# Massively Parallel Monte Carlo Methods for Discrete Linear and Nonlinear Systems

Stuart R. Slattery
Engineering Physics Department
University of Wisconsin - Madison

October 31, 2012



#### Introduction



- Predictive modeling and simulation enhances engineering capability
- Modern work focused on this task leverages multiple physics simulation (CASL, NEAMS)
- New hardware drives algorithm development (petascale and exascale)
- Monte Carlo methods have the potential to provide great improvements that permit finer simulations and better mapping to future hardware
- A set of massively parallel Monte Carlo methods is proposed to advance multiple physics simulation on contemporary and future leadership class machines

## Physics-Based Motivation



## Physics-Based Motivation



#### Predictive nuclear reactor analysis enables...

- Tighter design tolerance for improved thermal performance and efficiency
- Higher fuel burn-up
- High confidence in accident scenario models

## Physics-Based Motivation



#### Predictive nuclear reactor analysis enables...

- Tighter design tolerance for improved thermal performance and efficiency
- Higher fuel burn-up
- High confidence in accident scenario models

#### Multiple physics simulations are complicated...

- Neutronics, thermal hydraulics, computational fluid dynamics, structural mechanics, and many other physics
- Consistent models yield nonlinearities in the variables through feedback effects
- Tremendous computational resources are required with  $O(1 \times 10^9)$  element meshes and O(100,000)+ cores used in today's simulations.

## Physics-Based Motivation: DNB



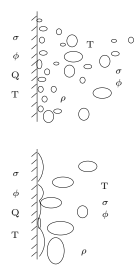
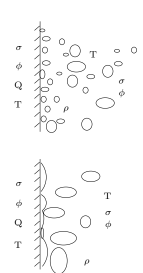


Figure: Departure from nucleate boiling scenario.

### Physics-Based Motivation: DNB





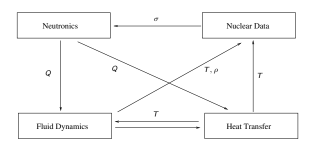


Figure: Multiphysics dependency analysis of departure from nucleate boiling.

Figure: Departure from nucleate boiling scenario.

#### Hardware-Based Motivation



- Modern hardware is moving in two directions:
  - Lightweight machines
  - Heterogeneous machines
  - Both characterized by low power and high concurrency
- Some issues:
  - Higher potential for both soft and hard failures
  - Memory restrictions are expected with a continued decrease in memory/FLOPS
- Potential resolution from Monte Carlo:
  - Soft failures buried within the tally variance
  - Hard failures are high variance events
  - Memory savings over conventional methods

#### Research Outline



- Parallelization of Monte Carlo methods for discrete systems
  - Parallel strategies taken from modern reactor physics methods
  - Research is required to explore varying parallel strategies
  - Scalability is of concern
- Development of a nonlinear solver leveraging Monte Carlo
  - Application to nonlinear problems of interest
  - Memory benefits
  - Performance benefits

# Linear Operator Equations



• We seek solutions of the general linear operator equation

$$\begin{aligned} \boldsymbol{A}\boldsymbol{x} &= \boldsymbol{b} \\ \boldsymbol{A} &\in \mathbb{R}^{N \times N}, \ \boldsymbol{A} : \mathbb{R}^N \to \mathbb{R}^N, \ \boldsymbol{x} \in \mathbb{R}^N, \ \boldsymbol{b} \in \mathbb{R}^N \end{aligned}$$

•  $\mathbf{r} = \mathbf{0}$  when an exact solution is found.

# Linear Operator Equations



• We seek solutions of the general linear operator equation

$$\begin{aligned} \mathbf{A}\mathbf{x} &= \mathbf{b} \\ \mathbf{A} &\in \mathbb{R}^{N \times N}, \ \mathbf{A} : \mathbb{R}^N \to \mathbb{R}^N, \ \mathbf{x} \in \mathbb{R}^N, \ \mathbf{b} \in \mathbb{R}^N \end{aligned}$$

•  $\mathbf{r} = \mathbf{0}$  when an exact solution is found.

#### A Requirement

Assert that **A** is *nonsingular*. The solution is then:

$$\mathbf{x} = \mathbf{A}^{-1}\mathbf{b}$$

# Stationary Methods



• General stationary methods are formed by splitting the linear operator

$$A = M - N$$
.

$$\mathbf{x} = \mathbf{M}^{-1}\mathbf{N}\mathbf{x} + \mathbf{M}^{-1}\mathbf{b} .$$

• We identify  $\mathbf{H} = \mathbf{M}^{-1}\mathbf{N}$  as the *iteration matrix* 

$$\mathbf{x}^{k+1} = \mathbf{H}\mathbf{x}^k + \mathbf{c}$$
 .

# Stationary Methods Convergence



- The qualities of the iteration matrix dictate convergence
- Define  $\mathbf{e}^k = \mathbf{x}^k \mathbf{x}$  as the error at the  $k^{th}$  iterate

$$e^{k+1} = He^k$$

We diagonalize H to extract its Eigenvalues

$$||\mathbf{e}^k||_2 = \rho(\mathbf{H})^k ||\mathbf{e}^0||_2$$
,

• We bound **H** by  $ho(\mathbf{H}) < 1$  for convergence

## Projection Methods



- Powerful class of iterative methods
- Provides theory that encapsulates most other iterative methods
- Leveraged in many modern physics codes at the petascale

## Projection Methods



- Powerful class of iterative methods
- Provides theory that encapsulates most other iterative methods
- Leveraged in many modern physics codes at the petascale

### Search Subspace ${\mathfrak K}$

Extract the solution from the search subspace:

$$\boldsymbol{\tilde{x}}=\boldsymbol{x}_0+\boldsymbol{\delta},\ \boldsymbol{\delta}\in\boldsymbol{\mathfrak{K}}$$

# Projection Methods



- Powerful class of iterative methods
- Provides theory that encapsulates most other iterative methods
- Leveraged in many modern physics codes at the petascale

