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ABSTRACT

This paper presents a sparsified Fourier neural operator for coupled time-dependent partial differential equations (ST-FNO) as an efficient machine learning surrogate for fluid and particle-based fusion codes such as NIMROD (Non-Ideal Magnetohydrodynamics with Rotation - Open Discussion) and GTC (Gyrokinetic Toroidal Code). ST-FNO leverages the structures in the governing equations and utilizes neural operators to represent Green's function-like numerical operators in the corresponding numerical solvers. Once trained, ST-FNO can rapidly and accurately predict dynamics in fusion devices compared with first-principle numerical algorithms. In general, ST-FNO represents an efficient and accurate machine learning surrogate for numerical simulators for multi-variable nonlinear time-dependent partial differential equations, with the proposed architectures and loss functions. The efficacy of ST-FNO has been demonstrated using quiescent H-mode simulation data from NIMROD and kink-mode simulation data from GTC. The ST-FNO H-mode results show orders of magnitude reduction in memory and central processing unit usage in comparison with the numerical solvers in NIMROD when computing fields over a selected poloidal plane. The ST-FNO kink-mode results achieve a factor of 2 reduction in the number of parameters compared to baseline FNO models without accuracy loss.

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I. INTRODUCTION

Simulation codes are indispensable tools in fusion plasma confinement research, enabling the study of plasma instabilities, prediction of plasma behaviors, design and control of fusion devices, and other critical aspects contributing to the advancement of fusion energy. Mainstream fusion codes use numerical approximations to solve physical models of varying fidelity and applicability with respect to temporal and spatial scales. For example, nonlinear extended magnetohydrodynamics (MHD) codes are routinely applied to study macroscale instability and its consequences in a broad class of magnetic fusion energy (MFE) devices such as tokamaks, stellarators, reversedfield pinches, and field-reversed configurations. Among many others, they include codes like NIMROD,¹ M3D-C1,² and JOREK.^{3,4} Understanding cross field transport due to microscale turbulence in MFE configurations requires a description of the kinetic deviations from a Maxwellian distribution function, due to important effects at the scale of particle gyroradii. Here, the particle gyro-averaged kinetic model, gyrokinetics (GK), is solved in both Eulerian form by codes such as GENE⁵ and GYRO⁶ and in Lagrangian particle-in-cell (PIC) form by codes like GTC^{7,8} and XGC.⁹ Despite their success, these "first-principles" fusion codes are computationally demanding for high-fidelity and whole-device simulation, and each practical run takes days using thousands of central processing unit (CPUs) or graphics processing unit (GPUs) on the leadership supercomputers.

Data-driven algorithms, particularly scientific machine learning (SciML) models for partial differential equations (PDEs), offer appealing complements and alternatives to the fusion codes, as they require much less computation and memory resources to predict an approximate solution once the models have been trained. Compared to other data-driven algorithms, SciML models incorporate physics knowledge and principles into the neural network design to reduce model sizes, improve training time, reduce training samples, and/or improve model 04 April 2025 21:54:50



accuracy. SciML models can potentially be used for fusion research in the following ways: (1) They can be used for the diagnosis of critical events in the fusion device by analyzing the snapshots generated by simulation and experiments. (2) They can be used to quickly predict long-range dynamics without running the simulation code with small time step sizes. (3) They can be used to produce initial guesses and preconditioners for solving linear systems arising in certain fusion codes. (4) They also allow training the ML model with one set of initial conditions or physics models, and testing the model with another set of initial conditions or physics models. Existing SciML models include physics-informed neural networks (PINN) that encode PDEs into the loss function,¹⁰ DeepOnet,¹¹ Fourier neural operators (FNO)¹² for operator learning, Gaussian process,13,14 and reduced-order models,^{15,16} to name a few. To date, most SciML models have been demonstrated in simplified configurations, and very few have been considered for production simulation in realistic geometry and with experimental data.¹⁷⁻²⁵ In this paper, we aim at developing new SciML models for production fusion computations based on the FNO model.

Inspired by the pseudo-spectral method,²⁶ the FNO model represents Green's functions in the spectral domain by nonlinear network operators.¹² Similar to the dealiased pseudo-spectral method that zeros out higher-frequency components, for example see Ref. 27, the FNO model keeps only the lowest few modes for efficiency and accuracy to achieve superior performance for PDEs with smooth solutions. FNO has been mainly applied to relatively simple time-dependent fluid dynamics simulations to predict solutions at future time from their past snapshots. We note that the FNO model is also well-suited for augmenting fusion codes, as many exploit Fourier transforms for periodic directions. Another consideration is that linearly unstable perturbations often exhibit alignment with the magnetic field, making the Fourier representation with field-line-following coordinates a compact representation for GK computation. Although FNO is parameter-efficient, further reducing the model size is critical for large-scale production fusion simulations, particularly due to the limited memory capacity of GPU devices. In addition to the baseline FNO model,¹² there exist a few FNO variants including Tensorized FNO (TFNO)²⁸ which leverages tensor compression of the network parameters, geo-FNO²⁹ that adapts FNO to arbitrary geometry and mesh, factorized FNO³⁰ that simplifies the FNO operations, and physics-informed FNOs^{31,32} that improve the model accuracy by incorporating PDE-based residuals.

Aside from these improved models, we remark that there is another largely open opportunity to improve FNO for fusion codes: production codes often solve a coupled system of equations involving a handful of variables such as temperature, pressure, magnetic field, current density, flow velocity, electrostatic and magnetic potentials, etc. One can leverage the sparse dependencies of these quantities, as indicated by the governing equations, to further simplify the FNO model architecture. We exploit this idea in this paper for two challenging applications of fusion codes that stretch their typical use. Our extended-MHD application models profile evolution from edge-driven fluctuations using the two-fluid description to reproduce electron-fluid dynamics that are not represented by MHD. Conversely, our GK application models macroscale evolution including kinetic effects over a relatively long time.

Our contribution is threefold:

 We bridge the gap between advanced SciML models with production fusion codes (those that solve multiple equations with large-scale and unstructured mesh). Our general methodology applies to fusion codes with linear and nonlinear mode modeling capabilities.

- We propose a parameter-efficient SciML model called sparsified FNO for coupled time-dependent partial differential equations (ST-FNO) that leverages the structure of the governing equations to simplify the architecture. ST-FNO can achieve up to 2× parameter reduction while maintaining similar inference accuracy as the baseline FNO model.
- We demonstrate the efficiency and accuracy of ST-FNO with two distinct fusion codes: NIMROD (extended-MHD) for a simulation of a saturated tokamak edge perturbation, and GTC (gyrokinetic PIC) for a kink mode simulation. Compared with the algorithm in NIMROD, which uses SuperLU_DIST to precondition the computation-intensive algebraic solves, ST-FNO can achieve over 100× memory reduction and significant speedup for generating approximate field solutions over a single poloidal plane, once the model has been trained.

II. METHODOLOGY

We first give a brief introduction of FNO and its building blocks in Sec. II A. We then explain in detail how to develop the proposed ST-FNO frameworks for two distinct examples of fusion simulation codes: NIMROD (Sec. II B) and GTC (Sec. II C) by essentially identifying the independent field variables, their governing equations and sparsifying connectivity in FNO. Finally, we present a short remark on how to develop ST-FNO for other fusion codes and, more generally, time-dependent PDEs in Sec. II D.

