

Spent Fuel and Waste Science and Technology (SFWST)









#### Machine Learning Surrogates for Fuel Degradation Processes in Nuclear Waste Repository Simulations

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Sandia ML DL Workshop Jul 18, 2023 SAND2023-05874C

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#### Acknowledgements and Disclaimers

#### Acknowledgements:

- Funding by the Department of Energy (DOE), Office of Nuclear Energy, Spent Fuel and Waste Science and Technology
- Disclaimer:
  - This presentation describes objective technical results and analysis. Any subjective views or opinions that might be expressed in the presentation do not necessarily represent the views of the U.S. Department of Energy or the United States Government.



- Computational Challenges in Handling Spent Nuclear Fuel
- Training Data for Machine Learning Surrogates
- ML Surrogates
  - kNNr
  - ANN
  - Neural ODEs
- Conclusions

#### Computational Challenges in Handling Spent Nuclear Fuel

### The US inventory of spent nuclear fuel is rapidly increasing

- 90,000 MTHM and increasing
- Pools have reached capacity limits
- Utilities have implemented dry storage
- Where facilities have shut down, some "stranded" fuel remains at independent spent fuel storage installations



**Storage Projections (2 models)** 

Adapted from Freeze et al. (2021, Figure 2-3)

## Our challenge is to provide realistic UO<sub>2</sub> degradation rates in underground nuclear waste repository simulations



Fuel Matrix Degradation Model (FMDM) adapted from Jerden et al. (2015)

Nuclear Waste Repository

#### Surrogate models provide a cheap-to-evaluate mapping between the model inputs and its outputs



Fuel surface boundary cell	Reactions, Diffusion	Bulk water boundary cell
-------------------------------------	-------------------------	--------------------------------

- Inputs are the environmental conditions along with the internal state at any point in time
  - Environmental Concentrations of CO<sub>3</sub><sup>2-</sup>, O<sub>2</sub>, Fe<sup>2+</sup>, and H<sub>2</sub>
  - Temperature T
  - Dose Rate, which is f(time, burnup)
  - Corrosion Layer Thickness
  - Internal concentration profiles
  - Relevant output is the UO<sub>2</sub> degradation rate (expressed as a flux)

#### Training Data for Machine Learning Surrogates

### Process model input parameters were sampled from expected ranges in reservoir simulations to generate training data

Parameter	Distribution	Min.	Max.
Init. Temp. (K)	Uniform	300	400
Burnup	Uniform	40	65
(Gwd/MTU)			
Env. CO <sub>3</sub> <sup>2-</sup> (mol/m <sup>3</sup> )	Log-uniform	10 <sup>-3</sup>	2x10 <sup>-2</sup>
Env. O <sub>2</sub> (mol/m <sup>3</sup> )	Log-uniform	10-7	10 <sup>-5</sup>
Env. Fe <sup>2+</sup> (mol/m <sup>3</sup> )	Log-uniform	10-3	10-2
Env. H <sub>2</sub> (mol/m <sup>3</sup> )	Log-uniform	10 <sup>-5</sup>	2x10 <sup>-2</sup>

- Same ranges used for training, validation, and testing data
- Ranges that span multiple orders of magnitude sampled with log-uniform distribution

### Training data is pulled from FMD Process model UO<sub>2</sub> Flux trajectories for randomly sampled initial conditions



### FMD surrogate model inputs aim to track the internal fuel cask state

![](_page_10_Figure_1.jpeg)

Fuel surface boundary cell	Reactions, Diffusion	Bulk water boundary cell
-------------------------------------	-------------------------	--------------------------------

- Inputs that do not require detailed knowledge of the fuel cask state
  - Environmental Concentrations of  $CO_3^{2-}$ ,  $O_2$ ,  $Fe^{2+}$ , and  $H_2$
  - Temperature T
  - Dose Rate, which is f(time, burnup)
  - Inputs that require detailed knowledge of the internal fuel cask state
    - Corrosion Layer Thickness
    - Internal concentration profiles

#### Prior results: kNNr and ANN

# kNNr has good but noisy predictions on the test data due to the local character of the representation

![](_page_12_Figure_1.jpeg)

- The inputs for each prediction are taken from test data (rather than from previously predicted points)
- More details in Debusschere et al. 2022 & 2023