#### Search Subspace ${\mathfrak K}$

Extract the solution from the search subspace:

$$\tilde{\mathbf{x}} = \mathbf{x}_0 + \boldsymbol{\delta}, \ \boldsymbol{\delta} \in \mathcal{K}$$

#### Constraint Subspace $\mathcal{L}$

Constrain the extraction with the constraint subspace by asserting orthogonality with the residual:

$$\langle \tilde{\mathbf{r}}, \mathbf{w} \rangle = 0, \ \forall \mathbf{w} \in \mathcal{L}$$

## The Orthogonality Constraint



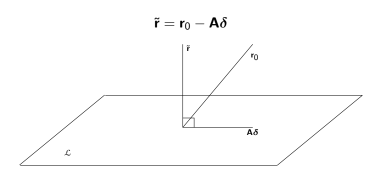


Figure: Orthogonality constraint of the new residual with respect to  $\mathcal{L}. \label{eq:local_local_local}$ 

# The Orthogonality Constraint



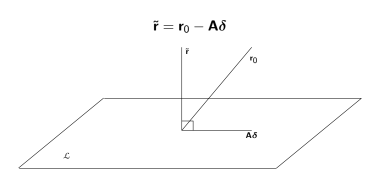


Figure: Orthogonality constraint of the new residual with respect to  $\mathcal{L}$ .

#### Minimization Property

The residual of the system will always be *minimized* with respect to the constraints

$$||\tilde{\mathbf{r}}||_2 \le ||\mathbf{r}_0||_2, \ \forall \mathbf{r}_0 \in \mathbb{R}^N$$

## Putting it All Together



ullet Choose  $oldsymbol{V}$  as a basis of  $\mathcal K$  and  $oldsymbol{W}$  as a basis of  $\mathcal L$ 

$$\boldsymbol{\delta} = \mathbf{V}\mathbf{y}, \ orall \mathbf{y} \in \mathbb{R}^N$$

$$\textbf{y} = (\textbf{W}^{T}\textbf{A}\textbf{V})^{-1}\textbf{W}^{T}\textbf{r}_{0}$$

# Putting it All Together



ullet Choose  $oldsymbol{V}$  as a basis of  $\mathcal K$  and  $oldsymbol{W}$  as a basis of  $\mathcal L$ 

$$\pmb{\delta} = \mathbf{V}\mathbf{y}, \ orall \mathbf{y} \in \mathbb{R}^{\mathcal{N}}$$

$$\mathbf{y} = (\mathbf{W}^T \mathbf{A} \mathbf{V})^{-1} \mathbf{W}^T \mathbf{r}_0$$

#### Projection Method Iteration

$$\mathbf{r}^k = \mathbf{b} - \mathbf{A}\mathbf{x}^k$$
 $\mathbf{y}^k = (\mathbf{W}^T \mathbf{A} \mathbf{V})^{-1} \mathbf{W}^T \mathbf{r}^k$ 
 $\mathbf{x}^{k+1} = \mathbf{x}^k + \mathbf{V} \mathbf{y}^k$ 
Update  $\mathbf{V}$  and  $\mathbf{W}$ 

# Krylov Subspace Methods



$$\mathcal{K}_m(\mathbf{A}, \mathbf{r}_0) = span\{\mathbf{r}_0, \mathbf{A}\mathbf{r}_0, \mathbf{A}^2\mathbf{r}_0, \dots, \mathbf{A}^{m-1}\mathbf{r}_0\}$$

$$\mathcal{L} = \mathbf{A}\mathcal{K}_m(\mathbf{A}, \mathbf{r}_0)$$

- Yields the normal system  $\mathbf{A}^T \mathbf{A} \mathbf{x} = \mathbf{A}^T \mathbf{b}$
- ullet Must generate an orthonormal basis  $oldsymbol{V}_m \in \mathbb{R}^{N imes m}$  for  $\mathfrak{K}_m(oldsymbol{\mathsf{A}}, oldsymbol{\mathsf{r}}_0)$
- $\mathbf{W}_m = \mathbf{AV}_m$
- Typically choose a Gram-Schmidt-like procedure such as Arnoldi or Lanzcos

#### **GMRES**



#### **Algorithm 1** GMRES Iteration

```
1: \mathbf{r}_0 := \mathbf{b} - \mathbf{A} \mathbf{x}_0
 2: \beta := ||\mathbf{r}_0||_2
 3: \mathbf{v}_1 := \mathbf{r}_0/\beta {Create the orthonormal basis for the Krylov subspace}
 4: for j = 1, 2, \dots, m do
 5: \mathbf{w}_i := \mathbf{A}\mathbf{v}_i
      for i = 1, 2, \dots, j do
 6:
      h_{ii} \leftarrow \langle \mathbf{w}_i, \mathbf{v}_i \rangle
          \mathbf{w}_i \leftarrow \mathbf{w}_i - h_{ii}\mathbf{v}_i
 8:
       end for
 9.
      h_{i+1,i} \leftarrow ||\mathbf{w}_i||_2
10:
       \mathbf{v}_{i+1} \leftarrow \mathbf{w}_i/h_{i+1,i}
11:
12: end for{Apply the orthogonality constraints}
13: \mathbf{y}_m \leftarrow \operatorname{argmin}_{\mathbf{v}} ||\beta \mathbf{e}_1 - \mathbf{H}_m \mathbf{y}||_2
14: \mathbf{x}_m \leftarrow \mathbf{x}_0 + \mathbf{V}_m \mathbf{y}_m
```

# Parallel Projection Methods



Parallel vector update

$$\mathbf{y}[n] \leftarrow \mathbf{y}[n] + a * \mathbf{x}[n], \ \forall n \in [1, N_g]$$
  
 $\mathbf{y}[n] \leftarrow \mathbf{y}[n] + a * \mathbf{x}[n], \ \forall n \in [1, N_l]$ 

• Parallel dot product

$$d_I = \mathbf{y}_I \cdot \mathbf{x}_I, \ d_g = \sum_{p} d_I$$

Parallel vector norm

$$||x||_{\infty,I} = \max_{n} \mathbf{y}[n], \ \forall n \in [1, N_I]$$
$$||x||_{\infty,g} = \max_{p} ||x||_{\infty,I}$$

## Parallel Matrix-Vector Multiplication



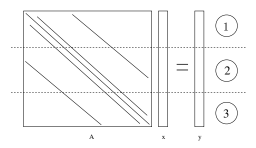


Figure: Matrix-vector multiply  $\mathbf{A}\mathbf{x} = \mathbf{y}$  operation on 3 processors.

