

# Evolving Gaussian Processes and Kernel Observers for Learning and Control in Spatiotemporally Varying Domains

with applications in agriculture, weather monitoring, and fluid  
dynamics

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## Introduction

Monitoring and modeling of large-scale stochastic phenomena with both spatial and temporal (spatiotemporal) evolution using a network of distributed sensors is a fundamental problem in many applications. Consider, for example, a team of mechanical weeding robots managing herbicide-resistant weeds on a farm (see Figure 1, also see “How can control engineers help agriculture?”). This team of robots needs to predict the weed growth across the whole farm to make intelligent decisions on robot coordination [1]. However, the robots can only observe a limited part of the field at any given time. This leads to the critical problem: How can a few robots that can only partially observe the field at any given time predict the full state of the spatiotemporally evolving weed growth over the entire field? The goal of this work is to take a step towards addressing this challenging problem. There are many examples in other fields, including modeling and monitoring ocean heat content and acidification in oceanography [2] using a network of satellites and surface sensors; prediction of traffic patterns using data from vehicles, cellphones, and traffic cameras; prediction of enemy movements through ground and aerial surveillance, and prediction of spatiotemporal evolution of extreme weather events [3] using data from weather stations and aerial drones. Indeed, with rapid advances in robotics and the capability to process large volumes of data on compact computational packages, the applications involving such distributed cyber physical systems are rapidly expanding.

allows one to solve very tough problems. SO its not the feature space that's the problem,

These types of problems are characterized by the need to estimate complex and stochastic dynamics that are distributed in space and time. The key constraint is that there are only a

few spatially distributed sensors available, and together, these few sensors are not enough to entirely cover the whole space at any given time. In the face of this constraint, one approach to solve the problem is to use a predictive model of the phenomena and use it to inform sensing strategy. While modeling such spatiotemporal phenomena has traditionally been in the province 5 of the field of geostatistics, it has in recent years gained more attention in the machine learning community [4]. The data-driven models developed through machine learning techniques provide a way to capture complex spatiotemporal phenomena which are not easily modeled by first-principles alone. However, machine learning models are limited by the data sets they are trained on, and the challenge with complex and distributed physical systems is that they have high 10 variability. For example, a model trained on weed growth over years of data in one field doesn't necessarily generalize to another, and a model trained on past few year's worth of data is still unreliable to predict weather variability in the following year. What is needed instead of just a *predict* system is a *predict-and-correct* system; we elucidate this point below in more detail.

In the machine learning community, kernel methods represent a class of extremely well- 15 studied and powerful methods for regression in spatial domains. In these techniques, correlations between the input variables are encoded through covariance kernels, and the model is formed through a linear weighted combination of the kernels [5]–[7]. In recent years, kernel methods have been applied to spatiotemporal regression problems with varying degrees of success [4], [5]. Many recent techniques in spatiotemporal modeling have focused on nonstationary covariance 20 kernel design and associated hyperparameter learning algorithms [8]–[10]. These methods, which focus on the careful design of covariance kernels, have been proposed as an alternative to the naive approach of simply including time as an additional input variable in the kernel [11]. The careful design/optimization of covariance kernel avoids an explosion in the number of parameters (kernels utilized) of the model which would be inevitable in a model that simply adds time as 25 an additional input variable, and has been shown to better account for spatiotemporal couplings.

However, there are two key challenges with existing kernel-based and other machine learning approaches for prediction over complex spatiotemporally-varying physical systems: The first is ensuring the scalability of the model to large scale phenomena, which manifests due to the fact that the problem of optimizing the covariance kernel (known as hyperparameter 30 optimization in the ML community) is not convex in general, leading to methods that are difficult to implement especially in online settings, susceptible to getting stuck at local minima, and computationally demanding for large datasets [12]. This has also been a key issue in utilizing deep learning models for this problem. The second key challenge is that no matter how much data the model is trained on, it cannot completely capture the variability in real world complex 35 systems with spatiotemporal dynamics. This is a problem that the controls community is quite aware of. Our solutions to this problem build upon feedback, leading to fundamental notions of

observability and controllability which can be used to build robust state estimators and controllers. To bring these notions to fruition in the spatiotemporal prediction problem, what is needed is a way to inform where sensing/control should be done to ensure that the state estimation problem can be made observable/controllable. It is however not clear how notions of observability  
5 and controllability could be utilized with existing kernel-based machine learning models, and how observers and controllers can be embedded within such models. While the computational challenge can be addressed with increasing computational power and large datasets, addressing the latter (and vastly more fundamental) challenge in designing robust observers/controllers is particularly important in the design of reliable engineering systems, such as distributed  
10 sensor/actuator networks intended for monitoring physical phenomena, autonomous soft-robots, or other physical systems with distributed sensing and actuation.

Within the computational flow dynamics (CFD) community, there has recently been considerable interest in methods inspired by Koopman operator theory. A Koopman operator is a linear but infinite-dimensional operator that is defined for an autonomous dynamical system  
15 and governs the evolution of its *observables* [13], rather than states. An observable is a function from the state space to a measurement space; the Koopman operator is in essence a composition operator on an observable and the state-transition operator, returning a new function which takes a state and gives a prediction of future measurements. By performing a spectral analysis of this linear operator, one reveals modes which show the spatial distribution, oscillation frequency, and  
20 growth rate/decay of the component dynamics of the system. Many advantages can be realized from this: the ability to transform the state space so the dynamics appear linear, to predict the temporal evolution of the linear system, to reconstruct the state of the original nonlinear system, and even to implement controller design. Dynamic Mode Decomposition (DMD) is the most widely used method for finding a finite-dimensional subspace of the Koopman operator's  
25 infinite-dimensional domain to work in. Williams et al., recently integrated DMD with the kernel trick, allowing the algorithm to be extended to systems with much larger dimensions [14]. Brunton et al., inspired by DMD, were able to generate governing equations from data by sparse identification of nonlinear dynamical systems [15]. However, these methods are restricted to approximating the Koopman operator given a fixed vector-valued observable, and have no way  
30 of effectively using measurements that vary in both size and location over time. Furthermore, the state of research into data-driven generalizing over similar systems with varying parameters is at best preliminary.

## Contributions

In this article, we present a new perspective on solving the spatiotemporal monitoring  
35 problem that brings together kernel-based modeling, systems theory, and Bayesian filtering.

We define the monitoring problem as follows: *Given an approximate predictive model of the spatiotemporal phenomena learned using historic data, estimate the current latent state of the phenomena in the presence of uncertainty using as few sensors as possible.* Ideally, the solution to the problem should also provide guidance on how many sensors are needed and where to place them. In this paper we argue that when it comes to predictive inference over spatiotemporal phenomena, a Kalman-filter type approach of predicting and correcting with feedback from a set of minimal sensors is a robust way of dealing with real-world uncertainties and inherent modeling errors. In the context of this specific problem, *our main contributions are two-fold*: first, we demonstrate that spatiotemporal functional evolution can be modeled using stationary kernels with a linear dynamical systems layer on their mixing weights. In particular, in contrast with existing work, this approach does not necessarily require the design of complex spatiotemporal kernels, and can accommodate positive-definite kernels on any domain on which it is possible to define them, which includes non-Euclidean domains such as Riemannian manifolds, strings, graphs and images [16]. Second, we show that such a model can be utilized to determine sensing locations that guarantee that the hidden states of functional evolution can be estimated using a Bayesian state-estimator (Kalman filter) that is *embedded in the feature space of the kernel model* with very few sensors. A benefit of our solution approach is that it provides guidance on how many sensors are needed and where to place them. Accordingly, we provide sufficient conditions on the number and location of sensor measurements required and prove non-conservative lower bounds on the minimum number of sampling locations by developing fundamental results on the observability of kernel-based models. **Thirdly, we show that this model can be utilized to generate paths for sensing agents to follow, that allows the agents to converge to a correct estimate of the system state as quickly as possible.** The validity of the presented model and sensing techniques is corroborated using synthetic and large real datasets.

25 *Broader Context*

The fundamental idea of building observers and controllers embedded in the feature spaces of machine learning models introduced in this paper is generalizable beyond the particular application of spatiotemporal monitoring. Indeed, the contributions of this paper demonstrate how machine learning theory can be combined with systems theory to address major challenges in distributed monitoring and control of complex spatiotemporally varying systems. We elucidate this point below.

Traditionally, the controls literature is strongest when the system dynamics can be represented as Ordinary Differential Equations (ODEs). As depicted in Figure 2, some of the major successes of controls have included results such as Linear Quadratic Gaussian control, reinforcement learning, and adaptive control in state spaces with well-defined, finite, and physi-

cally meaningful state variables. The problem of state estimation of a temporally evolving, finite-dimensional state-space system for example has been extensively studied in the context of Kalman filtering and observer design [17]. Here, fundamental results in observability/controllability provide sufficient conditions on the structure of the state transition and measurement matrix  
5 such that the latent state can be estimated in the presence of measurement and process noise. On the other hand, machine learning models operate in an abstract and learned function-space of *features* (for more details see “Feature Spaces in Machine Learning”). Kalman filters can be naively extended to the functional domain ([18]), but have not typically been studied in context of the spatiotemporal monitoring problem studied here.

10 On the other hand, recent advances in machine learning are providing different and highly powerful ways of modeling complex spatiotemporal dynamical systems. Yet, the main challenge with using machine learning approaches such as deep learning or kernel based methods has been that these approaches lack interpretability and analyzability, which makes it difficult to design robust engineered systems. This is because machine learning works in abstract feature  
15 spaces that are only relatable to physical quantities through complex functional operations (see “Feature Spaces in Machine Learning”). This leads to a major challenge in using machine learning models in control: For example, when kernel based models are used for spatiotemporal systems, how does one answer fundamental questions such as 1) the least number of sensors required to observe a distributed system, 2) the placement of sensors/actuators to guarantee  
20 observability/controllability of the system, and 3) the effect of random sensor placement on system observability/controllability.

In this paper we present an approach that can provide one formal way of addressing these and other questions about complex systems that are modeled with machine learning. In particular, we demonstrate how linear dynamical systems can be embedded in the reproducing kernel Hilbert  
25 space (RKHS) [6], [19], [20] generated by features used in Gaussian Process modeling [21], [22], and utilized to answer fundamental questions such as controllability and observability. The marriage of systems theory with machine learning pursued in this paper is exciting, and we expect that follow-up work will exploit the framework presented in this paper of utilizing linear models in RKHSs and feature spaces of other machine learning models to enable practical and  
30 analyzable data-driven engineering systems. To facilitate the development of the theory, we have focused this paper on the problem of monitoring spatiotemporal phenomena. However, the idea should be generalizable to any distributed cyber-physical system that is changing with space and time.

## Outline of the article and relationship to prior work by the authors

We begin the rest of the article by summarizing some related work in machine learning in this area. “Feature Spaces in Machine Learning” discusses the concept of feature spaces and its importance in machine learning in a broader context. We then formulate the problem, 5 introduce *Kernel Observers (KO)*, and develop the main theoretical and algorithmic results. This is followed by some results on the expected number of randomly placed sensors required to monitor a spatiotemporal process in the context of our model. An extension to the KO method called *Evolving Gaussian Processes (E-GP)* is presented that learns one model for multiple, similar spatiotemporal processes, the efficacy of which on real-world CFD data is presented 10 in Sidebar *Learning Fluid Flows with Evolving Gaussian Processes*. **Lastly, a method for separating invariant subspaces within our model and using them to generate paths for sensing agents is presented. Elements of the work presented in this paper first appeared in Neural Information Processing Systems (NIPS 2016) ([23], [24]), IEEE CDC 2015 conference [25], the Conference on Robot Learning (CoRL 2017) [26] and IEEE ACC 2018 conference [27].** This paper presents a comprehensive set of results and fills in the missing 15 links in a single encompassing publication, and introduces new results on observability in the presence of random sensor placement, connection to Koopman operators, and generating paths for single observing agents. As such, we have focused in this article mostly on the fundamental theory and practical algorithms for modeling, estimation, and control, 20 while the excruciating details of how to optimally implement the presented algorithms are omitted. Instead an open-source code-base is made available in MATLAB at <http://daslab.illinois.edu/software.html> or at <https://github.com/hkingravi/FunctionObservers> and in Python on GitHub at <https://github.com/hkingravi/funcobspy>.

## Related Work

25 **There is a very large amount of literature on Gaussian Processes and spatiotemporal modeling, a complete survey of this literature is beyond the scope of this paper. Since our contributions are in the area of creating a feedback based observer in the feature spaces of GP models, we discuss here related work in three related areas: Spatiotemporal modeling with GPs, Kalman filtering and GP connection, and sensor placement for inference in 30 spatiotemporal domains.**

There is a large body of literature on spatiotemporal modeling in geostatistics where specific process-dependent kernels can be used [4], [28], [29]. **Other approaches that utilize hierarchy or evolution of kernels have also been used for modeling spatiotemporal functions [30]–[32].** From the machine learning perspective, a naive approach is to utilize both spatial

and temporal variables as inputs to a Mercer kernel [33]. However, this technique leads to an ever-growing kernel dictionary. Furthermore, constraining the dictionary size or utilizing a moving window will occlude learning of long-term patterns. **A more clever approach is to use state-space representations of time-varying GPs [30], [34]. In this view point, each GP instance is viewed as a snapshot of an evolving set of weights. We follow in a similar vein here, with added emphasis on understanding the mathematical structures leading to observability and controllability.** Periodic or nonstationary covariance functions and nonlinear transformations have been proposed to address this issue [5], [9]. Work focusing on nonseparable and nonstationary covariance kernels seeks to design kernels optimized for environment-specific dynamics, and to tune their hyperparameters in local regions of the input space. Seminal work in [35] proposes a process convolution approach for space-time modeling. This model captures nonstationary structure by allowing the convolution kernel to vary across the input space. This approach can be extended to a class of nonstationary covariance functions, thereby allowing the use of a Gaussian process (GP) framework, as shown in [36]. However, since this model's hyperparameters are inferred using MCMC integration, its application has been limited to smaller datasets. To overcome this limitation, [10] proposes to use the mean estimates of a second isotropic GP (defined over latent length scales) to parameterize the nonstationary covariances. Finally, [8] considers non-isotropic variation across different dimension of input space for the second GP as opposed to isotropic variation by [10]. Issues with this line of approach include the nonconvexity of the hyperparameter optimization problem and the fact that selection of an appropriate nonstationary covariance function for the task at hand is a nontrivial design decision (as noted in [37]).

Apart from directly modeling the covariance function using additional latent GPs, there exist several other approaches for specifying nonstationary GP models. One approach maps the nonstationary spatial process into a latent space, in which the problem becomes approximately stationary [38]. Along similar lines, [39] extends the input space by adding latent variables, which allows the model to capture nonstationarity in original space. Both these approaches require MCMC sampling for inference, and as such are subject to the limitations mentioned in the preceding paragraph.

A geostatistics approach that finds dynamical transition models on the linear combination of weights of a parameterized model [4], [18] is advantageous when the spatial and temporal dynamics are hierarchically separated, leading to a convex learning problem. **This approach has been utilized previously in MRI imaging [40], [41].** As a result complex nonstationary kernels are often not necessary (although they can be accommodated). The approach presented in this paper aligns closely with this vein of work. A systems-theoretic study of this viewpoint enables the fundamental contributions of the paper, which are 1) allowing for inference on more

general domains with a larger class of basis functions than those typically considered in the geostatistics community, and 2) quantifying the minimum number of measurements required to estimate the state of the system.

5 **Kalman filtering in the context of Gaussian processes and Kernel models has also been quite widely studied [29], [42]–[45]. There is clearly a direct link between the Bayesian approach to inference taken in GPs and its natural extension to Kalman Filters. Our contributions here are in creating explicit connections between feedback observers and inference by deriving conditions of observability in the kernel space. This leads to explicit conditions on the number of sensors required and where to place them that we present in**  
10 **this paper.**

Lastly, sensor placement optimization is also a well-studied area. Examples include, but are not limited to 1) geometric approaches, which seek to provide a covering of the operating space without making assumptions about the spatiotemporal dynamics [46], and 2) information-theoretic approaches, which place their focus on sensor placement optimizing strategies based  
15 on mutual information and information entropy for Gaussian process models [47]. It should be noted that the contribution of the paper concerning sensor placement is to provide *sufficient conditions* for monitoring rather than optimization of the placement locations, and therefore a comparison with these approaches is not considered in the experiments.

## Kernel Observers

20 This section outlines our modeling framework and presents theoretical results associated with the number of sampling locations required for monitoring functional evolution.

### Problem Formulation

We focus on predictive inference of a time-varying stochastic process, whose mean  $f$  evolves temporally as  $f_{\tau+1} \sim \mathbb{F}(f_\tau, \eta_\tau)$ , where  $\mathbb{F}$  is a distribution varying with time  $\tau$  and  
25 exogenous inputs  $\eta$ . Our approach builds on the fact that in several cases, temporal evolution can be hierarchically separated from spatial functional evolution. A classical and quite general example of this is the *abstract evolution equation* (AEO), which can be defined as the evolution of a function  $u$  embedded in a Banach space  $\mathcal{B}$ :  $\dot{u}(t) = \mathcal{L}u(t)$ , subject to  $u(0) = u_0$ , and  $\mathcal{L} : \mathcal{B} \rightarrow \mathcal{B}$  determines spatiotemporal transitions of  $u \in \mathcal{B}$  [48]. This model of spatiotemporal  
30 evolution is very general (AEOs, for example, model many PDEs), but working in Banach spaces can be computationally taxing. A simple way to make the approach computationally feasible is to place restrictions on  $\mathcal{B}$ : in particular, we restrict the sequence  $f_\tau$  to lie in a RKHS, the theory of which provides powerful tools for generating flexible classes of functions with relative ease

[5]. In a kernel-based model,  $k : \Omega \times \Omega \rightarrow \mathbb{R}$  is a positive-definite Mercer kernel on a domain  $\Omega$  that models the covariance between any two points in the input space, and implies the existence of a smooth map  $\psi : \Omega \rightarrow \mathcal{H}$ , where  $\mathcal{H}$  is an RKHS with the property  $k(x, y) = \langle \psi(x), \psi(y) \rangle_{\mathcal{H}}$ . The key insight behind the proposed model is that spatiotemporal evolution in the input domain corresponds to temporal evolution of the mixing weights of a kernel model alone in the functional domain. Therefore,  $f_{\tau}$  can be modeled by tracing the evolution of its mean embedded in a RKHS using switched ordinary differential equations (ODE) when the evolution is continuous, or switched difference equations when it is discrete (Figure 3). The advantage of this approach is that it allows us to utilize powerful ideas from systems theory for deriving necessary and sufficient conditions for spatiotemporal monitoring.