# ANN has smoother predictions as it is a global functional approximation

![](_page_13_Figure_1.jpeg)

- The inputs for each prediction are taken from test data (rather than from previously predicted points)
- More details in Debusschere et al. 2022 & 2023

#### Neural ODE

# Neural ODEs approximate the derivative of the system state as a Neural Network

$$\frac{du}{dt} = f(u, \lambda) = NN$$

- Train NN based on data at equidistant timesteps<sup>1</sup>
- Predict with ODE Solver
- Hyperparameters to tune:
  - Number of layers
  - Number of nodes (neurons) per layer
  - Amount of training data

![](_page_15_Figure_8.jpeg)

<sup>1</sup> For more details, see Raissi et al. 2018

#### Data at regular time intervals is used to train a multi-step method

![](_page_16_Figure_1.jpeg)

• For more details, see Raissi 2018

#### Testing error plateaus by 20,000 epochs

![](_page_17_Figure_1.jpeg)

### Best results are obtained with 4 layers and 16 nodes per layer

![](_page_18_Figure_1.jpeg)

- Inputs: [CO<sub>3</sub><sup>2-</sup>], [H<sub>2</sub>], T, Dose Rate, CLT and UO2 Flux @ previous time, time
- Better accuracy when using more training data, but levels off at 500 runs
- Optimal accuracy for 4 layers, 16 nodes

# Neural ODE predicts the test data very well, and has fewer outliers than the regular ANN approach

![](_page_19_Figure_1.jpeg)

 Data integrated over a single time step only (all inputs taken from test data), to be consistent with kNNr and ANN results

#### Comparison kNNr – ANN – Neural ODE

# The neural ODE approach gives the lowest errors on the testing data

![](_page_21_Figure_1.jpeg)

Surroyate	mmse	inape_i	Surroyate	IIIIISe	паре
kNNr	0.26	1.4%	kNNr	0.11	29%
ANN	0.37	2.4%	ANN	0.12	14%
Neural ODE	0.18	0.80%	Neural ODE	0.086	1.9%

 Neural ODEs use the UO<sub>2</sub> flux at the current time step as input, whereas kNNr and ANN do not

#### Conclusions

#### Conclusions and Ongoing Work

- Machine Learning offers powerful ways to approximate the FMD process model outputs
- The Neural ODE formulation lends itself well to time advancement and gives
  very accurate results
- Adding more internal fuel surface state information may further improve accuracy
  - But will require additional surrogate predictions at each time step
- Surrogate models enable more detailed FMD dynamics in repository simulations
- Ongoing work focuses on determining an appropriate description of internal fuel cask state to balance accuracy and complexity

#### **Relevant References**

- 1. J. Jerden, K. Frey, and W. Ebert, "A Multiphase Interfacial Model for the Dissolution of Spent Nuclear Fuel," Journal of Nuclear Materials, 462, 135, <u>https://doi.org/10.1016/j.jnucmat.2015.03.036</u> (2015)
- 2. S. D. Sevougian et al., GDSA Repository Systems Analysis FY19 Update. SAND2019-11942R. Sandia National Laboratories, Albuquerque, New Mexico (2019).
- Freeze, G., E.J. Bonano, P. Swift, E. Kalinina, E. Hardin, L. Price, S. Durbin, R. Rechard, and K. Gupta, "Integration of the Back End of the Nuclear Fuel Cycle." SAND2021-10444. Sandia National Laboratories, Albuquerque, New Mexico (2021)
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- Bert J. Debusschere, D. Thomas Seidl, Timothy M. Berg, Kyung Won Chang, Rosemary C. Leone, Laura P. Swiler, and Paul E. Mariner, "Machine Learning Surrogate Process Models for Efficient Performance Assessment of a Nuclear Waste Repository," IHLRWM 22 (2022)
- J. Harvey et al.: Development of an Efficient Version of the Fuel Matrix Degradation Model, IHLRWM 22 (2022)
- 7. Maziar Raissi, Paris Perdikaris, George Em Karniadakis, "Multistep Neural Networks for Data-driven Discovery of Nonlinear Dynamical Systems," <u>https://doi.org/10.48550/arXiv.1801.01236</u> (2018).

