2.59
2	1.995258	80.16	3.98	4.32E-03	3.43	1.72E-02	3.41
3	1.995858	20.15	3.99	1.26E-03	3.88	5.04E-03	3.87
4	1.996009	5.05	4.00	3.25E-04	3.96	1.30E-03	3.96
5	1.996047	1.26	4.00	8.20E-05	3.93	3.28E-04	3.93
6	1.996057	0.32		2.09E-05		8.34E-05	
Ref.	1.996060						



# Two-Dimension, One-Group, Criticality

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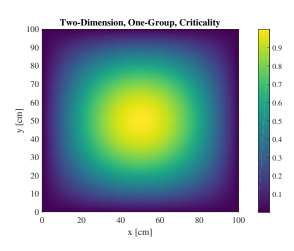
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$$\phi(x, y) = \phi_0 \sin\left(\frac{\pi}{L_x}x\right) \sin\left(\frac{\pi}{L_y}y\right)$$



### Three-Dimension, One-Group, Finite Cylinder

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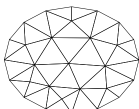
Coupled Results

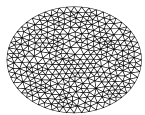
Conclusions

Refine	$k_{eff}$	k <sub>eff</sub> error [pcm]	$k_{eff}$ ratio	RMS	RMS ratio	$\ \mathbf{e}\ _{\infty}$	$\ e\ _{\!\scriptscriptstyle \infty}$ ratio
0	0.895108	10160.26	4.18	5.34E-02	2.57	2.12E-01	1.62
1	0.972412	2429.90	4.16	2.07E-02	3.19	1.31E-01	4.65
$2^{\dagger}$	0.990870	584.06	3.90	6.50E-03	1.85	2.81E-02	1.79
3	0.995215	149.61	3.99	3.51E-03	9.22	1.57E-02	8.28
4	0.996336	37.48		3.81E-04		1.90E-03	
Ref.	0.996711						

<sup>&</sup>lt;sup>†</sup> Refinement ratio ≈ 1 but next case ≈ 8.

This is due to the movement of mesh nodes in the process of circular mesh regeneration.





### Three-Dimension, One-Group, Finite Cylinder

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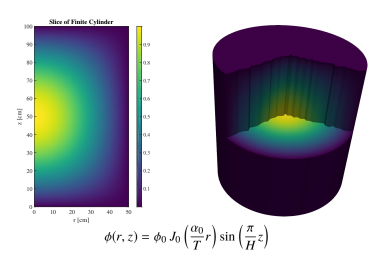
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### **Benchmark Solutions**

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• 9 benchmark problems.

- Two and three dimensional geometry.
- Varied energy group structure and neutron spectrum.

Benchmark	Dimensions	Groups	Reactor Type	Neutron Spectrum
VVER440	2	2	LWR	Thermal
SNR	2	4	SFR	Fast
HWR	2	2	HWR	Thermal
IAEA $(\times 4)$	2	2	PWR	Thermal
MONJU	3	3	SFR	Fast
KNK	3	4	SFR	Fast

### VVER440

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- Two-dimensional.
- Light Water Reactor (LWR).
- Two-group.

Refine	$k_{eff}$	k <sub>eff</sub> error [pcm]
0	1.005932	376.80
1	1.008980	72.00
2	1.009572	12.82
3	1.009666	3.35
4	1.009692	0.76
5	1.009698	0.22
Ref. <sup>†</sup>	1.009700	

<sup>†</sup> See [Cha95].



## VVER440 Benchmark Power Comparison

Fast Reactor and FEM

William Christopher Dawn

Introduction

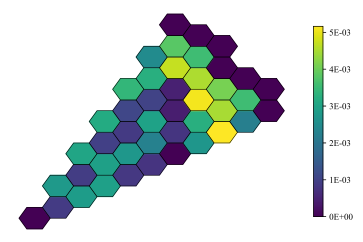
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VVER440 Benchmark Power Comparison for Most Refined Mesh.

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#### **MONJU**

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- Three-dimensional.
- Sodium-cooled Fast Reactor (SFR).
- Three-group.
- Case A. Control rods fully removed.
- Case B. Control rods partially inserted.
- Case C. Control rods fully inserted.

