



1	Efficient surrogate modeling methods for large-scale Earth system models based on
2	machine learning techniques
3	
4	Dan Lu ^{1,*} , Daniel Ricciuto ²
5	
6	
7	¹ Computational Sciences and Engineering Division, Climate Change Science Institute, Oak
8	Ridge National Laboratory, Oak Ridge, TN, USA;
9	² Environmental Sciences Division, Climate Change Science Institute, Oak Ridge National
10	Laboratory, Oak Ridge, TN, USA;
11	
12	
13	* Corresponding Author: Dan Lu, <u>lud1@ornl.gov</u>
14	
15	
16	
17	
18	
19	December 2018
20	For Publication in Geoscientific Model Development
21	





22	Abstract
23	Improving predictive understanding of Earth system variability and change requires data-model
24	integration. Efficient data-model integration for complex models requires surrogate modeling to
25	reduce model evaluation time. However, building a surrogate of a large-scale Earth system
26	model (ESM) with many output variables is computationally intensive because it involves a large
27	number of expensive ESM simulations. In this effort, we propose an efficient surrogate method
28	capable of using a few ESM runs to build an accurate and fast-to-evaluate surrogate system of
29	model outputs over large spatial and temporal domains. We first use singular value
30	decomposition to reduce the output dimensions, and then use Bayesian optimization techniques
31	to generate an accurate neural network surrogate model based on limited ESM simulation
32	samples. Our machine learning based surrogate methods can build and evaluate a large surrogate
33	system of many variables quickly. Thus, whenever the quantities of interest change such as a
34	different objective function, a new site, and a longer simulation time, we can simply extract the
35	information of interest from the surrogate system without rebuilding new surrogates, which
36	significantly saves computational efforts. We apply the proposed method to a regional ecosystem
37	model to approximate the relationship between 8 model parameters and 42660 carbon flux
38	outputs. Results indicate that using only 20 model simulations, we can build an accurate
39	surrogate system of the 42660 variables, where the consistency between the surrogate prediction
40	and actual model simulation is 0.93 and the mean squared error is 0.02. This highly-accurate and
41	fast-to-evaluate surrogate system will greatly enhance the computational efficiency in data-
42	model integration to improve predictions and advance our understanding of the Earth system.





43 1 Introduction

44	Improving predictive understanding of Earth system variability and change requires data-
45	model integration. For example, Bilionis et al. (2015) improved Community Land Model (CLM)
46	prediction of crop productivity after model calibration; Müller et al. (2015) improved the CLM
47	prediction of methane emission after parameter optimization; and Fox et al. (2009) and Lu et al.
48	(2017) improved the terrestrial ecosystem model predictive credibility of carbon fluxes after
49	uncertainty quantification. However, data-model integration methods are usually
50	computationally expensive involving a large ensemble of model simulations, which prohibits
51	their application to complex Earth system models (ESMs) with lengthy simulation time. To
52	reduce computational costs, surrogate modeling is widely used (Razavi et al., 2012; Gong et al,
53	2015; Ray et al., 2015; Huang et al., 2016, Lu et al., 2018; Ricciuto et al., 2018). The surrogate
54	model, which is a set of mathematical functions, approximates the actual simulation model based
55	on pairs of simulation model input-output samples, and then replaces the simulation model in the
56	data-model integration. As the ESMs evaluation is expensive, it is desired to use a limited
57	number of ESM simulation samples to build an accurate surrogate. As the surrogate model needs
58	to be calculated many times in data-model integration, it is required to build a fast-to-evaluate
59	surrogate. In this study, we use a very few simulation model runs to build an accurate and fast
60	evaluated surrogate system of a large scale problem based on advanced machine learning
61	methods.
62	In Earth system modeling, we usually need to build a surrogate system of many output

variables over large spatial and temporal domains. ESMs tend to be simulated in a regional or
global scale with many grid cells for several years, producing a large number of output variables.
In addition, ESMs are used to solve versatile scientific problems, so the quantities of interest





66	(QoIs) often change. Moreover, the development of a surrogate requires expensive ESM runs,
67	and a large number of runs are often needed to capture the complex model input-output
68	relationship. Therefore, it is reasonable to build a surrogate system for all possible model outputs
69	to reduce the efforts of rerunning ESMs for a new surrogate development when the QoIs change.
70	In this way, whenever we simulate the outputs in a new site or for additional sites, at a different
71	time or for a longer period, we can simply extract the information of interest from the large
72	surrogate system without spending extra efforts in building new surrogates, which significantly
73	saves the computational costs.
74	Building and evaluating a surrogate system of a large number of model outputs can be very
75	computationally intensive for almost all the surrogate methods. Polynomials and artificial neural
76	networks are widely used for surrogate modeling (Razavi et al., 2012; Viana et al., 2014).
77	Polynomial methods, such as polynomial regression and radial basis functions, need to solve
78	polynomial coefficients in the surrogate construction and to calculate matrix multiplications in
79	the surrogate evaluation. Using a p th-order polynomial to approximate a model with d
80	parameters, $M = (p+d)!/(p!d!)$ coefficients need to be solved, i.e., the number of coefficients
81	increases factorially fast with the parameter size and polynomial order. When $d=40$, a second-
82	order polynomial involves 861 coefficients and a third-order polynomial involves 12341
83	coefficients. ESMs have many uncertain parameters and a high-order polynomial is usually
84	needed to approximate complex ESMs, which can easily lead to a prohibitive number of model
85	evaluations, up to ~10 ⁵ , necessary to compute the polynomial coefficients. To reduce the
86	computational costs, some regularization techniques such as Bayesian compressive sensing have
87	been used (Sargsyan et al., 2014; Ricciuto et al., 2018). These regularization techniques can use a
88	few samples to solve a large number of coefficients (i.e., an underdetermined system) by





89	iteratively minimizing the L1 norm of the coefficient vector. But they usually perform
90	minimization once for one model output, so for a large model outputs problem, significant
91	computing effort is required. To reduce the computing burden in building polynomial-based
92	surrogates, we need to reduce the output dimensions.
93	Reducing the model output dimensions also improves computational efficiency in the
94	evaluation of the polynomial-based surrogates. For example, evaluating the third-order
95	polynomial-based surrogate of the model with 40 parameters and 300,000 outputs at 1 parameter
96	sample, we need to calculate two matrix multiplications where matrix A has the size $[1, M]$ and
97	B has the size [M, Nout] and $M = 12341$ and Nout=300,000. The surrogate evaluation takes about
98	90 seconds and most time is spent on loading the huge matrix. When Nout reduces to 20, the
99	surrogate evaluation quickly reduces to less than a second. Note that an ESM can easily have
100	more than 40 parameters and more than 300,000 model outputs. Even using the most advanced
101	supercomputers with GPUs, the data storage and loading are still a bottleneck. Thus, reducing
102	model output dimensions is necessary for both fast building and evaluating polynomial-based
103	surrogates.
104	Neural network (NN) assisted surrogate modeling also suffers from high computational
105	costs when applied to a large-scale problem with many QoIs. To approximate a complex ESM
106	with many outputs, a complicated NN with many wide hidden layers is usually needed to capture
107	the complex relationship between the model inputs and outputs, because each spatial and
108	temporal output variable is driven by different meteorological forcing such as air temperature,
109	humidity, wind speed, precipitation, and radiation. The full connections between nodes in the
110	input layer and the first hidden layer, between nodes of the hidden layers, and between nodes in
111	the last hidden layer and a large number of nodes on the output layer, involve a great amount of