#### **Additional Materials**

### US Department of Energy (DOE) Office of Nuclear Energy

- Spent Fuel and Waste Science and Technology (SFWST)
  - Research and Development (R&D) Campaign (2010 current)
- Mission
  - To identify alternatives and conduct scientific research and technology development to enable storage, transportation and disposal of used nuclear fuel and wastes generated by existing and future nuclear fuel cycles
- Mission work
  - Storage and transportation R&D
  - Disposal R&D

"Geological disposal remains the only long-term solution available."

National Research Council, 2001

#### Deep Geologic Disposal

- Several possible host rocks in US
- Investigating direct disposal of dual-purpose canisters (DPCs)

![](_page_27_Figure_3.jpeg)

![](_page_27_Picture_4.jpeg)

![](_page_27_Picture_5.jpeg)

28

overburden

salt dome

exploration level at a depth of 840m

backfilled drifts

emplacement drift

emplacement level at

a depth of 870m

(crushed salt)

#### Stages of a Deep Geologic Disposal Program

![](_page_28_Figure_1.jpeg)

#### Performance Assessment R&D

![](_page_29_Figure_1.jpeg)

#### **GDSA** Framework

- Geologic disposal safety assessment (GDSA) framework
- PFLOTRAN for multiphysics simulation
- Dakota for probabilistic performance assessment (PA)
- dfnWorks for DFN tools
- Open source
- Massively parallel
- Freely available (pa.sandia.gov)

#### The Need for Surrogate Models

# The Fuel Matrix Degradation (FMD) process model computes the degradation rate of spent nuclear fuel

- 1D reactive-transport model (diffusion only)
- Chemical (slow) and oxidative (fast) dissolution of UO<sub>2</sub> matrix
- Hydrogen peroxide production via alpharadiolysis
- Precipitation and dissolution of U(VI) (i.e., schoepite) corrosion layer at the fuel surface
- Arrhenius temperature dependence
- Complexation of uranium with carbonates
- Hydrogen as an oxidation sink (focused on fuel interface)
- Logarithmic spatial discretization for enhanced accuracy near the solid interfaces

![](_page_31_Figure_9.jpeg)

# Surrogate FMD models can alleviate cost of UO<sub>2</sub> flux computation in probabilistic repository assessments

![](_page_32_Figure_1.jpeg)

#### k-Nearest Neighbor regression (kNNr) interpolates between points in the training data closest to the query point

- Generalization of table lookup in higher dimensional setting
- Local approximation
- Inverse distance weighting means
  no training error
- Kd-Tree structure offers efficient table search
- Hyperparameters to tune:
  - Amount of training data
  - Number of nearest neighbors to use in interpolation

![](_page_33_Figure_8.jpeg)

# Artificial Neural Networks (ANN) approximate a function as a weighted combination of nonlinear functions

- Global functional
  approximation
- In each layer:
  - $\circ y_i = f(b_i + \sum w_{i,j} x_j)$
  - $\circ~$  ReLU activation function
- Prediction cost does not depend on amount of training data
- Hyperparameters to tune:
  - Number of layers
  - Number of nodes (neurons) per layer

![](_page_34_Figure_9.jpeg)

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#### Surrogate Models Based on Environmental and Global Inputs Only

### Hyperparameter tuning improves the performance for kNNr

![](_page_36_Figure_1.jpeg)

- Inputs: [CO<sub>3</sub><sup>2-</sup>], [H<sub>2</sub>], T, Dose Rate
- Accuracy improves with more training data
- Best accuracy with 9 subsamples per FMD process model run and 80 nearest neighbors

### ANN gives optimal results with 2 layers and 64 nodes per layer

![](_page_37_Figure_1.jpeg)

- Inputs: [CO<sub>3</sub><sup>2-</sup>], [O<sub>2</sub>], [Fe<sup>2+</sup>], [H<sub>2</sub>], T, Dose Rate
- Adding more layers or more nodes per layer does not significantly improve accuracy

## Despite minimal input, surrogates based on only environmental inputs approximate the actual $UO_2$ flux quite well

![](_page_38_Figure_1.jpeg)

- Overprediction between 10 and 1000 years
- kNNr approximations are noisier due to the nature of the local approximation