A. FNO

FNO¹² represents, arguably, one of the most successful machine learning tools for PDE simulations, and has been demonstrated on a variety of simple and production fluid dynamics simulation codes.^{17,31} FNO is an operator learning framework that represents the discretized Green's function in the spectral domain with parameter efficient neural network components. Suppose one wants to use FNO to predict a set of n_F physical quantities F_s , $s = 1, ..., n_F$ temporally discretized at time steps $k + 1, ..., k + k_o$ from their past values at time steps $k - k_i + 1, ..., k$. Assume that each physical quantity is spatially discretized into a $n_1 \times n_2$ array. In other words, the input of FNO for each physical quantity is of dimensions $n_1 \times n_2 \times k_i$ and the output is of dimensions $n_1 \times n_2 \times k_o$.

A typical FNO architecture consists of a lifting operator P_m, several FNO layers with FNO operator $F_{m,m}$ and a projection operator Q_m . Here, *m* denotes the number of input and output physical quantities of a FNO operator. In our baseline model, it is assumed that $m = n_F$, and the input and output numbers of physical quantities are the same. However, as will be seen later, our proposed FNO model builds upon F_{m_i,m_o} with $m_i, m_o \leq n_F$. Figures 1(a) and 1(b) show the baseline FNO models (similar to the idea of Ref. 24) for the NIMROD and GTC cases (see Secs. II B and II C for detailed explanation). The lifting operator P_m pads the input array with the normalized (x, y)spatial coordinates into a $n_1 \times n_2 \times (mk_i + 2)$ array and lifts the size of the last dimension to mW via linear transformation [see Fig. 1(c)]. W represents the internal width of each field quantity. Each FNO layer consists of one FNO operator $F_{m,m}$ [see Fig. 1(d)] that performs fast Fourier transforms (FFT) along the x and y dimensions and only keeps the lowest M Fourier components. These components are passed to a

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FIG. 1. (a) Baseline-FNO model with $k_i = 1$, $k_o = 1$ for the NIMROD case, " × 4" represents four Fourier layers $F_{m,n}$; (b) baseline-FNO model with $k_o = 1$, $k_o = 1$ for the GTC case; (c) input and output dimensions for the lifting operator P_m and projection operator Q_n ; (d) data dimensions for the Fourier layer $F_{m,n}$; (e) proposed ST-FNO model with $k_i = 1$, $k_o = 1$ for the MIMROD case; (f) the RMS-FC model to predict the magnitude information for the GTC case; and (g) proposed ST-FNO model with $k_i = 1$, $k_o = 1$ for the GTC case.

convolution operator R which learns the underlying Green's function of the PDE. Afterward, they are converted back to data of dimensions $n_1 \times n_2 \times nW$ via inverse fast Fourier transforms (IFFT). In addition, a convolution operator W with kernel size 1 is also used to regularize the learning process (see Refs. 12 and 24 for more explanation). Finally, the projection operator P_m converts the data to the output of dimensions $n_1 \times n_2 \times k_o$ [see Fig. 1(c)]. In $F_{m,m}$ and Q_m , the GeLU activation function is used.

Among the operators P_m , Q_m and $F_{m,m}$, the parameter count of FNO is dominated by that in $F_{m,m}$, each containing roughly $m^2 W^2 M^2$ parameters. This number is estimated by the fact that (1) the FNO operator is inspired by the pseudo-spectral method, which decouples the interactions among Fourier modes, and (2) the R operator performs convolution along the mW dimension. Therefore, the parameter count is linear with respect to the total number of retained Fourier modes M^2 and quadratic with respect to the total internal width mW. It is critical to choose the proper value for the internal width W and the number of Fourier modes M in each spatial coordinate to construct an efficient and effective FNO model. That said, the black-box way of using FNO for fusion simulation can still be prohibitively expensive given that (1) production fusion codes usually have multiple field variables leading to large parameter counts, and (2) production fusion codes are expensive to run, which may limit the amount of training data that can be collected. Therefore, we propose exploiting the algorithm structure of an existing fusion code to sparsify the connectivity of the FNO model and, hence, reduce the training and inference costs.

B. ST-FNO for NIMROD

1. Identification of field quantities and their dependencies

NIMROD^{1,33} is an extended-MHD code that uses a mixed spatial discretization of two-dimensional (2D) high-order finite-elements and

a spectral Fourier decomposition in the third (periodic) direction. For toroidal geometries, the finite-element method (FEM) is applied over the poloidal plane, and Fourier expansion is used for the toroidal direction. The mixed implicit/semi-implicit leapfrog time-advance algorithm in NIMROD exploits the structure of the governing MHD equations. We first discuss the extended-MHD equations and identify the independent field quantities and their dependencies in order to design an efficient ST-FNO model. In what follows, we assume that each field quantity *F* consists of both equilibrium and perturbed components $F = F_0 + \delta F$, but the ST-FNO model only operates on the perturbed quantities as input and output, and does not use the equilibrium quantities. Also, we use the notation F^k to denote the perturbed field quantity discretized at time step *k*.

We consider an extended-MHD model with separate electron dynamics and first-order finite-orbit-radius effects that also include both carbon and deuterium ion species. The center-of-mass velocity, **V**, is computed from the momentum equation,

$$\rho \frac{\partial \mathbf{V}}{\partial t} + \rho (\mathbf{V} \cdot \nabla) \mathbf{V} = \mathbf{J} \times \mathbf{B} - \nabla p - \nabla \cdot \mathbf{\Pi}, \tag{1}$$

where $\rho = m_d n_d + m_c n_c + m_e n_e$ is the total mass density with n_c , n_d , and n_e being the number densities for the carbon, deuterium, and electron species, respectively. The pressure is determined by the ideal-gas law, $p = k_B(n_e T_e + n_c T_c + n_d T_d)$, with a shared-ion temperature, $T_c = T_d = T_i$. The Π term is the stress tensor, which represents the traceless contributions to the pressure tensor and the exact form includes perpendicular, parallel, and gyro-viscosity as described in prior work³⁴ with straightforward modifications for multiple ion species. The current density, **J**, is computed with the pre-Maxwell's Ampère's law, $\mathbf{J} = \nabla \times \mathbf{B}/\mu_0$, as appropriate for low-frequency MHD dynamics. One of the objectives of this work is to leverage the variable dependencies of the equations when constructing the FNO models. However, the center-of-mass velocity equation depends on all of the evolved variables: density, velocity, temperature, and magnetic field. That said, our experiments indicate that dropping the dependency on density in our FNO model does not significantly affect the model accuracy.

The magnetic field is updated via the induction equation, which combines Faraday's law with a generalized Ohm's law

$$\frac{\partial \mathbf{B}}{\partial t} = \nabla \times \left[\mathbf{V} \times \mathbf{B} + \frac{1}{n_e e} (\mathbf{J} \times \mathbf{B} - \nabla p_e - \nabla \cdot \mathbf{\Pi}_e) - \eta \mathbf{J} \right].$$
(2)

Thus, the magnetic field is advected by the electron velocity, $\mathbf{v}_e = \mathbf{V} - \mathbf{J}/n_e e$. The resistivity, η , is density and temperature dependent as determined by a Spitzer expression with a multi-ion electron collision time. A neoclassical current drive is applied through Π_e^{35} where neoclassical poloidal flow damping is also included in the ion stress tensor. The ∇p_e and inverse electron dependencies are present in the induction equation. However, the dominant dependency is on the ideal-MHD center-of-mass velocity advective term, which involves only velocity and magnetic field.