112 NN weights and biases that need to be solved. For the same example discussed above, to 113 approximate the model with 40 parameters and 300,000 model outputs, an NN with two hidden 114 layers and each layer having 100 nodes has over 30 million weights and biases. Calculation of 115 these weights and biases requires many samples to train the NN for a good fit. Each training 116 sample involves one model evaluation. However, ESM simulation is time consuming, which 117 usually takes several hours or days and can be up to months or even years. A limited sample size 118 is not enough to train a deep and wide NN for convergence and a simple NN trained by a small 119 sample size may not capture underlying Earth systems accurately. Thus, reducing model output 120 dimensions is needed to advance the NN-based surrogate modeling. A small output size reduces 121 the width of the output layer and also simplifies the relationship between the model inputs and 122 outputs, so that a simple NN architecture can be appropriate and a small sample size can be 123 sufficient to accurately train the simple NN. In addition, a simple NN can also be fast evaluated 124 with small weight matrix multiplications. 125 In this work, we propose to use singular value decomposition (SVD) to reduce model

126 output dimensions, so as to improve the computational efficiency in both building and evaluating 127 the surrogates. ESM outputs usually show periodic changes along time and strong correlations 128 between locations, which promises a fast decay of singular values. So, we can use a small 129 number of singular value coefficients to capture a great amount of output information, enabling a 130 significant output dimension reduction. We use the NN for surrogate modeling, because 131 compared to polynomial methods, NNs have shown less difficulty in fitting highly nonlinear and 132 discontinuous functions which are usually observed in ESMs response surfaces. For example, 133 carbon flux state variables, such as gross primary productivity (GPP), are strongly affected by 134 vegetation related parameters. When the parameter samples cause zero vegetation growth, GPP





has zero values. Whereas when the parameter samples cause high vegetation growth, GPP has
large positive values. This leads to a discontinuous GPP response surface jumping from zeros to
nonzeros.

138 NNs theoretically can fit any functions, but their practical performance strongly depends 139 on the NN's architectures and hyperparameters. NN has many hyperparameters such as the 140 number of layers, number of nodes in each layer, type of activation functions, and learning rate 141 of the stochastic gradient descent optimization. A slight change in the hyperparameter value can 142 result in dramatically different NN performance. Development of a high-performing NN is time-143 intensive and usually requires trial-and-error tuning by machine learning experts. In this work, 144 we use Bayesian optimization techniques to optimize the NN architecture and hyperparameters 145 so as to produce an accurate NN model for the training data. Bayesian optimization searches the 146 hyperparameter space to iteratively minimize the validation errors of the NN by balancing 147 exploration and exploitation (Shahriari et al., 2016). Researches suggested that Bayesian 148 hyperparameter optimization of NNs is more efficient than manual, random, or grid search with 149 better overall performance on test data and less time required for optimization (Bergstra et al., 150 2011; Snoek et al., 2012). Bayesian optimization involves a large ensemble of NN fittings and it 151 is a sequential model-based optimization, thus, fast training of the NN models is important. Our 152 proposed SVD method can simplify the NN architecture so as to advance the NN training and 153 improve the Bayesian optimization performance. 154 In this effort, we propose an SVD-enhanced, Bayesian-optimized, and NN-based surrogate

method and aim to build an accurate and fast-to-evaluate surrogate system of a large-scale model using a few model runs, so as to improve computational efficiency in surrogate modeling and thus advance the data-model integration. We apply the method to a simplified land model in the





158	Energy Exascale Earth System Model (sELM) to improve the model predictive capability of
159	carbon fluxes. We build a surrogate system of 42660 model output variables which are annual
160	GPPs at 1422 locations simulated for 30 years. The sELM is a regional-scale terrestrial
161	ecosystem model that simulates terrestrial water, energy, and biogeochemical processes in
162	terrestrial surfaces. Simulation of sELM is important for improving our understanding of
163	ecosystem responses to climate change. However, sELM requires lengthy times for hydrologic
164	and carbon cycle equilibration, and these high computational costs limit the affordable number of
165	simulations in data-model integration thus resulting in poor model performance. The proposed
166	machine learning assisted surrogate method makes the sophisticated data-model integration
167	computationally feasible and promises an improvement of the sELM predictions.
168	The major contributions of this work are (1) using SVD to reduce model output
169	dimensions so as to improve computational efficiency in both building and evaluating an
170	accurate surrogate of a large-scale ESM; (2) using Bayesian optimization techniques to fast
171	generate an accurate NN-based surrogate; and (3) applying the proposed method to build a large
172	surrogate system of a regional-scale ESM to advance data-model integration. To our knowledge,
173	the method of using SVD to enhance surrogate modeling is novel and we have not seen the
174	application of Bayesian optimization to improve NN-based surrogates in Earth system modeling.
175	The paper is organized as follows. In section 2, we first describe the sELM, the model
176	parameters and the QoIs we build surrogates for; following that, we introduce the SVD, NNs,
177	and Bayesian optimization methods. In section 3, we apply the methods to the sELM and analyze
178	the surrogate accuracy. In section 4, we discuss strategies to improve surrogate accuracy and
179	investigate our method's performance in the application of these strategies. In section 5, we end
180	this paper by drawing our conclusions.





181 2 Materials and Methods

182 **2.1 Description of sELM and related parameters**

183 We developed a simplified version of Energy Exascale Earth System (E3SM) land model 184 (ELM), or sELM, to simulate carbon cycle processes relevant for Earth system models in a 185 computationally efficient framework. This framework allows us to perform large regional 186 ensembles that are computationally infeasible using offline land surface models such as ELM. 187 sELM is a combination of model elements from the Data Assimilation Linked Ecosystem 188 Carbon model (DALEC; Williams et al., 2005) and the Community Land Model version 4.5 189 (CLM4.5; Oleson et al., 2013). sELM consists of five process-based submodels that simulate 190 carbon fluxes between five major carbon pools using 49 overall parameters. Based on previous 191 sensitivity analysis using ELM (Ricciuto et al., 2018), this study considers the most sensitive 192 eight parameters associated with four out of the five submodels. We summarize all five process-193 based submodels and their interactions below and in Figure 1. 194 sELM consists of five major submodels: photosynthesis, autotrophic respiration, 195 allocation, deciduous phenology, and decomposition. Photosynthesis is driven by the aggregate 196 canopy model (ACM) from the DALEC, which itself is calibrated against the soil-plant-197 atmosphere model (Williams et al., 2005). ACM predicts GPP as a function of carbon dioxide 198 concentration, leaf area index, maximum and minimum daily temperature, and 199 photosynthetically active radiation. Here the GPP predicted by ACM is modified by BTRAN, 200 which reduces GPP when soil water is insufficient to support transpiration. Because sELM does not predict soil moisture, BTRAN is calculated in a full ELM simulation and is fed into sELM as 201 202 an input. ACM shares one parameter, the leaf carbon to nitrogen ratio (*leaf C:N*), with the





203 autotrophic respiration model and employs an additional parameter, the specific leaf area at the

top of the canopy (*slatop*).

