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## **RESEARCH ARTICLE**

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### **Key Points:**

- We propose a symbiotic ocean modeling framework in which models of different complexities benefit from each other
- Unresolved processes are represented through hybrid machine learning methods using data from the symbiotic framework
- Hybrid correction strategies with imperfect physics as control input improve the representation of key long-term flow properties

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# Symbiotic Ocean Modeling Using Physics-Controlled Echo State Networks

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**Abstract** We introduce a "symbiotic" ocean modeling strategy that leverages data-driven and machine learning methods to allow high- and low-resolution dynamical models to mutually benefit from each other. In this work we mainly focus on how a low-resolution model can be enhanced within a symbiotic model configuration. The broader aim is to enhance the representation of unresolved processes in low-resolution models, while simultaneously improving the efficiency of high-resolution models. To achieve this, we use a grid-switching approach together with hybrid modeling techniques that combine linear regression-based methods with nonlinear echo state networks. The approach is applied to both the Kuramoto–Sivashinsky equation and a single-layer quasi-geostrophic ocean model, and shown to simulate short-term and long-term behavior better than either purely data-based methods or low-resolution models. By maintaining key flow characteristics, the hybrid modeling techniques are also able to provide higher quality initial conditions for high-resolution models, thereby improving their efficiency.

**Plain Language Summary** Models of the ocean vary in complexity. Some are very detailed and manage to show oceanic vortices, whereas others are very efficient but coarse, and unable to compute such vortices. The idea in this paper is to let these different model types work together and benefit from each other, as if in a symbiosis. With knowledge of differences between the detailed and coarse model we can use machine learning techniques to improve the coarse model. In this way a coarse model can be used to provide good quality predictions and to aid a detailed model by taking over part of its computations. We apply our ideas to the Kuramoto–Sivashinsky (KS) model and a quasi-geostrophic (QG) ocean model, where we show that promising short-term KS results may generalize to models of the ocean. Long-term equilibrium experiments with QG show in addition how the correction strategies let a coarse model produce correct flow properties, where standalone physics- or data-based approaches fail. These improved coarse models are computationally cheap, yet good enough to give initial conditions for the fine model, showcasing the symbiotic modeling idea.

### 1. Introduction

One of the most important spatial scales in the ocean circulation is the internal Rossby radius of deformation  $L_D$ ; it ranges from 50 to 100 km at mid-latitudes to a few km in the polar regions (Hallberg, 2013). At this scale, perturbations are amplified on mean flows through mixed barotropic/baroclinic instability, giving rise to ocean eddies. Interactions between these eddies and the mean flow can lead to up-gradient momentum transport affecting the strength and separation of ocean western boundary currents such as the Kuroshio and Agulhas (Chassignet et al., 2020).

Most climate models, in particularly those used in CMIP5 and CMIP6, do not resolve ocean processes at the scale  $L_D$  as the spatial grid size used is too large; typically 1° (Eyring et al., 2016). The main reason is computational costs, as doubling the horizontal resolution increases these costs roughly by a factor 10. Effects of subgrid-scale processes are hence parameterized in these models. For example, the effect of ocean eddies on tracer transport is represented by the Gent–McWilliams (Gent et al., 1995) scheme, but such a scheme cannot capture, for example, the up-gradient momentum transport. Hence, western boundary flows are too weak and diffuse, and do not separate at the correct location (Chassignet et al., 2020).

Over the last few years, first simulations have been performed with global climate models, where the ocean model component has a resolution of  $0.1^{\circ}$ , which is smaller than  $L_D$  for many locations on the globe (Chang

et al., 2020; Jüling et al., 2021). We will refer to those models as high-resolution (HR) models to contrast them with the 1° models which we will call low-resolution (LR) models. But also the high-resolution models are not completely eddy-resolving as this requires an even higher spatial resolution. There is now a substantial amount of model data available to compare results on ocean-climate variability and climate change for both types of models. High-resolution models tend to reduce biases compared to observations, particularly in western boundary currents, sea surface temperature variability patterns and Southern Ocean mean flows (Chang et al., 2020; Jüling et al., 2021).

However, HR model simulations form a great drain on computational resources and hence there are still many efforts to represent the effects of unresolved processes in LR models. This parameterization process has been around for decades and approaches can be grouped into three types. First, semi-empirical parameterizations are used, where observation motivated schemes are implemented (Gargett, 1989; Viebahn et al., 2019). Second, theoretically derived schemes, where specific approximations are made in the underlying equations (Gent et al., 1995) have been used. Third, stochastic schemes derived from sampled high-resolution model simulations (Berloff, 2005; Mana & Zanna, 2014) have shown potential in representing unresolved processes in LR models (Hewitt et al., 2020).

Recently, a new approach has been added, where the subgrid-scale (SGS) model is based on machine learning (ML) techniques. In Bolton and Zanna (2019), a convolutional neural network (CNN) was trained with data from a high-resolution model of the mid-latitude ocean gyres. This CNN was shown to successfully capture the small-scale processes and the effects of those on the mean flow in the low-resolution version of the same model. Capturing up-gradient momentum transport in turbulent flows is a crucial test for ML-based SGS models and the quasi-geostrophic (QG) equations form an ideal testing ground for this problem. Effective learning strategies based on CNNs and applied to QG are for instance presented in Frezat et al. (2022) and Guan et al. (2023), where in the latter study physics-based augmentations and constraints are introduced.

CNNs are a special variant of the traditional feed-forward neural network architecture (FFNN), which has also been used for subgrid-scale representations in both ocean and atmospheric models (Irrgang et al., 2021; Rasp et al., 2018). Another ML architecture that shows promise in the modeling of climate physics is the reservoir computing approach, often referred to as an echo state network (ESN). An ESN is especially suited to simulate chaotic dynamics (Jaeger & Haas, 2004; Pathak et al., 2017) and is shown to be capable of emulating interactions between empirical orthogonal functions (EOFs) (Nadiga, 2021). ESNs are a type of recurrent neural network (RNN), which are in fact dynamical systems with internal states that are propagated according to recursive relations. This property sets them apart from FFNNs, which can be seen as functions (Lukoševičius & Jaeger, 2009). A relatively low training cost and a limited number of hyperparameters make ESNs stand out against similar RNN architectures such as Long Short-Term Memory (LSTM) and Gated Recurrent Units (GRU) (Vlachas et al., 2020). Furthermore, ESNs are shown to have close computational and theoretical connections with linear regression and models based on dynamic mode decomposition (DMD) (Bollt, 2021; Kutz et al., 2016; Schmid, 2010).

Recent "hybrid" (or physics-controlled) ESN advances (Pathak et al., 2017, 2018) provide a simple and effective approach to correct known model imperfections, such as those due to the lack of eddies in LR ocean models. With training data based on ground truths and imperfect model predictions, model tendencies and nonlinear model mismatches are encoded in an ESN. The result is an artificial dynamical system that can be controlled using an imperfect model. Combining an imperfect model with corrections from a trained ESN creates a hybrid dynamical system that greatly outperforms both the network and the imperfect model (Wikner et al., 2020). This approach was recently applied to an atmospheric model (SPEEDY) and shown to be able to improve the simulations of mean flow and variability considerably on short time scales (Arcomano et al., 2022).

In this paper we use the hybrid modeling framework as a key ingredient for a "symbiotic" ocean modeling approach. The idea is to couple models of different complexities and configure them to solve the same problem, where we distinguish between perfect and imperfect models in terms of differing resolution and parameterizations. This model coexistence can be made mutually beneficial using data-driven and ESN-based techniques. With the symbiotic approach we aim to improve the computational efficiency of HR models, while simultaneously enhancing the parameterizations of unresolved processes in LR models. Our approach shares similarities with the coupling techniques in Barthélémy et al. (2022) and Counillon et al. (2023). Yet, here our broader aim is not a synchronized solution, but to be able to alternate between LR and HR models during time integration, with



simultaneous transients only present during a training phase. Hence our concept of symbiotic modeling involves both synchronous and alternating propagations of HR and LR models.