Pattern	$k_{eff}$	Rod Worth $[\Delta k]$	Rod Difference $[\%\Delta k]$	
A	1.056816			
В	1.031623	$0.023 (2.51E-5)^{\dagger}$	2.52 (-0.07)	
C	1.006519	0.047 (1.77E-3)	5.03 (0.04)	

<sup>&</sup>lt;sup>†</sup> Value in parentheses is difference to reference value [Kom78].

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## Assembly Geometry

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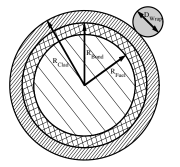
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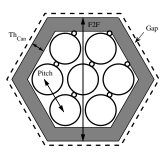
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## **Material Properties**

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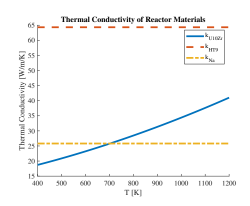
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- Functional sodium properties [Fin95].
- Clad and bond thermal conductivity assumed constant [Lei88].
- Fuel thermal conductivity assumed a function of temperature [Kim14].





#### Axial Convection Geometric Model

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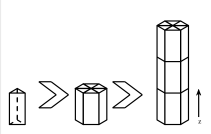
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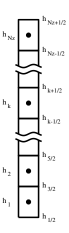
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Progression of Element (left), to Chunk (center), to Channel (right).



Nodalization for channel i.

# Channel Enthalpy

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Steady-state coolant enthalpy within the channel is given by an energy balance.

$$h_{i,k+1/2} = h_{in} + \frac{1}{\dot{m}_i} \sum_{k'=1}^k q_{i,k'}$$

Use a first-order approximation to estimate the chunk-average enthalpy.

$$h_{i,k} = \frac{1}{2}(h_{i,k-1/2} + h_{i,k+1/2})$$

 $T_{\infty,i,k}$  is then given by a state relationship [Fin95].

$$T_{\infty,i,k} = T(h_{i,k})$$



#### Radial Conduction Geometric Model

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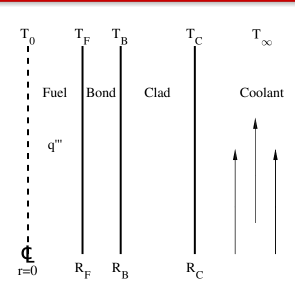
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# Clad Surface Temperature – Subbotin-Ushakov

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Using Newton's Law of Cooling.

$$q_{clad}^{\prime\prime}=H_c(T_C-T_\infty)$$

 $H_c$  is given by the Subbotin-Ushakov correlation [Pfr07] which relates the Nusselt and Péclet numbers for 1 < Pe < 4,000 and  $1.2 \le S/D \le 2.0$ .

$$Pe = Re Pr$$

$$Nu = 7.55 \frac{S}{D} - 20 \left(\frac{S}{D}\right)^{-13} + \frac{3.67}{90 \left(\frac{S}{D}\right)^2} Pe^{\left(0.56 + 0.19 \frac{S}{D}\right)}$$

$$H_c = \frac{Nu \, k}{D_c}$$

Then, the clad surface temperature,  $T_C$  follows.

## Fuel Centerline Temperature

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Conclusions

Define a conductivity integral.

$$K_F(T) = \int_0^T k_F(T') \ dT'$$

The value of the conductivity integral is given by the heat conduction equation.

$$K_F(T_0) = K_F(T_F) + \frac{q_{i,k}^{\prime\prime\prime}}{4}R_F^2$$

Then, a bisection method search is used to calculate  $T_0$  given a functional form of  $K_F(T)$ .

## Average Material Temperatures

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Average temperatures in the clad and bond are calculated analytically.

$$\overline{T_C} = T_B - \frac{q_{i,k}'''}{4k_C} R_F^2 \left( \frac{2R_C^2 \ln\left(\frac{R_C}{R_B}\right)}{R_C^2 - R_B^2} - 1 \right)$$

$$\overline{T_B} = T_F - \frac{q_{i,k}'''}{4k_B} R_F^2 \left( \frac{R_F^2 - R_B^2 + 2R_B^2 \ln\left(\frac{R_B}{R_F}\right)}{R_B^2 - R_F^2} \right)$$

# Average Fuel Temperature

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.