205	The remaining four submodules are based on ELM. The autotrophic respiration model
206	computes the growth and maintenance respiration components and is controlled by four
207	parameters, the <i>leaf C:N</i> , the fine root carbon to nitrogen ratio (<i>froot C:N</i>), the base rate of
208	maintenance respiration (<i>br_mr</i>), and temperature sensitivity for maintenance respiration
209	$(q10_mr)$. The allocation model partitions carbon to several vegetation carbon pools following
210	those in ELM: leaves, fine roots, live stem, dead stem, live coarse roots and dead coarse roots. In
211	the allocation model, we only consider one parameter, the fine root to leaf allocation ratio
212	(froot_leaf). The deciduous phenology model is used to predict the timing of budbreak and
213	senescence. It considers two parameters, the critical day length to initiate autumn senescence
214	(crit_dayl) and the number of accumulated growing degree days needed to initiate spring leaf-out
215	(crit_onset_gdd). The last submodel is a decomposition model that simulates heterotrophic
216	respiration and the decomposition of litter into soil organic matter using the converging trophic
217	cascade framework as in the CLM4.5 (Oleson et al., 2013). Because this study focuses on plant
218	carbon uptake, no uncertain parameters are considered in the decomposition model. In sELM,
219	nutrient feedbacks are not represented explicitly, however a constant nitrogen limitation factor is
220	included to downregulate photosynthetic uptake.
221	The sELM can simulate several carbon state and flux variables as shown in Figure 1 with
222	green shapes. GPP, which represents the total plant carbon uptake, is considered in this study.
223	Here we use sELM to predict annual GPP in deciduous forest systems in the eastern region of the
224	United States for 30 years between 1981-2010. The carbon state variables are spun up to steady
225	state by cycling the GSWP3 input meteorology (Kim et al., 2017) from 1981-2010 for 5 cycles,





226	and the 6 th cycle is used as the output for our surrogate modeling study. The region of interest
227	covers 1422 land grid cells (locations) as shown in Figure 2. Given 30 outputs at each location
228	(annual values over 30 years), a total of 42660 GPP variables are simulated. The model uses one
229	plant functional type and the phenological drivers such as air temperature, solar radiation, vapor
230	pressure deficit, and CO ₂ concentration are used as boundary conditions. One regional sELM run
231	takes about 24 hours on a single processor, which although much faster than ELM is still
232	computationally too expensive to be directly used in model-data integration studies. To improve
233	the computational efficiency in generating the sELM simulation samples to develop the surrogate
234	model, we use high performance computing to perform an ensemble of 2000 sELM model
235	simulations in parallel. The 2000 parameter input samples are randomly drawn from the
236	parameter space defined in Figure 3. The numerical ranges of these parameters are designed to
237	reflect their average values and broad uncertainties associated with the temperate deciduous
238	forest plant functional type. The output samples are sELM simulated GPPs at the 1422 locations
239	for 30 years. In the surrogate modeling, part of the 2000 input-output samples are used for
240	developing the surrogate and part of them are used to evaluate the surrogate accuracy, as
241	discussed in section 3.

242 **2.2 Efficient surrogate modeling methods**

In this section, we introduce our SVD-enhanced, Bayesian-optimized, and NN-based surrogate methods. We first describe the SVD for reducing data dimensionality, then introduce the NN techniques for building a surrogate model, and last depict the Bayesian optimization algorithm for producing a high-performing NN-based surrogate.





247 2.2.1 Singular value decomposition for data compression

248	We build a surrogate system of model outputs by fitting a data matrix whose columns are
249	output variables and rows are output samples. For a model with 100000 output variables, the
250	columns of this matrix span a 100000-dimensional space. Encoding this matrix on a computer
251	takes quite a lot of memory and evaluating this matrix takes a large number of calculations. We
252	are interested in approximating this matrix with some low-rank matrix but remaining its most
253	information, so as to reduce data transfer and accelerate matrix calculation.
254	Singular value decomposition (SVD) decomposes a matrix A with size $m \times n$ into three
255	other matrices, $\mathbf{A} = \mathbf{U}\mathbf{S}\mathbf{V}^T$, where \mathbf{U} is an $m \times m$ orthogonal matrix, \mathbf{V} is an $n \times n$ orthogonal
256	matrix, and S is an $m \times n$ diagonal matrix saving singular values in descending order on the
257	diagonal. Truncated SVD keeps the K largest singular values and corresponding K column
258	vectors of U and <i>K</i> row vectors of \mathbf{V}^T to form $\widetilde{\mathbf{A}} = \mathbf{U}_K \mathbf{S}_K \mathbf{V}_K^T$. The <i>K</i> -rank matrix $\widetilde{\mathbf{A}}$ has proven to
259	be the best approximation of \mathbf{A} in minimizing the Frobenius norm of the difference between \mathbf{A}
260	and $\widetilde{\mathbf{A}}$ under the constraint of rank($\widetilde{\mathbf{A}}$) = <i>K</i> . In addition, the total of the first <i>K</i> singular values
261	divided by the sum of all the singular values is the percentage of information that those singular
262	values contain. For example, if we want to keep 90% of the data information, we just need to
263	compute sums of K largest singular values until we reach 90% of the sum and discard the rest.
264	By dropping all but a few singular values and then recomputing the approximated matrix, the
265	SVD technique compresses the data information and reduces data dimensions. When the matrix
266	A shows strong correlations between columns (variables), a low-rank matrix \widetilde{A} can make a very
267	accurate approximation of A.
268	In this study, we use SVD to reduce training data dimensions. The training data matrix A

269 [m, n] for surrogate construction contains model output samples information. n columns are





- 270 output variables (e.g., the 42660 temporal and spatial GPPs in this work) and *m* rows are the
- samples of these variables (e.g., the sELM simulation results of the 42660 GPPs for the m
- 272 parameter samples), and usually $n \gg m$ for expensive ESMs with many outputs. In
- implementation, we first perform truncated SVD to get low-rank matrices $\mathbf{U}_{K}[m, K]$, $\mathbf{S}_{K}[K, K]$,
- and $\mathbf{V}_{K}^{T}[K, n]$ with $K \ll n$, we then use the low-dimensional dataset $(\mathbf{V}_{K}^{T} \mathbf{A}^{T})^{T}$ with reduced size m
- 275 $\times K$ as training data to build the surrogate model of the K largest singular value coefficients.
- 276 Next, we evaluate the surrogate model at q new data points to get results \mathbf{Y}_{new} with size $q \times K$.
- 277 Lastly, we transform the predicted values back to its original size $q \times n$ through $\mathbf{Y}_{new} \mathbf{V}_K^T$ to obtain
- the surrogate approximation of the *n* variables at the *q* new data points.

279 **2.2.2** Neural networks for surrogate modeling

280 Artificial neural networks (NNs) consist of fully connected hierarchical layers with nodes 281 which can be flexibly used for function approximation (Yegnanarayana, 2009). The first layer is 282 the input layer and each node in the input layer represents one model input variable. The last 283 layer is the output layer and each node in the output layer represents one model output variable. 284 The layers between input and output layers are hidden layers which are used to approximate the 285 relationship between model inputs and outputs. When the relationship is complex, a complicated 286 NN with many wide hidden layers is usually needed. The input layer first assigns model 287 parameter values to its nodes. Then each node in the first hidden layer takes multiple weighted 288 inputs, applies the activation function to the summation of these inputs, and calculates the node's 289 value. Next, the second hidden layer takes the values on the first hidden layer nodes as inputs 290 and calculates its nodes' values in the same way. This process moves forward till we get values 291 of all nodes in the output layer, i.e., obtaining NN predictions for the given model parameter 292 input values. The nodes in each layer are fully connected to all the nodes in its previous and





293 subsequent layers. Each of these connections has an associated weight and bias. A complicated 294 NN results in a large number of weights. By tuning these weights and biases based on some 295 training data, we improve the NN approximation of the underlying simulation model. 296 NN uses stochastic gradient descent (SGD) method to optimize its weights and biases 297 (Bottou, 2012). SGD optimizes variables by minimizing some loss function based on the 298 function's gradients to these variables. The loss function is usually defined as the mean squared 299 error (MSE) between the NN predictions and model simulations for the same set of model 300 parameter samples in the training data. SGD iteratively updates the optimized variables at the 301 end of each training epoch. In the process, the learning rate, which specifies how aggressively 302 the optimization algorithm jumps between iterations, greatly affects the algorithm's performance 303 and has to be tuned. A small learning rate will take a long time to reach the optimum causing a 304 slow convergence, whereas a big learning rate will bounce around the optimum causing unstable 305 results and a difficult convergence. Using SGD to optimize a complex NN with many weights 306 requires a great amount of computational efforts and has difficulty in convergence. First, many 307 training data are required to tune a large number of weights. Small training data can easily cause 308 over-fitting, i.e., the NN "perfectly" fits the training data but performs badly on new data, thus 309 deteriorating the NN prediction accuracy. In addition, a large number of weights involve massive 310 matrix calculations in evaluating the loss function, slowing down the training process. 311 Furthermore, a complicated NN has difficulty in convergence and can easily get stuck in local 312 minima. In this work, we use SVD to reduce the model output dimensions, so as to decrease the 313 number of nodes in the output layer and simplify the NN architecture, thus reducing the size of 314 the weights and enabling a reasonable NN training from small training data, and ultimately 315 improving the computational efficiency.