We will first focus on using the hybrid modeling strategy with data generated from both LR and HR models to correct imperfect model transients. Model corrections made can then be seen as modeling subgrid effects. To this end, we employ a linear grid-switching approach and introduce a correction framework that includes models based on linear regression, DMD, ESN and hybrid variants (Section 2). We apply the correction strategy to coupled LR and HR versions of the Kuramoto–Sivashinsky (KS) equations (Section 3) and a single-layer quasi-geostrophic (QG) ocean model (Section 4). With the ocean model we perform both short-term predictions and long-term equilibrium runs to compare the available corrective models. In the final part of Section 4 we test the symbiotic idea for the QG equations and lift LR predictions back to the HR grid where they serve as initial conditions for numerous HR simulations. A summary and discussion with the main conclusions is provided in Section 5.

### 2. Methods

In a general framework, the HR model is defined on a fine grid  $\Omega^{f}$  and is regarded as a *perfect* model. An LR model is considered as an *imperfect* model, and is defined on a coarse grid  $\Omega^{c}$ . The grids  $\Omega^{f}$  and  $\Omega^{c}$  have dimensions  $N_{f}$  and  $N_{c}$ , respectively, and cover the same domain. Both models attempt to solve the same problem, but apart from different grids we also allow differences in key parameters and forcings between the perfect and imperfect model. The physics resolved by the perfect model is then used as ground truth and the imperfect model results are considered to be in need of correction.

The perfect model is a system of coupled partial differential equations (PDEs), spatially discretized on  $\Omega^{f}$ , which leads to a large system of differential-algebraic equations (DAEs):

$$M_P \dot{\boldsymbol{\xi}} = F_P(\boldsymbol{\xi}), \text{ with } \boldsymbol{\xi} \in \mathbb{R}^{N_f}.$$
 (1)

Here,  $\boldsymbol{\xi} = \boldsymbol{\xi}(t)$  is a time dependent state vector and  $M_P \in \mathbb{R}^{N_f \times N_f}$  is a mass matrix that determines the dependence on temporal derivatives. The nonlinear operator  $F_P : \mathbb{R}^{N_f} \to \mathbb{R}^{N_f}$  is a spatial discretization of the perfect model physics. Similarly, the semi-discretized imperfect model has a coarse state  $\mathbf{x} = \mathbf{x}(t)$  that evolves according to

$$M_I \dot{\mathbf{x}} = F_I(\mathbf{x}), \text{ with } \mathbf{x} \in \mathbb{R}^{N_c},$$
 (2)

where  $M_I \in \mathbb{R}^{N_c \times N_c}$  and  $F_I : \mathbb{R}^{N_c} \to \mathbb{R}^{N_c}$  are again the mass matrix and spatial discretization operator.

Transfers between the solutions on the two grids  $\Omega^f$  and  $\Omega^c$  are made through a fully weighted restriction  $R \in \mathbb{R}^{N_c \times N_f}$  and a prolongation operator  $P \in \mathbb{R}^{N_f \times N_c}$ . We choose these operators for their convenient (variational) property that they are each other's transpose up to a constant factor:  $R = cP^{\top}$  (Briggs et al., 2000). The perfect model evolves according to  $\phi_P : \mathbb{R}^{N_f} \to \mathbb{R}^{N_f}$ . Similarly, the evolution of the imperfect model is given by  $\phi_I : \mathbb{R}^{N_c} \to \mathbb{R}^{N_c}$ . Hence  $\phi_P$  and  $\phi_I$  are time-propagation operators that represent time-discretized versions of Equations 1 and 2, respectively. We assume a fixed time step  $\Delta t$ ; an uncorrected, imperfect model state that has been propagated for a single  $\Delta t$  will be denoted as  $\tilde{\mathbf{x}}^{k+1} = \phi_I(\mathbf{x}^k)$ .

The imperfect spatial discretization  $F_I$  is incapable of capturing the physics resolved by the perfect model and we therefore attempt to improve the imperfect evolution  $\phi_I$  with a combination of linear and non-linear corrections. As these corrections are data-driven we divide our approach into a data gathering and a prediction phase.

### 2.1. Data Gathering

We gather data from a trajectory of  $\boldsymbol{\xi}(t)$  on  $\Omega^{f}$ . From this transient, associated restricted states, imperfect predictions and auxiliary states are computed. Starting at time  $t_0$ , we collect  $N_T + 1$  snapshots of the evolving state  $\boldsymbol{\xi}(t)$ :

$$\{\boldsymbol{\xi}^{0}, \boldsymbol{\xi}^{1}, \dots, \boldsymbol{\xi}^{N_{T}}\}, \quad \boldsymbol{\xi}^{k} = \boldsymbol{\xi}(t_{0} + k\Delta t),$$
(3)

at fixed time intervals  $\Delta t$  such that we cover the model time  $T = N_T \Delta t$ . The snapshots are restricted to the coarse grid and combined into two data matrices:

$$X = \begin{bmatrix} \mathbf{x}^0, \ \mathbf{x}^1, \ \dots, \ \mathbf{x}^{N_T - 1} \end{bmatrix} = \begin{bmatrix} R\xi^0, \ R\xi^1, \ \dots, \ R\xi^{N_T - 1} \end{bmatrix},$$
(4)



$$X' = \begin{bmatrix} \mathbf{x}^1, \ \mathbf{x}^2, \ \dots, \ \mathbf{x}^{N_T} \end{bmatrix} = \begin{bmatrix} R\xi^1, \ R\xi^2, \ \dots, \ R\xi^{N_T} \end{bmatrix}.$$
 (5)

Apart from the restricted data matrix  $X \in \mathbb{R}^{N_c \times N_T}$  and its shifted version  $X' \in \mathbb{R}^{N_c \times N_T}$ , we also create a collection of imperfect predictions  $\Phi(X)$ :

$$\Phi(X) = \left[\phi_I(\mathbf{x}^0), \ \phi_I(\mathbf{x}^1), \ \dots, \ \phi_I(\mathbf{x}^{N_T - 1})\right] \in \mathbb{R}^{N_c \times N_T}.$$
(6)

The matrices X and  $\Phi(X)$  contain perfect and imperfect model tendencies that serve as forcing to an auxiliary (surrogate) model *f*. The exact structure of this model is described in more detail in Section 2.3. The model *f* has an internal state **s** and is subjected to an input forcing **u**. We evolve *f* over the same time period *T* and obtain snapshots from it at the same fixed intervals  $\Delta t$ . Hence we can iterate according to

$$\mathbf{u}^{k} = h(\mathbf{x}^{k}, \phi_{I}(\mathbf{x}^{k})), \tag{7}$$

$$\mathbf{s}^{k+1} = f\left(\mathbf{s}^k, \mathbf{u}^k\right),\tag{8}$$

where the input forcing **u** is given by a mapping *h* and with initialization  $\mathbf{s}^0 = \mathbf{s}_0$  at  $t = t_0$ . For *h* we either use a selection, for example,  $h(\mathbf{x}^k, \mathbf{\tilde{x}}^{k+1}) = \mathbf{x}^k$  or combine the forcing such that  $h(\mathbf{x}^k, \mathbf{\tilde{x}}^{k+1}) = (\mathbf{x}^k; \mathbf{\tilde{x}}^{k+1})$ , where (;) denotes vertical stacking. These are the most straightforward choices and of course other options are possible here. The surrogate model *f* comes in the form of an Echo State Network (ESN) and is described in Section 2.3. From the evolution of *f* we gather  $N_T + 1$  state snapshots  $\mathbf{s}^0, \dots, \mathbf{s}^{N_T}$  and combine them into a data matrix, with the exception of the initialization  $\mathbf{s}^0$ :

$$S = \begin{bmatrix} \mathbf{s}^1, \ \mathbf{s}^2, \dots, \ \mathbf{s}^{N_T} \end{bmatrix} \in \mathbb{R}^{N_r \times N_T}.$$
(9)

#### 2.2. Prediction

The data gathered up until time  $t = t_0 + T$  is used to obtain linear best fit operators for a corrected prediction strategy. Given data X, X',  $\Phi(X)$ , and S, these operators optimally combine **x**,  $\phi_I(\mathbf{x})$  and **s** to improve the imperfect evolution given by  $\phi_I$  alone. Here we provide a general transient strategy that covers a number of different corrective methods.