In this paper, we restrict our attention to the class of functional evolutions  $\mathbb{F}$  defined by linear Markovian transitions in an RKHS. While extension to the nonlinear case is possible (and non-trivial), it is not pursued in this paper to help ease the exposition of the key ideas. The class of linear transitions in RKHS is rich enough to approximately model many real-world datasets, as suggested by our experiments.

Let  $y \in \mathbb{R}^N$  be the measurements of the function available from  $N$  sensors,  $\mathcal{A} : \mathcal{H} \rightarrow \mathcal{H}$  be a linear transition operator in the RKHS  $\mathcal{H}$ , and  $\mathcal{K} : \mathcal{H} \rightarrow \mathbb{R}^N$  be a linear measurement operator. The model for the functional evolution and measurement studied in this paper is:

$$f_{\tau+1} = \mathcal{A}f_{\tau} + \eta_{\tau}, \quad y_{\tau} = \mathcal{K}_{\tau}f_{\tau} + \zeta_{\tau}, \quad (1)$$

where  $\eta_{\tau}$  is a zero-mean stochastic process in  $\mathcal{H}$ , and  $\zeta_{\tau}$  is a Wiener process in  $\mathbb{R}^N$ . Classical treatments of kernel methods emphasize that for most kernels, the feature map  $\psi$  is unknown, and possibly infinite-dimensional; this forces practitioners to work in the dual space of  $\mathcal{H}$ , whose dimensionality is the number of samples in the dataset being modeled. This conventional wisdom precludes the use of kernel methods for most tasks involving modern datasets, which may have millions and sometimes billions of samples [49]. An alternative is to work with a feature map  $\widehat{\psi}(x) := [\widehat{\psi}_1(x) \dots \widehat{\psi}_M(x)]^T$  to an approximate feature space  $\widehat{\mathcal{H}}$ , with the property that for every element  $f \in \mathcal{H}$ ,  $\exists \widehat{f} \in \widehat{\mathcal{H}}$  and an  $\epsilon > 0$  s.t.  $\|f - \widehat{f}\| < \epsilon$  for an appropriate function norm. A few such approximations are listed below.

*Dictionary of atoms:* Let  $\Omega$  be compact. Given points  $\mathcal{C} = \{c_1, \dots, c_M\}$ ,  $c_i \in \Omega$ , we have a dictionary of atoms  $\mathcal{F}^{\mathcal{C}} = \{\psi(c_1), \dots, \psi(c_M)\}$ ,  $\psi(c_i) \in \mathcal{H}$ , the span of which is a strict subspace  $\widehat{\mathcal{H}}$  of the RKHS  $\mathcal{H}$  generated by the kernel. Here,

$$\widehat{\psi}_i(x) := \langle \psi(x), \psi(c_i) \rangle_{\mathcal{H}} = k(x, c_i). \quad (2)$$

*Low-rank approximations:* Let  $\Omega$  be compact, let  $\mathcal{C} = \{c_1, \dots, c_M\}$ ,  $c_i \in \Omega$ , and let  $K \in \mathbb{R}^{M \times M}$ ,  $K_{ij} := k(c_i, c_j)$  be the Gram matrix computed from  $\mathcal{C}$ . This matrix can be diag-

nalized to compute approximations  $(\widehat{\lambda}_i, \widehat{\phi}_i(x))$  of the eigenvalues and eigenfunctions  $(\lambda_i, \phi_i(x))$  of the kernel [50]. These spectral quantities can then be used to compute  $\widehat{\psi}_i(x) := \sqrt{\widehat{\lambda}_i} \widehat{\phi}_i(x)$ .

*Random Fourier features:* Let  $\Omega \subset \mathbb{R}^n$  be compact, and let  $k(x, y) = e^{-\|x-y\|^2/2\sigma^2}$  be the Gaussian RBF kernel. Then random Fourier features approximate the kernel feature map as  
 $\widehat{\psi}_\omega : \Omega \rightarrow \widehat{\mathcal{H}}$ , where  $\omega$  is a sample from the Fourier transform of  $k(x, y)$ , with the property that  
 $k(x, y) = \mathbb{E}_\omega[\langle \widehat{\psi}_\omega(x), \widehat{\psi}_\omega(y) \rangle_{\widehat{\mathcal{H}}}]$  [49]. In this case, if  $V \in \mathbb{R}^{M/2 \times n}$  is a random matrix representing the sample  $\omega$ , then  $\widehat{\psi}_i(x) := [\frac{1}{\sqrt{M}} \sin([Vx]_i), \frac{1}{\sqrt{M}} \cos([Vx]_i)]$ . Similar approximations exist for other radially symmetric kernels, as well as dot-product kernels.

In the approximate space case, we replace the transition operator  $\mathcal{A} : \mathcal{H} \rightarrow \mathcal{H}$  in (1) by  $\widehat{\mathcal{A}} : \widehat{\mathcal{H}} \rightarrow \widehat{\mathcal{H}}$ . This approximate regime, which combines the flexibility of a truly nonparametric approach with computational realizability, still allows for the representation of rich phenomena, as will be seen in the sequel, and in Figure 5. The finite-dimensional evolution equations approximating (1) in dual form are

$$w_{\tau+1} = \widehat{A}w_\tau + \eta_\tau, \quad y_\tau = Kw_\tau + \zeta_\tau, \quad (3)$$

where we have matrices  $\widehat{A} \in \mathbb{R}^{M \times M}$ ,  $K \in \mathbb{R}^{N \times M}$ , the vectors  $w_\tau \in \mathbb{R}^M$ , and where we have slightly abused notation to let  $y_\tau, \eta_\tau$  and  $\zeta_\tau$  denote their  $\widehat{\mathcal{H}}$  counterparts. Here  $K$  is the matrix whose rows are of the form  $K_{(i)} = \widehat{\Psi}(x_i) = [\widehat{\psi}_1(x_i) \ \widehat{\psi}_2(x_i) \ \dots \ \widehat{\psi}_M(x_i)]$ . In systems-theoretic language, each row of  $K$  corresponds to a *measurement* at a particular location, and the matrix itself acts as a measurement operator.  
 $\tau$

The equations in (1) suggest an immediate extension to functional control problems. Pick another basis for  $\mathcal{H}$  as  $\tilde{\psi}(x) := [\tilde{\psi}_1(x) \ \dots \ \tilde{\psi}_{\ell'}(x)]^T$ , where the functions  $\tilde{\psi}_j(x)$  are used to approximate the RKHS  $\mathcal{H}$  generated by the kernel. We denote the span of these functions as  $\widetilde{\mathcal{H}}$ . In the dictionary of atoms case, an example would be another set of atoms  $\mathcal{F}_D = [\psi(d_1) \ \dots \ \psi(d_{\ell'})]$ ,  $\psi(d_j) \in \mathcal{H}$ ,  $d_j \in \Omega$ , with  $\widetilde{\mathcal{H}}$  being a strict subspace of the RKHS  $\mathcal{H}$  generated by the kernel. The functional evolution equation is then as follows:

$$f_{\tau+1} = \mathcal{A}f_\tau + \mathcal{B}\delta_\tau + \eta_\tau, \quad y_\tau = \mathcal{K}_\tau f_\tau + \zeta_\tau, \quad (4)$$

where the control functions  $\delta_\tau$  evolve in  $\widetilde{\mathcal{H}}$ , and  $\mathcal{B} : \widetilde{\mathcal{H}} \rightarrow \widehat{\mathcal{H}}$ . To derive the finite-dimensional equivalent of  $\mathcal{B}$ , we have to work out the structure of the matrix  $B$ : since  $\widehat{\mathcal{H}}$  is not, in general, isomorphic to  $\widetilde{\mathcal{H}}$ , this imposes strict restrictions on  $B$ . We can derive  $B$  using least squares using the inner product of  $\mathcal{H}$ . An instructive example is where both  $\widehat{\mathcal{H}}$  and  $\widetilde{\mathcal{H}}$  are generated by dictionaries of atoms; recall that in this case,  $\mathcal{F}^C = [\psi(c_1) \ \dots \ \psi(c_M)]$  is the basis for  $\widehat{\mathcal{H}}$ ,

and let  $\delta = \sum_{j=1}^{\ell'} \hat{w}_j \psi(d_j)$ , and let  $\mathcal{F}^C = [\psi(c_1) \ \cdots \ \psi(c_M)]$  be the basis for  $\mathcal{H}^C$ . Then the projection of  $\delta$  onto  $\widehat{\mathcal{H}}$  can be derived as

$$\begin{bmatrix} \langle \delta, \psi(c_1) \rangle_{\mathcal{H}} \\ \vdots \\ \langle \delta, \psi(c_M) \rangle_{\mathcal{H}} \end{bmatrix} = \underbrace{\begin{bmatrix} \langle \psi(d_1), \psi(c_1) \rangle_{\mathcal{H}} & \cdots & \langle \psi(d_{\ell'}), \psi(c_1) \rangle_{\mathcal{H}} \\ \vdots & \ddots & \vdots \\ \langle \psi(d_1), \psi(c_M) \rangle_{\mathcal{H}} & \cdots & \langle \psi(d_{\ell'}), \psi(c_M) \rangle_{\mathcal{H}} \end{bmatrix}}_{K_{CD}} \begin{bmatrix} \hat{w}_1 \\ \vdots \\ \hat{w}_{\ell'} \end{bmatrix}. \quad (5)$$

Note that in the dictionary of atoms case, the entries of  $K_{CD}$  can be computed in closed form as  $K_{CDij} := k(d_i, c_j)$ , using the reproducing property. This derivation shows that the operator  $B$  is simply  $K_{CD} \in \mathbb{R}^{M \times \ell'}$ , the kernel matrix between the data  $C$  generating the atoms  $\mathcal{F}^C$  of  $\widehat{\mathcal{H}}$  and the data  $D$  generating the atoms  $\mathcal{F}_D$  of  $\widetilde{\mathcal{H}}$ . Thus, the finite-dimensional evolution equations equivalent to (4) are

$$w_{\tau+1} = \widehat{A}w_{\tau} + K_{CD}\hat{w}_{\tau}, \quad y_{\tau} = K_{\tau}w_{\tau}. \quad (6)$$

We define the *generalized observability matrix* [51] as  $\mathcal{O}_{\Upsilon} = \begin{bmatrix} K\widehat{A}^{\tau_1} \\ \vdots \\ K\widehat{A}^{\tau_L} \end{bmatrix}$  where  $\Upsilon = \{\tau_1, \dots, \tau_L\}$  are the set of instances  $\tau_i$  when we apply the operator  $K$ . A linear system is said to be *observable* if  $\mathcal{O}_{\Upsilon}$  has full column rank (i.e.  $\text{Rank}(\mathcal{O}_{\Upsilon}) = M$ ) for  $\Upsilon = \{0, 1, \dots, M-1\}$  [51]. Observability guarantees two critical facts: firstly, it guarantees that the state  $w_0$  can be recovered exactly from a finite series of measurements  $\{y_{\tau_1}, y_{\tau_2}, \dots, y_{\tau_L}\}$ ; in particular, defining  $y_{\Upsilon} = \begin{bmatrix} y_{\tau_1}^T, y_{\tau_2}^T, \dots, y_{\tau_L}^T \end{bmatrix}^T$ , we have that  $y_{\Upsilon} = \mathcal{O}_{\Upsilon}w_0$ . Secondly, it guarantees that a feedback based *observer* can be designed such that the estimate of  $w_{\tau}$ , denoted by  $\hat{w}_{\tau}$ , converges exponentially fast to  $w_{\tau}$  in the limit of samples. Note that all our theoretical results assume  $\widehat{A}$  is available: while we perform system identification in the experiments ([23]), it is not the focus of the paper.

We are now in a position to formally state the spatiotemporal modeling, control, and inference problems being considered: given a spatiotemporally evolving system modeled using (3), choose a set of  $N$  sensing locations such that even with  $N \ll M$ , the functional evolution of the spatiotemporal model can be estimated (which corresponds to *monitoring*), can be predicted robustly (which corresponds to *Bayesian filtering*), and which can be controlled (which corresponds to *functional control*). Our approach to solve the monitoring and prediction problem relies on the design of the measurement operator  $K$  so that the pair  $(K, \widehat{A})$  is observable: any Bayesian state estimator (e.g. a Kalman filter) utilizing this pair is denoted as a **kernel observer**. In the case where no measurements are taken, for the sake of consistency, we denote the state estimator as an *autonomous* kernel observer. In the controls case, given a spatiotemporally evolving system modeled using (6), we need to choose a set of  $N$  sensing locations and  $\ell'$  control locations, such that even with  $N \ll M$ ,  $\ell' \ll M$ , the functional evolution of the spatiotemporal model can be controlled; in this case, we must design both a measurement operator  $K$  and a

control operator  $K_{CD}$  such that the pair  $(K_{CD}, \widehat{A})$  is controllable: a controls system utilizing this pair and the measurement operator  $K$  is denoted as a **kernel controller**.

## Preliminaries on Rational Canonical Structures

We take a geometric approach towards the choice of sampling locations for inferring  $w_\tau$  in 5 (3); the extension for control is similar. We use the notation  $\mathcal{V}$ , with  $\dim(\mathcal{V}) = M$ , to emphasize the fact that these theorems hold for any finite-dimensional vector space. Consider the linear operator  $\mathcal{A} : \mathcal{V} \rightarrow \mathcal{V}$ , and recall that the definition of observability requires the construction of a linear operator  $\mathcal{K} : \mathcal{V} \rightarrow \mathcal{U}$ , with  $\dim(\mathcal{U}) = N$ , such that  $\text{rank} [\ (\mathcal{K})^T \dots (\mathcal{K}\mathcal{A}^{M-1})^T ]^T = M$ . In most applications, if  $N \geq M$ , and  $\text{rank } \mathcal{K} = N$ , it is reasonable to expect that observability 10 may be achieved. However, for our purposes,  $N$  must be *significantly* less than  $M$ . Therefore, we must design  $\mathcal{K}$  with as small a rank as possible. To do so, we require a series of vectors  $v_i$  that, under repeated iterations of  $\mathcal{A}$ , can generate a basis for  $\mathcal{V}$ . For this task, we will use a fundamental decomposition result from the theory of modules, known as the *rational canonical structure* of  $\mathcal{A}$  [52]. The intuition here is that if the sequence  $\{v_i\}_i$  can generate this basis, it 15 can be directly used to construct  $\mathcal{K}$ .

The linear operator  $\mathcal{A} : \mathcal{V} \rightarrow \mathcal{V}$  has a characteristic polynomial  $\pi(\lambda)$  such that  $\pi(\mathcal{A}) = 0$  by the Cayley-Hamilton theorem. The minimal polynomial (MP) of  $\mathcal{A}$  is the monic polynomial  $\alpha(\cdot)$  of least degree (denoted by  $\deg(\cdot)$ ) given as  $\alpha(\lambda) = a_0 + a_1\lambda + \dots + \lambda^{\deg(\alpha)} = 0$ , such 20 that  $\alpha(\mathcal{A}) = a_0I + a_1\mathcal{A} + \dots + \mathcal{A}^{\deg(\alpha)} = 0$ . The MP is unique and divides  $\pi(\lambda)$ , so that  $\deg(\alpha) \leq \deg(\pi)$ . The MP of a vector  $v \in \mathcal{V}$  relative to  $\mathcal{A}$  is the unique monic polynomial  $\xi_v$  of least degree such that  $\xi_v(\mathcal{A})v = a_0v + a_1\mathcal{A}v + \dots + \mathcal{A}^{\deg(\alpha)}v = 0$ . If  $\deg(\alpha) = M$ , then  $\mathcal{A}$  is *cyclic* and  $\exists v \in \mathcal{V}$ , such that the vectors  $\{v, \mathcal{A}v, \dots, \mathcal{A}^{M-1}v\}$  form a basis for  $\mathcal{V}$ ; this is the same 25 as saying that the pair  $(v^T, \mathcal{A}^T)$  is observable. A subspace  $\mathcal{V}_S \subset \mathcal{V}$  s.t.  $\mathcal{A}\mathcal{V}_S \subset \mathcal{V}_S$  is  $\mathcal{A}$ -*cyclic* if  $\mathcal{A}|_{\mathcal{V}_S}$ , the restriction of  $\mathcal{A}$  to the subspace  $\mathcal{V}_S$ , is cyclic. If  $\alpha(\lambda)$  is the minimal polynomial 30 of  $\mathcal{A}$  and  $\deg(\alpha) = m < M$ ,  $\exists v \in \mathcal{V}$  such that  $\{v, \mathcal{A}v, \dots, \mathcal{A}^{m-1}v\}$  span an  $m$ -dimensional  $\mathcal{A}$ -cyclic subspace  $\mathcal{V}_S$ , with  $v$  being the *cyclic generator* of  $\mathcal{V}_S$ . The subspace  $\mathcal{V}_S$  decomposes  $\mathcal{V}$  relative to  $\mathcal{A}$ . By the rational canonical structure theorem (Theorem 0.1 of [52]),  $\mathcal{A}$  can be successively decomposed into subspaces  $\mathcal{V}_i \subset \mathcal{V}$ ,  $i \in \{1, \dots, \ell\}$ , s.t.  $\mathcal{V} = \mathcal{V}_1 \oplus \dots \oplus \mathcal{V}_\ell$ ,  $\mathcal{A}\mathcal{V}_i \subset \mathcal{V}_i$ , and  $\mathcal{A}|_{\mathcal{V}_i}$ ,  $i \in \{1, \dots, \ell\}$ , are cyclic. In general, the subspaces  $\mathcal{V}_i$  are not unique for a fixed  $\mathcal{A}$ . The integer  $\ell$  is unique and is called the *cyclic index of  $\mathcal{A}$* .

One of our main results is to show that the cyclic index is a lower bound on the number of measurements required to reconstruct  $w_\tau$  (see Prop. 3 and Alg. 1 in [23]). The matrix transform associated to this theorem is known as the *Frobenius normal form* (denoted by  $C \in \mathbb{R}^{M \times M}$ ): for  $\mathcal{A} \in \mathbb{R}^{M \times M}$ ,  $\exists Q \in \mathbb{R}^{M \times M}$  invertible such that  $\mathcal{A} = QCQ^{-1}$ . We will also use the *Jordan*

*decomposition*, where for  $\mathcal{A} \in \mathbb{R}^{M \times M}$ ,  $\exists P \in \mathbb{R}^{M \times M}$  invertible such that  $\mathcal{A} = P\Lambda P^{-1}$ , where  $\Lambda$  is a unique block diagonal matrix with Jordan blocks with  $\lambda_i$  along the diagonal. If all the eigenvalues  $\lambda_i$  are nonzero and real, we say the matrix has a *full-rank Jordan decomposition*.