316 2.2.3 Bayesian optimization algorithm for NN hyperparameter optimization

317 NN involves a lot of hyperparameters that dramatically affect its performance such as the 318 number of layers, the number of nodes in each layer, and the learning rate of the SGD algorithm. 319 Hyperparameter optimization is needed to produce a high-performing NN. This requires 320 optimizing an objective function f(x) over a tree-structured configuration spaces $x \in X$, where 321 some leaf variables (e.g., the number of nodes in the third hidden layer of an NN) are only well 322 defined when branch variables (e.g., a discrete choice of how many layers to use) take particular 323 values. In addition, the optimization not only optimizes discrete and continuous variables, but 324 also simultaneously choose which variables to optimize. When the NN is used for surrogate 325 modeling, the objective function is the NN accuracy of predicting some validation data. In this 326 case, the f(x) does not have a simple closed form but can be evaluated at any arbitrary query 327 point x in the configuration space. For such optimization problem, a sequential search method is 328 needed, besides some inefficient grid search and random search approaches (Bergstra and 329 Bengio, 2012). The sequential search method starts with some random points in the search space, 330 and then iteratively evaluates new points based on NN predictions on previously evaluated 331 points. After N evaluations, we choose the optimal combination of the hyperparameters resulting 332 in the highest NN prediction accuracy. Among the sequential search algorithms, Bayesian 333 optimization is able to take advantage of full information provided by the history of the 334 optimization to improve the search efficiency. 335 Tree-structured Parzen estimator (TPE) and Gaussian process are two widely used 336 Bayesian optimization algorithms (Shahriari et al., 2016; Bardenet and Kegl, 2010; Niranjan et 337 al., 2010; Snoek et al., 2012). In comparison to the Gaussian process, TPE works well for all 338 types of NN hyperparameter variables, is robust to NN randomization, has a fast calculation and





- 339 a straightforward implementation without associated hyperparameters specification (Bergstra et
- al., 2011). In this work, we use the TPE algorithm for NN hyperparameter optimization.
- **341 3 Results**

342 In this section, we present the results of building the surrogate system of 42660 GPP 343 variables of sELM. First, we demonstrate that our method using SVD can efficiently build and 344 evaluate a large surrogate system by comparing the results with and without application of SVD. 345 We then investigate the influence of NN's architecture on surrogate performance and show that 346 our method using hyperparameter optimization can fast generate an accurate NN. Last, we 347 evaluate surrogate accuracy on the large-scale spatial and temporal GPPs. 348 We consider three sets of data, the training data for fitting the NN, the validation data to 349 detect overfitting in the NN training and to select the best-performing NN in the hyperparameter 350 optimization, and the test data to evaluate the NN prediction accuracy. Each data set contains 351 pairs of parameter and GPP samples. The parameter samples are randomly drawn from the 352 parameter space defined in Figure 3. To assess the effectiveness of our proposed surrogate 353 method for a small data set, we consider only 20 training data (Figure 3). The validation data is 354 chosen as 0.3 fractions of the training data. The NN model will not train on the validation data 355 but evaluate the loss function on them at the end of each epoch. In each epoch, the training data 356 is shuffled, and the validation data are always selected from the last 0.3 fraction. Precisely, we 357 only use 14 samples to tune NN weights. Attribute to shuffling, these 14 samples can be a 358 different subset from the 20 training data in each epoch, thus we sufficiently explore the limited 359 20 data information for building the surrogates. We use 1000 test data (Figure 3) to evaluate the 360 NN prediction accuracy, which makes a reasonable assessment of our proposed method within 361 an affordable computational cost. Note that the 1000 test data are not needed for building the





- surrogates but used to demonstrate the effectiveness and efficiency of our method. When using
 our method to build the surrogates of the 42660 GPPs, only 20 sELM model simulations are
 used.
- 365 We define the loss function as the mean squared error (MSE) between the NN predictions 366 and the sELM simulations based on the parameter samples for training. We use Adam algorithm 367 (Kingma and Ba, 2015) for stochastic optimization of NN and run it for 800 epochs to minimize 368 the loss function and update NN weights. Adam has been shown a superior stochastic 369 optimization algorithm in training NN (Basu et al., 2018). There is no right answer for the 370 optimal number of epochs. A small number of epochs could result in underfitting and a large 371 number of epochs may lead to overfitting. Here we consider a large number of epochs and in the 372 meantime use early stopping to avoid overfitting. During the training, when there is no 373 improvement of loss functions for the validation data in 100 epochs, we stop the training and 374 choose the weights at the epoch resulting in the smallest loss function of the validation data as 375 the optimal weights and the associated NN as the best trained NN under the given setting. 376 We then use the trained NN to predict the 1000 test data and compare the predictions with 377 the corresponding sELM simulation results to evaluate the NN accuracy. We define two metrics 378 for evaluation, the MSE and the coefficient of determination. The MSE computes the expected 379 value of the squared prediction errors; the small the MSE value is, the better the prediction. The 380 coefficient of determination, also called R^2 score, measures how well the unobserved data are 381 likely to be predicted by the NN model. Denote \hat{y}_i as the NN prediction of the *ith* sample and y_i 382 as the corresponding sELM simulation, the R^2 score estimated over N_s samples is defined as $R^2 = 1 - \frac{\sum_{i=1}^{N_s} (y_i - \hat{y}_i)^2}{\sum_{i=1}^{N_s} (y_i - \bar{y}_i)^2}$, where $\bar{y} = \frac{1}{N_s} \sum_{i=1}^{N_s} y_i$. Best possible value of R^2 score is 1.0, indicating 383 that the NN can perfectly predict the test data. R^2 score can be negative indicating the model is 384





arbitrarily poor. A constant model gets a R^2 score of 0.0. Compared to MSE, the R^2 score

386 considers the variability of the data which provides a more reasonable measure.

387 **3.1** SVD reduces data dimensionality and improves surrogate efficiency

388 We consider two scenarios when building the surrogate system of the 42660 GPP outputs;

389 Case I: building the surrogates of reduced data after SVD, and Case II: building the surrogates of

390 all GPPs directly. In Case I, we first apply SVD to reduce the training data dimensionality, then

391 build surrogates of the singular value coefficients, and last transfer the surrogate system back to

the original QoIs (i.e., the 42660 GPP variables).

The goal of this study is to develop a surrogate method that builds an accurate surrogate system with small training data, so as to reduce the computational costs in simulating the expensive ESMs. To demonstrate the effectiveness and efficiency of our method, we compare

the surrogate performance of the two cases in predicting the 1000 test data from two aspects: (1)

397 for the same number of training data, the predictive accuracy of the two surrogates, and (2) the

398 number of training data used to achieve the similar predictive accuracy.