A corrected imperfect transient is started at  $t_0 + T$ . Now, the models  $\phi_I$  and f operate in isolation from any perfect model data and f augments  $\phi_I$ . Using starting states  $\mathbf{x}^{N_T}$  and  $\mathbf{s}^{N_T}$ , the transient proceeds as follows:

$\widetilde{\mathbf{x}}^{k+1} = \boldsymbol{\phi}_{I}\left(\mathbf{x}^{k}\right)$	create an imperfect model prediction,	(10)
$\mathbf{u}^{k} = h\big(\mathbf{x}^{k}, \widetilde{\mathbf{x}}^{k+1}\big)$	construct a forcing,	(11)
$\mathbf{s}^{k+1} = f\left(\mathbf{s}^k, \mathbf{u}^k\right)$	evolve the auxiliary state,	(12)
$\mathbf{x}^{k+1} = A\mathbf{x}^k + B\widetilde{\mathbf{x}}^{k+1} + C\mathbf{s}^{k+1}$	create an improved prediction,	(13)

for  $k = N_T, N_T + 1, \dots$  Hence the trajectory of **x** is initialized with a restricted truth  $(\mathbf{x}^{N_T} = R\boldsymbol{\xi}^{N_T})$  but continues independently of the perfect model  $(\mathbf{x}^{N_T+1} \neq R\boldsymbol{\xi}^{N_T+1})$ .

With the general formulation in Equations 10-13 we aim to include several methods and their combinations in the same framework. The operators *A*, *B*, *C* in Equation 13 have separate interpretations. On its own, *A* is obtained as a linear best fit of the propagation from *X* to *X'*. Its eigendecomposition is known as a dynamic mode decomposition (DMD) (Kutz et al., 2016; Schmid, 2010) and *A* is often called a DMD-operator. The matrix *B* is the best direct correction of  $\Phi(X)$  to *X'* in the least squares sense. Lastly, as *f* is a neural net, the operator *C* is the optimal output layer, that is, the linear best fit translation of *S* to *X'*. Hence these different methods can be seen as special cases in Equations 10-13.

Combinations of the operators A, B, and C are fitted at  $t = t_0 + T$  using regularized linear regressions with the data matrices X, X',  $\Phi(X)$ , and S. Choices for the architecture of f and h and the use of operators A, B, C lead to a variety of predictive methods (Table 1). A *model only* approach uses B = I and ignores A and C. The transient



Table 1

Overview of Corrective Methods Based on Operator Configurations in Equation 13				
Method	$h(\mathbf{x}^k, \widetilde{\mathbf{x}}^{k+1})$	Known operators	Minimization to compute unknown operators	
Model only		A = 0, B = I, C = 0	No minimization necessary	
ESN	$\mathbf{x}^k$	A = 0, B = 0	$\min_{C} \left\  CS - X' \right\ _{F}$	
DMD		B = 0, C = 0	$\min_{A} \left\  AX - X' \right\ _{F}$	
ESNc	$\begin{bmatrix} \mathbf{x}^k \\ \widetilde{\mathbf{x}}^{k+1} \end{bmatrix}$	A = 0	$\min_{[B \ C]} \left\  \begin{bmatrix} B \ C \end{bmatrix} \begin{bmatrix} \Phi(X) \\ S \end{bmatrix} - X' \right\ _{F}$	
DMDc		<i>C</i> = 0	$\min_{[A \ B]} \left\  \begin{bmatrix} A \ B \end{bmatrix} \begin{bmatrix} X \\ \Phi(X) \end{bmatrix} - X' \right\ _{F}$	
Correction-only		A = 0, C = 0	$\min_{B} \left\  B\Phi(X) - X' \right\ _{F}$	
ESN + DMD	$\mathbf{x}^k$	B = 0	$\min_{[A  C]} \left\  \begin{bmatrix} A & C \end{bmatrix} \begin{bmatrix} X \\ S \end{bmatrix} - X' \right\ _{F}$	
ESN + DMDc	$\begin{bmatrix} \mathbf{x}^k \\ \widetilde{\mathbf{x}}^{k+1} \end{bmatrix}$		$\min_{\begin{bmatrix} A & B & C \end{bmatrix}} \begin{bmatrix} A & B & C \end{bmatrix} \begin{bmatrix} X \\ \Phi(X) \\ S \end{bmatrix} - X' \\ \end{bmatrix}_{E}$	

*Note.* The associated minimizations are linear regression problems for which we do not include the regularization here;  $\| \|_F$  is the Frobenius norm. Additional variations on these methods rely on the specific architecture chosen for *h* and *f*. The choices

(Equations 10–13) is reduced to only the imperfect model evolution. In an *ESN* prediction we translate from states of the neural net (ESN) to predictions using a best fit *C*. Here, *f* is forced with restricted states only:  $\mathbf{x}^k$ . A *DMD* prediction is based on the best linear approximation of the propagation from *X* to *X'*. When the operators *B* and *C* are combined and  $\mathbf{u}^k = (\mathbf{x}^k; \mathbf{\tilde{x}}^{k+1})$ , the auxiliary model *f* is subjected to a physics-based control  $\phi_I(\mathbf{x})$ , both internally through  $\mathbf{u}$  and externally through *B*. With *f* an ESN this is referred to as ESNc, which is equivalent to the hybrid scheme in Pathak et al. (2018). DMDc denotes DMD with control (Proctor et al., 2016) and is obtained by combining operators *A* and *B*. In DMDc the imperfect physics assist the DMD model which, on its own, generalizes poorly outside the training data. A basic *correction-only* approach follows from using only *B*, whereas additional combinations lead to the varieties ESN + DMD and ESN + DMDc. Connections between ESNs and DMD exist (Bollt, 2021) and within this framework it is straightforward to combine (and consequently isolate) both approaches.

The minimizations shown in Table 1 are computed using Tikhonov regularization, which introduces an additional penalty on the size of the fitted operator. Regularization is crucial as it reduces overfitting and improves the stability of a long-term transient (Lukosevicius, 2012). For instance, the DMD-operator actually minimizes

$$\min_{A} (\|AX - X'\|_{F} + \lambda^{2} \|A\|_{F}),$$
(14)

with  $\lambda > 0$  a regularization parameter.

we make for h are added as a separate column to this table.