## Main Results

In this section, we prove results concerning the observability of spatiotemporally varying functions modeled by the functional evolution and measurement equations (3). In particular, observability of the system states implies that we can recover the current state of the spatiotemporally varying function using a small number of sampling locations  $N$ , which allows us to 1) track the function, and 2) predict its evolution forward in time. We work with the approximation  $\widehat{\mathcal{H}} \approx \mathcal{H}$ : given  $M$  basis functions, this implies that the dual space of  $\widehat{\mathcal{H}}$  is  $\mathbb{R}^M$ . Proposition 1 shows that if  $\widehat{\mathcal{A}}$  has a full-rank Jordan decomposition, the observation matrix  $K$  meeting a condition called *shadedness* (Definition 1) is sufficient for the system to be observable. Proposition 2 provides a lower bound on the number of sampling locations required for observability which holds for any  $\widehat{\mathcal{A}}$ . Proposition 3 constructively shows the existence of an abstract measurement map  $\tilde{K}$  achieving this lower bound. Since the measurement map does not have the structure of a kernel matrix, a slightly weaker sufficient condition for the observability of any  $\widehat{\mathcal{A}}$  is in Theorem 1. Finally, since both  $K$  and  $K_{CD}$  are kernel matrices generated from a shared kernel, these observability results translate directly into controllability results. Proofs of all claims are in the appendix.

**Definition 1. (Shaded Observation Matrix)** Given  $k : \Omega \times \Omega \rightarrow \mathbb{R}$  positive-definite on a domain  $\Omega$ , let  $\{\widehat{\psi}_1(x), \dots, \widehat{\psi}_M(x)\}$  be the set of bases generating an approximate feature map  $\widehat{\psi} : \Omega \rightarrow \widehat{\mathcal{H}}$ , and let  $\mathcal{X} = \{x_1, \dots, x_N\}$  be the set of sampling (or sensing) locations, with each  $x_i \in \Omega$ . Let  $K \in \mathbb{R}^{N \times M}$  be the observation matrix, where  $K_{ij} := \widehat{\psi}_j(x_i)$ . For each row  $K_{(i)} := [\widehat{\psi}_1(x_i) \dots \widehat{\psi}_M(x_i)]$ , define the set  $\mathcal{I}_{(i)} := \{\iota_1^{(i)}, \iota_2^{(i)}, \dots, \iota_{M_i}^{(i)}\}$  to be the indices in the observation matrix row  $i$  which are nonzero. Then if  $\bigcup_{i \in \{1, \dots, N\}} \mathcal{I}^{(i)} = \{1, 2, \dots, M\}$ , we denote  $K$  as a shaded observation matrix (see Figure 4a).

This definition seems quite abstract, so the following remark considers a more concrete example.

**Remark 1.** let  $\widehat{\psi}$  be generated by the dictionary given by  $\mathcal{C} = \{c_1, \dots, c_M\}$ ,  $c_i \in \Omega$ . Note that since  $\widehat{\psi}_j(x_i) = \langle \psi(x_i), \psi(c_j) \rangle_{\mathcal{H}} = k(x_i, c_j)$ ,  $K$  is the kernel matrix between  $\mathcal{X}$  and  $\mathcal{C}$ . For the kernel matrix to be shaded thus implies that there does not exist an atom  $\psi(c_j)$  such that the projections  $\langle \psi(x_i), \psi(c_j) \rangle_{\mathcal{H}}$  vanish for all  $x_i$ ,  $1 \leq i \leq N$ . Intuitively, the shadedness property requires that the sensor locations  $x_i$  are privy to information propagating from every  $c_j$ . As an example, note that, in principle, for the Gaussian kernel, a single row generates a shaded kernel

matrix. However, in this case, the matrix can have many entries that are extremely close to zero, and will probably be very ill-conditioned.

**Proposition 1.** Given  $k : \Omega \times \Omega \rightarrow \mathbb{R}$  positive-definite on a domain  $\Omega$ , let  $\{\widehat{\psi}_1(x), \dots, \widehat{\psi}_M(x)\}$  be the set of bases generating an approximate feature map  $\widehat{\psi} : \Omega \rightarrow \widehat{\mathcal{H}}$ , and let  $\mathcal{X} = \{x_1, \dots, x_N\}$ ,  
5  $x_i \in \Omega$ . Consider the discrete linear system on  $\widehat{\mathcal{H}}$  given by the evolution and measurement equations (3). Suppose that a full-rank Jordan decomposition of  $\widehat{A} \in \mathbb{R}^{M \times M}$  of the form  $\widehat{A} = P\Lambda P^{-1}$  exists, where  $\Lambda = [\Lambda_1 \dots \Lambda_O]$ , and there are no repeated eigenvalues. Then, given a set of time instances  $\Upsilon = \{\tau_1, \tau_2, \dots, \tau_L\}$ , and a set of sampling locations  $\mathcal{X} = \{x_1, \dots, x_N\}$ , the system (3) is observable if the observation matrix  $K_{ij}$  is shaded according to Definition 1,  $\Upsilon$   
10 has distinct values, and  $|\Upsilon| \geq M$ .

When the eigenvalues of the system matrix are repeated, it is not enough for  $K$  to be shaded. In the next proposition, we take a geometric approach and utilize the rational canonical form of  $\widehat{A}$  to obtain a lower bound on the number of sampling locations required. Let  $r$  be the number of unique eigenvalues of  $\widehat{A}$ , and let  $\gamma_{\lambda_i}$  denote the geometric multiplicity of eigenvalue  
15  $\lambda_i$ . Then the *cyclic index* of  $\widehat{A}$  is defined as  $\ell = \max_{1 \leq i \leq r} \gamma_{\lambda_i}$  [52].

**Proposition 2.** Suppose that the conditions in Proposition 1 hold, with the relaxation that the Jordan blocks  $[\Lambda_1 \dots \Lambda_O]$  may have repeated eigenvalues (i.e.  $\exists \Lambda_i$  and  $\Lambda_j$  s.t.  $\lambda_i = \lambda_j$ ). Then there exist kernels  $k(x, y)$  such that the lower bound  $\ell$  on the number of sampling locations  $N$  is given by the cyclic index of  $\widehat{A}$ . In other words, the system in (3) is observable if  $N \geq \ell$ .

20 We will give a concrete example to build intuition regarding this lower bound below. For now, we note the following:

**Proposition 3.** Given the conditions stated in Proposition 2, it is possible to construct a measurement map  $\widetilde{K} \in \mathbb{R}^{\ell \times M}$  for the system given by (3), such that the pair  $(\widetilde{K}, \widehat{A})$  is observable.

The construction provided in the proof of Proposition 3 is utilized in Algorithm 1, which  
25 uses the rational canonical structure of  $\widehat{A}$  to generate a series of vectors  $v_i \in \mathbb{R}^M$ , whose iterations  $\{v_1, \dots, \widehat{A}^{m_1-1}v_1, \dots, v_\ell, \dots, \widehat{A}^{m_\ell-1}v_\ell\}$  generate a basis for  $\mathbb{R}^M$ . Unfortunately, the measurement map  $\widetilde{K}$ , being an abstract construction unrelated to the kernel, does not directly select  $\mathcal{X}$ . We will show how to use the measurement map to guide a search for  $\mathcal{X}$  in Remark 3 (in Appendix). For now, we state a sufficient condition for observability of a general system.

**Theorem 1.** Suppose that the conditions in Proposition 1 hold, with the relaxation that the Jordan blocks  $[\Lambda_1 \dots \Lambda_O]$  may have repeated eigenvalues. Let  $\ell$  be the cyclic index of  $\widehat{A}$ .

Define

$$\mathbf{K} = [K^{(1)^T} \dots K^{(\ell)^T}]^T \quad (7)$$

as the  $\ell$ -shaded matrix (see Figure 4b) which consists of  $\ell$  shaded matrices with the property that any subset of  $\ell$  columns in the matrix are linearly independent from each other. Then system (3) is observable if  $\Upsilon$  has distinct values, and  $|\Upsilon| \geq M$ .

While Theorem 1 is a quite general result, the condition that any  $\ell$  columns of  $\mathbf{K}$  be  
5 linearly independent is a very stringent condition. One scenario where this condition can be met with minimal measurements is in the case when the feature map  $\widehat{\psi}(x)$  is generated by a dictionary of atoms with the Gaussian RBF kernel evaluated at sampling locations  $\{x_1, \dots, x_N\}$  according to (2), where  $x_i \in \Omega \subset \mathbb{R}^d$ , and  $x_i$  are sampled from a non-degenerate probability distribution on  $\Omega$  such as the uniform distribution. For a semi-deterministic approach, when the  
10 dynamics matrix  $\widehat{A}$  is block-diagonal, we can utilize a simple heuristic:

**Remark 2.** Let  $\Omega$  be compact,  $\mathcal{C} = \{c_1, \dots, c_M\}$ ,  $c_i \in \Omega$ , and let the approximate feature map be defined by (2). Consider the system (3) with  $\widehat{A} = \Lambda$ , and let  $\Upsilon = \{0, 1, \dots, M - 1\}$ . Then the measurement map  $\widetilde{K}$ 's values lie in  $\{0, 1\}$ ; in particular, each row  $\widetilde{K}^{(j)}$ ,  $j \in \{1, \dots, \ell\}$ , corresponds to a subspace  $\widehat{\mathcal{H}}_j$ , generated by a subset of centers  $\mathcal{C}^{(j)} \subset \mathcal{C}$ . Generate samples  
15  $x_i^{(j)}$  to create a kernel matrix  $K^{(j)}$  that is shaded only with respect to centers  $\mathcal{C}^{(j)}$ . Once this is done, move on to the next subspace  $\widehat{\mathcal{H}}_{j+1}$ . When all  $\ell$  rows of  $\widetilde{K}$  are accounted for, construct the matrix  $\mathbf{K}$  as in (7). Then the resulting system  $(\mathbf{K}, \widehat{A})$  is observable.

This heuristic is formalized in Algorithm 2. Note that in practice, the matrix  $\widehat{A}$  needs to be inferred from measurements of the process  $f_\tau$ . If no assumptions are placed on  $\widehat{A}$ , it's clear  
20 that at least  $M$  sensors are required for the system identification phase. Future work will study the precise conditions under which system identification is possible with less than  $M$  sensors.

## Discussion of Theoretical Results

The systems-theoretic approach taken in this paper reveals something rather surprising: functions with complex dynamics (with a small cyclic index) can be recovered with less  
25 sensor placements than functions with simpler dynamics. Although seemingly counterintuitive, it becomes clear that this is because complex dynamics, which are characterized by a lower geometric multiplicity of the eigenvalues, ensure that the orbit  $\Theta := \{\widehat{A}w_\tau\}_{\tau \in \Upsilon}$  traverses a greater portion of  $\mathbb{R}^M \equiv \widehat{\mathcal{H}}$  and thus that fewer sensors can recover more geometric information. On the other hand, in ‘simpler’ functional evolution,  $\Theta$  evolves along strict subspaces of  $\mathbb{R}^M$ ,  
30 and so more independent sensors are required to infer the same amount of information.

In the case described in Remark 3, we have a set of centers  $\mathcal{C} = \{c_1, \dots, c_M\}$ , which generate the bases  $\mathcal{F}^{\mathcal{C}} = \{\psi(c_1), \dots, \psi(c_M)\}$ . Let the cyclic index be  $\ell$ : this implies that there exist  $\ell$  subsets  $\Psi^{(i)}$  of  $\mathcal{F}^{\mathcal{C}}$  with at least one element  $\psi(c_j)$  each, leading to  $\binom{M}{\ell}$  possible choices: Figure 8 represents these choices as hyperplanes separating the subsets. The measurement map 5 described in Alg. 1 in [23] induces this *decomposition of bases*  $\mathcal{F}^{\mathcal{C}} = \{\Psi^{(1)}, \dots, \Psi^{(\ell)}\}$  in polynomial time. Further, each subset  $\Psi^{(i)}$  is directly associated to a subset of centers  $\mathcal{C}^{(i)} \subset \mathcal{C}$ , which allows us to pick targeted sensor locations  $x_i \in \Omega$ . In particular, for radially symmetric 10 kernels such as the Gaussian, the centroid of the convex hull of  $\mathcal{C}^{(i)}$  is sufficient for generating a sensor placement. The measurement map is a significant theoretical insight into sensor placement for dynamically changing environments, because it directly takes into account the dynamics of 15 the process. Of course, in practice, this may be too expensive for approximate feature spaces with  $M$  very large, so one can use random sampling to generate the sensor locations instead, at the cost of  $N$  being larger than  $\ell$ . The advantage here though is that since random sampling is computationally inexpensive, different choices of sensor placements can be generated and evaluated relatively quickly.

Another point to note is that since the collection of bases  $\{\widehat{\psi}_i(x)\}_{i=1}^M$  determines the richness of the function space  $\widehat{\mathcal{H}} \approx \mathcal{H}$  we operate in, it determines the fidelity of the model approximation to the true time-varying function. As a consequence, observability of the system in  $\widehat{\mathcal{H}}$  refers to the best possible approximation in  $\widehat{\mathcal{H}}$ . The greater the number of bases, the higher 20 the dimensionality, which results in greater model fidelity, but which may require a much greater number of measurements for state recovery. This is where the lower bounds presented in the paper are particularly useful, because they show that for functional evolutions corresponding to certain  $\widehat{A}$ , *the number of sensor placements are essentially independent of the dimensionality  $M$* , but depend rather on the cyclic index of  $\widehat{A}$ .

25 Figure 7 gives an overall picture on the process of generating a kernel observer, while Figure 8 gives two approaches to sensor selection in our framework. The measurement map approach can generate a smaller set of sensors than the random placement approach, but comes at an additional computational cost.

## Random Sensor Placement

30 We now elaborate on how the challenging problem of sensor placement can be tackled through random selection. This process of random selection is a product of the kernel observer model described above. We present the theoretical background required to prove Theorem 2, which states the expected number of randomly placed sensors required to monitor a given spatiotemporal process, and Theorem 3, which determines the probability with which optimal

sensor placement is ensured given that,  $N$  number of sensors have been placed.

As discussed earlier, we work with an approximate feature space  $\widehat{\mathcal{H}}$ , with the corresponding transition operator  $\widehat{A} : \widehat{\mathcal{H}} \rightarrow \widehat{\mathcal{H}}$ , representing finite-dimensional functional evolution. To achieve observability for the pair  $(\widehat{A}, K)$ , row vectors of the corresponding observability matrix,  $\mathcal{O}$ ,  
5 should form the basis for the  $\mathbb{R}^M$ -dimensional space  $\widehat{\mathcal{H}}$ . According to the rational canonical structure Theorem [52],  $\widehat{A}$  can successively decompose the dual space  $\mathbb{R}^M$  into subspaces,  $\mathcal{V}_i \subset \mathcal{V}$ ,  $i \in \{1, \dots, \ell\}$ , with properties, i)  $\mathcal{V} = \mathcal{V}_1 \oplus \dots \oplus \mathcal{V}_\ell$ , ii)  $\widehat{A}\mathcal{V}_i \subset \mathcal{V}_i$ , and iii)  $\widehat{A}|_{\mathcal{V}_i}, i \in \{1, \dots, \ell\}$ , are cyclic. The integer  $\ell$  is unique and is called the *cyclic index of  $\widehat{A}$* . Each of these properties contribute towards the theorem on the number of random samples required to achieve  
10 observability. The first property shows that the space  $\mathbb{R}^M$  can be decomposed into  $\ell$  independent subspaces. The second property shows that the vector  $v_i \in \mathcal{V}_i$  stays in  $\mathcal{V}_i$  even when operated upon by  $\widehat{A}$ . Thus, to generate bases for  $\mathbb{R}^M$ , one needs at least  $\ell$  vectors, say,  $v_1, \dots, v_\ell$ , with respect to each subspace  $\mathcal{V}_1, \dots, \mathcal{V}_\ell$ . This holds due to the third property, but requires that the vectors  $v_1, \dots, v_\ell$ , are the cyclic generators of their corresponding subspaces. Our analysis is  
15 based on whether a randomly selected sensor can generate a cyclic generator. To examine this, recall that a row vector  $K_{(i)}$  generated by a randomly selected sensor location  $x_i$  takes the form,

$$K_{(i)} = \left[ k(x_i, c_1), \dots, k(x_i, c_M) \right]. \quad (8)$$

Here, for radial kernels for example, the entries corresponding to the centers closer to  $x_i$  tend to be non-zero, whereas the others tend to be zero. The rows  $K_{(i)}$  from random sensor placement  
20 must be able to generate a basis for a subspace  $\mathcal{V}_i$ , and thus must be cyclic generators. We will derive the expected number of random sensor placements sufficient for observability for the case where  $\widehat{A} = \Lambda$  and then attempt to generalize the result for any  $\widehat{A}$ . Note  $\Lambda$  is a block diagonal Jordan form. In this case, the cyclic generator for each subspace  $\mathcal{V}_i$ , is a vector  $v_i$  with non-zero entries corresponding to the leading entry of the Jordan blocks of  $\mathcal{V}_i$ .

Overall, our construction is as follows: for each subspace  $\mathcal{V}_i$ , let  $\mathcal{C}_{\mathcal{V}_i} \subset \mathcal{C}$  be the centers  
25 corresponding to those leading entry of Jordan blocks: then the minimum number of random samples required to generate the bases for  $\mathcal{V}_i$  is equal to the number of Jordan blocks comprising  $\mathcal{V}_i$ . Altogether, the minimum number of random samples required to generate a basis for  $\mathbb{R}^M$  is equal to the total number of Jordan blocks in  $\widehat{A}$ . Let  $\varsigma$  be the total number of Jordan blocks in  
30  $\widehat{A}$ , then

$$\varsigma = \sum_{\lambda \in \sigma(\widehat{A})} \gamma_{\widehat{A}}(\lambda) \quad (9)$$

where  $\sigma(\widehat{A})$  represents the spectrum of  $\widehat{A}$ , whose elements are the eigenvalues of  $\widehat{A}$ , and  $\gamma_{\widehat{A}}(\lambda)$  is the geometric multiplicity corresponding to the eigenvalue  $\lambda$ , which is also equal to the

total number of Jordan blocks corresponding to the eigenvalue  $\lambda$ . Define a set of centers  $\mathcal{C}_\varsigma$  with elements  $\{c_1, c_2, \dots, c_\varsigma\}$ , to be the centers corresponding to the leading entries of the Jordan blocks. For sensor location  $x \in \Omega$ , and  $\epsilon > 0$ , let  $k(x, c_j) > \epsilon$ , denote the region  $\Omega_j \subset \Omega$ , such that the kernel evaluation with respect to center  $c_j$  is greater than  $\epsilon$ , that is  
<sup>5</sup>  $\Omega_j \equiv \{x \in \Omega : k(x, c_j) > \epsilon\}$ . We define  $p_\epsilon$  as

$$p_\epsilon = \min_{c_j \in \mathcal{C}_\varsigma} \frac{\nu(k(x, c_j) > \epsilon)}{\nu(\Omega)}, \quad (10)$$

where  $\nu$  is a measure in the real analysis sense. Hence,  $p_\epsilon$  corresponds to a lower bound on the probability that a random sample lies within the  $\epsilon$ -shaded region of a particular center  $c_j$ . With all of this in place, we can prove the following theorem.