Figure 4 shows the singular value decay of decomposition of the training data matrix

400 having 20 samples and 42660 GPP variables. The figure indicates that the singular values decay

401 very fast. The first 2 singular values drop about 1 magnitude, and the first 5 singular values can

402 capture 97% information of the training data matrix. To choose a suitable number of singular

403 value coefficients (Nsvd) to compress the training data and build a surrogate for, we consider a

404 series of Nsvds, where Nsvd=1, 5, 10, 15, and 20, and investigate their impact on NN

405 performance. To make a fair comparison, the same NN architectures are used for all Nsvd cases.

406 We consider a simple NN with 2 hidden layers and each hidden layer has 10 nodes. Figure 5

407 shows the prediction performance of the NNs based on the 20 training data. The figure indicates





408	that with considering only 1 singular value coefficient, the averaged MSE of the predictions is
409	about 0.053, and the NN model can fit the sELM simulation data well with the R^2 score of 0.83.
410	When 5 singular value coefficients are considered, the NN prediction accuracy improves with the
411	MSE of 0.02 and the R^2 score of 0.93. After Nsvd=5, the MSE and R^2 score have minor changes,
412	suggesting that for the limited 20 training data, Nsvd=5 is a good choice to compress the GPPs
413	and build a surrogate for. At this time, the surrogate error becomes dominant compared to the
414	SVD approximation error and including more than 5 singular value coefficients would barely
415	improve the NN prediction unless more training data are included to reduce the surrogate error.
416	In the following, we consider Nsvd=5 in Case I and compare its surrogate prediction
417	performance with Case II which builds surrogates for all GPPs directly.
418	In Case I, our method is able to use 20 training data to build a highly accurate surrogate of
419	42660 GPP variables with a small MSE of 0.02 and a high R^2 score of 0.93. The detailed NN
420	performance is explained in Figure 6(a) where the training and validation loss decays in building
421	the surrogates of the 5 singular value coefficients are plotted. The figure indicates that the loss
422	functions of the two data sets have similar decay, decreasing dramatically at the first 10 epochs
423	and then slowly decreasing to the end of training. The closely overlapped two lines in Figure $6(a)$
424	suggest that the trained NN captures the relationship between sELM inputs and outputs pretty
425	well and can give reasonable predictions of GPPs for a given parameter sample.
426	To make a fair comparison, we use the same NN architecture in Case II as in Case I except
427	that the output layer of NN in Case II has all the 42660 GPPs and the output layer in Case I has
428	only 5 singular value coefficients. Figure 6(b) indicates that the simple NN with 20 hidden nodes
429	is not sophisticated enough to capture the complex relationship between the 8 inputs and 42660
430	outputs. As we can see in Figure 6(b), both training and validation losses are relatively high





431 suggesting an underfitting. The validation loss is always larger than the training loss suggesting 432 that the fitted NN does not generalize well and may result in poor performance in predicting new 433 data. Figure 7 shows R^2 scores of Case II in predicting the 1000 test data. The figure indicates 434 that the simple NN trained by 20 data in Case II has a very poor prediction accuracy with the R^2 score of only 0.05, close to a constant model's performance with a zero R^2 score. However, with 435 436 the same NN trained by the same 20 data, our SVD-based surrogate method can achieve a high 437 prediction accuracy with the R^2 score of 0.93. This demonstrates our method's capability in using 438 a few training samples to build an accurate surrogate model, greatly reducing the computational 439 costs in generating the expensive model simulation data. 440 On the other hand, the poor performance in Case II suggests that a wider and deeper NN is 441 needed when we consider the large outputs directly. We thus increase the nodes of each hidden 442 layer to 100 and use this complex NN with total 200 hidden nodes to approximate the 443 relationship of the 8 inputs and 42660 outputs in Case II. This complex NN blows up its 444 parameters (including weights and biases) to 4.3 million from 255 in Case I. To fit this wide NN 445 and calibrate its large parameters, 20 training data are way too small to get a reasonable fit. No 446 matter how we adjust the NN hyperparameters, we cannot get a stable solution in training. We 447 then increase the training data to 50, Figure 6(c) shows that the increased data greatly decrease 448 the training and validation losses and the validation loss is slightly higher than the training loss, 449 implying a good fit. Figure 7 indicates that the complex NN with 200 hidden nodes trained by 50 450 data in Case II significantly improves the prediction accuracy with the R^2 score of 0.73. 451 However, Case II's predictive performance is still worse than Case I which has the R^2 score of 452 0.93. We keep increasing the training data (Ntrain) to 100 and 200 in Case II. Figure 6(d) and (e) 453 indicate that the increase of training data brings the validation loss closer and closer to the





454	training loss making the fitted NN represent the underlying sELM better and better. Figure 7
455	shows that the nicely fitted NNs trained by large Ntrains lead to a high prediction accuracy. With
456	Ntrain=100, the R^2 score is about 0.89, and with Ntrain=200, the R^2 score is up to 0.95.
457	However, compared to Case I using 20 training data to get predictive R^2 score of 0.93, Case II
458	uses near 200 data to get the similar accuracy, increasing 10-fold computational costs. Note that,
459	each training data involves one sELM simulation and one regional sELM run takes about 24
460	hours on one processor. Thus, our SVD-based surrogate method greatly improves computational
461	efficiency in the accurate surrogate modeling.
462	Our method, in the means of simplifying NN architecture through data compression, not
463	only reduces the training data but also decreases the training time. Using 20 data to train a simple
464	NN with 255 parameters, our method takes about 4 seconds. In comparison, the traditional
465	surrogate method without data compression spends a great effort in training the complex NN
466	with 4.3 million parameters. As shown in Figure 7, Case II takes 270 seconds to fit the NN based
467	on 50 training data and 967 seconds for the 200 training data, showing a linear increase in
468	computing time. The long training time leads to high computational costs in NN hyperparameter
469	optimization where massive NN training are involved in searching the wide hyperparameter
470	space for a high-performing NN model, as discussed in the following section 3.2.
471	3.2 NN's hyperparameter optimization improves surrogate accuracy
472	NN has a large number of hyperparameters. Here we adjust 5 hyperparameters and use
473	Case I to investigate their influence on surrogate prediction accuracy. The 5 hyperparameters are,
474	the number of hidden layers (L) where we consider the most 3 hidden layers, the number of
475	nodes in hidden layer 1 (N1), in hidden layer 2 (N2), and in hidden layer 3 (N3), and the learning
476	rate (lr) of Adam optimization algorithm. We consider the following choices: L= $\{2, 3\}$, N1= $\{10, 10\}$





477	20, 40, 60, 80, 100}, N2={10, 20, 40, 60, 80, 100}, N3={0, 10, 20, 40, 60, 80, 100}, and
478	lr=U[0.001, 0.1]. The first four hyperparameters are discrete variables and the last one, lr, is a
479	continuous variable with uniform distribution. The choice of L determines the selection of N3
480	showing a tree-like structure. We use tree-structured Parzen estimator (TPE) to search the 5
481	hyperparameter space and find a set of values that gives the best-performing NN. We fix the
482	activation function as ReLU (Agarap, 2018) which has been widely used and shown to produce
483	good NN predictions.
484	We use TPE to evaluate 100 sets of hyperparameters and the one giving the best validation
485	score, i.e., the smallest MSE on validation data, is chosen as the optimal hyperparameters.
486	Results indicate that the combination of N1=10, N2=10, N3=0, and lr=0.08 gives the best
487	validation score. To investigate the impact of hyperparameters on NN prediction accuracy, we
488	show the 100 sets of hyperparameters and their resulting R^2 scores in predicting the 1000 test
489	data in Figure 8. The figure indicates that different hyperparameter values result in dramatically
490	different NN performance. The prediction R^2 scores range from 0.66 to 0.93 where 32
491	hyperparameter sets have the R^2 scores over 0.90. The selected optimal NN producing the
492	smallest MSE on the validation data also gives the best prediction performance on the test data
493	with the R^2 score of 0.93. It is desired that the best NN model chosen by validation data gives the
494	best predictions, however, in practice it is not always the case, especially when the prediction
495	data deviates a lot from the validation data. Extrapolation is always a difficulty in surrogate
496	modeling and several researches are going on to improve the extrapolation accuracy (Gal, 2014).
497	Although NNs perform significantly different with different combination of
498	hyperparameters, the TPE algorithm can efficiently find the high-performing NNs based on
499	previous samples information. As shown in Figure 8, good-performing NNs prefer simple