# Parallel Matrix-Vector Multiplication



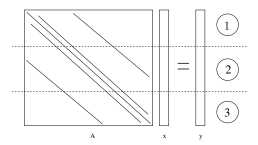


Figure: Matrix-vector multiply Ax = y operation on 3 processors.

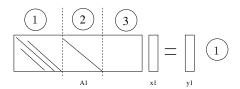


Figure: Components of multiply operation owned by process 1.

# **Projection Methods Summary**



- Used in practice everywhere
- Global reduction operations observed not to impede scalability
  - Dot product
  - Vector norms
- Nearest neighbor computations have poor algorithmic strong scaling
  - Matrix-vector multiply
  - Weak scaling is better

## Monte Carlo Methods for Discrete Linear Systems



- First proposed by J. Von Neumann and S.M. Ulam in the 1940's
- Earliest published reference in 1950
- General lack of published work
- Modern work by Evans and others has yielded new applications

### Monte Carlo Linear Solver Preliminaries



Split the operator

$$H = I - A$$

$$x = Hx + b$$

• Generate the Neumann series

$$A^{-1} = (I - H)^{-1} = \sum_{k=0}^{\infty} H^k$$

• Require  $\rho(\mathbf{H}) < 1$  for convergence

$$\mathbf{A}^{-1}\mathbf{b} = \sum_{k=0}^{\infty} \mathbf{H}^k \mathbf{b} = \mathbf{x}$$

### Monte Carlo Linear Solver Preliminaries



• Expand the Nuemann series

$$x_i = \sum_{k=0}^{\infty} \sum_{i_1}^{N} \sum_{i_2}^{N} \dots \sum_{i_k}^{N} h_{i,i_1} h_{i_1,i_2} \dots h_{i_{k-1},i_k} b_{i_k}$$

• Define a sequence of state transitions

$$\nu = i \rightarrow i_1 \rightarrow \cdots \rightarrow i_{k-1} \rightarrow i_k$$

• Define the Neumann-Ulam decomposition<sup>1</sup>

$$H = P \circ W$$

<sup>&</sup>lt;sup>1</sup>The Hadamard product  $\mathbf{A} = \mathbf{B} \circ \mathbf{C}$  is defined element-wise as  $a_{ij} = b_{ij}c_{ij}$ .

### Direct Method



• Compute row-normalized transition probabilities and weights

$$p_{ij} = \frac{|h_{ij}|}{\sum_j |h_{ij}|}, \ w_{ij} = \frac{h_{ij}}{p_{ij}}$$

• Generate an expectation value for the solution

$$W_m = \sum_{m=0}^k w_{i,i_1} w_{i_1,i_2} \cdots w_{i_{m-1},i_m}$$
  
 $X_{\nu}(i_0 = i) = \sum_{m=0}^k W_m b_{i_m}$ 

### Direct Method



• Compute the probability of a particular random walk permutation

$$P_{\nu} = p_{i,i_1} p_{i_1,i_2} \cdots p_{i_{k-1},i_k}$$

Generate the estimator

$$E\{X(i_0=i)\}=\sum_{\nu}P_{\nu}X_{\nu}$$

Check that we recover the exact solution

$$E\{X(i_0=i)\} = \sum_{k=0}^{\infty} \sum_{i_1}^{N} \sum_{i_2}^{N} \dots \sum_{i_k}^{N} p_{i,i_1} p_{i_1,i_2} \dots p_{i_{k-1},i_k} w_{i,i_1} w_{i_1,i_2} \dots w_{i_{k-1},i_k} b_{i_k}$$

$$= x_i,$$





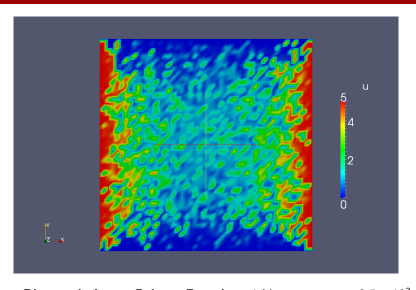


Figure: **Direct solution to Poisson Equation.** 1 history per state,  $2.5 \times 10^3$  total histories. 0.785 seconds CPU time.



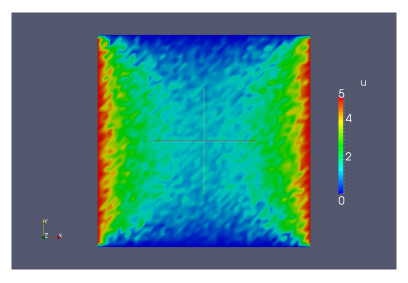


Figure: **Direct solution to Poisson Equation.** 10 histories per state,  $2.5 \times 10^4$  total histories. 5.9 seconds CPU time.



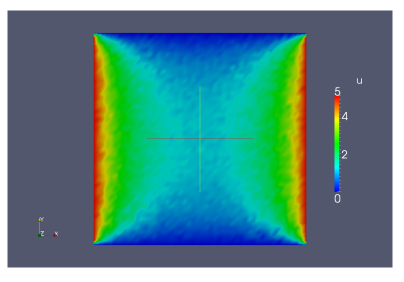


Figure: Direct solution to Poisson Equation. 100 histories per state,  $2.5 \times 10^5$  total histories. 54.7 seconds CPU time.



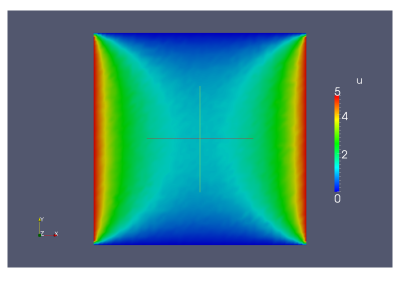


Figure: Direct solution to Poisson Equation. 100 histories per state,  $2.5 \times 10^6$  total histories. 644 seconds CPU time.