**Theorem 2.** *Given the spatiotemporal function  $f(x, \tau)$  with  $x \in \Omega \subseteq \mathbb{R}^D, \tau \in \mathbb{Z}^+$  its kernel observer model (3), and a tolerance parameter  $\epsilon > 0$ , the expected number of randomly placed sensor locations required to achieve observability for the pair  $(K, \hat{A})$  is  $\varsigma/p_\epsilon$  where  $\varsigma$  is the summation over geometric multiplicities of each  $\lambda \in \sigma(\hat{A})$  given by (9).*

**Theorem 3.** *Given the spatiotemporal function  $f(x, \tau)$  with  $x \in \Omega \subseteq \mathbb{R}^D, \tau \in \mathbb{Z}^+$ , its kernel observer model (3), a tolerance parameter  $\epsilon > 0$ , summation over geometric multiplicities of each  $\lambda \in \sigma(\hat{A})$  denoted by  $\varsigma$  as in (9), and a constant  $\delta \in (0, 1]$ , the probability that pair  $(K, \hat{A})$  is unobservable after the selection of  $N$  random sensors is at most  $e^{-\frac{1}{2}(Np_\epsilon - 2\varsigma)}$ , where  $p_\epsilon$  is given by (10) and  $N > 2\varsigma/p_\epsilon$ .*

For the case when  $\hat{A} \neq \Lambda$ , a change of basis can be used to obtain  $\Lambda = P^{-1}\hat{A}P$ , where  $P$  is the projection map. There are two challenges in performing the above analysis for  $\Lambda$  so obtained: first, the leading entries of Jordan blocks do not directly correspond to the centers  $\{c_1, \dots, c_M\}$  which was the case for  $\hat{A} = \Lambda$ . Second, although we can obtain the transformation of the row vector (8) using the projection map  $P$ , we can no longer arrive at the definition of the probability  $p_\epsilon$  as in (10). The existence of the similarity transform hints that the results in Theorems 2-3 should hold for any  $\hat{A}$ , but the mathematical tools utilized in the paper seem to be insufficient to prove them. However, we present some empirical evidence for these claims for when  $\hat{A} \neq \Lambda$  in the empirical results section.  
<sup>20</sup> <sup>25</sup>

## Generalizing Across Similar Spatiotemporally Evolving Systems

Building on the Kernel Observers method, let us introduce Evolving Gaussian Processes (E-GP). The primary novelty in this method of generating a model is learning an  $\hat{A}$  matrix  
<sup>30</sup> for *multiple* systems. The ultimate goal of this research would be to generate highly efficient machine learning models that can be used instead of the costly numerical simulations for design

and autonomy purposes. This would be a major success for the design and control of complex physical systems, such as soft robotics, as they would significantly reduce the cost and resources required in simulations. The ability to generalize across different physical situations, is critical. This is a difficult problem, as it requires that the model have the capability to actually learn  
5 the underlying physics and not just input-output relationships. For example, in the context of fluid flows, these models must be able to predict fluid dynamics at different conditions (e.g. Reynolds number) than the training data. E-GP, as far as the authors know, was the first machine learning method to generalize across spatiotemporally evolving systems of such complexity using end-to-end data.

<sup>10</sup> We found that the class of functional evolutions  $\mathbb{F}$  defined by linear Markovian transitions in a RKHS is still sufficient to model the nonlinear Navier Stokes equations which govern fluid dynamics, since the unknown map  $\psi$  allows us to model highly nonlinear dynamics in the input space. However, we do expect that phenomena such as bifurcation or turbulence will require nonlinear mappings  $\mathcal{H}$ . There are three steps to generate an E-GP model:

- <sup>15</sup> 1) After picking the kernel and estimating the bandwidth hyperparameter  $\sigma$  (we utilize the maximum likelihood approach, although other approaches can be used), find an optimal basis vector set  $\mathcal{C}$  using the algorithm in [53].
- 2) Use Gaussian process inference to find weight vectors for each time-step in the training set(s), generating the sequence  $w_\tau, \tau = 1, \dots, T$  for each system. A uniform time-step makes next step easier but can be worked around for non-uniform data sets  
<sup>20</sup>
- 3) Using the weight trajectory, use matrix least-squares with the equation  $\hat{A}[w_1, w_2, \dots, w_{T-1}] = [w_2, w_3, \dots, w_T]$  to solve for  $\hat{A}$ .
- 4) To generate a multi-system model, concatenate the weight trajectories from each similar system in the least-squares computation of  $\hat{A}$ . That is, let  $W_\theta = [w_1^{(\theta)}, w_2^{(\theta)}, \dots, w_{n-1}^{(\theta)}]$  and  $W'_\theta = [w_2^{(\theta)}, w_3^{(\theta)}, \dots, w_n^{(\theta)}]$  be the weight trajectory and next weight trajectory for some parameter  $\theta$ . Then we solve the least-squares problem  $\hat{A} = [W_{\theta_1}, \dots, W_{\theta_n}] = [W'_\theta, \dots, W'_{\theta_n}]$   
<sup>25</sup>

For the sake of defining when it is appropriate to expect this method to be able to generalize across different spatiotemporally evolving systems, we shall define what it means for two fluid flows to be *similar*. In configuring a fluid dynamics simulation, a set of quantifiable parameters  
<sup>30</sup> are defined. Two dynamical fluid systems  $S_1$  and  $S_2$  are considered *similar* if they have the same configuration of parameters and differ only in the value of at most one parameter. Furthermore, we require that the parameter be continuously variable, and that any observable quantity in the domain of the system vary smoothly as that parameter varies from its value in  $S_1$  to its value in  $S_2$ . For example, for fluids flowing past identical cylinders, the Reynolds number associated  
<sup>35</sup> with the free stream velocity may be varied to produce similar systems. However, to replace

the system's cylinder with a triangle would be to qualitatively change the configuration of the system parameters, and thus would produce a non-similar system.

Unlike neural networks, the weights in an E-GP do not exist in some abstract, difficult-to-comprehend space, but are associated with kernel centers in specific locations in the domain. We refer to this attribute of E-GPs as the *spatial encoding* property. This property is an extremely valuable tool for gaining insight into the learned model works:

- 1) By plotting which kernel centers are associated with which invariant subspaces in the transition matrix, one can visualize where the eigenfunctions are found and how the dynamic modes are separated spatially
- 10 2) By plotting arrows from center  $c_j$  to  $c_i$  for each of the largest elements  $\hat{a}_{ij}$  of  $\hat{A}$ , one can visualize how different areas of the domain influence each other's evolution.
- 3) By performing an eigendecomposition of the  $\hat{A}$  matrix, and transforming the eigenvectors back from the weight space to the function space, one can obtain the Koopman modes (and associated eigenvalues) of the system (see next section)

## 15 Spectral Analysis of Evolving Gaussian Process Model and Resulting Algorithms

For a general dynamical system  $f_{\tau+1} = \mathbb{F}(f_\tau)$  defined on a state space (for us, an RKHS, i.e.  $f \in \mathcal{H}$ ), we can define an arbitrary, vector-valued observable  $g : \mathcal{H} \rightarrow \mathbb{R}^N$ . Note that the space of observables  $g$  is a vector space. The Koopman operator  $U$  is defined to be the operator on the space of observables such that

$$Ug(f_\tau) = g(\mathbb{F}(f_\tau)) = g(f_{\tau+1}) \quad (11)$$

This operator is clearly linear from its definition, and thus it is reasonable to examine its spectral properties. The special observables  $\phi : \mathcal{H} \rightarrow \mathbb{C}$  that have the property

$$U\phi(f_\tau) = \phi(\mathbb{F}(f_\tau)) = \phi(f_{\tau+1}) = \lambda\phi(f_\tau) \quad (12)$$

are the eigenfunctions of  $U$ , and the associated  $\lambda$  are the eigenvalues. Suppose now we have a vector-valued observable  $u(f, x)$ , where  $x \in \Omega$  and  $f \in \mathcal{H}$ .

**Definition 2.** The Koopman mode  $s(x)$  at isolated eigenvalue  $\lambda$  of algebraic multiplicity 1 is the projection of  $u(f, x)$  onto the eigenfunction  $\phi_\lambda(f)$  of  $U$  at  $\lambda$ . [54]

20 As previously shown by Rowley in 2009 [55], the modes produced by the Dynamic Mode Decomposition (DMD) algorithm constitute a subset of Koopman modes. Mezic (2013) showed that there exists, in principle if not in practice, a rigorous method for computing the full set of Koopman modes by a method known as generalized Laplace

analysis (GLA). We have generated a number of results interpreting the E-GP model in terms of Koopman operator theory, culminating in the proof that the eigenvalues and eigenvectors of  $\hat{A}$  are related to the Koopman eigenvalues and modes.

**Proposition 4.** Let  $\mathcal{H}$  be a RKHS with an approximate feature space  $\hat{\mathcal{H}}$ , and let  $\mathbf{u}(f, \mathbf{x})$  be an observable in the Koopman sense with respect to the dynamical system  $f_{\tau+1} = \mathbb{F}(f_\tau)$  in  $\mathcal{H}$ . Then  $\hat{\mathbf{u}}(f, \mathbf{x}) := \mathbf{u}(\hat{f}, \mathbf{x})$ , where  $\hat{f}$  is the projection of  $f \in \mathcal{H}$  onto  $\hat{\mathcal{H}}$ , is also an observable.

This follows from the fact that the projection from the function space to its subspace is well-defined when the  $\hat{\psi}_i$  are independent. Now, Koopman modes of observables are of interest because they are akin to the eigenvector expansions utilized in linear dynamics. If we separate the Koopman operator into  $U = U_s + U_r$  where  $U_s$  has a pure point spectrum and  $U_r$  has a pure continuous spectrum, then we can write

$$U^t \mathbf{u}(f, \mathbf{x}) = \mathbf{u}^*(f, \mathbf{x}) + \sum_{j=1}^k \lambda_j^t \phi_j(f) \mathbf{s}_j(\mathbf{x}) + U_r^t \mathbf{u}(f, \mathbf{x}) \quad (13)$$

where  $\mathbf{u}^*(\mathbf{x})$  represents the time-averaged part of the field, which corresponds with  $\lambda = 1$  (see GLA in [54]). The continuous part is usually discarded/neglected since it represents the part of the field that is genuinely aperiodic (or chaotic) in time. Now, from this expansion we can prove that:

**Proposition 5.** Let  $\mathbf{u}(f, \mathbf{x})$  and  $\hat{\mathbf{u}}(f, \mathbf{x})$  be observables as in the Proposition 4, and  $\mathbf{s}_j(\mathbf{x})$  be the Koopman modes associated with the projection of the former onto the eigenfunctions  $\phi_j$  of  $U$  at  $\lambda_j$ . Then the Koopman modes associated with  $\hat{\mathbf{u}}$  are

$$\hat{\mathbf{s}}_j(\mathbf{x}) = \frac{\phi_j(\hat{f})}{\phi_j(f)} \mathbf{s}_j(\mathbf{x}) \quad (14)$$

*Proof.* Using the spectral expansion and the definition of the observables, we have

$$\begin{aligned} U^t \hat{\mathbf{u}}(f, \mathbf{x}) &= U^t \mathbf{u}(\hat{f}, \mathbf{x}) = \mathbf{u}^*(\hat{f}, \mathbf{x}) + \sum_{j=1}^k \lambda_j^t \phi_j(\hat{f}) \mathbf{s}_j(\mathbf{x}) + U_r^t \mathbf{u}(\hat{f}, \mathbf{x}) \\ &= \hat{\mathbf{u}}^*(f, \mathbf{x}) + \sum_{j=1}^k \lambda_j^t \phi_j(f) \hat{\mathbf{s}}_j(\mathbf{x}) + U_r^t \hat{\mathbf{u}}(f, \mathbf{x}) \end{aligned}$$

It is easy to see that this must be the spectral expansion of  $\hat{\mathbf{u}}(f, \mathbf{x})$   $\square$

Note that this means the modes of the two observables are identical in shape (only differing by a multiplicative constant). In the exceptional case that  $\phi_j(f) = 0$ , the GLA method is unable

to compute  $s_j(\mathbf{x})$  anyway, so that is not an issue for this proposition. The final step in our analysis is to connect the Koopman modes with the spectral decomposition of our model's  $\widehat{A}$  operator in the dual space.

**Theorem 4.** *In an E-GP or KO model of the dynamical system  $f_{\tau+1} = \mathcal{A}f_\tau$ , the eigenvalues of  $\widehat{A}$  correspond to the eigenvalues of the Koopman operator. Furthermore, if the observable  $\mathbf{u}(f, \mathbf{x})$  is the evaluation operator,  $\mathbf{u}(f, \mathbf{x}) = f(\mathbf{x})$ , then the eigenvectors  $v_j$  of  $\widehat{A}$  are related to the Koopman modes of the observable  $s_j(\mathbf{x}) = \frac{\phi_j(f)}{\phi_j(f)} \widehat{\Psi}(\mathbf{x}) v_j$ .*

*Proof.* It should be clear that, as long as the projection onto the approximate feature space is well defined, that  $w(f_\tau) := w_\tau$  is an observable. Then if we let  $\phi_j(f) = \langle w(f), q_j \rangle$ , where  $q_j$  are eigenvectors of the adjoint  $\widehat{A}_*$ , then  $\phi_j$  is an eigenfunction of the Koopman operator:  
 $U\phi_j(f_\tau) = \langle w(f_{\tau+1}), q_j \rangle = \langle w_{\tau+1}, q_j \rangle = \langle \widehat{A}w_\tau, q_j \rangle = \langle w_\tau, \widehat{A}^*q_j \rangle = \lambda_j \langle w_\tau, q_j \rangle = \lambda_j \phi_j(f_\tau)$   $\square$

in an rkhs, f and fhat being close means they are pointwise close.

It is well known that in the case in which the dynamical system is linear, such as  $w_{\tau+1} = \widehat{A}w_\tau$ , its matrix eigenvalues are eigenvalues of the associated Koopman operator. The associated Koopman eigenfunctions

We can show that there exists a relationship between the

### Invariant Subspaces

One way of viewing the invariant subspaces concept (as described earlier) is say that information contained in an invariant subspace never leaves that invariant subspace.  
 $20$  We hypothesize that the kernel centers associated with the invariant subspaces of a spatiotemporally evolving system are generally associated with spatial regions in the domain (and not just homogeneously spread all over and or mixed with the other invariant subspaces). This hypothesis makes sense both physically and mathematically. In physics, the principle of locality states that an object is only directly influenced by its immediate surroundings [56]. If this is the case (and there is no reason to believe that it is not, apart from certain quantum dynamics situations), and the E-GP model accurately captures the physics of the system, then information (measurable phenomena) may only travel continuously from one point in the domain to another. Mathematically, since a value at any one point in the domain is influenced by the weights of multiple nearby centers, we  
 $25$  would expect nearby centers to be connected dynamically, unless separated by “plains” where the values are indistinguishable from noise.

When a square matrix is perfectly formed, the Jordan form of a  $n \times n$  matrix  $\hat{A}$  is block diagonal, and therefore gives a decomposition of the  $n$  dimensional Euclidean space into invariant subspaces of  $\hat{A}$ . The cyclic index, which can be found by counting the geometric multiplicities of eigenvalues in  $\hat{A}$ , gives the number of invariant subspaces. In reality, data-driven approximations of  $\hat{A}$  rarely give such easily interpretable properties. Hence, the need for an algorithm can divide the system into invariant subspaces even when the boundaries between them are “fuzzy”.

Each block in the Jordan normal form has a set of corresponding eigenvectors and an eigenvalue with geometric multiplicity. When we transform the former back into the domain space, we obtain complex-valued functions which are the Koopman modes of the system. These provide an image of what kind of structures we see in the dynamics. The eigenvalues describe the frequency with which these structures oscillate between their real and imaginary forms, as well as the exponential growth or decay of their magnitudes.

### *k*-Invariant Subspaces Clustering

The intuition behind our  $k$ -invariant subspaces clustering algorithm is to replace the Euclidean distance in the  $k$ -means algorithm with a different metric of “nearness”, namely one corresponding with the dynamic connections in the space. The  $\hat{A}$  matrix provides easy access to these: its rows  $\hat{A}_{i*}$  indicate which centers inform the  $i^{\text{th}}$  value of  $w_{t\text{index}+1}$ , and its columns  $\hat{A}_{*j}$  indicate what centers will be informed by the  $j^{\text{th}}$  value of  $w_\tau$ . However, these values do tend to be biased towards the eigenmodes with higher frequencies (i.e. those whose eigenvalues have a greater polar angle) and those which grow exponentially. To control for this, we chose modify the eigenvalues of the matrix in the following way: (1) We zeroed any eigenvalue that is clearly inside (not on) the unit circle, (2) We unitized the remaining eigenvalues, and (3) We adjusted the frequency of the remaining eigenvalues on the unit circle to either  $\pm\frac{\pi}{4}$ . If the eigen-decomposition of the original was  $\hat{A} = UDU^{-1}$ , then we can reconstruct a new matrix with our modified eigenvalues  $\bar{D}$  as  $\bar{A} = U\bar{D}U^{-1}$ .

Much like the pairwise-squared deviations of points formulation of the  $k$ -means clustering problem, we can now write our problem as

$$\arg \max_S \sum_{i=1}^k \frac{1}{2|S_i|} \sum_{\substack{x_i \neq x_j \\ x_j, x_i \in S_i}} \bar{A}_{ij}^2$$

This problem can be solved with the following algorithm:

Note the differences between this and the  $k$ -means clustering algorithm: first, we

are *maximizing* since the terms represent influence rather than distance. Secondly, in the possible case that  $|S_k| = 0$ , we make the total value zero in order to avoid division by zero. Thirdly, we exclude centers' influence on themselves from the score by taking  $S_k \setminus \{i\}$ .