500	architectures with 2 hidden layers, e.g., most blue lines have N3 of 0. After TPE finds a good
501	architecture of N1=10 and N2=10, it samples around this architecture in the hyperparameter
502	space to fine tune the learning rate till finds the most suitable lr of 0.08. This work considers 5
503	hyperparameters with limited choices, increasing the dimensions and possible choices of the
504	hyperparameters would make the search more thorough and could produce a better-performing
505	NN. Our surrogate method with SVD can accelerate the optimization process by reducing the
506	NN training time.
507	3.3 Evaluation of surrogate accuracy on large-scale spatial and temporal data
508	We, using only 20 expensive sELM runs, fast build an accurate surrogate system of 42660
509	GPPs at 1422 locations for 30 years. Therefore, for a data-model integration problem with the
510	QoIs within the spatial and temporal ranges, we can directly extract the information of interest
511	from the surrogate system to advance the analysis. The best-performing NN generated from our
512	method gives an overall accurate prediction of the 42660 GPPs with averaged MSE of 0.02 and
513	R^2 scores of 0.93. When using the subset of the surrogate system for data-model integration
514	studies, it is desired to analyze the surrogate accuracy at individual locations for specific times.
515	Figure 9 shows averaged R^2 scores over 30 years at 1422 locations. The figure indicates
516	that the surrogate accuracy is not uniformly good for all the locations. We observe that most
517	locations have R^2 scores above 0.9 with the best R^2 score of 0.96, and about 100 locations have
518	R^2 scores below 0.90 with the smallest R^2 score of 0.79. We highlight the locations having zero
519	GPP simulations in blue circles and find that these locations generally have poor predictions with
520	low R^2 scores. Connecting to Figure 2 where we label the locations in column-wise from south to
521	north and from west to east, we identify that those locations with zero GPPs are mostly located in





522 the north where the temperature is relatively low and annual GPPs tend to be zero for parameter

523 samples.

524 We pick 3 locations to closely evaluate the surrogate accuracy (Figure 9). Location 1046 has the best prediction with the highest R^2 score, location 1345 has the worst prediction 525 526 accuracy, and location 428 performs best among the locations with zero GPP simulations. Figure 527 10 shows annual GPP simulations based on sELM and NN-based surrogate in evaluating the 528 1000 test data for 30 years at the 3 locations. It can be seen that NN has difficulty in fitting zero 529 GPP data. At location 1046 where the annual GPPs are relatively high with positive values, NN produces a great fit with a high R^2 score of 0.96 and a small MSE of 0.013. Location 1046 530 531 (Figure 2) is close to the lake where the variance in atmospheric drivers (e.g., temperature) is moderated. This reduced variance leads to a smooth response surface of GPP for which NN can 532 533 easily build an accurate surrogate. In contrast, location 1345 has a large number of simulated 534 GPPs less than 1.0 including many zero GPPs. NN shows difficulty in predicting these small GPPs resulting in a relatively poor performance with the R^2 score of 0.79. Location 1345 is 535 sitting in the north and has the lowest mean annual temperature, so the most parameter samples 536 537 cause low vegetation growth and small GPP values. Moreover, location 1345 is far away from 538 the lakes and has a large variation in atmospheric drivers. Since this location has a climate that is 539 at the extreme end of the range for deciduous forests, the model response is expected and 540 reasonable. However, this leads to a strong nonlinear response surface that casts difficulty in 541 surrogate modeling. In comparison, although location 428 is located in the north with some small 542 GPPs including zero values, it is also close to the lake which has a small variance in the 543 atmospheric drivers. Thus, the NN prediction performance in location 428 is not bad with the R^2 544 score of 0.91.





545	Figure 11 plots the averaged R^2 scores over all locations for 30 years. The R^2 scores have
546	small fluctuations between 0.93 and 0.94, displaying a uniformly good fit among the simulated
547	years. So, when using the surrogate model at any specific year for a data-model analysis, we
548	should be able to obtain a good approximation. In this study, we are considering annual GPPs.
549	Although the variation of atmospheric drivers between years has an impact on surrogate
550	accuracy, its influence is less strong compared to monthly GPPs, so a uniformly good fit among
551	years is expected.
552	Building a surrogate of the discontinuous response surface, e.g., vegetation turns from
553	alive to dead representing as the GPP jumps from nonzero to zero, is a difficulty for almost all
554	the state-of-the-art surrogate methods. Nevertheless, NNs, attribute to the layered architecture
555	and the nonlinear activation function, usually show better performance compared to other
556	surrogate approaches. To improve the surrogate accuracy for strong nonlinear and discontinuous
557	problems, one strategy is using physics-informed domain decomposition methods to build
558	surrogate models separately in different response surface regimes. This strategy requires the
559	surrogate methods strongly connecting to the simulation model, and the methods are generally
560	problem-specific requiring experts' interaction. Another strategy is increasing the training data to
561	explore complex problems. This strategy requires an increase in computational costs for extra
562	expensive model simulations. In the following section 4, we investigate these two strategies and
563	discuss their influence on surrogate accuracy.

564 4 Discussion

ESMs are complex whose response surfaces always display strong nonlinearity and discontinuity, casting a challenge to surrogate modeling. In this section, we consider the strategies of physics-informed learning and increase of training data to improve the surrogate





568	accuracy. We conduct two corresponding experiments to investigate our method's performance
569	in application of these two strategies. In experiment I, we divide the parameter space into two
570	parts producing zero GPPs and nonzero GPPs, and we use 20 training data to build surrogates of
571	the 42660 GPPs in the regime generating nonzero GPP samples. In experiment II, we build the
572	surrogates of the 42660 GPPs in the original parameter domain (Figure 3), but with increasing
573	training data of 200 and 1000.
574	We use the results of Case I as a baseline to investigate our method's performance in the
575	two experiments. Figure 12 shows averaged R^2 scores over 30 years at the 1422 locations in
576	experiment I. The figure indicates that without zero GPPs our method can produce a very
577	accurate surrogate at all locations with a uniformly high R^2 score of 0.98. Building the surrogates
578	in the subdomain without zero GPPs not only significantly improves the prediction accuracy in
579	locations originally having poor fit in Case I, but also further improves the prediction accuracy in
580	locations which already have a good fit in Case I. For example, the R^2 score is dramatically
581	improved from 0.79 to 0.97 at location 1345, from 0.96 to 0.99 at location 1046, and from 0.91
582	to 0.98 at location 428. As shown in Figure 13, the NN almost perfectly reproduces sELM
583	simulations at these 3 locations. Experiment I indicates that physics-informed domain
584	decomposition can be a good strategy to improve surrogate accuracy. For smooth problems (e.g.,
585	no sharp jumps from non-zeros to zeros in response surfaces), our method can build a very
586	accurate surrogate model based on a few training data.
587	Figure 14 shows averaged R^2 scores over 30 years at 1422 locations based on 200 and
588	1000 training data in experiment II. The figure indicates that an increase of training data greatly
589	enhances NN prediction accuracy. Adding 10 folds additional data from Ntrain=20 to
590	Ntrain=200, the overall R^2 score improves from 0.93 to 0.98; further increasing Ntrain to 1000,