The density equations for only the ion species are evolved and the electron density is determined by the quasineutrality relation, $n_e = n_d + Z_c n_c$. The ion number densities are updated by the continuity equation,

$$\frac{\partial n_{\alpha}}{\partial t} + \nabla \cdot (n_{\alpha} \mathbf{V}) = \nabla \cdot D_{n_{\alpha}} \nabla n_{\alpha} + \nabla \cdot D_{n_{\alpha}, hypd} \frac{1}{R^2} \frac{\partial^2}{\partial \phi^2} \nabla n_{\alpha}, \quad (3)$$

where α is a species index. The last two terms are numerical density diffusion terms added for computational practicality. The last term, a ϕ -directed hyperdiffusivity, is effective to resolve electro-static turbulence which may become prevalent when using an extended-MHD model.³⁶ The density has a simple dependence on only itself and the velocity.

Finally, the temperature T_{α} , $\alpha = i, e$ is updated from the energy equation,

$$\frac{n_{\alpha}}{\gamma - 1} \left(\frac{\partial T_{\alpha}}{\partial t} + \mathbf{V}_{\alpha} \cdot \nabla T_{\alpha} \right) = -\nabla \cdot \mathbf{q}_{\alpha} - p_{\alpha} \nabla \cdot \mathbf{V}_{\alpha}, \qquad (4)$$

where $p_i = k_B(n_c + n_d)T_i$ and $p_e = k_B n_e T_e$ are the ion and electron pressures, respectively. \mathbf{q}_{α} is the conductive heat flux that includes perpendicular, parallel and cross contributions as described in prior work^{34,37} again with straightforward modifications for multiple ion species. Heating terms are not included in this present NIMROD modeling but in principle can be included in future modeling in a straightforward manner. The temperature/energy/pressure equations depend on all of the variables, density, velocity, temperature, and magnetic field where the magnetic field is important for anisotropic thermal conduction and electron advection. However, we choose to drop the dependency of ion temperature on the magnetic field in our FNO model. Also, we drop the weak dependency on density in our FNO model.

Overall, the system is composed of 10 evolved field quantities: 3 for **V**, 3 for **B**, 2 for the ion number density n_c and n_d , and 2 for the temperature, T_e and T_i , and we use their sparse dependencies to design the proposed ST-FNO architecture.

2. Network design

In this paper, we focus on a tokamak simulation case where the inner core region of the poloidal plane is not meshed [see Fig. 2(a) for



FIG. 2. (a) NIMROD training data: coordinate transformation from the finite-element grid in the R - Z coordinate to the uniform grid in the $r - \theta$ coordinate and (b) GTC training data: data sampled in the radial-poloidal ($\psi - \theta$) coordinate can be directly viewed as 2D arrays.

the FEM mesh]. The dynamics in this region are not expected to be significant for this particular application. To prepare training data for the FNO models, we first perform a coordinate transformation from the *R* – *Z* coordinate to the *r* – θ coordinate, where *r* = 0 at the inner boundary and r = 1 at the outer boundary [see Fig. 2(a)]. We generate 2D arrays for each field quantity in a fixed toroidal plane using a structured grid in the $r - \theta$ coordinate. These arrays are evaluated by evaluating the spectral basis functions of the field data on the FEM mesh. Let F denote one of the 10 field quantities. The training data are labeled $F^k \in \mathbb{R}^{n_r \times n_\theta}$ for each time step k. Note that for this simulation case, there is no dramatic change in the magnitude of each field quantity across time steps, but a significant scale difference exists across different quantities. To address such a scale difference, we normalize each field data F by its mean $\mu_F = \sum_{i,j,k} F^k(i,j)/(n_t n_\theta n_r)$ and standard deviation σ_F as $F \leftarrow (F - \mu_F) / \sigma_F$. In this study, the purpose of the networks is to predict all 10 (unnormalized) field quantities in the $\phi = 0$ poloidal plane at time steps (or snapshots if the fields are collected very few time steps) $k + 1, ..., k + k_o$ from their history at time steps (or snapshots) $k - k_i + 1, \dots, k$.

Our baseline FNO model is shown in Fig. 1(a) consisting of 1 lifting operator P₁₀, 1 projection operator Q₁₀ and 4 FNO layers F_{10,10}. The parameter count of the baseline FNO model is dominated by the FNO layers as about $4 \times 10^2 M^2 W^2 = 400M^2 W^2$. In comparison, the proposed ST-FNO model is shown in Fig. 1(e) consisting of lifting and projection operators in 4 groups, as well as four FNO layers each having multiple FNO operators. Each FNO operator F_{m,n} corresponds to one governing equation in Sec. II B 1 and exploits the same structure of the MHD equations that is exploited by the semi-implicit leapfrog algorithm within NIMROD.¹ Due to sparser connectivity, the parameter count can be estimated as $4 \times (8 \times 3 + 5 \times 2 + 7 \times 1 + 4 \times 1 + 6 \times 3)M^2W^2 = 252M^2W^2$, which is about 37% smaller than the baseline-FNO model. We note that the typical mean squared errors (MSE) in machine learning community is not a suitable error metric or loss function for multi-variable fusion simulation data, as it cannot distinguish the magnitude difference between variables. For both the baseline and the ST-FNO models, we propose the following loss function:

$$loss_{F} = \frac{1}{n_{F}|\mathbb{K}|} \sum_{k \in \mathbb{K}} \sum_{s=1}^{n_{F}} \sqrt{\frac{\sum_{i=1}^{k_{o}} |F_{s}^{k+i} - \bar{F}_{s}^{k+i}|_{2}^{2}}{\sum_{i=1}^{k_{o}} |F_{s}^{k+i}|_{2}^{2}}},$$
(5)

where \bar{F}_s and F_s denote the sth predicted and ground truth field quantity, and $n_F = 10$. $|F|_2$ denotes the L2 norm of the vectorization of the matrix F. \mathbb{K} denotes the set of training or testing snapshot indices, $|\mathbb{K}| = n_{train}$ for the training error, $|\mathbb{K}| = n_{test}$ for the test error, and $|\mathbb{K}| = 1$ for the error of each individual test index. Unlike many other existing loss functions,²⁴ this formula can faithfully represent the average relative difference between simulation and FNO outputs. Here \bar{F}_s and F_s are the normalized field data, but Eq. (5) can be used for unnormalized field data as well.

C. ST-FNO for GTC

1. Identification of field quantities and their dependencies

GTC⁷ is an advanced particle-in-cell code for simulating plasma turbulence in fusion reactors capable of modeling kinetic electrons, kinetic ions, electromagnetic waves, etc. In this paper, we consider a GTC-generated dataset for simulating the internal kink mode in the low frequency, long-wavelength limit. To develop a ST-FNO architecture leveraging the algorithm structure of GTC, we first summarize the governing equations used to derive the GTC algorithm. To be consistent with the kink mode dataset, we assume that electrons are treated as fluid, and (thermal) ions are treated as gyrokinetic particles.³⁸ Our ST-FNO design for GTC, just like ST-FNO for NIMROD, only operates on perturbed physical quantities, and we focus on the explanation of the perturbed quantities in what follows.