### Adjoint Method



• Solve the adjoint linear system

$$\mathbf{A}^T \mathbf{y} = \mathbf{d}$$

$$\mathbf{y} = \mathbf{H}^T \mathbf{y} + \mathbf{d}$$

• Set the adjoint constraint

$$\langle \mathbf{A}^T \mathbf{x}, \mathbf{y} \rangle = \langle \mathbf{x}, \mathbf{A} \mathbf{y} \rangle$$

$$\langle \mathbf{x}, \mathbf{d} \rangle = \langle \mathbf{y}, \mathbf{b} \rangle$$

### Adjoint Method



• Generate the Neumann series for the adjoint operator

$$\mathbf{y} = (\mathbf{I} - \mathbf{H}^T)^{-1} \mathbf{d}$$

$$\mathbf{y} = \sum_{k=0}^{\infty} (\mathbf{H}^T)^k \mathbf{d}$$

Expand the series

$$y_i = \sum_{k=0}^{\infty} \sum_{i_1}^{N} \sum_{i_2}^{N} \dots \sum_{i_k}^{N} h_{i_k, i_{k-1}} \dots h_{i_2, i_1} h_{i_1, i} d_{i_k}$$

• Pick another constraint to yield the original solution

$$\mathbf{d} = \boldsymbol{\delta}_i, \ \langle \mathbf{y}, \mathbf{b} \rangle = \langle \mathbf{x}, \boldsymbol{\delta}_i \rangle = x_i$$

### Adjoint Method



• Use the adjoint Neumann-Ulam decomposition

$$\mathbf{H}^T = \mathbf{P} \circ \mathbf{W}$$

$$p_{ij} = \frac{|h_{ji}|}{\sum_j |h_{ji}|}, \quad w_{ij} = \frac{h_{ji}}{p_{ij}}$$

Build the estimator and expectation value

$$X_{\nu} = \sum_{m=0}^{k} W_{m} b_{i_{0}} \delta_{i,i_{m}}$$

$$E\{X_{j}\} = \sum_{k=0}^{\infty} \sum_{i_{1}}^{N} \sum_{i_{2}}^{N} \dots \sum_{i_{k}}^{N} b_{i_{0}} h_{i,i_{1}} h_{i_{1},i_{2}} \dots h_{i_{k-1},i_{k}} \delta_{i_{k},j}$$

$$= x_{j},$$





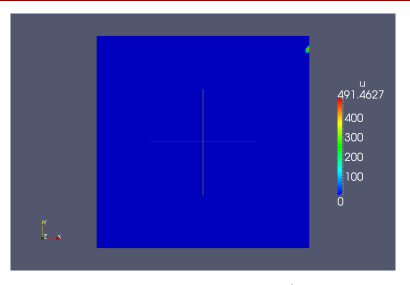


Figure: Adjoint solution to Poisson Equation.  $1 \times 10^0$  total histories, 0.286 seconds CPU time.





Figure: Adjoint solution to Poisson Equation.  $1 \times 10^1$  total histories, 0.278 seconds CPU time.





Figure: Adjoint solution to Poisson Equation.  $1 \times 10^2$  total histories, 0.275 seconds CPU time.



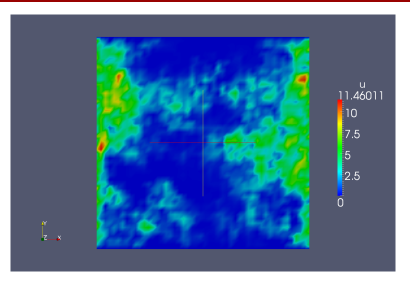


Figure: Adjoint solution to Poisson Equation.  $1 \times 10^3$  total histories, 0.291 seconds CPU time.



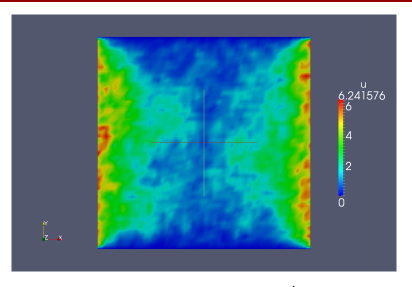


Figure: Adjoint solution to Poisson Equation.  $1 \times 10^4$  total histories, 0.428 seconds CPU time.





Figure: Adjoint solution to Poisson Equation.  $1 \times 10^5$  total histories, 1.76 seconds CPU time.



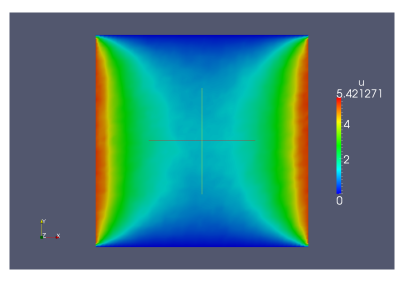


Figure: Adjoint solution to Poisson Equation.  $1 \times 10^6$  total histories, 15.1 seconds CPU time.





Figure: Adjoint solution to Poisson Equation.  $1 \times 10^7$  total histories, 149 seconds CPU time.