Calculate an effective thermal conductivity in the fuel.

$$\overline{k_F} = \frac{q_{i,k}^{\prime\prime\prime} \, R_F^2}{4(T_0 - T_F)}$$

Assume thermal conductivity is constant  $\overline{k_F}$ . Calculate an analytic value for the average fuel temperature.

$$\overline{T_F} = T_0 - \frac{q_{i,k}^{\prime\prime\prime}}{8\overline{k_F}}R_F^2$$

 $\overline{T_F}$  is used to calculate fuel cross sections.

Due to self-shielding, an effective fuel temperature would weight the surface temperature more.



## Radial Temperatures for Typical Fuel Rod

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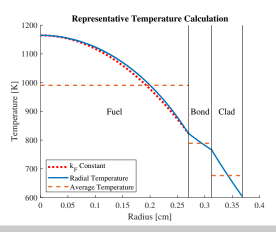
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Difference less than 15 [K].

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### Cross Section Treatment - Coolant & Bond

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• Number density and microscopic cross sections are functionalized and updated based on  $T_{\infty,i,k}$ .

Linear interpolation for microscopic cross sections.

Number density functionalization.

$$M_{Na} = 22.989769 \left[ \frac{\text{gram}}{\text{mol}} \right]$$

$$N_{Na}(T) = \frac{\rho_{Na}(T)\,N_A}{M_{Na}}$$

Microscopic cross section functionalization for  $T_n < T_{\infty,i,k} < T_{n+1}$ .

$$\Sigma_{x,i,k,g} = N_{Na}(T_{\infty,i,k}) \left( \frac{T_{\infty,i,k} - T_n}{T_{n+1} - T_n} (\sigma_{x,Na,g,n+1} - \sigma_{x,Na,g,n}) + \sigma_{x,Na,g,n} \right)$$

Bond is assumed to have the same macroscopic cross section as coolant. Consistent with homogenization approximation.

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#### Cross Section Treatment - Clad

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- Macroscopic cross section updated based on  $\overline{T_{C,i,k}}$ .
- Linear interpolation.

Macroscopic cross section functionalization for  $T_n < \overline{T_{C,i,k}} < T_{n+1}$ .

$$\Sigma_{x,i,k,g} = \frac{T_{C,i,k} - T_n}{T_{n+1} - T_n} (\Sigma_{x,g,n+1} - \Sigma_{x,g,n}) + \Sigma_{x,g,n}$$

#### Cross Section Treatment – Fuel

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- Macroscopic cross section update based on  $\overline{T_{F,i,k}}$ .
- Square-root interpolation due to Doppler effect.

Macroscopic cross section functionalization for  $T_n < \overline{T_{F,i,k}} < T_{n+1}$ .

$$\Sigma_{x,i,k,g} = \frac{\sqrt{\overline{T_{F,i,k}}} - \sqrt{T_n}}{\sqrt{T_{n+1}} - \sqrt{T_n}} (\Sigma_{x,g,n+1} - \Sigma_{x,g,n}) + \Sigma_{x,g,n}$$

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## Thermal Expansion Motivation

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- Strong feedback.
- Metallic fuels.
- Small active fuel region with high leakage ( $\mathcal{L} \approx 20\%$ ).
- Experimental Breeder Reactor II (EBR-II) designed and built by Argonne National Laboratory (ANL) [Til11].
  - ► Full-power demonstrations from April 1986 [Pla87].
  - Unprotected Loss-Of-Flow (ULOF).
  - ► Unprotected Loss-Of-Heat-Sink (ULOHS).



### **Material Properties**

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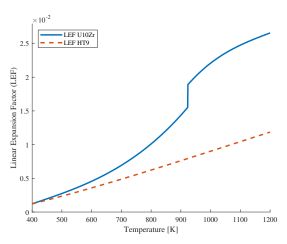
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Linear Expansion Factor for HT9 Steel and U10Zr Fuel.



# Simplified Thermal Expansion Model

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Coupled Results

- User input expansion temperatures  $T_{exp,fuel}$  and  $T_{exp,struct}$ .
- Leakage effects.
  - ► Finite Elements.
    - Radial (x and y) directions expanded as structural material, HT9 stainless steel.
    - Axial (z) direction expanded as fuel material, U10Zr.
  - Area fractions.
    - Fuel radius expanded as U10Zr.
    - All other material expanded as HT9 stainless steel.
- Density Effects.
  - ► Material densities decreased to conserve quantity of material.
  - Cross sections decrease proportionally according to  $\Sigma = N \sigma$ .