591	the averaged R^2 score is up to 0.993 with the worst value of 0.96. Although we observe similar
592	nonuniform performance among locations in Figure 14 as in Figure 9, where the locations with
593	zero GPPs have smaller R^2 scores than others, increasing Ntrain significantly improves the
594	accuracy at all locations, especially those originally having poor fits in Case I. For example,
595	when Ntrain=200, most blue-circled locations have R^2 scores above 0.95 and for Ntrain=1000,
596	the R^2 scores at these blue-circled locations are above 0.985 in comparison to the values below
597	0.9 when Ntrain=20. In the examination of the 3 individual locations by comparing Figure 10
598	and Figure 15, we see that at the location of 1046, an increase of Ntrain enables the NN to
599	perfectly predict sELM simulations with negligible MSEs. Even for the location 428 with zero
600	GPPs, more training data can capture the discontinuous behavior better with R^2 score of 0.99 and
601	MSE of 0.003 when Ntrain=1000. The worst location happens at 1345 for all cases due to its
602	highly changed atmospheric drivers. Even so, the increase of Ntrain can still dramatically
603	enhance the NN's capability in simulating the difficult response surface. Experiment II indicates
604	that increasing training data is able to significantly improve the surrogate accuracy. Our method
605	scales well with the increase of training data and greatly improves prediction accuracy as Ntrain
606	increases.
607	The analysis of the two experiments suggests that our method is data-efficient for
608	continuous problems. To improve the surrogate accuracy in discontinuous and highly nonlinear

609 problems, we can use the physical-informed domain decomposition to focus on the continuous

610 and smooth regions of the response surface. If the discontinuity is the inherent feature of the

- 611 underlying function that we need to surrogate, an increase of training data would be a good
- 612 solution for surrogate accuracy improvement.





613	Having built a surrogate system of many GPP variables over large spatial and temporal
614	domains provides great flexibility and possibility for subsequent predictive analytics tasks. For
615	example, the surrogate model can be used for analyzing sensitivities of model parameters to any
616	set of spatial and temporal GPP variables, and for parameter optimization and uncertainty
617	quantification based on a single-site or multiple-site, a single-year or multiple-year GPP
618	observations using any defined objective functions. In addition, with the newly collected
619	observations from additional sites or further time periods, we can use the same surrogate system
620	for analysis as long as the QoIs are within the surrogate simulation ranges. In the future study,
621	we will pursue the data-model integration using the constructed surrogate system.
622	5 Conclusions
623	In this work, we develop an SVD-enhanced, Bayesian-optimized, and NN-based surrogate
624	method to improve the computational efficiency of large-scale surrogate modeling, so as to
625	advance model-data integration studies in Earth system model simulations. Our method is data
626	efficient in the fact that only 20 model simulations are needed to build an accurate surrogate
627	system. This is a promising result because large Earth system model ensembles are always

628 computationally infeasible, and 20 is a reasonable and affordable number of simulations to

629 consider. In addition, our method is general purpose and can be efficiently applied to a wide

range of Earth system problems with different spatial scales (local, regional, or global) at

631 different simulation periods. It is super effective for smooth problems and scaled well for highly

632 nonlinear and discontinuous problems.

633 We apply our surrogate method to a regional ecosystem model. The results indicate that 634 using only 20 model runs, we can build an accurate surrogate system of 42660 spatially- and 635 temporally-varied GPPs with the R^2 score of 0.93 and MSE of 0.02. For locations with robust





- 636 vegetation growth across the ensemble, our method can almost perfectly predict the model
- 637 simulations with the R^2 score of 0.96. For locations with low vegetation growth for some
- 638 parameter samples and large variation in atmospheric drivers that cause discontinuous response
- 639 surfaces, using physics-informed domain decomposition or the increase of training samples, our
- 640 method can produce accurate predictions with the R^2 score of 0.97 and 0.96, respectively. This
- application demonstrates our method's capability in accurately reproducing expensive model
- 642 simulations based on a few parallel model runs.
- 643 Data availability
- All the data used in this study are model simulation data, which can be generated by running
- 645 the sELM.

646 Code availability

sELM is presented in its 1.0 version, which is realized in the Python language. It is an open-647 648 use computer code which can be accessed freelv from 649 https://github.com/dmricciuto/OSCM SciDAC/tree/master/models/simple ELM. The source 650 code of surrogate modeling using machine learning techniques can be provided upon request via 651 lud1@ornl.gov.

652 Author contribution

Dan Lu developed the methods and carried them out. Daniel Ricciuto developed the model
code and performed the model simulations. Dan Lu prepared the manuscript with contributions
from all coauthors.

656 Acknowledgments

Primary support for this work was provided by the Scientific Discovery through Advanced
Computing (SciDAC) program, funded by the U.S. Department of Energy (DOE), Office of





- 659 Advanced Scientific Computing Research (ASCR) and Office of Biological and Environmental
- 660 Research (BER). Additional support was provided by BER's Terrestrial Ecosystem Science
- 661 Scientific Focus Area (TES-SFA) project. The authors are supported by Oak Ridge National
- 662 Laboratory, which is supported by the DOE under Contract DE-AC05-00OR22725.
- 663 **References**
- Agarap, A. F. M.: Deep learning using Rectified Linear Units (ReLU),
- 665 <u>https://arxiv.org/pdf/1803.08375</u>, 2018.
- 666 Bardenet, R. and Kegl, B.: Surrogating the surrogate: accelerating Gaussian Process optimization
- 667 with mixtures, In ICML, 2010.
- Basu A., De, S., Mukherjee, A., and Ullah, E.: Convergence guarantees for rmsprop and adam in
- 669 nonconvex optimization and their comparison to nesterov acceleration on autoencoders,
- 670 arXiv preprint arXiv:1807.06766, 2018.
- 671 Bergstra J. S., Bardenet, R., Bengio, Y., and Kegl, B.: Algorithms for hyperparameter
- 672 optimization, NIPS 24, 2546-2554, 2011.
- Bergstra J., and Bengio, Y.: Random search for hyper-parameter optimization, Journal of
- 674 Machine Learning Research, 13(1): 281-305, 2012.
- 675 Bergstra, J., Yamins, D., and Cox, D. D.: Hyperopt: A Python library for optimizing the
- hyperparameters of machine learning algorithms, In Proceedings of the 12th Python in
 Science Conference, 13-20, 2013.
- Bottou, L.: Stochastic gradient descent tricks, Neural networks: tricks of the trade: 2nd edition,
 Springer Berlin Heidelberg, 2012.
- 680 Bilionis, I., Drewniak, B. A., and Constantinescu, E. M.: Crop physiology calibration in the
- 681 CLM, Geosci. Model Dev., 8, 1071-1083, 2015.