The perturbed ion particle distribution function δf_i or equivalently ion particle weight *w* is updated from the Vlasov equation

$$\frac{dw}{dt} = (1 - w) \left[-\left(\nu_{||} \frac{\delta \mathbf{B}_{\perp}}{B_{0}} + \mathbf{v}_{E} \right) \cdot \nabla \ln f_{i0} - \frac{\mu \nu_{||} \widehat{\mathbf{b}}_{0} \cdot \nabla \langle \langle \delta B_{||} \rangle \rangle}{T_{i}} + \frac{Z_{i}}{T_{i}} \left(\nu_{||} E_{||} - \mathbf{v}_{d} \cdot \nabla \left(\phi + \frac{\mu}{Z_{i}} \langle \langle \delta B_{||} \rangle \rangle \right) \right) \right].$$
(6)

Here, the perturbed perpendicular magnetic field $\delta \mathbf{B}_{\perp}$ is computed from the parallel magnetic potential δA_{\parallel} as $\delta \mathbf{B}_{\perp} = \nabla \times \delta A_{\parallel} \hat{\mathbf{b}}_0$. \mathbf{v}_E $= \frac{\hat{\mathbf{d}}_0 \times \nabla \phi}{B_0^*}$ is the E × B drift velocity with ϕ being the electrostatic potential, \mathbf{v}_d is the sum of the magnetic curvature drift current and the magnetic gradient drift current. The parallel particle velocity v_{\parallel} and the gyrocenter **R** are updated from the equations of motion. After temporal discretization, one can realize that the distribution function δf_i^{k+1} at time step k + 1 only depends on δf_i^k , ϕ^k , and A_{\parallel}^k . Note that δf_i is a particle quantity in GTC, but our ST-FNO model only operates on field quantities. Therefore, we use ion particle density n_i , ion flow velocity u_i , ion perpendicular pressure $\delta P_{\perp i}$, and ion parallel pressure $\delta P_{\parallel i}$ to approximately represent δf_i

$$n_i = \frac{2\pi B_0^*}{m} \int d\nu_{||} d\mu \delta f_i, \tag{7}$$

$$u_i = \frac{2\pi B_0^*}{m} \int d\nu_{||} d\mu \nu_{||} \delta f_i, \qquad (8)$$

$$\delta P_{\perp i} = \frac{2\pi B_0^*}{m} \int d\nu_{||} d\mu \mu B_0^* \delta f_i, \tag{9}$$

$$\delta P_{\parallel i} = \frac{2\pi B_0^*}{m} \int dv_{\parallel} d\mu m v_{\parallel}^2 \delta f_i.$$
⁽¹⁰⁾

Although these quantities cannot be directly used in the GTC, ST-FNO can learn to predict these low-order fluid moments of, e.g., the kink-mode evolution based on the same fluid moments extracted from GTC results, which effectively provides closure information for the kinetic system.

The perturbed parallel magnetic potential $A_{||}$ is updated via Faraday's law,

$$\frac{\partial A_{\parallel}}{\partial t} = \widehat{\mathbf{b}}_0 \cdot \nabla(\phi_{eff} - \phi), \qquad (11)$$

where the effective electrostatic potential ϕ_{eff} is computed from the perturbed electron number density δn_e and the perturbed parallel magnetic field $\delta B_{||}$ by

$$\frac{e\phi_{eff}}{T_e} = \frac{\delta n_e}{n_{0e}} + \frac{\delta B_{||}}{B_0} - \frac{\partial \ln n_0}{\partial \psi_0} \delta \psi - \frac{\partial \ln n_0}{\partial \alpha_0} \delta \alpha + \frac{e}{T_e} \frac{\partial \phi_{eq}}{\partial \psi_0} \delta \psi. \quad (12)$$

Here, $\delta \psi$ and $\delta \alpha$ represent the Clebsch representation of $\delta \mathbf{B}_{\perp}$, and δB_{\parallel} can be directly computed from $\delta P_{\perp i}$. Therefore, one can conclude that the parallel magnetic potential at time step k + 1, A_{\parallel}^{k+1} , only depends on A_{\parallel}^k , δn_e^k , and $\delta P_{\perp i}^k$.

The perturbed electron number density δn_e can be updated by the electron continuity equation,

$$\frac{\partial \delta n_e}{\partial t} = -\nabla \cdot \left[n_{0e} u_{\parallel e} \left(\frac{\mathbf{B}_0 + \delta \mathbf{B}_\perp}{B_0} \right) + n_e \mathbf{v}_E - \frac{P_{\parallel e} \widehat{\mathbf{b}}_0 \times \kappa}{eB_0} - \frac{P_{\perp e} \widehat{\mathbf{b}}_0 \times \nabla B_0}{eB_0^2} - \frac{P_{\perp e} \widehat{\mathbf{b}}_0 \times \nabla \delta B_{\parallel}}{eB_0^2} \right], \quad (13)$$

where $n_e = n_{0e} + \delta n_e$, $\boldsymbol{\kappa} = (\hat{\mathbf{b}}_0 \cdot \nabla) \hat{\mathbf{b}}_0$ (field line curvature), $P_{\parallel e} = P_{e0} + \delta P_{\parallel e}, P_{\perp e} = P_{e0} + \delta P_{\perp e}$. In Eq. (13) δu_e is the perturbed electron flow velocity from Ampère's law $\frac{4\pi}{c} en_e u_{\parallel e} = \nabla_{\perp}^2 A_{\parallel}$ $+ \frac{4\pi}{c} Z_i n_i u_{\parallel i}, \mathbf{v}_E$ depends on ϕ , the perturbed diamagnetic drift velocity $\mathbf{v}_* = \frac{1}{n_0 m_e \Omega_e} \hat{\mathbf{b}}_0 \times \nabla (\delta P_{\parallel e} + \delta P_{\perp e})$ with perturbed pressures $\delta P_{\parallel e}$ and $\delta P_{\perp e}$ depending on ϕ_{eff} and $\delta \mathbf{B}_{\perp}$. δB_{\parallel} depends on $\delta P_{\perp i}$. Recall their dependencies explained above, the perturbed electron number density at time step k + 1, δn_e^{k+1} , only depend on δn_e^k , $A_{\parallel i}^k$, δu_i^k , $\delta P_{\perp i}^k$, and ϕ^k .

In addition to the above-described governing equations and quantity dependencies, the perturbed electrostatic potential at time step k + 1, ϕ^{k+1} , can be computed from δn_e^{k+1} and δn_i^{k+1} via the Poisson equation

$$\frac{Z_i^2 \rho_i^2}{T_i} \sum_{s \neq e} \frac{n_{0s} m_s}{m_i} \nabla_\perp^2 \phi = -(1 - \rho_i^2 \nabla_\perp^2) \sum_s Z_s \bar{n}_s.$$
(14)

Overall, we choose 7 independent field quantities δn_e , δn_i , δu_i , $\delta P_{\perp i}$, $\delta P_{\parallel i}$, A_{\parallel} , and ϕ , and use their sparse dependencies to design the proposed ST-FNO architecture.

2. Network design

In principle, we can apply the same methodology as the case of NIMROD to design ST-FNO for GTC. Similar to the ST-FNO for the NIMROD case, here we aim at predicting all 7 field quantities at time steps (or snapshots) $k + 1, ..., k + k_o$ from their history at time steps (snapshots) $k - k_i + 1, ..., k$.