This algorithm can be repeated as many times as desired with different random

5 initial conditions, and the result which produces the highest score retained.

### *Scoring Paths for a Moving Agent*

Consider the problem of predicting the current state of a spatiotemporally-evolving function with a set of robots, each one of whom can only observe the phenomena partially. When there are sufficient such robots to cover the entire domain, there are 10 many approaches in the coverage literature available for solving this problem [?], [?]. In those approaches, the robots figure out the best trajectories to create a network of connected sensors that can simultaneously observe the whole environment. However, when the number of robots is limited, full coverage of the domain is not possible. How a few robots whose combined sensors can only observe a small part of the domain can predict 15 the state of the spatiotemporally-evolving function remains an open problem. Here we consider a limiting case of this problem where only a single robot is available to observe the whole environment, but the robot has available to it an approximate model of the function learned through past experience Evolving Gaussian Processes.

We will be using a Kalman filter in the weight space of the E-GP as described in 20 previous sections. In order to determine where a moving agent should go in order to gather the measurements which will help it converge to the true state of the system quickest, we use information from the spectral analysis of the E-GP model. We postulate that the value of taking a measurement at a point in the domain with respect to a particular eigenvector is proportional to the following factors:

- 1) The spatial extent of the eigenvector. This can be taken to be equivalent to the area under the curve of the magnitude of the complex function corresponding to that eigenvector (normalized by the function peak)

$$SE_i = \frac{1}{\sup_x |v_i \hat{\Phi}(x)|} \int_X |v_i \hat{\Phi}(x)| dx$$

25 This Riemann integral can of course be approximated by discretely partitioning the space. From this point of view, we can see that this expression represents what proportion of the domain is subject to variance due to the oscillation of this mode.

- 2) The expected size of the measurement itself, which we take to be proportional to  $|v_i \hat{\Phi}(x)|$ . This is included because greater measurements mean relatively greater correction

to the state estimate.

- 3) The ratio of the frequency with which that Koopman mode is visited to the frequency of its eigenvalue. This factor is included due to the aliasing effect, and is related to the concept of the Nyquist frequency. Modes with higher frequencies need to be visited more often than modes with lower frequencies.  
5
- 4) A discount factor equal to a decaying exponential of the number of times that eigenmode has been visited. This factor is included due to the exponential convergence of a Kernel Observer. The information gained by taking a measurement within the active area of a particular dynamic mode decreases exponentially the more measurements are taken.

10 **This scoring method allows one to decide the likeliest candidate amongst a large family of paths.**

#### *Generating High-Scoring Random Paths*

**Leveraging the above insights, the following procedure can be utilized for high-scoring paths:**

- 15 1) Cluster the centers into invariant subspaces according to the  $k$ -invariant subspaces algorithm, with the best reasonable guess(es) for  $k$ . This is necessary in order that the robot may be sure it is keeping track of each of the independent subsystems.
- 2) Within each cluster, select waypoints at local maxima of the sum of the Koopman modes. These are where the best measurements may be taken.
- 20 3) Generate a path by selecting waypoints randomly according to a weighting scheme. The weights are equal to the spatial extent of the clusters, multiplied by an exponential decay factor for the number of times each has been visited (see item 4 in the previous section)

## **Empirical Results**

25 We begin with numerical results on determining sensing location and application of Kernel Observers to predict the spatiotemporal functions. The highlight here is results on sensing global ocean temperature with information from just a few location using the AVHRR satellite data. This is followed by application of E-GP in predicting spatiotemporal functions. We include results in prediction weed growth in simulated agricultural fields. Application of E-GP to model  
30 flow past a cylinder in the Reynold's number range 100-1000 are explored in "Learning Fluid Flows with Evolving Gaussian Processes". We conclude the section with some results on control of linear Partial Differential Equation.

## Sensing Locations for Synthetic Data Sets

The goal of this experiment is to investigate the dependency of the observability of system  
5 (3) on the shaded observation matrix and the lower bound presented in Proposition 2. The domain  
is fixed on the interval  $\Omega = [0, 2\pi]$ . First, we pick sets of points  $\mathcal{C}^{(\ell)} = \{c_1, \dots, c_{M_\ell}\}$ ,  $c_j \in \Omega$ ,  
 $M = 50$ , and construct a dynamics matrix  $A = \Lambda \in \mathbb{R}^{M \times M}$ , with cyclic index 5. We pick the  
RBF kernel  $k(x, y) = e^{-\|x-y\|^2/2\sigma^2}$ ,  $\sigma = 0.02$ . Generating samples  $\mathcal{X} = \{x_1, \dots, x_N\}$ ,  $x_i \in \Omega$   
randomly, we compute the  $\ell$ -shaded property and observability for this system. Figure 9a shows  
10 how shadedness is a necessary condition for observability, validating Proposition 1: the slight  
gap between shadedness and observability here can be explained due to numerical issues in  
computing the rank of  $\mathcal{O}_Y$ .

Next, we consider a system with a cyclic index  $\ell = 18$  to verify random sensor placement  
results. We constructed the measurement operator  $K$  using the heuristic in Remark 3 (Algorithm  
15 8), and random sensor selection to generate the sampling locations  $\mathcal{X}$ . These results are presented  
in Figure 9b. The plot for random sampling which has been averaged over 100 runs, resembles a  
c.d.f function of an exponential distribution  $F(X = x) = 1 - \exp(-\lambda x)$ . This verifies the claim  
made in Theorem 3, as the probability of becoming unobservable decays exponentially with the  
number of sensors placed. Also, fitting an exponential distribution we found that the mean  $\lambda^{-1}$   
20 comes close to the ratio  $s/p$ , which is the expected number of randomly placed sensors required  
for observability as per Theorem 2. Note that observability is not achieved if the number of  
samples  $N < \ell$  verifying the result in Proposition 2.

## Comparison With Nonstationary Kernel Methods on Real-World Data

We use three real-world datasets to evaluate and compare the kernel observer with the two  
25 different lines of approach for non-stationary kernels discussed in section on related work. For  
the Process Convolution with Local Smoothing Kernel (PCLSK) and Latent Extension of Input  
Space (LEIS) approaches, we compare with NOSTILL-GP [8] and [39] respectively, on the Intel  
Berkeley, Irish Wind and Ozone data-sets.

Model inference for the kernel observer involved three steps: 1) picking the Gaussian  
30 RBF kernel  $k(x, y) = e^{-\|x-y\|^2/2\sigma^2}$ , a search for the ideal  $\sigma$  is performed for a sparse Gaussian  
Process model (with a fixed basis vector set  $\mathcal{C}$  selected using the method in [53]). For the data  
set discussed in this section, the number of basis vectors were equal to the number of sensing  
locations in the training set, with the domain for input set defined over  $\mathbb{R}^2$ ; 2) having obtained  
 $\sigma$ , Gaussian process inference is used to generate weight vectors for each time-step in the  
training set, resulting in the sequence  $w_\tau, \tau \in \{1, \dots, T\}$ ; 3) matrix least-squares is applied to  
this sequence to infer  $\hat{A}$  (Algorithm 9). For prediction in the autonomous setup,  $\hat{A}$  is used to

propagate the state  $w_\tau$  forward to make predictions with no feedback, and in the observer setup, a Kalman filter (Algorithm 10) with  $N$  determined using Proposition 2, and locations picked 5 randomly, is used to propagate  $w_\tau$  forward to make predictions. We also compare with a baseline GP (denoted by ‘original GP’), which is the sparse GP model trained using all of the available data.

Our first dataset, the Intel Berkeley research lab temperature data, consists of 50 wireless temperature sensors in indoor laboratory region spanning 40.5 meters in length and 31 meters in 10 width (<http://db.csail.mit.edu/labdata/labdata.html>). Training data consists of temperature data on March 6th 2004 at intervals of 20 minutes (beginning 00:20 hrs) which totals to 72 timesteps. Testing is performed over another 72 timesteps beginning 12:20 hrs of the same day. Out of 50 15 locations, we uniformly selected 25 locations each for training and testing purposes. Results of the prediction error are shown in box-plot form in Figure 10a and as a time-series in Figure 10b, note that ‘Auto’ refers to autonomous set up. Here, the cyclic index of  $\hat{A}$  was determined to be 2, so  $N$  was set to 2 for the kernel observer with feedback. Note that here, even the autonomous kernel observer outperforms PCLSK and LEIS overall, and the kernel observer with feedback with  $N = 2$  significantly outperforms all other methods, which is why we did not include results with  $N > 2$ .

20 The second dataset is the Irish wind dataset, consisting of daily average wind speed data collected from year 1961 to 1978 at 12 meteorological stations in the Republic of Ireland (<http://lib.stat.cmu.edu/datasets/wind.desc>). The prediction error is in box-plot form in Figure 11a and as a time-series in Figure 11b. Again, the cyclic index of  $\hat{A}$  was determined to be 2. In this case, the autonomous kernel observer’s performance is comparable to PCLSK and LEIS, while 25 the kernel observer with feedback with  $N = 2$  again outperforms all other methods.

Finally, the Ozone dataset measures ozone concentration (in parts per billion) measured at 60 stations by the United States Environmental Protection Agency [57] across USA. Due to missing measurements, we only selected data from year 1997 to 2013 for training and evaluation. For each station, we averaged ozone concentration over a period of three months, resulting in 30 four quarters per year. Out of 60 sensor locations, we uniformly selected 30 for training and the remaining locations for testing purposes. The prediction error results are presented in box-plot form in Figure 12a and as a time-series in Figure 12b. Here, the cyclic index of  $\hat{A}$  was determined to be 1. In this case, the performance of autonomous kernel observer is comparable to PCLSK and LEIS, with kernel observer with feedback with  $N = 1$  performing the best. Table 1 reports 35 the total training and prediction times associated with PCLSK, LEIS, and the kernel observer. We observed that, 1) the kernel observer is an order of magnitude faster than the competing methods, and 2) even for small sets, competing methods did not scale well.

## Prediction of Global Ocean Surface Temperature

We analyzed the feasibility of our approach on a very large dataset from the National Oceanographic Data Center: the 4 km AVHRR Pathfinder project, which is a satellite monitoring global ocean surface temperature (Figure 13a shows raw satellite data). This dataset is challenging, with measurements at over 37 million possible coordinates, but with only around 3-4 million measurements available per day, leading to a lot of missing data. The goal was to learn the day and night temperature models on data from the year 2011, and then to monitor thereafter for 2012. Success in monitoring would demonstrate two things: 1) the modeling process can capture spatiotemporal trends that generalize across years, and 2) the observer framework allows us to infer the state using a number of measurements that are an order of magnitude fewer than available. Note that due to the size of the dataset and the high computational requirements of the nonstationary kernel methods, a comparison with them was not pursued. To build the autonomous kernel observer and general kernel observer models, we followed the same procedure outlined in Section , but with  $\mathcal{C} = \{c_1, \dots, c_M\}$ ,  $c_j \in \mathbb{R}^2$ ,  $|\mathcal{C}| = 300$ . The Kalman filter for the general kernel observer model used  $N \in \{250, 500, 1000\}$  at random locations to track the system state given a random initial condition  $w_0$ . As a fair baseline, the observers are compared to training a sparse GP model (labeled ‘original’) on approximately 400,000 measurements per day. Figure 13b is an estimate of global ocean surface temperatures obtained using autonomous kernel observer. Figures 13c and 13d compare the autonomous and feedback approach with 1,000 samples to the baseline GP; here, it can be seen that the autonomous does well in the beginning, but then incurs an unacceptable amount of error when the time series goes into 2012, i.e. where the model has not seen any training data, whereas FKO does well throughout. Figures 13e and 13f show a comparison of the RMS error of estimated values from the real data. This figure shows the trend of the observer getting better and better state estimates as a function of the number of sensing locations  $N$ . Note that we checked the performance of training a GP with only 1,000 samples as a control, but the average error was about 10 Kelvins, i.e. much worse than FKO. Time required for kernel observer is much lesser than retraining the model every time step, see Figures 13g-13h.

**Weather Anomaly in 2012:** We further investigated the poor performance of autonomous kernel observer in the year 2012 as observed in Figures 13c and 13d. Clearly, the error in prediction blows up at the start of May in the year 2012. Indicating that the autonomous model trained using the data of the year 2011 does well in capturing the annual weather dynamics up to the month of May 2012. We turned our attention towards the weather in May 2012, as changes in ocean temperatures are directly related to the weather. Surprisingly, severe weather onset was reported on the east coast of United States in May 2012, and this anomaly continued over the period of May to June 2012. Thus, the apparent poor performance of autonomous kernel observer

can actually be a useful indicator in detecting the anomalous behavior, as was the case with the ocean temperature in May 2012 which had deviated from the nominal weather dynamics observed  
5 in the year 2011 in which no severe weather anomalies were reported. Further, we identified the locations at which the prediction error was two standard deviations above the mean error. These error locations are plotted in Figure 14. Error locations on 6-7th May coincide with the severe weather onset on the east coast of United states and that of 28-29th May were along the track of storm Beryl approaching the eastern coastline. This shows that the autonomous kernel observer  
10 model captures the dynamics of spatiotemporal evolution, and has the potential of identifying current behavior as anomalous to that observed in the training set.

### Predicting evolution of weed density in agricultural fields

As a proof-of-concept in applying our methods for learning dynamics and/or inferring the state of spatiotemporally-evolving systems, we consider the problem of predicting weed growth  
15 in agricultural fields.

Past work has presented detailed analysis of weed growth models, in which measurements of seed bank density for various species of weeds were conducted [58]–[61]. In order to generate data upon which to test our methods, we implemented a weed growth simulation model whose rate of seedling emergence agrees with that found in the above research [1]. A Poisson process  
20 is utilized to simulate the temporal evolution of emergence events. This assumption is reasonable over the short time scales in which robots may fully weed a field.

Other work in the field of crop science [62], [63] has shown that the spatial variation in the seed bank density for some species of weeds may be modeled via the Gini Coefficient of Concentration (GCC). This model is accurate for common waterhemp (*Amaranthus tuberculatus*),  
25 and thus we found it useful in our simulation of the spatial distribution of the seed bank density. For more reading. see [1], [64], [65] regarding the relationship between seed bank emergence patterns and environmental conditions such as temperature and moisture.

The weed growth simulation is based on Bernoulli random variables, operating on a matrix of cells, each  $0.8 \text{ m}^2$ , comprising a gridded field of 0.4 hectares, or a cellular automata  
30 model [66]. Seeds emerge from a limited seed bank, forming a binomial distribution over time. The parameters used, summarized in Table 2, are aligned with the growth model for common waterhemp determined in [62]. The initial density of the seed bank in each cell is  $S_0$  on average (between 600 and 1560 seeds per cell). However, at the start of the simulation, the seed bank density in each cell,  $S_0(x, y)$ , is chosen so that Gini Coefficient of Concentration (GCC) between all the cells, used to ensure the relative density of weeds aligns with that seen in real experiments,

is from 0.31 to 0.35, as was determined experimentally in [62]. The field is first divided into fifty patches of weeds, with centers chosen uniformly at random, and sizes from zero to twenty 5 cells in each dimension. Each cell has an initial density between zero and twenty percent of  $S_0$ , chosen uniformly at random. Finally, for each patch of weeds, up to an additional  $S_0$  weeds are added to each cell within each patch, so that the density of each patch follows a normal distribution.

Upon initialization of the simulation, a certain number of days,  $d_0$ , are allowed to elapse 10 before weeding starts. The number of emerging weeds in each cell,  $N_{\text{emerge}}$ , is a randomly generated Poisson variable with mean,  $\lambda(x, y, t)$ , such that 90 percent of the seed bank,  $S(x, y, t)$ , emerges in  $T_{\text{total}}$ , which is two months. This emergence rate is aligned with past work [58]–[61], [65], which all present measurements of the seed bank densities for various species of weeds, and provide an analysis of weed growth models for these species.

15 The weed density in each cell,  $\zeta(x, y, t)$ , grows as seeds emerge from the seed bank. The maximum weed height at each cell,  $\delta(x, y, t)$ , increases from zero height at a fixed rate  $\Gamma$  inches per day.

The main challenge with weeding robots is that they only have access to sparse data about 20 weed density and height, and no data about the seedbank. In fact, the robots can only sense the row that they are in and potentially the adjacent rows on either side. They have no information 25 about the field in locations that have not been visited. Hence, the robots have to try and predict the global weed density given sparse information. This is a great application for the Kernel observer techniques studied in this paper. To demonstrate the proof of concept, we have trained a Kernel observers model on the weed density data generated by our simulations, so that robots in the field can quickly infer the state of the field globally from partial measurements from a few rows.

We used a Radial Basis Function (RBF) kernels with a bandwidth approximately 5 cells (4.5 meters) wide. These were centered throughout the domain using the algorithm in [67]. After training a GP on each snapshot of the data, we obtained a weight vector trajectory. We 30 used linear least squares do determine the best linear transition in the weight space for this trajectory. Below we have included images of system as predicted by feeding forward the initial condition through this linear transition, as a comparison with the original data. The model is able to approximate the weed density growth data very well, with percent error averaging 5%. These results show the feasibility of modeling weed growth using E-GPs. Ongoing work involves 35 massive field campaign to collect the weed density data required to learn these models. Such datasets are currently not available, and the TerraSentia robots discussed in Sidebar: Key control problems in agriculture are being used to collect these data.

## Control of a linear PDE

We then employed kernel controllers for controlling an approximation to the scalar diffusion equation  $u_t = bu_{xx}$  on the domain  $\Omega = [0, 1]$ , with  $b = 0.25$ . The solution to this equation is infinite-dimensional, so we chose a kernel  $k(x, y) = e^{-(\|x-y\|^2/2\sigma^2)}$ , and a set of atoms  $\mathcal{F}^C = \{c_1, \dots, c_M\}$ ,  $c_i \in \Omega$ , with  $M = 25$  generating  $\mathcal{H}^C$ , the space approximating  $\mathcal{H}$ , and another set of atoms  $\mathcal{F}_D = \{\psi(d_1), \dots, \psi(d_{\ell'})\}$ ,  $d_j \in \Omega$ ,  $\ell' = 13$ , generating the control space  $\tilde{\mathcal{H}}$ . The number of, and the location of the observations was chosen to be the same as that of the actuation locations  $d_j$ . First, tests (not reported here) were conducted to ensure that the solution to the diffusion equation is well approximated in  $\mathcal{H}^C$ . Matrix least-squares was used to infer  $\hat{A}$ . Figure 16a shows an example of an initial function  $f_{\text{init}}$  evolving according to the PDE. A reference function  $f_{\text{ref}} \in \mathcal{H}^C$  was chosen to drive  $f_{\text{init}}$  to  $f_{\text{ref}}$  under the action of the PDE. Finally, Algorithm 11 was used to control the PDE, driving  $f_{\text{init}}$  to  $f_{\text{ref}}$ ; Figure 16b shows the absolute value of the error between  $f_k$  and  $f_{\text{ref}}$  as a function of time.