### Sequential Monte Carlo



- Neumann-Ulam methods bound by the Central Limit Theorem
- Halton proposed an iterative residual method
- Iteration error decoupled from Monte Carlo error
- Exponential convergence

$$\mathbf{r}^k = \mathbf{b} - \mathbf{A}\mathbf{x}^k$$
 $\mathbf{A}\delta^k = \mathbf{r}^k$ 
 $\mathbf{x}^{k+1} = \mathbf{x}^k + \delta^k$ 

### Monte Carlo Synthetic-Acceleration



• Split the operator to yield Richardson's iteration

$$\mathbf{x} = (\mathbf{I} - \mathbf{A})\mathbf{x} + \mathbf{b}$$
  
 $\mathbf{x}^{k+1} = (\mathbf{I} - \mathbf{A})\mathbf{x}^k + \mathbf{b}$ 

Define the iteration error

$$\delta \mathbf{x}^k = \mathbf{x} - \mathbf{x}^k$$
$$\delta \mathbf{x}^{k+1} = (\mathbf{I} - \mathbf{A}) \delta \mathbf{x}^k$$

# Monte Carlo Synthetic-Acceleration



• Subtract  $(\mathbf{I} - \mathbf{A})\delta \mathbf{x}^{k+1}$ 

$$\mathbf{A}\delta\mathbf{x}^{k+1} = (\mathbf{I} - \mathbf{A})(\mathbf{x}^{k+1} - \mathbf{x}^k)$$
$$= \mathbf{r}^{k+1}$$

• The following converges in one iteration with exact inversion of A:

$$\mathbf{x}^{k+1} = (\mathbf{I} - \mathbf{A})\mathbf{x}^k + \mathbf{b}$$
$$\mathbf{A}\delta\mathbf{x}^{k+1} = \mathbf{r}^{k+1}$$
$$\mathbf{x} = \mathbf{x}^{k+1} + \delta\mathbf{x}^{k+1}$$

### Monte Carlo Synthetic-Acceleration



#### MCSA Iteration

$$\mathbf{x}^{k+1/2} = (\mathbf{I} - \mathbf{A})\mathbf{x}^{k} + \mathbf{b}$$

$$\mathbf{r}^{k+1/2} = \mathbf{b} - \mathbf{A}\mathbf{x}^{k+1/2}$$

$$\hat{\mathbf{A}}\delta\mathbf{x}^{k+1/2} = \mathbf{r}^{k+1/2}$$

$$\mathbf{x}^{k+1} = \mathbf{x}^{k+1/2} + \delta\mathbf{x}^{k+1/2}$$

- Adjoint Neumann-Ulam solver computes the correction
- Decouples Monte Carlo error from solution error
- Exponential convergence
- Demonstrated by Evans and colleagues to be competitive with Krylov methods

## Preconditioning Monte Carlo Methods



- No symmetry requirements
- Require ρ(**H**) < 1</li>
- Choose Jacobi preconditioning at a minimum

$$\mathbf{M} = diag(\mathbf{A})$$
 $\mathbf{M}^{-1}\mathbf{A}\mathbf{x} = \mathbf{M}^{-1}\mathbf{b}$ 

Yields a preconditioned MCSA iteration with no in-state transitions

$$\mathbf{x}^{k+1/2} = (\mathbf{I} - \mathbf{M}^{-1}\mathbf{A})\mathbf{x}^k + \mathbf{b}$$
 $\mathbf{r}^{k+1/2} = \mathbf{b} - \mathbf{M}^{-1}\mathbf{A}\mathbf{x}^{k+1/2}$ 
 $\mathbf{M}^{-1}\mathbf{A}\delta\mathbf{x}^{k+1/2} = \mathbf{r}^{k+1/2}$ 
 $\mathbf{x}^{k+1} = \mathbf{x}^{k+1/2} + \delta\mathbf{x}^{k+1/2}$ 

## Direct vs. Adjoint Analysis



- Analysis needed to select Monte Carlo method
- Time-dependent 2-dimensional Poisson equation
- Spectral radius fixed
- Sparsity varied with 2 Laplacian stencils

$$\nabla_5^2 = \frac{1}{\Delta^2} [u_{i-1,j} + u_{i+1,j} + u_{i,j-1} + u_{i,j+1} - 4u_{i,j}]$$

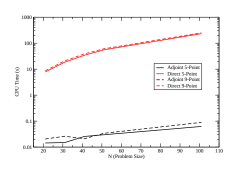
$$\nabla_9^2 = \frac{1}{6\Delta^2} [4u_{i-1,j} + 4u_{i+1,j} + 4u_{i,j-1} + 4u_{i,j+1} + u_{i-1,j-1} + u_{i-1,j+1} + u_{i+1,j-1} + u_{i+1,j+1} - 20u_{i,j}]$$

Implicit Euler time differencing

$$\mathbf{A}\mathbf{u}^{n+1}=\mathbf{u}^n$$

### Direct vs. Adjoint Analysis





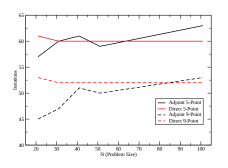


Figure: CPU Time (s) to converge vs. Figure: Iterations to converge vs. mesh).

Problem Size (N for an  $N \times N$  square Problem Size (N for an  $N \times N$  square mesh).

## Direct vs. Adjoint Analysis





Figure: Infinity norm of the solution residual vs. iteration number for a problem of fixed size.

- CPU time dominating factor in method selection
- Significant speedup with adjoint method
- Does not affect convergence behavior
- Use adjoint with MCSA and Sequential Monte Carlo

### Generalization of MCSA for Linear Problems



- Work to date has used a physics-based formulation
  - Radiation transport equations used as model system
  - Transition probabilities built from problem-specific parameters
  - Probabilities and weights must be rederived for each new equation set
- Desire a generalization for all linear operator equations
  - Requires a general parallel framework
  - Requires implementation with a general linear algebra framework
  - Operator, vector, and graph abstractions
- Neumann-Ulam solvers and MCSA implemented using the Trilinos Petra frameworks
- Can be leveraged in modern physics implementations

### Parallelization of Monte Carlo Methods



- No literature observed for parallel Neumann-Ulam solvers
- Numerous references for modern parallel Monte Carlo methods in reactor physics
- Build a strategy for applying modern methods to the Neumann-Ulam method
- MCSA iteration-level parallelism comes from parallel matrix/vector operations

# Domain Decomposition



# Multiple-Set Overlapping-Domain Decomposition



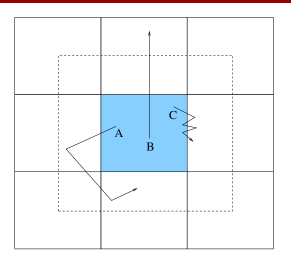


Figure: Overlapping domain example illustrating how domain overlap can reduce communication costs.