However, there are three significant differences. First, the NIMROD extended-MHD simulations considered here only consider fluid dynamics, but GTC is a PIC code that evolves both field and particle data. As explained in Sec. II C 1, we can use n_i , u_i , $\delta P_{\perp i}$, and $\delta P_{\parallel i}$ to describe the lowest-order moments of the particle distribution data δf_i . Second, unlike NIMROD whose simulation data in poloidal planes is represented with FEM mesh, GTC's simulation data in poloidal planes is directly sampled on the field-aligned coordinate with radial flux coordinate ψ and poloidal coordinate θ , leading to 2D arrays [see Fig. 1(b)] as the training data. Let $F = \delta n_e, \delta n_i, \delta u_i, \delta P_{\perp i}, \delta P_{\parallel i}, A_{\parallel}, \phi$ be one of the seven field quantities for ST-FNO, $F \in \mathbb{R}^{\overline{n_{\psi}} \times n_{\theta}}$. In principle, a coordinate transformation to a field-aligned coordinate could be performed with the NIMROD data but was not considered in this work. Third, our GTC dataset is generated from a kink mode simulation case, which consists of the linear and nonlinear phases. In the linear phase, several field-quantity magnitudes grow exponentially as time increases while the field patterns remain stable; in contrast, in the nonlinear phase, the field magnitudes stay stable, but the field patterns can evolve dramatically. The following design addresses this last challenge efficiently.

To design an efficient network surrogate for the kink mode simulation, we normalize each field quantity at time step k, F^k , by its root mean square (RMS) value $[F^k]$ as $F^k \leftarrow F^k/[F^k]$ and use the normalized data to train the ST-FNO model, as depicted in Fig. 1(g). Compared to the baseline FNO model in Fig. 1(b), ST-FNO's sparse connectivity can significantly reduce the network parameter count. To preserve the RMS information for both ST-FNO and baseline FNO, we create a separate network consisting of two fully connected (FC) layers called RMS-FC model. When $k_i = k_o = 1$, RMS-FC predicts log $[F^{k+1}]$ from log $[F^k]$ as shown in Fig. 1(f). We use the following loss function for RMS-FC:

$$loss_{RMS} = \frac{1}{n_F} \sum_{k \in \mathbb{K}} \sum_{i=1}^{k_o} \sqrt{\frac{\sum_{s=1}^{n_F} \left(\log \left[F_s^{k+i} \right] - \log \left[\bar{F}_s^{k+i} \right] \right)^2}{\sum_{s=1}^{n_F} \left(\log \left[F_s^{k+i} \right] \right)^2}}, \quad (15)$$

with \mathbb{K} the same as that in Eq. (5).

Our baseline FNO model is shown in Fig. 1(b) consisting of one lifting operator P₇, one projection operator Q₇, and four FNO layers F_{7,7}. The parameter count of the baseline FNO model is dominated by the FNO layers as about $4 \times 7^2 M^2 W^2 = 196M^2 W^2$. In comparison, the proposed ST-FNO model is shown in Fig. 1(g) consisting of seven lifting and seven projection operators, as well as four FNO layers each

having seven FNO operators. Each FNO operator $F_{m,n}$ corresponds to one governing equation in Sec. II C 1, except that those for n_i , u_i , $\delta P_{\perp i}$, and $\delta P_{\parallel i}$ approximately representing the Vlasov Eq. (6) (the connectivity is shown in red). Due to sparser connectivity, the parameter count of ST-FNO can be estimated as $4 \times (5 + 3 + 3 + 3 + 3 + 4 + 2)M^2W^2 = 92M^2W^2$, which is significantly smaller than the baseline-FNO model.

For both the baseline and the ST-FNO models, we use the loss function in Eq. (5) with $n_F = 7$. Recall that F_s^k in Eq. (5) represent the normalized perturbed field quantities.

D. A note on generalizing ST-FNO to other codes

Although in this paper we only demonstrate how to design ST-FNO for two simulation codes, NIMROD and GTC, it shall be clear by now that ST-FNO can be easily adapted to other fusion codes and, more generally, codes solving systems of time-dependent PDEs. Assume that we have a set of n_F field variables $\mathbf{F}(x, y) = (F_1, ..., F_{n_F})$ satisfying

$$\frac{\partial \mathbf{F}}{\partial t} = \mathscr{L}(\mathbf{F}, \nabla \mathbf{F}, \nabla^2 \mathbf{F}, \dots), \tag{16}$$

where ${\mathscr L}$ is a vector of differential operators involving only spatial derivatives.

One can identify the dependency of F_s based on the right-hand side of Eq. (16) and naturally design a FNO model whose connectivity respects such dependency. The training data can be generated using a regular grid in (*x*, *y*), regardless of the numerical algorithms being used, e.g., FEM, finite-difference, finite-volume, spectral method, etc. If particle-based methods, e.g., PIC, are used, one can convert the particle data to field data, just like Eqs. (7)–(10) for GTC. In addition, if the field-dependency is weak, it can be neglected in the FNO connectivity model. Introducing further sparsity into the NIMROD ST-FNO model may be possible.

III. NUMERICAL RESULTS

In this section, we provide numerical results using two distinct fusion simulations, an extended-MHD result from NIMROD and a gyrokinetic PIC result from GTC, to demonstrate the efficiency and accuracy of the proposed ST-FNO methodology.

A. NIMROD results

The dataset is generated by a nonlinear simulation using the model described in Sec. II B 1 using the NIMROD code. This case extends prior work^{39,40} that studies the quiescent H-mode regime in the DIII-D tokamak. The 3D tokamak-plasma volume is discretized with a 96 × 256 grid of bi-quartic elements over the poloidal plane and with 22 toroidal Fourier modes over the toroidal angle. For the FNO application, we use a set of 257 snapshots from the NIMROD simulation steps numbered 12 000–63 200 with one FNO snapshot per 200 NIMROD steps. The NIMROD time steps are of size $\Delta t_m = 5 \times 10^{-9}$ s, and at this time step size and resolution, the compressional Alfvén wave Courant–Friedrichs–Lewy (CFL) condition is approximately 800 while the flow-speed CFL is approximately two. Eleven Perlmutter CPU nodes are used, which requires approximately 7 s per step. On a fixed poloidal plane at $\phi = 0$, we generate 2D data with $n_r = 64$ and $n_{\theta} = 64$ as data for the ST-FNO model.

For both the baseline-FNO and ST-FNO models, we set the interior widths W = 20, the number of Fourier modes in each of r and θ directions M = 12, and the number of FNO layers to 4. Recall that n_{train} and n_{test} are the number of snapshots in the training and test sets, respectively. Here, we consider two ways of splitting the time steps. (1) Random splitting: the snapshots are randomly split into the training set with $n_{train} = 128$ and the test set with $n_{test} = 129$. (2) Sequential splitting: the snapshots are split into two halves, the first n_{train} snapshots as the training set and the rest as the test set. In other words, we only train the models from data in the past and test the models with data in the future. We train both models for 500 epochs, i.e., 500 network training passes through the data, with the training and testing errors defined by Eq. (5). The training is performed on 1 NVIDIA A100 GPU of Perlmutter.

1. Random splitting

We first compare the convergence of the test error as a function of the epoch number with $k_i = 1, 5$ and $k_o = 1, 5$ in Fig. 3. We remark that the convergence of the baseline-FNO and ST-FNO models are very similar. The training and test errors at epoch 500 listed in Table I confirm that the two models have very similar inference accuracy.