## Paths for Moving Agents

To begin our investigation into the problem of finding best paths for sensing agents in motion, we constructed a number of simple synthetic spatiotemporally evolving systems for the agents to explore. One of these is displayed below 17a. This is a system with two invariant subspaces, in each of which, two hills oscillate up and down in sync at a different frequency than the other invariant subspace. This is ideal for testing our methods. If successful, our methods should be able to (1) divide the space evenly between the two invariant subspaces, (2) be able to evaluate the spatial extent of the subspaces as roughly equivalent, (3) be able to identify the centers of each of the four hills as valuable waypoints, and (4) choose paths that travel between each of the hills equitably. In the example below, we used 100 centers with an RBF kernel with a bandwidth of 0.2 to train our E-GP model, though in testing we could also go as low as 20 centers.

The  $k$ -means algorithm divides the system nicely through the middle, as expected.

We found that the naive method of generating paths purely at random was largely futile, due to the large plain of zero values in-between the two invariant subspaces, which offer little in the way of meaningful data to correct the state estimate. By working with these synthetic data sets, we were able to test the three algorithms described in the previous section and demonstrate that they achieved the four criteria we laid out for them.

To test our methods on challenging, realistic data sets, we used CFD methods to generate the states for a canonical fluid mechanics problem: flow past a cylinder at various Reynolds numbers, namely  $\text{Re} = 100, 300, 600, 800$  and  $1000$ . This deterministic, high-dimensional spatiotemporal dynamical system is well-studied in the fluid dynamics literature, both experimentally and numerically [68]–[70]. In our CFD simulation, we used a 4th-order polynomial expansion with spectral element method on the incompressible Navier-Stokes equation to generate the cylinder flow data. The spatial domain is  $[-2, 10] \times [-3, 3]$ , excluding the diameter-1 cylinder at the origin. Neumann boundary conditions are applied to the far-field of the cylinder in the  $y$ -direction and the outlet of the flow field; and a Dirichlet boundary condition is applied to the inlet. The flow is unstable, periodic, and clearly nonlinear.

For training the E-GP we used 600 centers with an RBF kernel with a bandwidth of 0.4. This output a  $600 \times 600$  matrix  $\hat{A}$  which accurately captures the dynamics of the whole system. (Note that this reduces the size of the model by almost two orders of magnitude from the original CFD data).

Since the CFD set has dynamics almost everywhere, most paths are able to converge to the state eventually, as long as they avoid wandering at the edges where the fluid velocity is semi-constant. However, we were able to demonstrate (as shown in the figures below) that our method of generating paths from valuable waypoints performs statistically much better than either purely random paths or random paths selected by the scoring algorithm.

## Conclusion

Machine learning techniques such as Gaussian Process regression and deep learning are providing increasingly more powerful ways of learning predictive models. However, these techniques are limited by the datasets that they are trained with. Hence, their predictions are not always reliable when predicting real-world complex spatiotemporally-evolving phenomena with lots of variability. Indeed, years worth of weather data cannot be a reliable predictor of weather, nor can data from one farm directly relate to another. Hence, when engineering complex cyber-physical systems such as team of mechanically weeding robots, engineers have to think of ways to supplement the predictions of the machine learning models with real measurements. However, this is not always immediate when one works in the abstract feature spaces utilized in machine learning models. Here we presented Evolving Gaussian Processes, which are formed by embedding a linear dynamical system in the reproducing kernel Hilbert spaces generated by the Gaussian Process model. The parameters of the linear dynamical system can be trained to approximately predict the evolution. In itself this can lead to powerful and generalizable models,

as our results demonstrated in predicting solutions to the Navier-Stokes equations. Furthermore, a Kalman filter can be embedded with the linear dynamical system in the RKHS that can keep  
5 the predictions on track with real sensor measurements. This creates a very powerful prediction system that is capable of predicting nonlinear evolution using very few sensor measurements. We showed that the geometric properties of the linear dynamical system can be utilized to determine sensor numbers and locations. Looking forward, it would be interesting to extend the idea to include nonlinear dynamical systems in the feature space. Such embeddings could improve the capability  
10 of the E-GP model. However, it seems from our results with complex datasets that the feedback with sensors creates a robust monitoring system with the predictions with a simple linear model. The theoretical limits of the tradeoff between improving the model versus increasing the sensing in the context of E-GPs is interesting to study. Finally, also we note that deep neural networks have also been applied to the spatiotemporal modeling problem with significant empirical success  
15 [71], but because these methods are 1) nonlinear in their parameters, and 2) seem to lack the spatial-encoding properties of some types of kernels, how to generalize the analysis considered here for those systems is an interesting open question. Indeed, as sidebar ‘Feature Spaces in Machine Learning’ indicates, there could be very interesting future work that generalizes the idea of embedding controllers and observers in features spaces of machine learning models.

20

## Appendix

### Proofs of Main Theorems

Before we prove Proposition 1, recall that a Jordan decomposition of a matrix  $\widehat{A} \in \mathbb{R}^{M \times M}$  with no repeated eigenvalues can be composed of  $O$  Jordan blocks, where  $O$  can be strictly less than  $M$ . This is in direct contrast to the case where the matrix  $\widehat{A}$  has an eigenvalue decomposition,  
25 in which case no repeated eigenvalues implies that  $O = M$ .

*Proof. (Proposition 1)* To begin, consider a system where  $\widehat{A} = \Lambda$ , with Jordan blocks  $\{\Lambda_1, \Lambda_2, \dots, \Lambda_O\}$  along the diagonal. Then  $\widehat{A}^{\tau_i} = \text{diag}(\begin{bmatrix} \Lambda_1^{\tau_i} & \Lambda_2^{\tau_i} & \dots & \Lambda_O^{\tau_i} \end{bmatrix})$ . We have that

$$\mathcal{O}_Y = \underbrace{\begin{bmatrix} K\widehat{A}^{\tau_1} \\ \vdots \\ K\widehat{A}^{\tau_L} \end{bmatrix}}_{\mathcal{O}_Y \in \mathbb{R}^{NL \times M}}$$

We need to prove that the column rank of  $\mathcal{O}_Y$  is  $M$ , which is not immediately obvious since typically  $N \ll M$ . To prove the statement, we will show that computing the rank of  $\mathcal{O}_Y$  is equivalent to the rank computation of the product of two simple matrices. In what follows, we use the notation  $\mathbf{0}_{\mathbb{R}^{I \times J}}$  to denote an  $I \times J$  matrix of all zeros.

In the first step, we write the above matrix as the product of two matrices. Then it can be shown that  $\mathcal{O}_Y$  is the product of two block matrices

$$\mathcal{O}_Y = \underbrace{\begin{bmatrix} K & \cdots & \mathbf{0}_{\mathbb{R}^{N \times M}} \\ \vdots & \ddots & \vdots \\ \mathbf{0}_{\mathbb{R}^{N \times M}} & \cdots & K \end{bmatrix}}_{\widehat{\mathbf{K}} \in \mathbb{R}^{NL \times ML}} \cdot \underbrace{\begin{bmatrix} \Lambda_1^{\tau_1} & \cdots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \cdots & \Lambda_O^{\tau_1} \\ \hline \vdots & \ddots & \vdots \\ \Lambda_L^{\tau_L} & \cdots & 0 \\ \vdots & \ddots & \vdots \\ 0 & \cdots & \Lambda_O^{\tau_L} \end{bmatrix}}_{\widehat{\mathbf{A}} \in \mathbb{R}^{ML \times M}}.$$

We need to simplify  $\widehat{\mathbf{K}}$  even further. Recall that a matrix's rank is preserved under a product with an invertible matrix. Design a matrix of elementary row operations  $U \in \mathbb{R}^{N \times N}$  such that  $\breve{K} := UK$  is a matrix with at least one row vector of nonzeros; this can be achieved by having an elementary matrix that adds rows together. By the shadedness assumption, such a matrix exists. We can write this operation as

$$UK = \begin{bmatrix} \breve{K}_{11} & \breve{K}_{12} & \cdots & \breve{K}_{1M} \\ \breve{K}_{21} & \breve{K}_{22} & \cdots & \breve{K}_{2M} \\ \vdots & \vdots & \ddots & \vdots \\ \breve{K}_{N1} & \breve{K}_{N2} & \cdots & \breve{K}_{NM} \end{bmatrix}$$

Without loss of generality, and abusing notation slightly, let this multiplication lead to one nonzero row, with the rest of the elements of the matrix being zero, as

$$UK = \begin{bmatrix} k_{11} & k_{12} & \cdots & k_{1M} \\ 0 & 0 & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & 0 \end{bmatrix}.$$

Since elementary matrices are full-rank, we then have that  $\text{rank}(UK) = \text{rank}(K)$ .

To analyze the rank of  $\mathcal{O}_Y$ , we apply these elementary matrices to every  $K \in \widehat{\mathbf{K}}$ . To do so, consider the block-diagonal matrix  $\mathcal{U} \in \mathbb{R}^{NL \times NL}$  with  $U \in \mathbb{R}^{N \times N}$  along the diagonal, and zeros everywhere else, i.e.

$$\mathcal{U} := \begin{bmatrix} U & \mathbf{0}_{\mathbb{R}^{N \times N}} & \cdots & \mathbf{0}_{\mathbb{R}^{N \times N}} \\ \mathbf{0}_{\mathbb{R}^{N \times N}} & U & \cdots & \mathbf{0}_{\mathbb{R}^{N \times N}} \\ \vdots & \vdots & \ddots & \vdots \\ \mathbf{0}_{\mathbb{R}^{N \times N}} & \mathbf{0}_{\mathbb{R}^{N \times N}} & \cdots & U \end{bmatrix}. \quad (15)$$

It can be shown that  $\mathcal{U}$  is full-rank, i.e. has rank  $NL$ . Going back to the observability matrix, we have that

$$\begin{aligned} \mathcal{U}\mathcal{O}_Y &= \mathcal{U}\widehat{\mathbf{K}}\widehat{\mathbf{A}} \\ &= \underbrace{\begin{bmatrix} UK & \mathbf{0}_{\mathbb{R}^{N \times M}} & \cdots & \mathbf{0}_{\mathbb{R}^{N \times M}} \\ \mathbf{0}_{\mathbb{R}^{N \times M}} & UK & \cdots & \mathbf{0}_{\mathbb{R}^{N \times M}} \\ \vdots & \vdots & \ddots & \vdots \\ \mathbf{0}_{\mathbb{R}^{N \times M}} & \mathbf{0}_{\mathbb{R}^{N \times M}} & \cdots & UK \end{bmatrix}}_{\mathcal{U}\widehat{\mathbf{K}} \in \mathbb{R}^{NL \times ML}} \underbrace{\widehat{\mathbf{A}}}_{\in \mathbb{R}^{ML \times M}}, \end{aligned}$$

since  $\mathbf{0}_{\mathbb{R}^{N \times N}}\mathbf{0}_{\mathbb{R}^{N \times M}} = \mathbf{0}_{\mathbb{R}^{N \times M}}$ . Due to the fact that  $\text{rank}(\mathcal{U}\mathcal{O}_Y) = \text{rank}(\mathcal{O}_Y)$ , we can therefore perform our rank analysis on the simpler matrix  $\text{rank}(\mathcal{U}\mathcal{O}_Y)$ . Note that

$$\begin{aligned} UK\widehat{A}^{\tau_j} &= \begin{bmatrix} k_{11} & k_{12} & \cdots & k_{1M} \\ 0 & 0 & \cdots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \cdots & 0 \end{bmatrix} \widehat{A}^{\tau_j} \\ &= \begin{bmatrix} k_{11}\lambda_1^{\tau_j} & \binom{\tau_j}{1}\lambda_1^{\tau_j-1} + k_{12}\lambda_1^{\tau_j} & \cdots & k_{1M}\lambda_O^{\tau_j} \\ 0 & 0 & \cdots & 0 \\ \vdots & \vdots & \ddots & 0 \\ 0 & 0 & \cdots & 0 \end{bmatrix}. \end{aligned}$$

Therefore, following some more elementary row operations encoded by  $V \in \mathbb{R}^{ML \times ML}$ , we have that

$$\begin{aligned} V\mathcal{U}\mathcal{O}_Y &= \begin{bmatrix} k_{11}\lambda_1^{\tau_1} & \cdots & k_{1M}\lambda_O^{\tau_1} \\ k_{11}\lambda_1^{\tau_2} & \cdots & k_{1M}\lambda_O^{\tau_2} \\ \vdots & \ddots & 0 \\ k_{11}\lambda_1^{\tau_L} & \cdots & k_{1M}\lambda_O^{\tau_L} \\ \mathbf{0}_{\mathbb{R}^{M(L-1) \times 1}} & \cdots & \mathbf{0}_{\mathbb{R}^{M(L-1) \times 1}} \end{bmatrix} \\ &= \begin{bmatrix} \Phi \\ \mathbf{0}_{\mathbb{R}^{M(L-1) \times M}} \end{bmatrix}. \end{aligned}$$

If the individual entries  $k_{1i}$  are nonzero, and the Jordan block diagonals have nonzero eigenvalues, the columns of  $\Phi$  become linearly independent. Therefore, if  $L \geq M$ , the column rank of  $\mathcal{O}_Y$  is  $M$ , which results in an observable system.

To extend this proof to matrices  $\widehat{A} = P\Lambda P^{-1}$ , note that

$$\begin{aligned}\mathcal{O}_Y &= \begin{bmatrix} K\widehat{A}^{\tau_1} \\ \vdots \\ K\widehat{A}^{\tau_L} \end{bmatrix} \\ &= \begin{bmatrix} KP\Lambda^{\tau_1}P^{-1} \\ \vdots \\ KP\Lambda^{\tau_L}P^{-1} \end{bmatrix} \\ &= \widehat{\mathbf{K}}\mathbf{P}\Lambda^t\mathbf{P}^{-1},\end{aligned}$$

where  $\mathbf{P} \in \mathbb{R}^{ML \times ML}$ ,  $\Lambda^t \in \mathbb{R}^{ML \times ML}$ , and  $\mathbf{P}^{-1} \in \mathbb{R}^{ML \times ML}$  are the block diagonal matrices associated with the system. Since  $\mathbf{P}$  is an invertible matrix, the conclusions about the column rank drawn before still hold, and the system is observable.  $\square$

*Proof. (Proposition 2) By Contrapositive.* We will show that if the number of sampling locations are  $N = \ell - 1$  (i.e.  $N < \ell$ ), then the system is not observable. Pick the Gaussian kernel in the dictionary of atoms framework, with sampling locations  $x_i \in \mathcal{X}$  and centers  $c_j \in \mathcal{C}$ , with the additional property that  $x_i \neq x_j \forall i, j \in \{1, \dots, N\}, i \neq j$ . In this case,  $\mathbf{K}$  has  $\ell - 1$  nonzero, linearly independent rows, and can be written as

$$\mathbf{K} = \begin{bmatrix} k_{11} & k_{12} & \cdots & k_{1M} \\ \vdots & \vdots & \cdots & \vdots \\ k_{(\ell-1)1} & k_{(\ell-1)2} & \cdots & k_{(\ell-1)M} \end{bmatrix}.$$

Since the cyclic index is  $\ell$ , this implies that at least one eigenvalue, say  $\lambda$ , has  $\ell$  Jordan blocks. Define indices  $j_1, j_2, \dots, j_\ell \in \{1, 2, \dots, M\}$  as the columns corresponding to the leading entries of the  $\ell$  Jordan blocks corresponding to  $\lambda$ . WLOG, let  $j_1 = 1$ . Using ideas similar to the last proof, we can write the observability matrix as

$$\mathcal{O}_Y := \begin{bmatrix} k_{11}\lambda^{\tau_1} & \cdots & k_{1j_\ell}\lambda^{\tau_1} & \cdots \\ \vdots & \ddots & \vdots & \ddots \\ k_{11}\lambda^{\tau_L} & k_{1j_\ell}\lambda^{\tau_L} & \cdots & \cdots \\ \vdots & \ddots & \vdots & \ddots \\ k_{(\ell-1)1}\lambda^{\tau_1} & \cdots & k_{(\ell-1)j_\ell}\lambda^{\tau_1} & \cdots \\ \vdots & \ddots & \vdots & \ddots \\ k_{(\ell-1)1}\lambda^{\tau_L} & \cdots & k_{(\ell-1)j_\ell}\lambda^{\tau_L} & \cdots \end{bmatrix}.$$

Define  $\boldsymbol{\lambda} := [\lambda^{\tau_1} \ \lambda^{\tau_2} \ \dots \ \lambda^{\tau_L}]^T$ . Then the above matrix becomes

$$\mathcal{O}_\Upsilon := \begin{bmatrix} k_{11}\boldsymbol{\lambda} & \cdots & k_{1j_2}\boldsymbol{\lambda} & \cdots & k_{1j_\ell}\boldsymbol{\lambda} & \cdots \\ \vdots & \ddots & \vdots & \ddots & \vdots & \ddots \\ k_{(\ell-1)1}\boldsymbol{\lambda} & \cdots & k_{(\ell-1)j_2}\boldsymbol{\lambda} & \cdots & k_{(\ell-1)j_\ell}\boldsymbol{\lambda} & \cdots \end{bmatrix}.$$

We need to show that one of the columns above can be written in terms of the others. This is equivalent to solving the linear system

$$\begin{bmatrix} k_{1j_1} \\ k_{2j_1} \\ \vdots \\ k_{(\ell-1)j_1} \end{bmatrix} = \begin{bmatrix} k_{1j_2} & \cdots & k_{1j_\ell} \\ k_{2j_2} & \cdots & k_{2j_\ell} \\ \vdots & \ddots & \vdots \\ k_{(\ell-1)j_2} & \cdots & k_{(\ell-1)j_\ell} \end{bmatrix} \begin{bmatrix} c_1 \\ c_2 \\ \vdots \\ c_{(\ell-1)} \end{bmatrix}.$$

Since the kernel matrix on the RHS is generated from the Gaussian kernel, from [72], it's known that every principal minor of a Gaussian kernel matrix is invertible, which implies that  $\mathcal{O}_\Upsilon$  cannot be observable.  $\square$

*Proof. (Proposition 3)* The construction of the measurement map  $\tilde{K}$  is based on the rational canonical structure of  $\hat{A}^T$ , which decomposes  $\mathcal{V}$  into  $\hat{A}^T$ -cyclic direct summands such that  $\mathcal{V} = \mathcal{V}_1 \oplus \dots \oplus \mathcal{V}_\ell$ , where  $\ell$  is the cyclic index of  $\hat{A}$ . Let  $\xi_v$  be the minimal polynomial (m.p.) of  $v$  (relative to  $\hat{A}^T$ ): it is then the unique monic polynomial of least degree such that  $\xi_v(\hat{A}^T)v = 0$ .