### Domain-to-Domain Communication



### Load Balancing



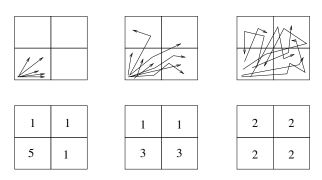


Figure: Example illustrating how domain decomposition can create load balance issues in Monte Carlo.

### Reproducability



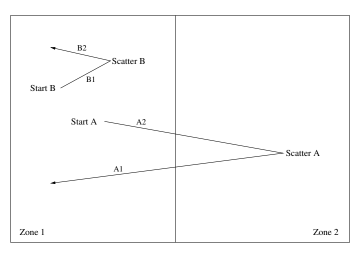


Figure: Gentile's example illustrating how domain decomposition can create reproducibility issues in Monte Carlo.

# Parallel Adjoint Method



### Parallel MCSA



#### MCSA Iteration

$$\mathbf{x}^{k+1/2} = (\mathbf{I} - \mathbf{A})\mathbf{x}^k + \mathbf{b}$$
 $\mathbf{r}^{k+1/2} = \mathbf{b} - \mathbf{A}\mathbf{x}^{k+1/2}$ 
 $\hat{\mathbf{A}}\delta\mathbf{x}^{k+1/2} = \mathbf{r}^{k+1/2}$ 
 $\mathbf{x}^{k+1} = \mathbf{x}^{k+1/2} + \delta\mathbf{x}^{k+1/2}$ 

- Richardson iteration and residual computation require parallel matrix-vector multiply and parallel vector update
- This work will generate a parallel adjoint Neumann-Ulam solver
- Application of correction requires parallel vector update
- Convergence checks through parallel norm computation

### Monte Carlo Solution Methods for Nonlinear Problems

### Monte Carlo Solution Methods for Nonlinear Problems

- · Many multiphysics problems of interest are nonlinear
- Segregated methods lack the consistency of fully implicit methods
- Newton methods often leverage Krylov solvers
  - Robust implementations
  - No operator required
  - Memory intensive
- Monte Carlo methods need the full operator
- Automatic construction of the linear operator available
  - Ideal for Monte Carlo
  - Relaxes memory requirements
  - Potential scaling improvements
  - Resiliency benefits

### Nonlinear Preliminaries



• We seek solutions of the general nonlinear problem

$$\begin{aligned} & F(u) = 0 \\ & u \in \mathbb{R}^n, \ F: \mathbb{R}^N \rightarrow \mathbb{R}^N \end{aligned}$$

- We interpret the exact solution  $\mathbf{u}$  to be the roots of  $\mathbf{F}(\mathbf{u})$
- Taylor expand the residuals at the k+1 iterate about the k iterate

$$\mathbf{F}(\mathbf{u}^{k+1}) = \mathbf{F}(\mathbf{u}^k) + \mathbf{F}'(\mathbf{u}^k)(\mathbf{u}^{k+1} - \mathbf{u}^k) + \frac{\mathbf{F}''(\mathbf{u}^k)}{2}(\mathbf{u}^{k+1} - \mathbf{u}^k)^2 + \cdots$$

• Assert  $\mathbf{u}^{k+1}$  is the exact solution

$$-\mathsf{F}(\mathsf{u}^k) = \mathsf{F}'(\mathsf{u}^k)(\mathsf{u}^{k+1} - \mathsf{u}^k)$$

### Nonlinear Preliminaries



•  $\mathbf{F}'(\mathbf{u}^k)$  is the *Jacobian*  $\mathbf{J}(\mathbf{u})$ 

$$J_{ij} = \frac{\partial F_i(\mathbf{u})}{\partial u_j}$$

•  $(\mathbf{u}^{k+1} - \mathbf{u}^k)$  is the solution update from the k iterate to the k+1 iterate

$$\delta \mathbf{u}^k = \mathbf{u}^{k+1} - \mathbf{u}^k$$

Form Newton's method

$$J(\mathbf{u})\delta\mathbf{u}^k = -\mathbf{F}(\mathbf{u}^k)$$
$$\mathbf{u}^{k+1} = \mathbf{u}^k + \delta\mathbf{u}^k$$

## Newton-Krylov Methods



A form of inexact Newton methods

$$||\mathbf{J}(\mathbf{u}^k)\delta\mathbf{u}^k + \mathbf{F}(\mathbf{u}^k)|| \le \eta^k ||\mathbf{F}(\mathbf{u}^k)||, \ \eta^k \in [0, 1)$$

- Choose a Krylov method to solve for the Newton correction
- GMRES with a long recurrence relation observed as more robust
- Generates a monotonically decreasing residual from maintaining the optimization and orthogonality conditions

## Newton-Krylov Methods



A form of inexact Newton methods

$$||\mathbf{J}(\mathbf{u}^k)\delta\mathbf{u}^k + \mathbf{F}(\mathbf{u}^k)|| \le \eta^k ||\mathbf{F}(\mathbf{u}^k)||, \ \eta^k \in [0, 1)$$

- Choose a Krylov method to solve for the Newton correction
- GMRES with a long recurrence relation observed as more robust
- Generates a monotonically decreasing residual from maintaining the optimization and orthogonality conditions

Where does the Jacobian come from?

## Newton-Krylov Methods



A form of inexact Newton methods

$$||\mathbf{J}(\mathbf{u}^k)\delta\mathbf{u}^k + \mathbf{F}(\mathbf{u}^k)|| \le \eta^k ||\mathbf{F}(\mathbf{u}^k)||, \ \eta^k \in [0, 1)$$

- Choose a Krylov method to solve for the Newton correction
- GMRES with a long recurrence relation observed as more robust
- Generates a monotonically decreasing residual from maintaining the optimization and orthogonality conditions

#### Where does the Jacobian come from?

- The Jacobian can come from hand-coded derivatives
  - Tedious and error prone
  - Repeated for each equation set and hard to do for multiphysics

# Matrix-Free Approximation



- Krylov methods only need the action of the linear operator
- The action of the Jacobian can be approximated using a forward difference

$$\mathsf{J}(\mathsf{u})\mathsf{v} = \frac{\mathsf{F}(\mathsf{u} + \epsilon \mathsf{v}) - \mathsf{F}(\mathsf{u})}{\epsilon}$$