Next, we remark that ST-FNO with a parameter count of 67.5 $\times 10^6$ is more efficient compared to baseline-FNO with a parameter count of 92.8 $\times 10^6$, leading to a 30% model size reduction (see Table I). It is worth noting that the FNO parameter count is insensitive to k_i and k_o as the FNO operators dominate the parameter count instead of the lifting/projection operators. For both models, the training time per epoch is about three to tens and the inference time per time step is about 5–20 ms using 1 Perlmutter A100 GPU. The memory requirement for ST-FNO is about 280 MB. In comparison, the approximate solutions in NIMROD are computed with the SuperLU_DIST⁴¹ direct solver used to precondition the algebraic solves, which requires 34 GB memory, 9s factorization time, and 340 ms apply time (summing up the numbers for all physical quantities) for each NIMROD toroidal Fourier mode (or equivalently, for each poloidal plane) using 64 Perlmutter CPU cores.



FIG. 3. Test error using Eq. (5) vs epoch number for ST-FNO and baseline FNO for the NIMROD case with $k_i = 1, 5$ and $k_o = 1, 5$.

TABLE I. Comparison of accuracy and parameter counts of the ST-FNO and the baseline FNO models for the NIMROD case (random splitting). k_i represents the number of past snapshots used as the model input, and k_o represents the number of future snapshots as the module output.

Model	k _i	k _o	Train error (10 ⁻³)	Test error (10 ⁻³)	Parameter count (M)
Baseline-FNO	1	1	8.76	9.19	92.8
Baseline-FNO	3	1	5.61	5.8	92.8
Baseline-FNO	5	1	5.65	6.94	92.8
ST-FNO	1	1	7.97	8.7	58.2
ST-FNO	3	1	5.79	6.09	58.2
ST-FNO	5	1	6.13	7.39	58.2

reused for multiple time steps and their computational cost can be amortized, we can just compare the memory requirement and inference/apply time. To this end, ST-FNO is capable of achieving 120× memory reduction compared with SuperLU_DIST-based solution in predicting an estimate of the field quantities over a single poloidal plane. If one wants to use ST-FNO to provide an initial guess for the algebraic solve at every time step, that requires only 20 ms × 44 ≈ 1 s. Compared with the 340 ms × 22 ≈ 7.5 s of SuperLU_DIST-based preconditioner per Krylov iteration, the ST-FNO-based initial guess is computationally inexpensive. Note that the factor of 44 indicates that prediction from 44 poloidal planes are required to resolve the 22 Fourier modes used in NIMROD. If one wants to get an estimate of the fields every 200 time steps, e.g., to perform long-term diagnostics/prediction, ST-FNO only takes 1s. In contrast, the first-principle numerical solver takes 200 × 7 ≈ 1400 s.

Next, we compare the field plots predicted by ST-FNO and simulated by NIMROD. Figure 4 plots the NIMROD simulation results,



FIG. 4. Color contour plots of perturbed fields from (top) NIMROD simulation results, (middle) ST-FNO results and (bottom) their relative difference using Eq. (17) (random splitting) at time step 58 400 (snapshot k = 233) for 4 out of the total 10 field quantities: V_{R} , B_{R} , n_{d} , and T_{i} . $k_{i} = 1$, $k_{o} = 1$.

ST-FNO results, and their relative difference for 4 out of the 10 field quantities at time step 58 400. For a NIMROD field quantity F^k and its ST-FNO output \bar{F}^k , the relative difference is computed as

relative diff =
$$(F^k - \overline{F}^k) / \max |F^k(i,j)|.$$
 (17)

Note that ST-FNO can attain $10^{-2} - 10^{-3}$ relative difference for most field quantities. Furthermore, Fig. 5 shows the ST-FNO results for V_R and the relative difference with respect to the NIMROD results for 5 different time steps (from the test set), a relative difference of 10^{-2} can be observed. Notable dynamics are developed near the edge of the inner core, and the largest error appears near the magnetic x-point. This point is a stagnation point of the poloidal magnetic field where field-lines on the last-closed magnetic-flux surface will asymptotically approach the x-point. This makes this spatial location unique within the simulated domain and thus exhibits differing dynamics relative to the rest of the domain.

Next, we examine the inference error per test sample in the random splitting setting. Figure 6(a) plots the relative error of the predicted data $\bar{F}_s^k = M_{FNO}(F_s^{k-1})$ for each sample snapshot k. Here $y = M_{FNO}(x)$ denotes the ST-FNO model with input x and output y. In other words, this represents a single forward pass of the input x through the ST-FNO model. As the reference, we also plot the relative error for directly using F_s^{k-1} as an estimate for F_s^k (dubbed "previous iterate (PI)"), i.e., $\bar{F}_s^k = F_s^{k-1}$. As we can see, ST-FNO inference can outperform PI by up to 5× more accuracy for most test snapshots.

2. Sequential splitting

In this subsection, we examine the inference error per test sample in the sequential splitting settings [see Fig. 6(b)], we set n_{train} to different values and try to investigate how many more snapshots ST-FNO can predict a valuable solution without retraining the model with newer snapshots. First, we consider $k_i = 1$ and $k_o = 1$. The FNO prediction is $\bar{F}_s^k = M_{FNO}(\bar{F}_s^{k-1})$ with $n_{train} = 130, 180, 220, 240$. In other words, we always predict the next snapshot \bar{F}_s^k , using the model M_{FNO} , with the NIMROD simulation data at previous snapshot F_s^{k-1} as the input. Note that for each curve, the ST-FNO model is trained only once. In comparison, we also plot results using PI $\bar{F}_s^k = F_s^{k-1}$. We remark that the ST-FNO prediction can always outperform PI for the first 3–4 snapshots, then a retraining of the model becomes necessary.



FIG. 5. Color contour plots of perturbed fields from the (top) ST-FNO results and (bottom) their relative difference using Eq. (17) (random splitting) with respect to the NIMROD simulation results at time steps 19 600, 28 600, 38 600, 48 800, and 58 400 for the field quantity V_{R} , $k_i = 1$, $k_o = 1$.



FIG. 6. Relative error using Eq. (5) of the estimated fields \bar{F}_s^k for each test sample in the NIMROD case. Each snapshot *k* represents 200 time steps in the NIMROD code, and snapshot 1 represents time step 12 000: (a) Random splitting setting: \bar{F}_s^k is computed from ST-FNO $\bar{F}_s^k = M_{ENO}(F_s^{k-1})$ with $n_{train} = 128$ and the previous iterate (PI) $\bar{F}_s^k = F_s^{k-1}$. $k_i = k_o = 1$; (b) sequential splitting setting: \bar{F}_s^k is computed from ST-FNO $\bar{F}_s^k = M_{ENO}(F_s^{k-1})$ with $n_{train} = 130$, 180, 220, 240 and PI $\bar{F}_s^k = F_s^{k-1}$. $k_i = k_o = 1$; (c) sequential splitting setting: Same as (b) but with $k_i = k_o = 5$. In other words, $\bar{F}_s^{k,\dots,k+4} = M_{ENO}(F_s^{k-5,\dots,k-1})$; and (d) sequential splitting setting: The ST-FNO operates in an auto-regressive fashion to predict snapshot *k* from $F_s^{k-k_0}$: $\bar{F}_s^k = M_{ENO}(\bar{F}_s^{k-1})$, $\bar{F}_s^{k-1} = M_{ENO}(\bar{F}_s^{k-2})$, ..., $\bar{F}_s^{k-k_0+1} = M_{ENO}(\bar{F}_s^{k-k_0})$.

Note that the PI estimate quality drops significantly for snapshot number k > 250, which justifies the need for ST-FNO prediction (with more frequent retraining).