### 2.3. Echo State Network

An echo state network (Jaeger, 2001; Jaeger & Haas, 2004) will act as the auxiliary predictive model *f*. Here we will roughly outline the organization of an ESN. For a detailed explanation we refer to Pathak et al. (2018), which we follow closely. An ESN is a recurrent neural network that can be viewed as an artificial nonlinear dynamical system with a state  $\mathbf{s} \in \mathbb{R}^{N_r}$  of sufficient dimension  $N_r$ . Typically, better prediction results are achieved for



larger  $N_r$ , at the cost of increased training time. The size of the ESN state **s** is therefore one of the key tunable hyperparameters in our framework. The components of **s** interact through a sparse, random linear operator  $W : \mathbb{R}^{N_r} \to \mathbb{R}^{N_r}$  that is not altered after initialization. The average degree of the adjacency graph associated with W is denoted with  $\overline{d}$ . Input data  $\mathbf{u} \in \mathbb{R}^{N_u}$  is standardized (every unknown has zero mean and unit variance) and is fed as forcing to the system, where it is combined with the state using a fixed linear operator  $W_{in} : \mathbb{R}^{N_u} \to \mathbb{R}^{N_r}$ . The input operator  $W_{in}$  is random and sparse, with only a single element per row that is drawn from a uniform distribution on [-1, 1]. The internal state evolves according to

$$\mathbf{s}^{k+1} = f\left(\mathbf{s}^{k}, \mathbf{u}^{k}\right) = (1 - \alpha)\mathbf{s}^{k} + \alpha \tanh\left(W\mathbf{s}^{k} + W_{in}\mathbf{u}^{k}\right), \quad \mathbf{s}^{0} = \mathbf{s}_{0}$$
(15)

with initialization  $\mathbf{s}_0$  and a relaxation parameter  $\alpha \in (0, 1]$  (also known as the leaking rate) that controls the "speed" of the artificial dynamics (Lukoševičius & Jaeger, 2009). Hence the state  $\mathbf{s}$  evolves according to a deterministic iteration with internal interactions given by a random (but fixed) W and forcing provided by the input data. The tanh(·) activation function introduces a nonlinearity that is controlled by the weights in  $W_{in}$ . The spectral radius of W,  $\rho(W)$ , determines the amplification or damping of  $\mathbf{s}^k$  due to the recursive application of Equation 15 and thereby controls the memory of system. From Equation 15 it is apparent that  $\alpha$  allows a matching of time scales between the network and the variability in the training data, which is beneficial to the network's predictive performance (Lukoševičius & Jaeger, 2009). The presence of  $\alpha \in (0, 1]$  is the only significant difference between our formulation of f and the original hybrid ESN in Pathak et al. (2018), which can be viewed as having  $\alpha = 1$ . ESN prediction results are in fact sensitive to  $\alpha$  and we treat it as a key hyperparameter that requires tuning.

Starting at  $t = t_0$  with  $\mathbf{s}_0$ , the recursion (Equation 15) generates  $N_T$  new states that are combined into a data matrix S, as described in Section 2.1. A linear operator C provides output predictions by translating the auxiliary state to a prediction. In the standard ESN approach the output operator  $C : \mathbb{R}^{N_r} \to \mathbb{R}^{N_c}$  is computed from a regularized minimization problem using  $S^*$  and X', see Section 2.2. Here  $S^*$  is an adapted version of S. As in Pathak et al. (2018), we take the square of the even elements in each state  $\mathbf{s}^k \in S$ . The motivation for this is largely empirical but related to problems that may originate with capturing symmetry in the model equations (Lu et al., 2017).

### 3. Results: Kuramoto-Sivashinsky Model

In Pathak et al. (2018) a hybrid ESN was applied to the Kuramoto–Sivashinsky (KS) equation. Here we will begin with a test of our framework by replicating these results. We will first consider equal grids ( $N_f = N_c$ ) and introduce an imperfection through a perturbation in one of the parameters. Later we explore a perfect/imperfect model setup with  $N_f = 2N_c$  and no parameter perturbation in the KS-equation.

The KS-equation is capable of displaying rich spatiotemporal dynamics and is used to study a variety of phenomena such as flame front dynamics (Sivashinsky, 1977) and reaction-diffusion dynamics (Kuramoto, 1984). In one dimension it is given by

$$\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} + (1+\epsilon) \frac{\partial^2 u}{\partial x^2} + \frac{\partial^4 u}{\partial x^4} = 0,$$
(16)

with  $u \in [0, L]$ , initial value  $u(x, 0) = u_0(x)$  and periodic boundaries u(x, t) = u(x + L, t). The domain size *L* is also the bifurcation parameter of the problem. In Pathak et al. (2018), the domain size is chosen at L = 35, for which the KS-equation has a positive maximum Lyapunov exponent  $\lambda_{max}$  and produces chaotic behavior (Hyman & Nicolaenko, 1986). A perturbation  $\epsilon \ge 0$  is introduced to create an imperfection. With  $\epsilon = 0$  we obtain the true, "perfect" evolution whereas our "imperfect" model will have  $\epsilon > 0$ .

The KS-equation is discretized on an equidistant grid:  $x_i = i/N_f$  with  $i = 1, 2, ..., N_f = N_c = 64$ . We use a fully implicit time stepping scheme with  $\Delta t = 0.25$  and initialize with

$$u_0(x_i) = \begin{cases} 1, & i = 1, \\ 0, & i > 1. \end{cases}$$

Starting at  $t = t_0$ , a transient is computed up to T = 6,000 from which we select a large number of training and testing intervals. In the remaining experiments we follow a similar procedure with long transients to sample



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**Figure 1.** Results for the replication of the experiments in Pathak et al. (2018) where the imperfect model is a perturbed version of the KS-equation with perturbation parameter  $\epsilon$ . "Valid time" is the time it takes until the error threshold is passed:  $E(\mathbf{x}^k, \mathbf{y}^k) > 0.4$ . These timings are in Lyapunov units ( $\lambda_{max}t$ ). The experiment is repeated for 100 different training intervals and network realizations. For each  $N_r$  a box plot is depicted showing the first, second and third quartile.

training periods from. This approach is efficient from a data-management perspective and will be necessary to consistently train in the same dynamical regime, but it does not guarantee uncorrelated data and might introduce biases. However, in our comparison with Pathak et al. (2018) we do not encounter such problems.

The ESN used closely follows that in Pathak et al. (2018). The spectral radius is set at  $\rho(W) = 0.4$ , the average degree is  $\overline{d} = 3$ , we use training intervals of size T = 5,000 and ignore any relaxation with  $\alpha = 1$ . The KS-equation and its discretization are also equivalent to Pathak et al. (2018) so, for a coherent interpretation of the predictions, we scale the obtained timings with the same Lyapunov exponent  $\lambda_{max} = 0.07$ .

The methods summarized in Table 1 are compared in a scaling experiment where the auxiliary state size  $N_r$  is doubled several times (see Figure 1). Only those methods based on an ESN depend on this parameter which leads to constant results for the other predictions. For each method we use 100 different training intervals and hence network realizations, as we do not reuse W. We fix the regularization parameter at  $\lambda = 1 \cdot 10^{-5}$ . The pure DMD-based methods (DMD and ESN + DMD) are not shown as they did not produce meaningful results. This is likely caused by DMD generalizing poorly to unseen data and hence showing only valid predictions for a short period after  $t_0 + T$ .

The short-term prediction accuracy is measured with the normalized error used in Pathak et al. (2018). We compare the *k*th prediction  $\mathbf{x}^k$  with the restricted truth  $\mathbf{y}^k = R\boldsymbol{\xi}^k$  through

$$E(\mathbf{x}^{k}, \mathbf{y}^{k}) = \frac{\|\mathbf{x}^{k} - \mathbf{y}^{k}\|}{\sqrt{\langle \|\mathbf{y}^{k}\|^{2} \rangle}},$$
(17)





**Figure 2.** Grid experiment with the KS-equation. The imperfect model consists of the same equations but discretized on a grid half the resolution of the perfect model. Solutions are valid until  $E(\mathbf{x}^k, \mathbf{y}^k) > 0.4$ . As in Figure 1, we repeat the experiment for 100 different training sets and network realizations.

with  $\langle \cdot \rangle$  the mean over a time window up until *k*.