- Forms the basis of Jacobian-Free Newton-Krylov (JFNK) methods
  - Sensitive to scaling and discretization error
  - Still have to form part of Jacobian periodically for preconditioning
  - Eventually break even with generating and storing the full Jacobian

#### Automatic Differentiation



- Automatically generate Jacobians from nonlinear function evaluations
  - Overload math operators and apply the chain rule (FAD)
  - Yields evaluations as accurate as function discretization
- Modern packages take an element-level assembly approach
  - Function evaluations are local, communication builds global data

$$\mathbf{J}(\mathbf{u}) = \sum_{i=1}^{N} \mathbf{Q}_{i}^{T} \mathbf{J}_{k} \mathbf{P}_{i}$$

$$e_{k} : \mathbb{R}^{n_{k}} \to \mathbb{R}^{m_{k}}, \ \mathbf{J}_{k_{i}} = \partial e_{k_{i}} / \partial P_{i} u, \ \mathbf{P} \in \mathbb{R}^{m_{k_{i}} \times N}$$

 Performance studies give acceptable results for use in large-scale, production physics codes

#### The FANM Method



Forward-Automated Newton-MCSA

#### The FANM Method



#### Forward-Automated Newton-MCSA

#### **Algorithm 3** FANM

- 1: k := 0
- 2: while  $||\mathbf{F}(\mathbf{u}^k)|| > \epsilon ||\mathbf{F}(\mathbf{u}^0)||$  do
- 3:  $\mathbf{J}(\mathbf{u}^k) \leftarrow AD(\mathbf{F}(\mathbf{u}^k))$  {Automatic differentiation}
- 4:  $\mathbf{J}(\mathbf{u}^k)\delta\mathbf{u}^k = -\mathbf{F}(\mathbf{u}^k)$  {Solve for the Newton correction with MCSA}
- 5:  $\mathbf{u}^{k+1} \leftarrow \mathbf{u}^k + \delta \mathbf{u}^k$
- 6:  $k \leftarrow k + 1$
- 7: end while

#### The FANM Method



#### Forward-Automated Newton-MCSA

#### **Algorithm 4** FANM

- 1: k := 0
- 2: while  $||\mathbf{F}(\mathbf{u}^k)|| > \epsilon ||\mathbf{F}(\mathbf{u}^0)||$  do
- 3:  $J(\mathbf{u}^k) \leftarrow AD(\mathbf{F}(\mathbf{u}^k))$  {Automatic differentiation}
- 4:  $\mathbf{J}(\mathbf{u}^k)\delta\mathbf{u}^k = -\mathbf{F}(\mathbf{u}^k)$  {Solve for the Newton correction with MCSA}
- 5:  $\mathbf{u}^{k+1} \leftarrow \mathbf{u}^k + \delta \mathbf{u}^k$
- 6:  $k \leftarrow k + 1$
- 7: end while
  - Robustness of Newton's method (inexact)
  - Accuracy and convenience of FAD
  - · Parallelism, memory, and resiliency benefits of MCSA
  - Requires only nonlinear function evaluations



 Jacobian will be in a compressed row storage format

$$\mathbf{A} = \begin{bmatrix} 2 & 0 & 8 & 0 & 0 & 0 \\ 4 & 5 & 0 & 1 & 0 & 0 \\ 0 & 2 & 1 & 0 & 1 & 0 \\ 0 & 0 & 3 & 7 & 0 & 2 \\ 0 & 0 & 0 & 4 & 9 & 0 \\ 0 & 0 & 0 & 0 & 9 & 1 \end{bmatrix}$$



• Jacobian will be in a compressed row storage format

$$\mathbf{A} = \begin{bmatrix} 2 & 0 & 8 & 0 & 0 & 0 \\ 4 & 5 & 0 & 1 & 0 & 0 \\ 0 & 2 & 1 & 0 & 1 & 0 \\ 0 & 0 & 3 & 7 & 0 & 2 \\ 0 & 0 & 0 & 4 & 9 & 0 \\ 0 & 0 & 0 & 0 & 9 & 1 \end{bmatrix}$$

62 / 71



 Jacobian will be in a compressed row storage format

$$\mathbf{A} = \begin{bmatrix} 2 & 0 & 8 & 0 & 0 & 0 \\ 4 & 5 & 0 & 1 & 0 & 0 \\ 0 & 2 & 1 & 0 & 1 & 0 \\ 0 & 0 & 3 & 7 & 0 & 2 \\ 0 & 0 & 0 & 4 & 9 & 0 \\ 0 & 0 & 0 & 0 & 9 & 1 \end{bmatrix}$$

**A** values: 2 8 4 5 1 2 1 1 3 7 2 4 9 9 1 column: 1 3 1 2 4 2 3 5 3 4 6 4 5 5 6 row start: 1 3 6 9 12 14 16

- m Krylov iterations require (m+1) subspace vectors
- Storage requirement is  $\lceil (m+1)N \rceil$  for  $\mathbf{x} \in \mathbb{R}^{N \times N}$



• Jacobian will be in a compressed row storage format

$$\mathbf{A} = \begin{bmatrix} 2 & 0 & 8 & 0 & 0 & 0 \\ 4 & 5 & 0 & 1 & 0 & 0 \\ 0 & 2 & 1 & 0 & 1 & 0 \\ 0 & 0 & 3 & 7 & 0 & 2 \\ 0 & 0 & 0 & 4 & 9 & 0 \\ 0 & 0 & 0 & 0 & 9 & 1 \end{bmatrix}$$

**A** values: 2 8 4 5 1 2 1 1 3 7 2 4 9 9 1 column: 1 3 1 2 4 2 3 5 3 4 6 4 5 5 6 row start: 1 3 6 9 12 14 16

- m Krylov iterations require (m+1) subspace vectors
- Storage requirement is  $\lceil (m+1)N \rceil$  for  $\mathbf{x} \in \mathbb{R}^{N \times N}$