Let  $\alpha_1(\lambda)$  be the m.p. of  $\hat{A}_{|\mathcal{V}_1}^T$ : then  $\deg(\alpha_1(\lambda)) < M$ . By the rational canonical structure theorem [52], there exists a vector  $\hat{v}_1$ , such that  $\xi_{v_1}(\lambda) = \alpha_1(\lambda)$ . Similarly there exists a vector  $\hat{v}_2$ , such that  $\xi_{v_2}(\lambda) = \alpha_2(\lambda)$ , where  $\alpha_2(\lambda)$ , is the minimal polynomial of  $\hat{A}_{|\mathcal{V}_2}^T$  and so on. Thus we can obtain  $\ell$  such vectors that form the measurement map  $\tilde{K} = [\hat{v}_1, \hat{v}_2, \dots, \hat{v}_\ell]^T$ . Construction of these vectors  $\hat{v}_i$ , can be simplified by first performing the Jordan decomposition as  $\hat{A}^T = P\Lambda P^{-1}$ .

Then the vectors  $\tilde{v}_i$ ,  $i \in \{1, \dots, \ell\}$  for  $\Lambda$ , can be constructed such that the entries corresponding to the leading entries of Jordan blocks of  $\Lambda_{|\mathcal{V}_i}$  are nonzero. Such a construction ensures that the m.p. of vector  $\tilde{v}_i$  w.r.t  $\Lambda_{|\mathcal{V}_i}$ , is also the corresponding m.p. of  $\Lambda_{|\mathcal{V}_i}$ . Finally, the required map can be obtained as  $\tilde{K} = [\tilde{v}_1, \tilde{v}_2, \dots, \tilde{v}_\ell]^T P^{-1}$ .  $\square$

*Proof. (Theorem 1)* A cyclic index of  $\ell$  for this system implies that there exists an eigenvalue  $\lambda$  that's repeated  $\ell$  times. We prove the theorem for repeated eigenvalues of dimension 1: the same statement can be proven for repeated eigenvalues for Jordan blocks using the ideas in the proof of Proposition 1. WLOG, let  $\mathbf{K}$  have  $\ell$  fully shaded, linearly independent rows, and,

assume that the column indices corresponding to this eigenvalue are  $\{1, 2, \dots, \ell\}$ . Define  $\boldsymbol{\lambda}_i := \begin{bmatrix} \lambda_i^{\tau_1} & \lambda_i^{\tau_2} & \cdots & \lambda_i^{\tau_L} \end{bmatrix}^T$ . Then

$$\mathcal{O}_Y := \begin{bmatrix} k_{11}\boldsymbol{\lambda}_1 & k_{12}\boldsymbol{\lambda}_2 & \cdots & k_{1M}\boldsymbol{\lambda}_M \\ \vdots & \vdots & \ddots & \vdots \\ k_{\ell 1}\boldsymbol{\lambda}_1 & k_{\ell 2}\boldsymbol{\lambda}_2 & \cdots & k_{\ell M}\boldsymbol{\lambda}_M \end{bmatrix}.$$

Let  $\boldsymbol{\lambda}_1 = \boldsymbol{\lambda}_2 = \cdots = \boldsymbol{\lambda}_\ell := \boldsymbol{\lambda}$ . Focusing on these first  $\ell$  columns of this matrix, this implies that we need to find constants  $c_1, c_2, \dots, c_{\ell-1}$  such that

$$\begin{bmatrix} k_{11} \\ \vdots \\ k_{\ell 1} \end{bmatrix} = c_1 \begin{bmatrix} k_{12} \\ \vdots \\ k_{\ell 2} \end{bmatrix} + \cdots + c_{\ell-1} \begin{bmatrix} k_{1\ell} \\ \vdots \\ k_{\ell\ell} \end{bmatrix}.$$

However, these columns are linearly independent by assumption, and thus no such constants exist, implying that  $\mathcal{O}_Y$  is observable.  $\square$

<sup>5</sup> While Theorem 1 is a quite general result, the condition that any  $\ell$  columns of  $\mathbf{K}$  be linearly independent is a very stringent condition. One scenario where this condition can be met with minimal measurements is in the case when the feature map  $\widehat{\psi}(x)$  is generated by a dictionary of atoms with the Gaussian RBF kernel evaluated at sampling locations  $\{x_1, \dots, x_N\}$  according to (2), where  $x_i \in \Omega \subset \mathbb{R}^d$ , and  $x_i$  are sampled from a non-degenerate probability distribution on  $\Omega$  such as the uniform distribution. For a semi-deterministic approach, when the dynamics matrix  $\widehat{A}$  is block-diagonal, we can utilize a simple heuristic:

<sup>10</sup> **Remark 3.** Let  $\Omega$  be compact,  $\mathcal{C} = \{c_1, \dots, c_M\}$ ,  $c_i \in \Omega$ , and let the approximate feature map be defined by (2). Consider the system (3) with  $\widehat{A} = \Lambda$ , and let  $\Upsilon = \{0, 1, \dots, M-1\}$ . Then the measurement map  $\widetilde{K}$ 's values lie in  $\{0, 1\}$ ; in particular, each row  $\widetilde{K}^{(j)}$ ,  $j \in \{1, \dots, \ell\}$ , corresponds to a subspace  $\widehat{\mathcal{H}}_j$ , generated by a subset of centers  $\mathcal{C}^{(j)} \subset \mathcal{C}$ . Generate samples  $x_i^{(j)}$  to create a kernel matrix  $K^{(j)}$  that is shaded only with respect to centers  $\mathcal{C}^{(j)}$ . Once this is done, move on to the next subspace  $\widehat{\mathcal{H}}_{j+1}$ . When all  $\ell$  rows of  $\widetilde{K}$  are accounted for, construct the matrix  $\mathbf{K}$  as in (7). Then the resulting system  $(\mathbf{K}, \widehat{A})$  is observable.

<sup>15</sup> This heuristic is formalized in Algorithm 2. Note that in practice, the matrix  $\widehat{A}$  needs to be inferred from measurements of the process  $f_\tau$ . If no assumptions are placed on  $\widehat{A}$ , it's clear that at least  $M$  sensors are required for the system identification phase. Future work will study the precise conditions under which system identification is possible with less than  $M$  sensors.

*Proof.* (**Theorem 2**) For each random sample, the probability that it lies within the  $\epsilon$ -shaded region of a particular center  $c_j \in \mathcal{C}_\zeta$  is at least  $p_\epsilon$ . The series of random samples can be considered

as Bernoulli trials in which  $p_\epsilon$  is the probability of a successful outcome. Note this is assuming worst case scenario that the intersection between any two  $\epsilon$ -shaded region of centers belonging  
<sup>5</sup> to the set  $\mathcal{C}_\varsigma$  is empty. Observability for the pair  $(K, \widehat{A})$  is achieved after  $\varsigma$  successful outcomes are obtained because each success ensures a row vector with non-zero entry corresponding to the leading entry of the Jordan block.

Let  $X_1, X_2, \dots, X_N$  be i.i.d. random variables whose common distribution is the Bernoulli distribution with parameter  $p_\epsilon$ . The random variable  $X = X_1 + X_2 + \dots + X_N$  denotes the number of success after  $N$  random samples. Since each  $X_i$  has the Bernoulli distribution,  $X$  will have binomial distribution,

$$P(X = h) = \binom{N}{h} p_\epsilon^h (1 - p_\epsilon)^{N-h},$$

in which  $h$  is the number of success. The expectation of the binomial distribution, that is the expected number of success is  $Np_\epsilon$ , and thus the expected number of trials required will be  
<sup>10</sup>  $N = \varsigma/p_\epsilon$ .  $\square$

*Proof.* (**Theorem 3**) The random variable  $X$  from the proof of Theorem 1 has a binomial distribution, which enables the application of a Chernoff-type bound on its tail probabilities. A well known result on multiplicative Chernoff bound [73] is directly applied to establish this Theorem. If  $X$  is binomially distributed,  $\delta \in (0, 1]$ , and  $\mu = \mathbb{E}[X]$ , then  $P[X \leq (1 - \delta)\mu] \leq \exp(-\mu\delta^2/2)$ , in which we let  $\delta = 1 - \frac{\varsigma}{Np_\epsilon}$ . The expression in the exponent can be simplified to  $-\frac{1}{2}Np_\epsilon + \varsigma - \frac{\varsigma^2}{2Np_\epsilon}$ , using  $\mu = Np_\epsilon$ . Note that  $e^{\frac{-\varsigma^2}{2Np_\epsilon}} \leq 1$ . This implies that,

$$\exp(-\mu\delta^2/2) = e^{-\frac{1}{2}(Np_\epsilon - 2\varsigma)} \cdot e^{\frac{-\varsigma^2}{2Np_\epsilon}} \leq e^{-\frac{1}{2}(Np_\epsilon - 2\varsigma)}$$

Note,  $(1 - \delta)\mu = \varsigma$ , hence we obtain that  $P[X \leq \varsigma] \leq e^{-\frac{1}{2}(Np_\epsilon - 2\varsigma)}$ .  $\square$

## Software

To facilitate implementation of the material presented in this paper we have made an open-source code-base. A MATLAB version is available at <http://daslab.illinois.edu/software.html> or at  
<sup>15</sup> <https://github.com/hkingravi/FunctionObservers> with examples and documentation. In addition, a Python version is available on GitHub at <https://github.com/hkingravi/funcobspy>.

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Table 1: Total training and prediction times for Figs. 10 and 11

	<b>Intel</b> Berkeley	<b>Irish</b> Wind	<b>Ozone</b>
<i>Data Size (bases-timesteps)</i>	25-72	12-36	30-68
Kernel Observer	2.1 sec	0.1 sec	1.61 sec
PCLSK	121.4 sec	7.0 sec	91.90 sec
LEIS	43.8 sec	2.8 sec	37.41 sec

Table 2: Seed Bank Density Parameters

<b>Parameter</b>	GCC	$S_0$ (seeds/cell)	Num. of Patches	Patch Size (Cells in X and Y)
<b>Range</b>	[0.31,0.35]	[600,1560]	50	[0,20]

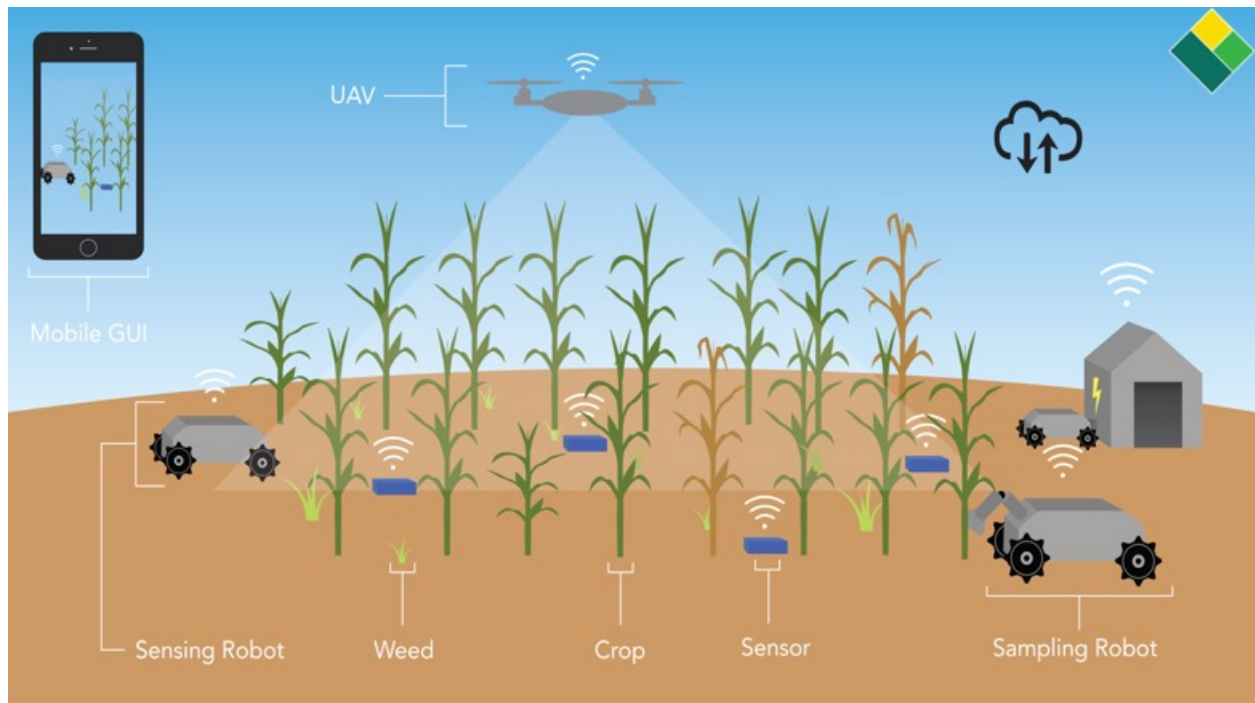


Figure 1: A Cyber Physical system consisting of a distributed team of robots for mechanical weed management on a farm. Image courtesy EarthSense inc.

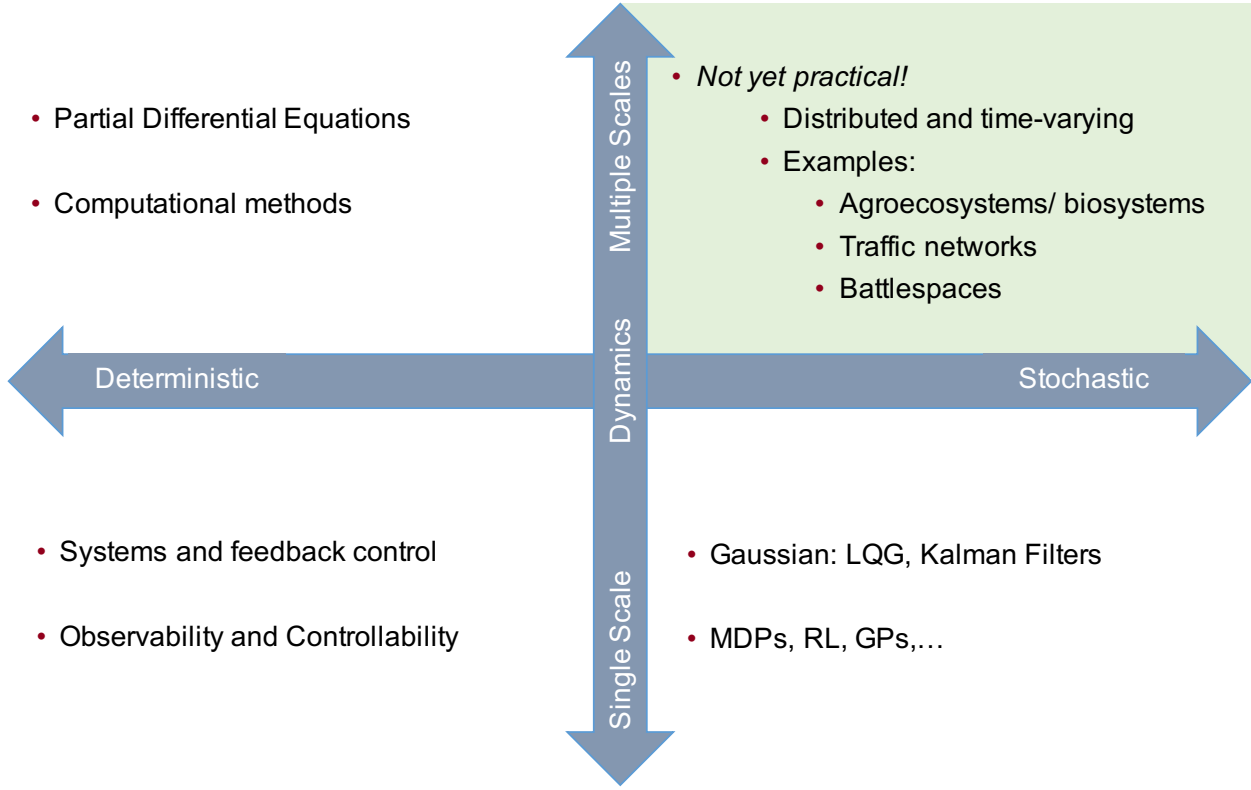
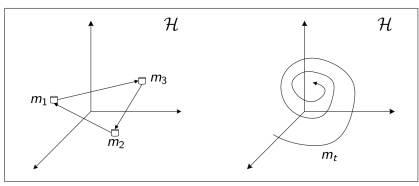
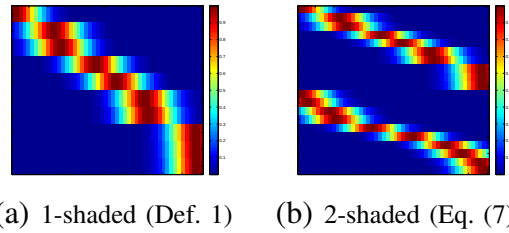


Figure 2: Modeling, monitoring, and controlling dynamical systems with complex and uncertain dynamics, such as agricultural, traffic, or weather monitoring systems, presents exciting open challenge for the controls community. The bottom left quadrant describes linear and time-invariant systems with single- scale dynamics, for which the theory of feedback control of dynamical systems is often sufficient. The bottom right quadrant shows stochastic single-scaled systems, where approaches such as Kalman Filters and Gaussian optimization have marked several successes of control systems enabling endeavors from Lunar landings to GPS navigation. The top left quadrant denotes systems with dynamics at multiple scales, where efficient computational solutions to Partial Differential Equations (PDEs) is an highly active area of research. However, fundamental theoretical advances and practical algorithms are needed to enable autonomous decision making for distributed stochastic Cyber Physical Systems with dynamics at multiple scales shown in the top right quadrant such as distributed agricultural robotic systems, traffic networks, and weather monitoring systems with mobile and stationary sensors.