The results in Figure 1 show a strong resemblance with the scaling experiment in Pathak et al. (2018) (Figure 8 in that paper). There, similar valid times are obtained for ESN, ESNc and the standalone imperfect model. However, our ESN appears to suffer more from a stagnation at higher  $N_r$  values, which is likely due to differences in regularization. The imperfect model performs poorly on its own and the ESN-based methods improve the predictions as expected. A standalone ESN is able to achieve decent predictions for  $\epsilon = 1$  and  $\epsilon = 0.1$ . For  $\epsilon = 0.01$ , however, it appears impossible for a standalone ESN to perform better than the imperfect model. In all studied cases it is remarkable how the hybrid variant ESNc stands out. By combining the imperfect model physics with the ESN a significant improvement is achieved.

The three additional models in Table 1 further explain the advantage of the hybrid ESNc over the standalone ESN. The correction-only and DMDc predictions do not depend on  $N_r$  and show up here as constant solutions. These two regression-based corrections outperform the standalone ESN for  $\epsilon = 0.1$  and  $\epsilon = 0.01$ . The third approach, ESN + DMDc, follows the ESNc performance but with an overall slight advantage for the two largest perturbations  $\epsilon$ . This advantage is explained by the performance of DMDc and correction-only, as these are the linear components of ESN + DMDc and ESNc, respectively. In experiments where DMDc outperforms the correction

we find a similar overall gain between ESN + DMDc and ESNc. From the experiments in Figure 1 it is apparent that ESN + DMDc and ESNc reduce to their linear components for low  $N_r$ , which is what would be expected from the correction Equation 13. Hence the performance of the linear models can be seen as a departure point for hybrid variants that add a nonlinear ESN. This largely explains the performance gain of, for example, ESNc over the standalone ESN.

In a different perfect/imperfect model setup, illustrating the symbiotic modeling approach, the models both use  $\epsilon = 0$  and have different spatial resolutions instead. The perfect model is discretized on a grid with twice the resolution,  $N_f = 2N_c$ . The domain size, ESN parameters and regularization remain unchanged. As explained in Section 2.1, fine grid information is restricted to the coarse grid and any data-driven corrections are made to the imperfect, coarse model evolution. Hence, instead of a model perturbation, it is now the difference in truncation errors and resolved scales between two resolutions that causes a model mismatch. With this setup the approach given by Equations 10–13 can be seen as a subgrid scale (SGS) modeling technique.

The coarse model is capable of a good prediction in this setup (Figure 2). DMDc, the correction-only and the standalone ESN are all unable to improve the coarse model. However, the hybrid variants ESNc and ESN + DMDc do show an overall improvement and an increase in predictive skill for larger  $N_r$ , similar to the parameter perturbation results (Figure 1). For large values of  $N_r$  the hybrid methods double the predictive performance. This, again, shows the benefit of introducing the imperfect physical predictions to both force and control the artificial ESN. Hence the hybrid approach in Pathak et al. (2018) shows promise as a nonlinear subgrid modeling technique.

### 4. Results: Quasi-Geostrophic Model

The barotropic quasi-geostrophic (QG) vorticity equation for a square (length *L*, constant depth *D*) ocean basin is solved on a  $\beta$ -plane. The ocean flow is driven by an idealized and deterministic wind-stress forcing pattern. Typical horizontal length and velocity scales are denoted *L* and *U*, from which the time scale follows as L/U. Using  $L = 10^6$  m and  $U = 3.17 \cdot 10^{-2}$  ms<sup>-1</sup>, we obtain a time scale of approximately one year. The equations are solved on a square domain,  $x \in [0, 1], y \in [0, 1]$ , with periodic boundaries in both directions.

The QG equations in non-dimensional form are given by

$$\left[\frac{\partial}{\partial t} - \frac{\partial\psi}{\partial y}\frac{\partial}{\partial x} + \frac{\partial\psi}{\partial x}\frac{\partial}{\partial y}\right](\omega + \beta y) = \frac{1}{\mathrm{Re}}\nabla^2\omega + \alpha_{\mathrm{r}}C_{\mathrm{r}}(x, y),\tag{18}$$





**Figure 3.** Snapshots of the vorticity fields (in day<sup>-1</sup>) at the end of the transient depicted in Figure 5. (a) Perfect model vorticity snapshot from a statistical equilibrium with  $N_f = 2 \cdot 256^2$  and  $\text{Re}_f = 2,000$ . (b) Imperfect model vorticity, also in a statistical equilibrium,  $N_c = 2 \cdot 32^2$  and  $\text{Re}_c = 500$ .

$$\omega = \nabla^2 \psi, \tag{19}$$

with  $\omega$  the vertical component of the vorticity vector and streamfunction  $\psi$ . The Reynolds number is Re =  $LU/A_H$ , where  $A_H$  is the horizontal mixing coefficient and  $\beta = \beta_0 L^2/U$ , with  $\beta_0 = 1.6 \cdot 10^{-11} \text{ (ms)}^{-1}$ . Wind forcing enters through the nondimensional parameter  $\alpha_\tau = \tau_0 L/(\rho D U^2)$ , with forcing amplitude  $\tau_0 = 0.3 \text{ Nm}^{-2}$ , density  $\rho = 1 \cdot 10^3 \text{ kg m}^{-3}$  and layer depth  $D = 6 \cdot 10^2 \text{ m}$ . We use a constant idealized wind-stress curl forcing in the form of a stirring pattern with stirring wavenumber  $k_f = 5$  in both directions:

$$C_{\tau}(x, y) = \cos(2k_f \pi x) \cos(2k_f \pi y).$$
(20)

This problem setup is a variant of the approach in Edeling and Crommelin (2019), but here a rotating situation is considered. With doubly periodic boundaries, sufficient resolution, and a high enough Reynolds number, the rotating barotropic flow will organize into a zonal jet pattern, similar to the structure of Jupiter's atmosphere (Farrell & Ioannou, 2007). Different zonal patterns are possible under the same forcing conditions due, not only to translational symmetries, but also to turbulent self-organization processes (Bouchet et al., 2019).

### 4.1. Approach

Following the perfect/imperfect modeling approach we discretize the QG equations on two different grids. The perfect model uses a fine discretization on  $\Omega^{f}$  with  $N_{f} = 2 \cdot 256^{2}$  unknowns and the imperfect variant is discretized on  $\Omega^{c}$  with  $N_{c} = 2 \cdot 32^{2}$  unknowns ( $N_{f} = 64N_{c}$ ). Furthermore, for both grids we model a flow with a Reynolds number that does not cause any numerical artifacts. With the perfect model we can run with  $\text{Re}_{f} = 2,000$ . A stable flow for this Reynolds number and forcing amplitude  $\tau_{0}$  is impossible to achieve on the coarse grid and we therefore choose to use  $\text{Re}_{c} = 500$  for the imperfect model.

For the discretization in time we use a fully implicit time stepping scheme that allows the use of the same time step for both models. In our experiments we will use  $\Delta t = 1$  day which is stable on both grids and choices for Re. The perfect QG state ( $\omega, \psi$ ) is randomly initialized and we run the model into a statistical steady state. From the steady state we select training periods of size  $N_T = 10,000$  days and follow the data gathering process described in Section 2.1. We ensure that the perfect QG reference trajectory contains no transitions between zonal jet patterns and hence all training intervals are part of the same statistical equilibrium. To get an idea of the perfect and imperfect flows we restart the imperfect model from a restricted fine state and run it into a steady regime. Snapshots from the two different statistical steady states are shown in Figure 3. The imperfect model solution in Figure 3b is highly diffusion dominated and shows a flow that strongly reflects the forcing pattern. The "perfect" solution in Figure 3a is—with 256<sup>2</sup> grid points—a moderately high-resolution flow and the difference in resolved features with the imperfect model is substantial, which makes this setup an ideal testing ground for the corrective approaches in Table 1. Without explicitly formulating a residual subgrid term, this setup can still be seen





**Figure 4.** Short-term prediction experiments with the imperfect QG equations in a setup similar to Figure 2. The experiments are repeated for 50 different network realizations and training sets. "Accurate days" marks the time steps  $(\Delta t = 1 \text{ day})$  it takes until the error threshold is passed:  $E(\mathbf{x}^k, \mathbf{y}^k) > 0.2$ .

as an SGS model and shares several similarities with the approach in Frezat et al. (2022).