In addition, we visualize the fields predicted by ST-FNO (in the sequential splitting settings with $k_i = 1$ and $k_o = 1$) and simulated by NIMROD. Figure 7 plots the NIMROD simulation results, ST-FNO



FIG. 7. Color contour plots of perturbed fields from (top) ST-FNO results and (bottom) their relative difference using Eq. (17) with respect to the NIMROD simulation results (sequential splitting) at time steps 48 000, 56 000 with different sized training data for the field quantity V_R . $k_i = 1$, $k_o = 1$.

results, and their relative difference defined by Eq. (17) for the field quantity V_R at two time steps. For time step 48 000 (snapshot 181), we consider with ST-FNO model with $n_{train} = 130$ and $n_{train} = 180$. For time step 56 000 (snapshot 131), we consider the ST-FNO model with $n_{train} = 130$ and $n_{train} = 180$. It is clear from both Figs. 6(b) and 7 that retraining the ST-FNO model with more available snapshot data can significantly improve the model prediction accuracy.

Next, we consider $k_i = 5$ and $k_o = 5$. In other words, we try to predict the field at a timescale 5 times larger than the above experiment. The ST-FNO prediction is $\bar{F}_s^{k,...,k+4} = M_{FNO}(F_s^{k-5,...,k-1})$ with $n_{train} = 130, 180, 220, 240$ [see Fig. 6(c)]. In comparison, we also plot results using PI $\bar{F}_s^k = F_s^{k-5}$. Note for snapshot k + 4, ST-FNO uses all five snapshots k - 5, ..., k - 1, while PI uses only snapshot k - 1. Therefore, ST-FNO can significantly outperform PI for almost all the data points with $n_{train} = 180, 220, 240$.

Finally, we consider using ST-FNO in an auto-regressive fashion. In other words, we want to predict the field \overline{F}_s^k from input quantities that are k_0 snapshots back in time: $\overline{F}_s^k = M_{FNO}(\overline{F}_s^{k-1}), \overline{F}_s^{k-1} = M_{FNO}(\overline{F}_s^{k-2}), \dots, \overline{F}_s^{k-k_0+1} = M_{FNO}(F_s^{k-k_0})$. We trained the ST-FNO model with $k_i = k_o = 1$ using $n_{train} = 130$ snapshots, and applied the trained model with k_o varying from 1 to 5. First, the error increases over time for all curves, just like the other subfigures, in Fig. 6(d). Note that the " $k_0 = 1$ " curve is the same as the " $n_{train} = 130$ " curve in Fig. 6(b). Next, one can clearly see that increasing k_0 will quickly increase the prediction error due to error accumulation of autoregression. This suggests that both a proper fine-tuning/retraining of the ST-FNO model over time and a proper choice of number of autoregression steps are needed.

B. GTC results

The dataset is generated by a nonlinear gyrokinetic simulation using GTC for the DIII-D discharge #141 216 at t = 1750 ms. The DIII-D tokamak geometry is discretized with a $100 \times 250 \times 24$ mesh in radial, poloidal, and parallel directions; the time step size is set to $\Delta t = 1.483 \times 10^{-8}s$. We run the simulation for 20 000 time steps and keep both n = 0 and n = 1 modes in the simulation. This simulation requires 3.9 h on 6 Perlmutter GPU nodes. The simulation generates one snapshot per 100 time steps, and we use time steps 4000 to 18 800 as the dataset for training and testing ST-FNO. This is due to the fact that the first 3999 time steps consist of significant initial noises, and the last 200 time steps become physically unreliable due to numerical instabilities. Note that there is a transition point near time step 15 300 (snapshot number k = 114) that separates the linear phase and nonlinear kink mode phase.

For both the baseline-FNO and ST-FNO models, we set the interior widths W = 20, the number of Fourier modes in each direction M = 12, and the number of FNO layers to 4. Here, we consider two ways of splitting the time steps. (1) Random splitting: the snapshots are randomly split into the training set with $n_{train} = 74$ and the test set with $n_{test} = 75$. (2) Sequential splitting: the snapshots are split into two halves, the first n_{train} snapshots as the training set and the rest as the test set. In other words, we only train the models from data in the past and test the models with data in the future. We train the models for 500 epochs with the errors defined by Eq. (5) for the ST-FNO and baseline-FNO model and Eq. (15) for the RMS-FC model. The training and testing are performed with 1 NVIDIA A100 GPU of Perlmutter.





FIG. 8. Test error using Eq. (5) for the normalized field data vs epoch number for ST-FNO and baseline FNO for the GTC case with $k_i = 1, 5$ and $k_o = 1, 5$.

1. Random splitting

We first compare the convergence of the test error in the random splitting setting as a function of the epoch number with $k_i = 1, 5$ and $k_o = 1, 5$ in Fig. 8. We remark that the convergence of the baseline-FNO and ST-FNO models are very similar. The training and test errors at epoch 500 listed in Table II confirm that the two models have very similar inference accuracy. Next, we remark that ST-FNO with a parameter count of 21.2×10^6 is more efficient compared to baseline-FNO with a parameter count of 45.4×10^6 , leading to a 2X model size reduction (see Table II). For ST-FNO models, the training time per epoch is about 6.78 s and the inference time per snapshot (i.e., per 100 time steps) is about 2.00 ms. In comparison, the average time per time step in GTC is about 0.7 s.

Next, we compare the field plots predicted by ST-FNO and simulated by GTC in the random splitting setting. Figure 9 plots the GTC simulation results, ST-FNO results, and their relative difference for 4 field quantities at time step 16 000. Note that these plots show the combination of the ST-FNO results and the RMS-FC results. Note that ST-FNO can attain $10^{-2} - 10^{-3}$ relative difference for all field quantities. Furthermore, Fig. 10 shows the ST-FNO results for $A_{||}$ and the relative difference with respect to the GTC results for five different time steps (from the test set), a relative difference of $10^{-2} - 10^{-3}$ can be observed.

TABLE II. Comparison of accuracy and parameter counts of the ST-FNO and the baseline FNO models for the GTC case (random splitting). k_i represents the number of past snapshots used as the model input, and k_o represents the number of future snapshots as the module output.

Model	k _i	k _o	Train error (10 ⁻³)	Test error (10 ⁻²)	Parameter count (M)
Baseline-FNO	1	1	9.96	1.78	45.4
Baseline-FNO	3	1	9.51	1.77	45.4
Baseline-FNO	5	1	8.23	1.79	45.4
ST-FNO	1	1	10.8	1.77	21.2
ST-FNO	3	1	7.93	1.81	21.2
ST-FNO	5	1	7.95	1.85	21.2

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FIG. 9. Color contour plots of perturbed fields from (top) GTC simulation results, (middle) ST-FNO results and (bottom) their relative difference using Eq. (17) (random splitting) at time step 16 000 for 4 out of the total 7 field quantities: ϕ , $A_{||}$, u_i and n_e . $k_i = 1$, $k_o = 1$.

Note that these time steps cover both the linear and nonlinear phases. The RMS values range from $10^{-5} - 10^0$ and the kink mode exhibits dramatic pattern changes, but our model can still predict very well.