## Finite Element Expansion

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Define radial and axial expansion factors.

$$F_r(T_{exp,struct}) = 1 + \left(\frac{\Delta L}{L}\right)_{HT9}$$
  
 $F_a(T_{exp,fuel}) = 1 + \left(\frac{\Delta L}{L}\right)_{U10Zr}$ 

Expand all coordinates in the finite element mesh.

$$x^{H} = x^{C} F_{r}(T_{exp,struct})$$
$$y^{H} = y^{C} F_{r}(T_{exp,struct})$$
$$z^{H} = z^{C} F_{a}(T_{exp,fuel})$$

• Elements will not overlap or intersect due to uniform expansion assumptions.



## **Arbitrary Volume Expansion**

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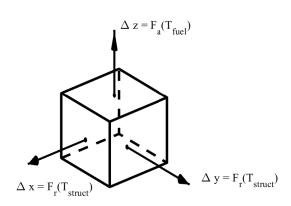
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$$\frac{V^C}{V^H} = \frac{1}{(F_r(T_{exp,struct}))^2(F_a(T_{exp,fuel}))}$$



## Area Fraction Expansion

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- Dimensions within a hexagonal assembly are expanded.
- Area fractions are used for cross section homogenization.
- Fuel radius,  $R_F$ , expanded as U10Zr.
- All other dimensions expanded as HT9 stainless steel.
- No general formula for expansion of area fractions, calculated directly.

# Conservation of Material & Cross Section Effects

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Conservation of number of atoms of species i.

$$n_i^H = n_i^C$$

Rewrite the number of atoms using number density and volume.

$$N_i^H \, V_i^H = N_i^C \, V_i^C$$

Volume  $V_i$  can be expressed using element volume and area fraction.

$$N_i^H = N_i^C \frac{a_j^C V_e^C}{a_i^H V_e^H}$$

Recall the volume ratio.

$$N_i^H = N_i^C \frac{a_j^C}{a_j^H} \frac{1}{(F_r(T_{exp,struct}))^2 F_a(T_{exp,fuel})}$$

Macroscopic cross sections can be updated directly.



## Demonstration of Reactor Thermal Expansion

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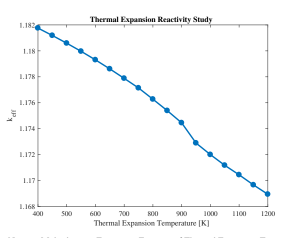
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Effective Neutron Multiplication Factor as a Function of Thermal Expansion Temperature.

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## Remember why we are here.

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## Remember why we are here.

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Model a nuclear reactor.



#### Advanced Burner Reactor (ABR) – MET-1000

Fast Reactor and FFM

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- Benchmark published February 2016 [OEC16].
- Four designs including MET-1000.
- 31 independent solutions submitted so far including DIF3D.
- Cross sections generated independently.

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#### **Benchmark Results**

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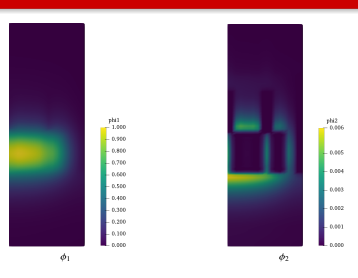
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$$k_{eff} = 1.006694$$
 (DIF3D -700 [pcm])

# **Reactivity Coefficients**

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The reactivity of a reactor can be defined.

$$\rho_i = \frac{k_{eff,i} - 1}{k_{eff,i}}$$

Reactivity coefficient is a derivative with respect to a variable of interest.

$$\alpha_x(x_i) = \left. \frac{\partial \rho}{\partial x} \right|_{x=x_i}$$
$$\Delta \rho \approx \alpha_x(x_i) \, \Delta x$$

## Reactivity Coefficient Formulae

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Consider a series of reactor powers  $Q_{Rx,i} = \{0\%, ..., 100\%\}$ . Define the following reactivity coefficients.