For the QG flow problem we will investigate the performance of the corrected transients, following Equations 10–13, in three different ways. We will make short-term predictions with the methods in Table 1 and compare with the truth using a normalized error, similar to the KS results in Figures 1 and 2. Then we let the different corrective models run into a statistical equilibrium and compare probability density functions (PDFs) of key flow properties with the perfect model equilibrium using the Kullback–Leibler divergence  $D_{KL}$ ; a measure of the similarity between two distributions (explained in more detail below). Finally we perform a coupling experiment that tests the long-term corrective model predictions by using them as initial conditions on the fine grid.

### 4.2. Short-Term Predictions

In Figure 4 we present a short-term prediction experiment using the methods in Table 1. From here on we exclude the standalone DMD and DMD + ESN corrections for their lack of meaningful results, which was also noticed for the KS problem. For the ESN operators we again use  $\rho(W) = 0.4$  and  $\overline{d} = 3$ , but with T = 10,000 days and  $\Delta t = 1$  day we use half the amount of training data. For this problem we find that, after scanning the range (0,1], the optimal relaxation parameter lies around  $\alpha = 0.2$  which roughly corresponds to the

eddy turnover time scale in the model (~73 days). The regularization is increased to  $\lambda = 1 \cdot 10^{-4}$  and the number of accurate days is measured using a stricter tolerance  $E(\mathbf{x}^k, \mathbf{y}^k) < 0.2$ , allowing only a small departure from the true trajectory.

The poor performance of the imperfect QG model shown in Figure 4 is improved by all studied methods. The standalone ESN needs at least  $N_r = 1,600$ , while the other methods show a significant improvement for all chosen  $N_r$ . From Equation 13 and Table 1 we see that ESNc is the optimal (in the least squares sense) hybrid between ESN and the correction-only approach. In the short-term QG predictions we find that these methods coincide for small  $N_r$ . A similar observation can be made for DMDc and the combination ESN + DMDc, which also coincide for low  $N_r$ . Controlled DMDc has better short-term predictive power than the correction-only variant, which is



**Figure 5.** Spinup and long-term transient dynamics indicated by mean kinetic energy  $K_m$ . (a) A 100 years spinup with the perfect QG equations using a time step  $\Delta t = 1$  day is followed by a training period of  $N_T \Delta t = T = 10,000$  days. After the training period, 100 years predictions with imperfect QG, ESN and ESNc are shown, using  $N_r = 3,200$  and  $\lambda = 1$ . (b) Approximations of the probability density functions (PDFs) associated with the equilibrium transients, using 20 bins and excluding spinup/transition periods. Vorticity snapshots at the end of the depicted trajectories are shown in Figures 3 and 6. The ESN results are with single realizations and serve as a demonstration of the corrected dynamics.



also reflected in the behavior of ESN + DMDc and ESNc at low  $N_r$ . With this domain setup we expect DMDc to perform reasonably well on short time scales and it can therefore be viewed as a linear benchmark. The nonlinear ESN + DMDc hybrid improves on it immediately but it takes at least  $N_r = 3,200$  for the other ESN-based methods to outperform DMDc. For large  $N_r$  both hybrid methods (ESN + DMDc and ESNc) almost coincide and any positive influence of the DMD component is negligible.

The  $N_r$  doubling results are reminiscent of the findings with the KS-equation here and in Pathak et al. (2018). Similar to the KS scaling results, increasing  $N_r$  improves the short-term predictions of ESN-based methods for the QG problem. Based on the experiments with the KS-equation we expect that also here a plateau or a maximum will be reached for  $N_r > 12,800$ . For ESN state sizes ranging between 200 and 1,600 the ESN + DMDc combination gives the best results, where ESNc shows a slight deterioration in performance. After  $N_r = 1,600$ , the ESN component begins to dominate the results and ESNc becomes comparable to ESN + DMDc. Note, however, that also the standalone ESN is doing remarkably well for large state sizes. Hence, we again find that the linear components provide a positive offset to the ESN prediction skill at low  $N_r$ . For larger  $N_r$  the differences are less pronounced compared to the KS results, yet still present.

### 4.3. Long-Term Dynamical Regime

For the short-term results in the previous subsection, we used a normalized error based on the full fields  $(\omega, \psi)$  for a comparison of the "hybrid" model results with the (restricted) perfect model truth. Failure in terms of this measure does not imply the predictions are invalid, only that the exact truth is not reproduced. We are therefore also interested in reproducing ergodic properties of long-term time series as in Pathak et al. (2017). In this fashion we will continue here and investigate three flow properties for long-term transient runs: mean kinetic energy  $K_m$ , eddy kinetic energy  $K_e$  and enstrophy Z.

Horizontal velocities u, v follow from the streamfunction  $\psi$ , with  $u = -\partial \psi/\partial y$ ,  $v = \partial \psi/\partial x$ , and are decomposed into a (time) mean and transient component:  $u = \langle u \rangle + u'$ ,  $v = \langle v \rangle + v'$  with the mean  $\langle \cdot \rangle$  taken over a window of 50 days. The quantities  $K_m$ ,  $K_e$ , and Z are then given by

$$K_m = \int_{\Omega} \left( \langle u \rangle^2 + \langle v \rangle^2 \right) d\Omega, \qquad (21)$$

$$K_{e} = \int_{\Omega} \left( \left\langle u^{\prime 2} \right\rangle + \left\langle v^{\prime 2} \right\rangle \right) d\Omega = \int_{\Omega} \left( \left\langle u^{2} \right\rangle - \left\langle u \right\rangle^{2} + \left\langle v^{2} \right\rangle - \left\langle v \right\rangle^{2} \right) d\Omega, \tag{22}$$

$$Z = \int_{\Omega} \omega^2 \ d\Omega, \tag{23}$$

where the integral is approximated with a Riemann sum over the coarse domain  $\Omega^c$ .

A switch from the perfect ( $N_f = 2 \cdot 256^2$ ,  $\text{Re}_f = 2,000$ ) to the imperfect ( $N_c = 2 \cdot 32^2$ ,  $\text{Re}_f = 500$ ) QG model solution will inevitably lead to a different statistical steady state. An example of this process is presented in Figure 5. The perfect QG model is randomly initialized and runs into a statistical equilibrium. Predictions using imperfect QG, a standalone ESN and the hybrid ESNc then start from a restricted perfect QG state and run for 100 years. For stable long-term transients with the ESN-based methods we need a significantly larger regularization parameter ( $\lambda = 1$ ) compared to the short-term experiments. Vorticity snapshots of the perfect and imperfect model depicted in Figure 3 are taken at the end of the trajectories in Figure 5. In Figure 6 we present vorticity snapshots at the end of the ESN and ESNc trajectories. By examining these vorticity snapshots we observe that the imperfect model enters a new regime and loses track of the zonal jet pattern that is present in the perfect QG solution. The two ESN-based methods, however, appear to maintain this structure to some degree after 100 years.