Surrogate	nrmse	mape
kNNr	0.48	44%
ANN	0.52	25%

# Model inputs that do not impact the fuel degradation rate much can be dropped

- Correlation between fuel degradation rate and O<sub>2</sub>, Fe<sup>2+</sup> is very small
- Training kNNr without these species gave better accuracy
- Fewer inputs also speeds up table lookup
- ANN not as impacted by extra inputs

![](_page_39_Figure_5.jpeg)

#### Most of the errors are very small

![](_page_40_Figure_1.jpeg)

- Histogram shows MSE averaged over each FMD process model run
- Some outliers with very low probability have MSE greater than 2 10<sup>-5</sup> (mol/m<sup>2</sup>/yr)<sup>2</sup>

#### Except for outliers, the agreement with test data is adequate

![](_page_41_Figure_1.jpeg)

#### Surrogate Models with CLT added

#### The corrosion layer thickness can be predicted with high accuracy

![](_page_43_Figure_1.jpeg)

- Prediction of CLT based on test data
- No time integration used

Surrogate	nrmse	mape_f
kNNr	0.26	1.4%
ANN	0.37	2.4%

### Adding the corrosion layer thickness as input gives dramatically better accuracy in fuel degradation rate prediction

![](_page_44_Figure_1.jpeg)

- Prediction of UO<sub>2</sub> flux based on test data
- No time integration used

Surrogate	nrmse	mape
kNNr	0.11	29%
ANN	0.12	14%

#### **Application to Repository Simulation**

### The surrogate models are demonstrated in a PFLOTRAN simulation of a generic shale repository reference case

- 2 x 41 drifts at a depth of 405m
- 10M grid cells
- 2000 4-PWR packages
  - 65 GWd/MTHM burn-up
  - 100 year Out of Reactor storage
- Sevougian et al. 2019

![](_page_46_Figure_7.jpeg)

## Surrogate models give more realistic prediction of fuel degradation rate than constant approximation

- Degradation starts after waste package is breached
- Fractional Degradation Rate (FDR) model assumes constant fractional rate of degradation
- ANN and kNNr surrogates provide higher fidelity by considering environmental inputs and changes in dose rate and temperature over time

![](_page_47_Figure_4.jpeg)

### Surrogate models are comparable in computational cost to constant fractional rate approximation

- Simulations use 1024 processors
- Transport is more expensive in ANN and kNNr runs to model transport of environmental species
- Running the full FMD process model on 2000 waste packages for 1M years would not be feasible in probabilistic (UQ) setting

![](_page_48_Figure_4.jpeg)

#### kNNr has best results with 8 – 12 nearest neighbors

![](_page_49_Figure_1.jpeg)

- Inputs: [CO<sub>3</sub><sup>2-</sup>], [H<sub>2</sub>], T, Dose Rate, CLT @ previous time, time step
- Better accuracy when using more training data
- Fewer nearest neighbors results in faster table lookup

#### Data conditioning improves the quality of the training data

- Remove FMD process model runs that are physically unrealistic
  - Runs that do not finish
  - Runs that stagnate at late time
  - Runs with Corrosion Layer Thicknesses that exceed physical domain size
- Log-transform data
- Subsample FMD process model runs
  - Random subset of points to reduce clustering in training data

![](_page_50_Picture_8.jpeg)

![](_page_50_Picture_9.jpeg)

### A variety of metrics evaluate different elements of the surrogate model accuracy

- (Normalized Root) Mean Squared Error
  - Good metric for engineering purposes

$$mse = \frac{1}{N} \sum_{i=1}^{N} (y_{pred,i} - y_{true,i})^{2}$$

$$nrmse = \frac{\sqrt{\frac{1}{N}\sum_{i=1}^{N}(y_{pred,i} - y_{true,i})^{2}}}{\frac{1}{N}\sum_{i=1}^{N}y_{true,i}}$$

- Mean Absolute Percentage Error
  - Highlights errors in small values

$$mape = \frac{1}{N} \sum_{i=1}^{N} \left| \frac{y_{pred,i} - y_{true,i}}{y_{true,i}} \right| \times 100$$

- Mean Absolute Error
  - Not as sensitive to outliers

$$mae = \frac{1}{N} \sum_{i=1}^{N} \left| y_{pred,i} - y_{true,i} \right|$$