- 682 Fox, A., Williams, M., Richardson, A. D., Cameron, D., Gove, J. H., Quaife, T., Ricciuto, D.,
- 683 Reichstein, M., Tomelleri, E., Trudinger, C. M., and Van Wijk, M. T.: The REFLEX
- 684 project: Comparing different algorithms and implementations for the inversion of a
- terrestrial ecosystem model against eddy covariance data, Agric. For. Meteorol., 149,
- 686 1597-1615, 2009.
- 687 Gong, W., Duan, Q., Li, J., Wang, C., Di, Z., Dai, Y., Ye, A., and Miao, C.: Multiobjective
- parameter optimization of community land model using adaptive surrogate modeling,
 Hydrol. Earth Syst. Sci., 19, 2409-2425, 2015.
- Huang, M., Ray, J., Hou, Z., Ren, H., Liu, Y., and Swiler, L.: On the applicability of surrogate-
- 691 based Markov chain Monte Carlo-Bayesian inversion to the Community Land Model:
- 692 Case studies at flux tower sites, J. Geophys. Res. Atmos., 121, 7548-7563,
- 693 doi:10.1002/2015JD024339, 2016.
- 694 Kim, H.: Global Soil Wetness Project Phase 3 Atmospheric Boundary Conditions (Experiment
- bata Integration and Analysis System (DIAS), https://doi.org/10.20783/DIAS.501, 2017.
- Kingma, D. P., and Ba, J.: Adam: a Method for Stochastic Optimization, International
 Conference on Learning Representations, 1-13, 2015.
- 699 Lu, D., Ricciuto, D., Walker, A., Safta, C., and Munger W.: Bayesian calibration of terrestrial
- 700 ecosystem models: a study of advanced Markov chain Monte Carlo methods,
- 701 Biogeosciences, 14, 4295-4314, 2017.
- 702 Lu, D., Ricciuto, D., Stoyanov, M., and Gu, L.: Calibration of the E3SM land model using
- surrogate-based global optimization. Journal of Advances in Modeling Earth Systems,
- 704 10. <u>https://doi.org/10.1002/2017MS001134</u>, 2018.





705	Müller, J., Paudel, R., Shoemaker, C. A., Woodbury, J., Wang, Y., and Mahowald, N.: CH4
706	parameter estimation in CLM4.5bgc using surrogate global optimization, Geosci. Model
707	Dev., 8, 3285-3310, 2015.
708	Niranjan, S., Krause, A., Kakade, A., and Seeger, M.: Gaussian process optimization in the
709	bandit setting: No regret and experimental design. In Proceedings of the 27th
710	International Conference on Machine Learning, 2010.
711	Oleson, K. W., et al.: Technical description of version 4.5 of the Community Land Model
712	(CLM). (NCAR Tech. Note NCAR/TN-5031STR, 420 pp. Boulder, CA: National
713	Center for Atmospheric Research, <u>https://doi.org/10.5065/D6RR1W7M</u> , 2013.
714	Ray, J., Hou, Z., Huang, M., Sargsyan, K., and Swiler, L.: Bayesian calibration of the
715	Community Land Model using surrogates, SIAM/ASA J. Uncertain. Quantif., 199-233,
716	doi:10.1137/140957998, 2015.
717	Razavi, S., Tolson, B. A., and Burn, D. H.: Review of surrogate modeling in water resources,
718	Water Resour. Res., 48, W07401, doi:10.1029/2011WR011527, 2012.
719	Ricciuto, D., Sargsyan, K., and Thornton, P.: The impact of parametric uncertainties on
720	biogeochemistry in the E3SM land model. Journal of Advances in Modeling Earth
721	Systems, 10, 297-319, 2018.
722	Sargsyan, K., Safta, C., Najm, H. N., Debusschere, B., Ricciuto, D. M., and Thornton, P.E.:
723	Dimensionality reduction for complex models via Bayesian compressive sensing, Int. J.
724	Uncert. Quant., 4, 63-93, 2014.
725	Shahriari, B., Swersky, K., Wang, Z., Adams, R. P., and de Freitas, N.: Taking the Human Out of
726	the Loop: A Review of Bayesian Optimization, Proc. IEEE 104 (1): 148-175,
727	doi:10.1109/jproc.2015.2494218, 2016.





- 728 Snoek, J., Larochelle, H., and Adams, R. P.: Practical Bayesian optimization of machine learning
- algorithms, in 26th Annual Conference on Neural Information Processing Systems,
- 730 2960-2968, 2012.
- 731 Williams, M., Schwarz, P. A., Law, B. E., Irvine, J., and Kurpius, M.: An improved analysis of
- forest carbon dynamics using data assimilation, Global Change Biol., 11, 89-105, 2005.
- 733 Viana, F. A., Simpson, T.W., Balabanov, V., and Toropov, V.: Metamodeling in
- multidisciplinary design optimization: How far have we really come?, AIAA J., 52(4),
- 735 670-690, 2014.
- 736 Yegnanarayana B.: Artificial neural networks, PHI Learning Pvt. Ltd, 2009.





737



738

Figure 1. Schematic of sELM, where processes are shown using blue boxes with dependencies
on environmental data, 8 uncertain parameter inputs are listed in orange ovals, and model state
variables are indicated by green shapes. Parameters are input to one or more processes as

indicated by blue arrows. Model state variables may be outputs for some processes and input for

743 other processes as indicated by red arrows.

744







746

Figure 2. Locations of interest for which we build surrogates of GPP (gC/m^2/day) variables;

total 1422 locations are considered. The figure shows the sELM simulated annual GPP based onone parameter sample.









753 20 training and 1000 test data are randomly drawn from the parameter space.

754







755

Figure 4. Singular value decay and the information contained in the first largest singular values.

The top 5 singular values contain 97% information of training data matrix with 42660 GPP

variables and 20 samples.







760

Figure 5. Performance of the NNs trained by 20 data with considering the different number of

762 singular value coefficients after SVD.







Figure 6. Changes of loss function values along epochs for training and validation data (a) in

766 Case I which builds surrogates of the 5 singular value coefficients with a simple NN (two hidden

767 layers and each layer has 10 nodes, N1=N2=10) based on 20 training data (Ntrain=20), and (b-e)

in Case II which builds surrogates of all outputs with different NN architectures and different

769 training data size.







771

Figure 7. Comparison of NN performance between Case I: building surrogates of 5 singular

value coefficients (Nsvd=5) after SVD based on 20 training data (red dashed line) and Case II:

building surrogates for all outputs directly with different numbers of training data (red solid

175 line). Each training data represents one sELM simulation. The right y-axis shows the time in

training the NN in Case II. The time for training the NN in Case I is 4 seconds.









Figure 8. Different sets of NN hyperparameters result in different R^2 score in evaluating the 1000

test data. N*l* is the number of nodes in hidden layer *l*, where l=1, 2, and 3. It is the learning rate

781 of Adam algorithm for NN weights optimization.







783

Figure 9. Averaged R^2 scores over 30 years at 1422 locations in evaluating the 1000 test data based on 20 training samples, where the blue circles identify the locations having zero GPP simulations.









Figure 10. Simulations of annual GPPs (gC/m²/day) from sELM and NN-based surrogate

model in evaluating 1000 test data for 30 years at 3 locations, where the NN is trained by 20 data using our method.







793

Figure 11. Averaged R^2 scores over 1422 locations at 30 years in evaluating the 1000 test data.









Figure 12. Averaged R^2 scores over 30 years at 1422 locations in evaluating the 1000 test data

- based on 20 training data in experiment I where the samples are generated in a subdomain of the
- parameter space without zero GPP simulations. The averaged R^2 score is 0.98.









Figure 13. Simulations of annual GPPs (gC/m^2/day) from sELM and NN-based surrogate

model in evaluating 1000 test data for 30 years at 3 locations in experiment I where the samples are generated in a subdomain of the parameter space without zero GPP simulations.







Figure 14. Averaged R^2 scores over 30 years at 1422 locations in evaluating the 1000 test data based on 200 and 1000 training samples, where the blue circles identify the locations having zero GPP simulations.







Figure 15. Simulations of annual GPPs (gC/m²/day) from sELM and NN-baed surrogate model



816 data.