The imperfect model reaches a very different statistical equilibrium after a transition period of approximately 10 years. A corrected transient based on Equations 10–13 should stay closer to the perfect model's dynamical regime and the presented ESN and ESNc trajectories show that this is feasible. Especially the hybrid ESNc shows a significantly better reproduction of the perfect model's  $K_m$  PDF, compared to imperfect model (Figure 5b).

In Figure 7 the average energy spectrum over the final 80 years in Figure 5a is shown. The spectrum provides another demonstration of the improved dynamics given by the standalone ESN and the hybrid ESNc. The imperfect QG solution strongly reflects the forcing, which is also noticeable in the vorticity snapshot (Figure 3b). In



Figure 6. Snapshots of the vorticity fields (in day<sup>-1</sup>) at the end of the transient in Figure 5. (a) Standalone ESN prediction with  $N_r = 3,200$  and  $\lambda = 1$ , (b) hybrid ESNc prediction with  $N_r = 3,200$  and  $\lambda = 1$ .

an incompressible 2D flow we expect energy to be transferred from the stirring wavelength to the larger scales, whereas enstrophy is transferred from the stirring wavelength to the smaller scales and dissipated (Vallis, 2019). Both the energy and the enstrophy transfer are poorly represented in the imperfect model. The ESN-based methods are a lot better at producing the correct energy transfer and achieve a good correspondence for the lowest frequencies. Around the stirring frequency ESNc still performs well, whereas the standalone ESN is overestimating. The enstrophy transfer appears even more difficult to capture correctly but still the hybrid ESNc shows a great improvement over the standalone ESN at these scales.

The transients shown in Figure 5 are specific examples and provide only information for a single realization of the ESN and a single training range. For a more rigorous approach we compute transients for 50 training periods (and hence network realizations). We turn to all models studied in the short-term experiment (Figure 4) and, to maintain a stable iteration, need to increase the regularization parameter  $\lambda$ . For the ESN-based methods we use  $\lambda = 1$ , for correction-only we will use  $\lambda = 5$  and with DMDc we use  $\lambda = 10$  to compute stable evolutions. Later in this section we explore how these methods perform for various other  $\lambda$  choices.



**Figure 7.** Average equilibrium energy spectrum based on the final 80 years of the trajectories in Figure 5a. A dashed line is added to mark the frequency of the forcing. The spectrum is normalized with the largest amplitude in the perfect QG spectrum.





**Figure 8.** Long-term (100 years) transient results for 50 different training intervals.  $D_{KL}$  results from methods that do not depend on an ESN are shown in (a, c, and e), for  $K_m$ ,  $K_e$ , and Z, respectively. In (b, d, and f) the respective scalings with  $N_r$  are depicted for models with an ESN-dependence. Missing values in the plots are caused by unstable configurations.

From the trajectories we compute flow properties  $K_m$ ,  $K_e$ , and Z, as defined in Equations 21–23, and compare their PDFs to the perfect model using their Kullback–Leibler (KL) divergence (Cover & Thomas, 2006): for two discrete distributions P and Q, the divergence of Q from P is given by

$$D_{KL}(P,Q) = \sum_{i} P_{i} \ln\left(\frac{P_{i}}{Q_{i}}\right).$$
(24)

Here, the distribution *P* is obtained from the perfect (reference) model. The PDFs are approximated using a domain that ranges beyond the perfect model's PDF with twice the standard deviation. This domain is divided into 100 bins and every transient is truncated to exclude initial spinup effects. For each flow property the divergence of its PDF from the "truth" is computed and combined into boxplots for different ESN state sizes  $N_r$  (Figure 8). We avoid division by zero in Equation 24 by substituting zero-values with machine precision. This leads to large but finite divergences for non-overlapping distributions (~32). The KL–divergence  $D_{KL}$  is a time-integrated measure that is here only used for evaluation, but is also well-suited for a-posteriori training strategies (Frezat et al., 2022).

The imperfect model shows a poor representation of the variability, which should be expected from the transient example in Figure 5. The PDFs for all flow properties show no resemblance with the true PDF, giving  $D_{KL}$  results that remain at the maximum divergence value. Controlled DMDc and correction-only methods are better at capturing the variability, although this is highly dependent on the stabilizing regularization. Especially for the correction-only approach it is possible to find a configuration such that PDFs give a reasonable correspondence.

The KL-divergences for ESN-based methods in Figure 8 are partly missing. For low  $N_r$ , ESNc and ESN + DMDc are unstable when  $\lambda = 1$ . The remaining results show an overall improvement for increasing ESN state size  $N_r$  (cf. Figure 1), although not very clear for all flow properties. Both mean and eddy kinetic energy KL-divergences are somewhat irregular with optima at moderate  $N_r$  values. For enstrophy, the ESN-based methods gradually improve with ESN state size. From the energy spectrum in Figure 7 (and related results in Frezat et al. (2022)) we know that the enstrophy transfer is difficult to capture and here a similar effect is visible in the correspondence between PDFs. ESNc requires at least  $N_r = 1,600$  to obtain small KL-divergences from the enstrophy PDF, further improving for larger  $N_r$ .

Diverged trajectories show up as non-overlapping with either a maximal KL-divergence or a missing value in the  $D_{KL}$  results. Poor performing methods are hence indistinguishable from unstable ones. Especially the combination ESN + DMDc appears to suffer from stability issues for small  $N_r$ , leading to missing  $D_{KL}$  values. We find that the ESN stabilizes regression-based corrective methods, as already noted in Arcomano et al. (2022). When the regression-based methods run on their own we choose a regularization that stabilizes sufficiently such that divergent trajectories are rare.

To provide an idea of how regularization affects the long-term performance of various methods we perform numerous equilibrium runs for different  $\lambda$ . In Figure 9 we present the results for enstrophy Z. The correction-only approach gives remarkably good results within a narrow optimal region for  $\lambda$ . It is also only slightly enhanced by the combination with an ESN (i.e., ESNc). The hybrid ESNc and ESN + DMDc are, however, much more robust and overall better at reproducing the correct enstrophy variability. From the regularization parameter study it is clear that DMDc needs a stronger regularization than the correction-only approach. The KL-divergences in Figure 8 show a related problem for the models that incorporate an ESN, where the ESN that combines with DMDc needs a much larger state size  $N_r$  to achieve sufficient stabilization. Hence stabilization is achieved through both regularization  $\lambda$  and ESN complexity  $N_r$ .







### 4.4. Symbiotic Modeling

The symbiotic modeling idea proposes a two-way coupling between HR and LR models that benefits both model types. In the experiments above we have only studied a one-way coupling, with data-driven subgrid modeling schemes that make use of HR data on the LR grid. In this section we set out to complete a symbiotic setup by coupling the LR model configurations to the HR reference model and compute HR restart (spinup) trajectories.

We limit our investigation to the LR QG model configurations that have successfully produced long-term predictions. Following the 100-year prediction experiments (Figures 8 and 9), the coarse predictions on  $\Omega^c$  are lifted to the fine grid  $\Omega^f$  using the prolongation operator *P*. From there an ensemble of 40-year restarts are performed with the high-resolution QG version ( $N_f = 2 \cdot 256^2$ , Re<sub>f</sub> = 2,000), using all available (50) coarse predictions. The results are shown in Figure 10.

The QG solution in a doubly periodic domain, with our chosen set of model parameters and wind stress pattern, exhibits multi-stability and hence the flow can organize into different zonal jet patterns under the same forcing conditions. Hence the statistical equilibrium of the HR QG setup that has been used to generate training data (Figure 5) is not unique. In Figure 10a it is clear that the imperfect QG predictions fail at maintaining the correct dynamics. The majority of HR restarts (48 out of 50) from imperfect QG end up in different statistical steady states with markedly higher  $K_m$ . Moreover, the spinup times into the different regimes are significant.