Next, we examine the inference error per test sample in the random splitting settings. Figure 11(left) plots the relative error of the normalized data, RMS, and the de-normalized data. Roughly speaking, let $[\bar{F}] = [F](1 + \varepsilon_{RMS})$ be the model-generated RMS data and $\bar{F} = F(1 + \varepsilon_{norm})$ be the model-generated normalized field data. In the random splitting setting (left column), the RMS values can be very computed as $[\bar{F}_s^k] = M_{RMS}([F_s^{k-1}])$ with M_{RMS} denoting the RMS-FC model. The inference errors are $\varepsilon_{RMS} \approx 10^{-4} - 10^{-2}$ given the presence of both linear and nonlinear phase data in the training set [see Fig. 11(a)]. Similarly, the normalized field data can be accurately computed as $\bar{F}_s^k = M_{FNO}(F_s^{k-1})$ with at least $|\varepsilon_{norm}| \approx 6 \times 10^{-2}$ error [see Fig. 11(b)]. The de-normalized field $[\bar{F}]\bar{F} \approx [F]F(1 + \varepsilon_{RMS} + |\varepsilon_{norm}|)$ shows an inference error about 6×10^{-2} [see Fig. 11(c)]. Note that for both the RMS and the field data, the inference error is larger toward snapshots 4000 and 18 800 due to the presence of more out-of-distribution data in these regions. In comparison, we also show the PI



FIG. 11. Relative error using Eq. (5) of the estimated fields \overline{F}_s^k for each test sample in the GTC case with $k_i = k_o = 1$. Each snapshot *k* represents 100 time steps in the GTC code, and snapshot 1 represents time step 4000. The vertical dashed line represents the linear-to-nonlinear transition point at time step 15 300 (i.e., snapshot k = 114). Left (random splitting setting): $n_{train} = 74$ and the previous iterate (PI) $|\overline{F}_s^k|\overline{F}_s^k = [F_s^{k-1}]F_s^{k-1}$. Right (sequential splitting setting): $n_{train} = 80, 100, 110, 120, 130$ and PI results. Top: Relative error of the RMS values using Eq. (15) for (a) sequential splitting. The quantities are computed as $[\overline{F}_s^k] = M_{RMS}([F_s^{k-1}])$ with M_{RMS} denoting the RMS-FC model. Middle: Relative error using Eq. (5) for normalized fields for (b) random splitting and (e) sequential splitting. The quantities are computed as $\overline{F}_s^k = M_{ENO}(F_s^{k-1})$ with M_{ENO} denoting the ST-FNO model. Bottom: Relative error using Eq. (5) for fields de-normalized with the RMS data as $[\overline{F}_s^k]\overline{F}_s^k$ for (c) random splitting and (f) sequential splitting.

results for the de-normalized fields: $[\bar{F}_s^k]\bar{F}_s^k = [F_s^{k-1}]F_s^{k-1}$ in Fig. 11(c). The proposed ST-FNO + RMS-FC model can yield up to 5× better inference accuracy compared with PI, but in the nonlinear phase their difference becomes less significant.



FIG. 10. Color contour plots of perturbed fields from the (top) ST-FNO results and (bottom) their relative difference using Eq. (17) with respect to the GTC simulation results (random splitting) at time steps 7500, 10 000, 12 500, 15 000, 17 500 for the field quantity $A_{||}$. $k_i = 1$, $k_o = 1$.

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FIG. 12. Color contour plots of perturbed fields from (top) ST-FNO results and (bottom) their relative difference using Eq. (17) with respect to the GTC simulation results (sequential splitting) at time steps 14 000 and 17 000 with different sized training data for the field quantity $A_{||}$, $k_i = 1$, $k_o = 1$.

2. Sequential splitting

Finally, we examine the prediction error per test sample in the sequential splitting settings [see Fig. 11(right column)]. We set n_{train} to different values and try to investigate how many more snapshots ST-FNO can predict a valuable solution (e.g., with the error below approximately 0.1) without retraining the model with newer snapshots. As expected, the errors go up as the snapshot index increases. That said, for RMS [Fig. 11(d)], one can accurately predict the results in the linear phase for the next 30–40 snapshots with only $n_{train} = 80$ training samples. For the normalized data [Fig. 11(e)] and denormalized data [Fig. 11(f)], the linear phase can be accurately predicted with $n_{train} = 80, 100, 110$. However, in the nonlinear phase, the ST-FNO can only predict the next snapshot more accurately than PI, suggesting that retraining is needed for every snapshot.

In addition, we visualize the fields predicted by ST-FNO (in the sequential splitting settings with $k_i = 1$ and $k_o = 1$) and simulated by GTC. Figure 12 plots the GTC simulation results, ST-FNO results, and their relative difference defined by Eq. (17) for the field quantity $A_{||}$ at two time steps. For time step 14 000 (snapshot 101), we consider the ST-FNO model with $n_{train} = 80$ and $n_{train} = 101$. For time step 17 000 (snapshot 131), we consider the ST-FNO model with $n_{train} = 120$ and $n_{train} = 130$. It is clear from both Figs. 11(f) and 12 that retraining the ST-FNO model becomes necessary in the sequential splitting setting.

IV. CONCLUSION

This paper proposed an efficient and accurate SciML model, called ST-FNO, which leverages the sparse connectivity indicated by the governing equations of fusion codes. ST-FNO has been applied to an extended-MHD code NIMROD and a gyrokinetic PIC code GTC to demonstrate its inference accuracy, memory efficiency, and CPU efficiency. We remark that for fusion simulation codes, or more generally multi-variable time-dependent PDEs with multiple variables, explicitly exploiting the sparsity dependency indicated by the governing equations can effectively reduce the sizes of SciML models without sacrificing the interference accuracy, which is particularly useful for data-scarce scientific applications.

The limitation of ST-FNO is the expensive training cost (just like most other ML models), and we plan to explore the idea of finetuning³² as well as further increasing the number of time steps per snapshot to reduce such cost. It is also likely that one can further sparsify ST-FNO by dropping weaker dependencies using more domain knowledge. A more systematic study of the prediction confidence, particularly for a large number of output time steps, is highly desirable for ST-FNO to be used as a reliable diagnostic tool for critical plasma events. Future work also includes integrating ST-FNO into fusion codes as an ML-based initial high-accuracy guess for preconditioners. Taking NIMROD for an example, this would require (1) conversion of the model output from uniform mesh in $r - \theta$ coordinate to the FEM mesh, (2) extension of the ST-FNO model to predict fully 3D data in the complex Fourier representation used by NIMROD, and (3) an automatic mechanism to determine when to retrain or fine-tune the ST-FNO model on-the-fly.

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AUTHOR DECLARATIONS

Conflict of Interest

The authors have no conflicts to disclose.

Author Contributions

Mustafa Mutiur Rahman: Data curation (equal); Investigation (equal); Methodology (lead); Software (lead); Validation (equal); Visualization (lead); Writing - original draft (equal); Writing - review & editing (equal). Zhe Bai: Data curation (equal); Methodology (equal); Validation (equal); Visualization (equal); Writing - review & editing (equal). Jacob Robert King: Data curation (equal); Methodology (equal); Validation (equal); Visualization (equal); Writing - review & editing (equal). Carl R. Sovinec: Validation (equal); Writing - review & editing (equal). Xishuo Wei: Data curation (equal); Methodology (equal); Validation (equal); Visualization (equal); Writing - review & editing (equal). Samuel Williams: Project administration (equal); Writing - review & editing (equal). Yang Liu: Conceptualization (lead); Data curation (equal); Funding acquisition (lead); Methodology (equal); Project administration (equal); Supervision (lead); Writing - original draft (equal); Writing - review & editing (equal).

DATA AVAILABILITY

The machine learning code supporting the findings of this study is available in the GitHub repository at https://github.com/liuyangzhuan/STFNO. A selected dataset is available at https://zenodo.org/ records/13901806. The other datasets are available from the corresponding author upon reasonable request. 04 April 2025 21:54:50

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