#### If we need 10 Krylov iterations to converge...

- 66 elements required for subspace vectors
- 72 total elements required for Jacobian and probability matrix

#### Parallel FANM Method



#### **Algorithm 5** FANM

- 1: k := 02: while  $||\mathbf{F}(\mathbf{u}^k)|| > \epsilon ||\mathbf{F}(\mathbf{u}^0)||$  do
- 2: While  $||\mathbf{F}(\mathbf{u}^*)|| > \epsilon ||\mathbf{F}(\mathbf{u}^*)||$  do
- 3:  $J(\mathbf{u}^k) \leftarrow AD(\mathbf{F}(\mathbf{u}^k))$  {Automatic differentiation}
- 4:  $\mathbf{J}(\mathbf{u}^k)\delta\mathbf{u}^k = -\mathbf{F}(\mathbf{u}^k)$  {Solve for the Newton correction with MCSA}
- 5:  $\mathbf{u}^{k+1} \leftarrow \mathbf{u}^k + \delta \mathbf{u}^k$
- 6:  $k \leftarrow k + 1$
- 7: end while
  - Modern FAD packages are parallelized using element-level assembly
  - This work will generate a parallel MCSA solver
  - Application of the Newton correction requires a parallel vector update
  - Convergence checks through parallel norm comptutation

## Research Proposal



- Experimental framework
- Methods verification
- Numerical experiments
- Challenge problem

## Experimental Framework



- Need a framework to generate linear and nonlinear systems in parallel
  - Linear algebra data structures
  - Parallel communication framework
  - Mesh generation and partitioning
  - Automatic differentiation and more...
- Choose the finite element method to leverage a finite element assembly engine

$$\begin{aligned} \mathbf{A}(\mathbf{u}) &= \mathbf{0}, \ \forall \mathbf{u} \in \Omega \\ \mathbf{B}(\mathbf{u}) &= \mathbf{0}, \ \forall \mathbf{u} \in \Gamma \\ \int_{\Omega} \mathbf{v}^{T} \mathbf{A}(\mathbf{u}) d\Omega + \int_{\Gamma} \bar{\mathbf{v}}^{T} \mathbf{B}(\mathbf{u}) d\Gamma &= \mathbf{0} \\ \int_{\Omega} \mathbf{C}(\mathbf{v}^{T}) \mathbf{D}(\mathbf{u}) d\Omega + \int_{\Gamma} \mathbf{E}(\bar{\mathbf{v}}^{T}) \mathbf{F}(\mathbf{u}) d\Gamma &= \mathbf{0} \end{aligned}$$

#### Monte Carlo Methods Verification



- Analytic solution to the heat equation for the linear methods
- Sequence of Navier-Stokes benchmarks for the nonlinear methods
  - Thermal convection cavity problem (De Vahl Davis, 1983)
  - Lid driven cavity problem (Ghia et al., 1982)
  - Backward-Facing step problem (Gartling, 1990)
- Tuning benchmark parameters varies the strength of nonlinearities

$$\rho \mathbf{u} \cdot \nabla \mathbf{u} - \nabla \cdot \mathbf{T} - \rho \mathbf{g} = \mathbf{0}$$

$$\nabla \cdot \mathbf{u} = 0$$

$$\rho C_{\rho} \mathbf{u} \cdot \nabla T + \nabla \cdot \mathbf{q} = 0$$

$$\mathbf{T} = -P\mathbf{I} + \mu [\nabla \mathbf{u} + \nabla \mathbf{u}^{T}]$$

$$\mathbf{q} = -k \nabla T$$





Domain Overlap Studies for the Parallel Neumann-Ulam Method

- Correlate domain overlap behavior to linear system properties
- Analyze random walk transport with respect to the operator:
  - Eigenvalues
  - Sparsity
  - Asymmetry
- Evaluate communication cost
- Evaluate memory cost



Domain Overlap Studies for the Parallel Neumann-Ulam Method

- Correlate domain overlap behavior to linear system properties
- Analyze random walk transport with respect to the operator:
  - Eigenvalues
  - Sparsity
  - Asymmetry
- Evaluate communication cost
- Evaluate memory cost

Domain Replication Studies for the Parallel Neumann-Ulam Method

- Feasibility from a memory perspective
- Impact on scalability
- Impact on load balancing



# Parallel Performance and Numerical Accuracy Studies for the MCSA Method

- Characterize Neumann-Ulam parameters required for good correction
- Feasibility of MCSA with full-clip Neumann-Ulam
- Performance for assymmetric systems
- Memory usage compared to Krylov methods



# Parallel Performance and Numerical Accuracy Studies for the MCSA Method

- Characterize Neumann-Ulam parameters required for good correction
- Feasibility of MCSA with full-clip Neumann-Ulam
- Performance for assymmetric systems
- Memory usage compared to Krylov methods

# Parallel Performance and Numerical Accuracy Studies for the FANM Method

- Feasibility for problems of interest
- Memory usage vs. Newton-Krylov methods
- Scalability vs. Newton-Krylov methods

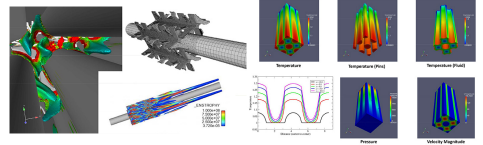
## Proposed Challenge Problem



- Problems of interest are coupled large scale problems in reactor physics
- The Consortium for Advanced Simulation of LWRs (CASL) is a modeling and simulation program aimed at:
  - Higher power uprates and efficiency
  - Higher burn-up
  - Predictive accident scenario analysis
- CASL and industry partners identified challenge problems:
  - Departure from nucleate boiling
  - Grid-to-rod-fretting
- Our challenge problem should reflect how this work aids the solution of these problems

# Proposed Challenge Problem





- CASL has leveraged the Drekar multiphysics code (SNL)
- Coupled fluid flow and heat transfer helps characterize many phenomena
- Drekar is massively parallel and leverages Newton-Krylov methods
- Propose using the largest Drekar problem to date as a challenge problem for the new Monte Carlo methods

Images source: www.casl.gov

## Conclusion