From the restarts using DMDc predictions (Figure 10b), 41 out of 50 trajectories end up in the correct regime after 40 model years. The correction-only approach performs worse, with 34 out of 50 trajectories approximating the desired dynamical range after 40 years. Both these linear methods are able to provide suitable initializations for HR QG, but not in a robust way. Many predictions have greatly diverged from the original training equilibrium. The states that do provide a good initialization yield only a short spinup time for the HR model, indicating the quality of these predictions.

The initializations with ESN-based predictions do show a consistent and reliable return to the training equilibrium. No significant differences in spinup behavior are observed between predictions for this particular choice of ESN state size ( $N_r = 3,200$ ) and regularization ( $\lambda = 8$ ). Throughout the prediction phase the ESN-based methods are able to maintain key flow properties, thereby creating ideal conditions for HR QG restarts with a minimal spinup period. Hence this result shows how LR QG with an ESN-based subgrid model can be used to improve the performance of the HR QG configuration.

Evolving HR QG for one year takes ~3000s (on average, for runs on a single Intel i7 CPU at 1.80 GHz). Taking ESNc as an example, with  $\lambda = 8$  and  $N_r = 3,200$ , it requires ~150s for training and an additional ~20s to compute one year. Switching for one year to the coarse grid using ESNc constitutes a speedup of a factor ~17. For 100 years we reach speedup factors ~140 as training is needed only once.



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**Figure 10.** Density heat maps of high-resolution QG restart trajectories. Restarts are based on the final result of 100-year long predictions using (a) imperfect QG, (b) DMDc ( $\lambda = 10$ ), (c) correction-only ( $\lambda = 5$ ), (d) ESN ( $\lambda = 8$ ,  $N_r = 3,200$ ), (e) ESNc ( $\lambda = 8$ ,  $N_r = 3,200$ ), (f) ESN + DMDc ( $\lambda = 8$ ,  $N_r = 3,200$ ). Each method produces 50 long-term predictions from which perfect QG is restarted. The choice for  $\lambda = 8$  in the ESN-based methods is informed by the results in Figure 9. For the heat maps, the (time,  $K_m$ )-domain is rasterized using 160 × 130 bins. A measure of density is then obtained by counting the number of trajectory points per bin.

### 5. Summary and Discussion

In this paper we have proposed a symbiotic ocean modeling approach, that is, a framework in which models with different complexities are coupled in order to benefit from each other. We distinguish between perfect and imperfect models in terms of differing spatial resolutions and key parameterizations, and mainly focus on how an imperfect model can benefit from a symbiotic setup. With data generated from both model types we seek to correct imperfect model transients, which can be viewed as a subgrid-scale (SGS) modeling effort. To this end, we make use of hybrid modeling techniques that combine linear regression-based methods with nonlinear echo state networks (ESNs). Furthermore, we illustrate how perfect models may benefit from corrected imperfect models in terms of performance, thus demonstrating the symbiotic concept.

Comparisons with similar ML methods show that ESNs perform remarkably well when tasked with predicting complex spatiotemporal dynamics, such as those generated by the Kuramoto–Sivashinsky (KS) equation (Vlachas et al., 2020). With the KS equation as benchmark problem we establish that our physics-controlled ESNc implementation reproduces short-term predictions that are consistent with earlier work in Pathak et al. (2017). Our framework furthermore allows a straightforward comparison with purely regression-based methods. We show how corrections based on linear regression contribute to the success of the hybrid machine learning combinations and serve as a departure point for hybrid methods. When we apply these techniques to a subgrid modeling version of the KS problem, we observe a similar scaling behavior with ESN complexity and departure points rooted in the linear regression techniques. Here it is worth noting that our framework is model-agnostic, making it applicable to a wide range of problems. It therefore generalizes easily from the KS equation to the rotating flow problem (QG) with minimal hyperparameter adjustments.

For the subgrid modeling problem with the QG equations, short-term predictions give results that are comparable to the findings with the KS-equation. A scaling behavior is found with the size of the ESN. When the ESN complexity is negligible, the hybrid methods reduce to their linear regression components. For the long-term flow development, our comparison of statistical steady states shows that the hybrid combinations are robust and perform well for various flow parameters. The parameter study with long-term statistics also shows how the ESN-based methods improve with ESN state size, reminiscent of the short-term full-field reproductions. For our purposes, however, the comparison of long-term flow characteristics is more informative than an error norm on state differences.

The experimental setup with long-term (100-year) transient experiments is an ideal testing ground to study the stability of ML-based SGS models. Our parameter studies with equilibrium simulations show that subgrid



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models based on only an ESN or regression often diverge and are difficult to stabilize. The problem of stabilization is in fact a major issue for ML-based subgrid-scale models. In Guan et al. (2023) a stable CNN-based SGS model is obtained through adding physics constraints, while in Frezat et al. (2022) stability is achieved with a time-integrated loss-function for the training of a CNN. In our ESN-based framework training is limited to a simple linear regression problem and hence we stabilize through regularization of this problem. Moreover, for purely linear correction strategies, the regularization parameter  $\lambda$  is the only available hyperparameter. In our long-term experiments with varying  $\lambda$  we observe that hybrid combinations are more stable. Here it is the ESN that stabilizes a correction-only strategy, which was also mentioned in Arcomano et al. (2022), whereas pure DMD is stabilized by the correction-only controlling term. Another important factor is the role of ESN complexity. Hence we find that, for a single architecture, stabilization is achieved through both regularization  $\lambda$  and ESN state size N<sub>r</sub>. However, other configuration decisions such as the spectral radius  $\rho(W)$  and the fully implicit time-discretization for the physics-based control term were not studied in this context, yet these are both expected to have a dampening effect on the prediction transient. The combinations with a DMD model yield interesting comparisons, especially in the short-term QG experiments. Benefits of adding a DMD model are visible for moderate ESN state sizes. For long-term transient runs the advantage of hybrid DMD-ESN models is less pronounced, which is possibly due to the DMD model being valid for only a short period and hence it should be (partially) rebuilt in an online fashion (Pendergrass et al., 2016).

Obviously, the QG ocean model used here is highly idealized compared to state-of-the-art ocean models. Still, we think that these ideas are applicable to the general problem of correcting large scale flows, that is, improving coarse versions of the flow problem at hand. For models with a higher dimension than studied here a reduced order version of the corrective transient framework, as defined by Equations 10-13, is worth investigating. Here the best choice of reduced coordinates (POD, Fourier, wavelets) in combination with an ESN remains uncertain. Projecting with global POD modes, for instance, greatly reduces the ESN's predictive skill (Vlachas et al., 2020). A localized representation as used in Wan et al. (2021) shows more promise. Another way to tackle high-dimensional problems is through parallelization. A parallel hybrid ESNc based on a local domain decomposition is used in Wikner et al. (2020) and Arcomano et al. (2022). It would be interesting to apply this approach as a subgrid model and reproduce long-term flow characteristics, comparing especially its ability to correctly capture energy and enstrophy transfer at low wave numbers.

The full, two-way coupling between high and low-resolution QG setups constitutes an essential test for the data-driven subgrid models. With a reference equilibrium that is not unique, it is interesting to observe the return to a high resolution steady regime from prolongated low-resolution predictions. It turns out that only the ESN-based methods are able to consistently maintain the zonal flow pattern that is present in the training data. Considering both its computational efficiency and adequate prediction capabilities we therefore conclude that the physics-controlled ESNc is a promising candidate to facilitate our envisioned model symbiosis.

### **Data Availability Statement**

The software developed for this paper is archived at Zenodo (Mulder & Baars, 2023). Included here are configurations for the KS and QG models with initial and boundary conditions as described in the paper. The experimental framework that handles transient computations, ESN/DMD configurations and error metrics is available here as well.

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