$$\begin{split} \alpha_{power}(Q_{Rx,i}) &= \frac{\rho(Q_{Rx,i}) - \rho(Q_{Rx,i} + \Delta Q_{Rx})}{\Delta Q_{Rx}} \\ \alpha_{thexp}(Q_{Rx,i}) &= \frac{\rho(T_{exp}(Q_{Rx,i})) - \rho(T_{exp}(Q_{Rx,i} + \Delta Q_{Rx}))}{\Delta Q_{Rx}} \\ \alpha_{CTC}(Q_{Rx,i}) &= \frac{\rho(Q_{Rx,i}) - \rho(T_{cool} + \Delta T_{cool})}{\Delta T_{cool}} \\ \alpha_{Doppler}(Q_{Rx,i}) &= \frac{\rho(Q_{Rx,i}) - \rho_i(T_{fuel} + \Delta T_{fuel})}{\Delta T_{fuel}} \end{split}$$



### Eigenvalue Feedback

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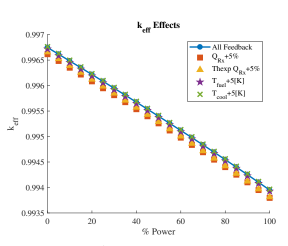
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 $\textit{k}_{\it{eff}}$  Feedback Effects.



### Temperature Reactivity Coefficients

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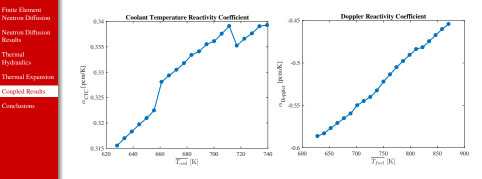
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## Power Reactivity Coefficients

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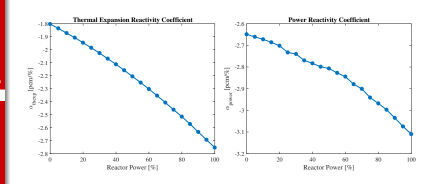
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## Multiphysics Contributions to Total Power Defect

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- −559.64 [pcm] due to thermal expansion effects.
- -29.85 [pcm] due to thermal hydraulics effects.
- Cancellation of error due to  $\alpha_{Doppler}$  and  $\alpha_{CTC}$ .

Case	Thermal Expansion Power	Thermal Hydraulic Power	$k_{eff}$	Reactivity [pcm]
1	0%	0%	0.999808	
2	100%	0%	0.994246	-559.64
3	100%	100%	0.993950	-589.49

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# Summary

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# Summary

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Modeled a nuclear reactor.



## Summary

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- Solved multigroup neutron diffusion equation via FEM.
- Developed thermal hydraulics models.
- Developed thermal expansion model.
- Demonstrated multiphysics simulation based on ABR.
- Estimated multiphysics reactivity coefficients.



### **Future Improvements**

Fast Reactor and FEM

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Code Enhancements and New Features.

- Depletion with Chebyshev Rational Approximation Method (CRAM) [Pus13].
- ► Higher order finite elements (e.g. quadratic) [Hos13].
- Simplified  $P_N$  (SP<sub>N</sub>) [Ryu13].
- Encouraging Code Usage.
  - ▶ Should be a tool for core design optimization.
  - ► More users encourage more feedback.
  - Unique reactor designs encourage feature additions.



#### Thank You!

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Thank you all for coming this morning!

I would like to thank my advisor, Dr. Scott Palmtag, and my committee, Dr. J. Michael Doster and Dr. Ralph Smith.



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## Acronyms I

Fast Reactor and FEM

William Christopher Dawn

Introduction

Finite Element Neutron Diffusion

Results Thermal

Hydraulics Thermal Expansion

Coupled Results

Conclusions

**ABR** Advanced Burner Reactor.

**ANL** Argonne National Laboratory.

**CG** Conjugate Gradient.

**CRAM** Chebyshev Rational Approximation Method.

**EBR-II** Experimental Breeder Reactor II.

FEM Finite Element Method.

LWR Light Water Reactor.

**RCM** Reverse Cuthill-McKee.

RMS Root-Mean-Squared.

SFR Sodium-cooled Fast Reactor.

**SPD** Symmetric Positive Definite.



#### Source Codes

Fast Reactor and FEM William

Christopher Dawn

Introduction

Finite Element Neutron Diffusion

Neutron Diffusion Results

Thermal Hydraulics

Thermal Expansion

Coupled Results

Conclusions

Defense Slides & Thesis.

 $\verb|https://github.com/wcdawn/WilliamDawn-thesis||$ 

Thesis Code.

https://github.ncsu.edu/wcdawn/masters\_thesis

Note: Not currently open-source. Contact the author for access.