**Figure 3:** Two types of Hilbert space evolutions. Left: discrete switches in RKHS  $\mathcal{H}$ ; Right: smooth evolution in  $\mathcal{H}$ .



**Figure 4:** Shaded observation matrices for dictionary of atoms. Each row represents a sensing location with the color map indicating the evaluation of kernel function w.r.t the others points in the domain.

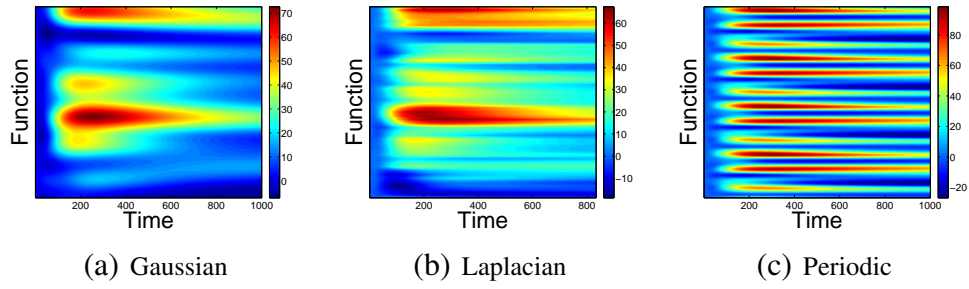


Figure 5: One-dimensional function evolution over a fixed transition matrix  $A$ , initial condition  $w_0$  and centers  $\mathcal{C}$ , but with different kernels  $k(x, y)$ . Each  $y$ -vector at a given value of  $x$  represents the output of the function, which evolves from left to right. As seen, changing the kernel creates quite different dynamic behaviors.

$$\begin{array}{ccc}
\widehat{\mathcal{H}} & \xrightarrow{\mathcal{A}} & \widehat{\mathcal{H}} \\
\downarrow \mathcal{W} & & \uparrow \mathcal{W}^{-1} \\
\mathbb{R}^M & \xrightarrow{A} & \mathbb{R}^M
\end{array}
\qquad
\begin{array}{ccc}
\widehat{\mathcal{H}} & \xrightarrow{\mathcal{K}} & \mathbb{R}^N \\
\downarrow \mathcal{W} & \nearrow \mathcal{W}^{-1} & \searrow K \\
\mathbb{R}^M & &
\end{array}$$

(a) Relationship between  $\mathcal{A}$  and  $A$  (b) Relationship between  $\mathcal{K}$  and  $K$

$$\begin{array}{ccc}
\widetilde{\mathcal{H}} & \xrightarrow{\mathcal{B}} & \widehat{\mathcal{H}} \\
\downarrow \mathcal{W} & & \uparrow \mathcal{W}^{-1} \\
\mathbb{R}^{\ell'} & \xrightarrow{B} & \mathbb{R}^M
\end{array}$$

(c) Relationship between  $\mathcal{B}$  and  $B$

Figure 6: Commutative diagrams between primal and dual spaces

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**Algorithm 1** Measurement Map  $\tilde{K}$ 

---

**Input:**  $\hat{A} \in \mathbb{R}^{M \times M}$

Compute Rational Canonical Form, s.t.  $C = Q^{-1}\hat{A}^TQ$ . Set  $C_0 := C$ , and  $M_0 := M$ .

**for**  $i = 1$  **to**  $\ell$  **do**

    Obtain MP  $\alpha_i(\lambda)$  of  $C_{i-1}$ . This returns associated indices  $\mathcal{J}^{(i)} \subset \{1, 2, \dots, M_{i-1}\}$ .

    Construct vector  $v_i \in \mathbb{R}^M$  such that  $\xi_{v_i}(\lambda) = \alpha_i(\lambda)$ .

    Use indices  $\{1, 2, \dots, M_{i-1}\} \setminus \mathcal{J}^{(i)}$  to select matrix  $C_i$ . Set  $M_i := |\{1, 2, \dots, M_{i-1}\} \setminus \mathcal{J}^{(i)}|$

**end for**

Compute  $\mathring{K} = [v_1^T, v_2^T, \dots, v_\ell^T]^T$

**Output:**  $\tilde{K} = \mathring{K}Q^{-1}$

---

---

**Algorithm 2** Sampling locations set  $\mathcal{X}$ 

---

**Input:**  $\widehat{A} = C$ , lower bound  $\ell$

Decompose  $C$  to generate invariant subspaces  $\widehat{\mathcal{H}}_j$ ,  $j \in \{1, 2, \dots, \ell\}$  (see section )

**for**  $j = 1$  **to**  $\ell$  **do**

    Obtain centers  $\mathcal{C}^{(j)}$  w.r.t subspace  $\widehat{\mathcal{H}}_j$ ,

    Generate samples  $x_i^{(j)}$  to create a kernel matrix  $K^{(j)}$  that is shaded only with respect to centers  $\mathcal{C}^{(j)}$

**end for**

**Output:** Sampling locations set  $\mathcal{X} = \{x^{(1)}, x^{(2)} \dots, x^{(\ell)}\}$ .

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**Algorithm 3** Kernel Observer (Transition Learning)

---

**Input:** Kernel  $k$ , basis centers  $\mathcal{C}$ , final time step  $T$ .

**while**  $\tau \leq T$  **do**

- 1) Sample data  $\{y_\tau^i\}_{i=1}^M$  from  $f_\tau$ .
- 2) Estimate  $\hat{w}_\tau$  via standard kernel inference procedure.
- 3) Store weights  $\hat{w}_\tau$  in matrix  $\mathcal{W} \in \mathbb{R}^{M \times T}$ .

**end while**

To infer  $\hat{A}$ , define matrix  $\Phi = \mathcal{W}^T \mathcal{W}$ . Then:

**for**  $i = 1$  **to**  $M$  **do**

At step  $i$ , solve system

$$\hat{A}^{(i)} = ((\Phi + \lambda I)^{-1} (\mathcal{W}^T \mathcal{W}^{(i)}))^T, \quad (16)$$

where  $\hat{A}^{(i)}$ , and  $\mathcal{W}^{(i)}$  are the  $i$ th columns of  $\hat{A}$  and  $\mathcal{W}$  respectively.

**end for**

Compute the covariance matrix  $\hat{B}$  of the observed weights  $\mathcal{W}$ .

**Output:** estimated transition matrix  $\hat{A}$ , predictive covariance matrix  $\hat{B}$ .

---

---

**Algorithm 4** Kernel Observer (Monitoring and Prediction)

---

**Input:** Kernel  $k$ , basis centers  $\mathcal{C}$ , estimated system matrix  $\hat{A}$ , estimated covariance matrix  $\hat{B}$ .  
**Compute Observation Matrix:** Compute the cyclic index  $\ell$  of  $\hat{A}$ , and compute  $K$ .  
**Initialize Observer:** Use  $\hat{A}$ ,  $\hat{B}$ , and  $K$  to initialize a state-observer (e.g. Kalman filter (KF)) on  $\hat{\mathcal{H}}$ .  
**while** measurements available **do**  
    1) Sample data  $\{y_\tau^i\}_{i=1}^N$  from  $f_\tau$ .  
    2) Propagate KF estimate  $\hat{w}_\tau$  forward to time  $\tau + 1$ , correct using measurement feedback with  $\{y_{\tau+1}^i\}_{i=1}^N$ .  
    3) Output predicted function  $\hat{f}_{\tau+1}$  of KF.  
**end while**

---

---

**Algorithm 5** Kernel Controller

---

**Input:** Kernel  $k$ , basis points  $\mathcal{C}$ , estimated system matrix  $\hat{A}$ , estimated covariance matrix  $\hat{B}$ , and function  $f_{\text{ref}}$  to drive initial function to.

**Initialize Observer:** (see Algorithm 10).

**Initialize Controller:** Use Jordan decomposition of  $\hat{A}$  to obtain no. of control locations  $\mathcal{D}$ , compute kernel matrix  $K_{CD} \in \mathbb{R}^{\ell \times M}$  between  $\mathcal{D}$  and  $\mathcal{C}$ , and initialize controller (e.g. LQR) utilizing  $(\hat{A}, \hat{B})$ .

**while** measurements available **do**

- 1) Sample data  $\{y_k^i\}_{i=1}^N$  from  $f(x, \tau)$ .
- 2) Utilize observer to estimate  $\hat{w}_{\tau+1}$ .
- 3) Use  $\hat{w}_{\tau+1}$  and  $f_{\text{ref}}$  as input to controller to get feedback.

**end while**

---

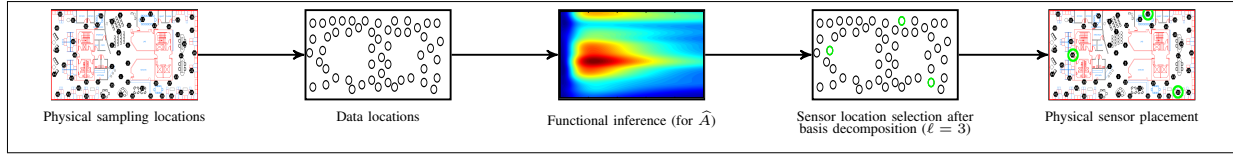


Figure 7: Overall description of how the kernel observer fits in the sensing framework. Physical locations are mapped to data locations, over which historical data is collected as a time series. Functional inference is performed over  $\hat{\mathcal{H}}$  to solve for  $\hat{A}$ . The measurement operator  $K$  is then computed (see Figure 8), leading to sensor placement.

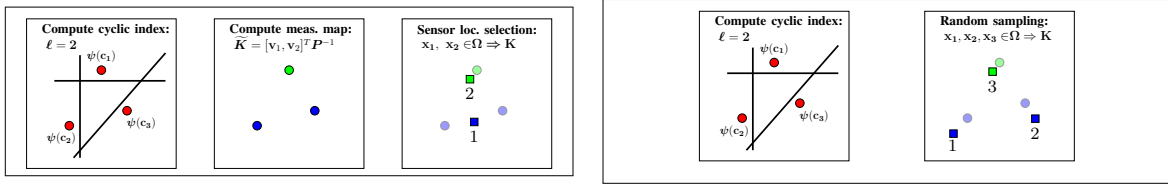


Figure 8: Diagram demonstrating sensor placement using the measurement map or random sampling approaches. The circles represent data locations associated to bases (e.g.  $c_j \Leftrightarrow \psi(c_j)$ ) and the squares represent sensor locations (e.g.  $x_i \Leftrightarrow \psi(x_i)$ ). The cyclic index ( $\ell = 2$ ) indicates how many possible couplings of bases exist, which can be represented as a choice of  $\binom{M}{\ell}$  hyperplanes in  $\Omega$ . If the measurement map is computed (left), the correct couplings are chosen (green vs. blue), and a smaller number of sensors (2) can be placed. Alternatively, random sampling (right) is more computationally efficient, but generally requires more sensors (3).

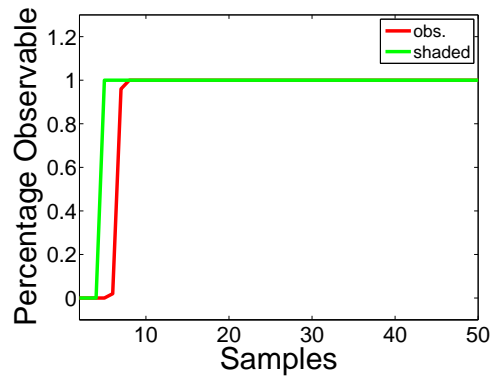
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**Algorithm 6** *k*-Invariant Subspaces Algorithm

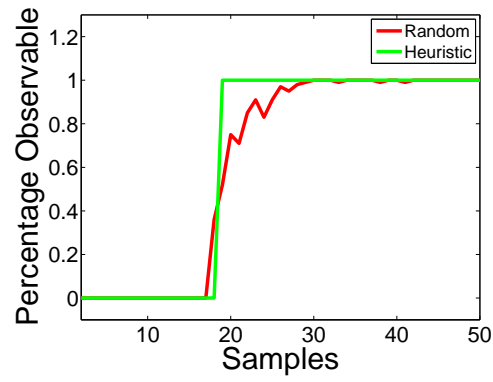
---

```
while clusters have changed do
  for each center  $i$  do
    Find cluster  $k$  which maximizes the score  $\frac{1}{|S_k|} (\|\bar{A}_{i,S_k \setminus \{i\}}\|^2 + \|\bar{A}_{S_k \setminus \{i\}, i}\|^2)$ 
    Reassign center  $i$  to cluster with highest score
  end for
end while
return clusters
```

---



(a) Shaded vs. observability



(b) Heuristic vs. random

Figure 9: Kernel observability results.

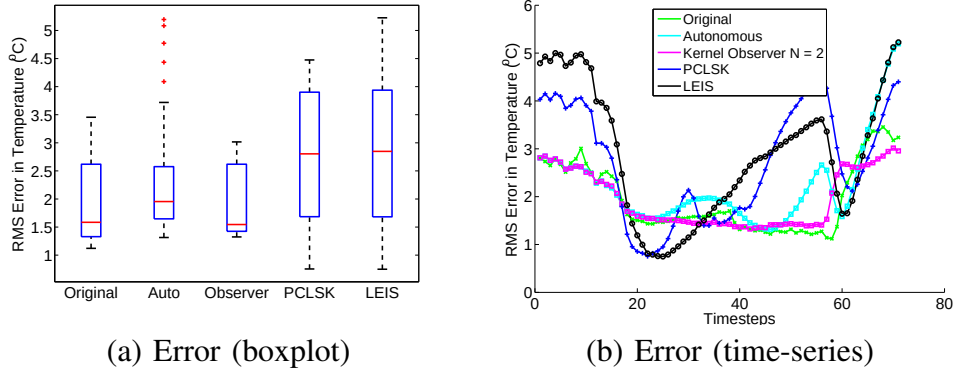


Figure 10: Comparison of kernel observer to PCLSK and LEIS methods on Intel Berkeley dataset.

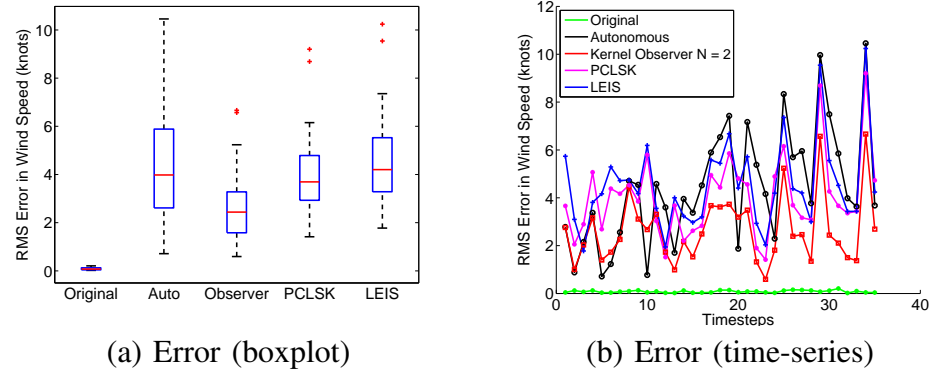


Figure 11: Comparison of kernel observer to PCLSK and LEIS methods on Irish Wind dataset.

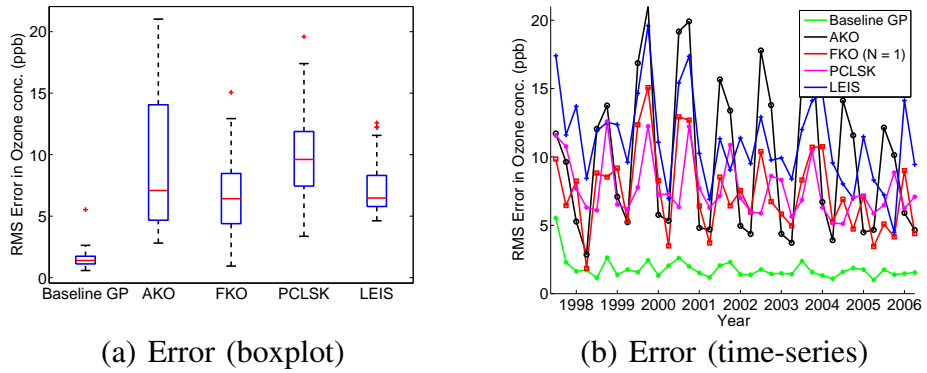
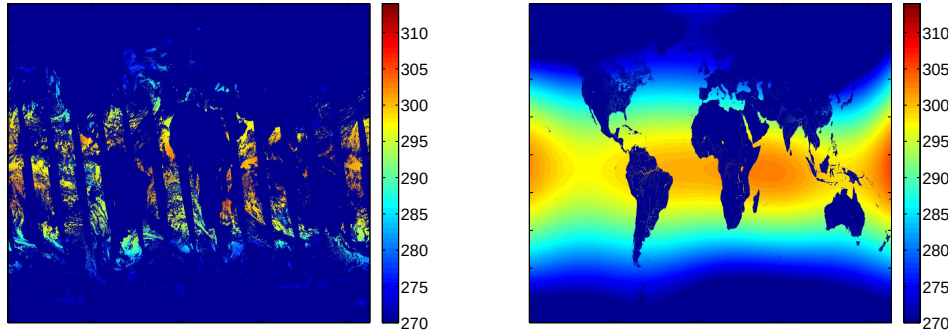
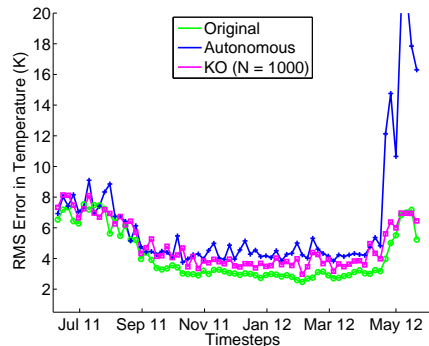


Figure 12: Comparison of kernel observer to PCLSK and LEIS methods on Irish Wind dataset.

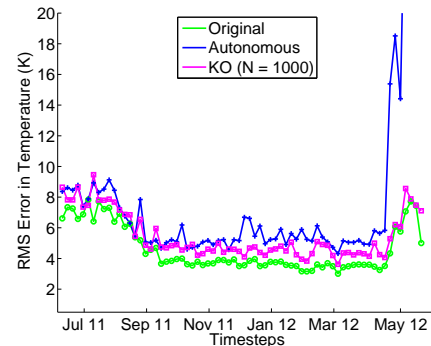


(a) Raw Satellite Data

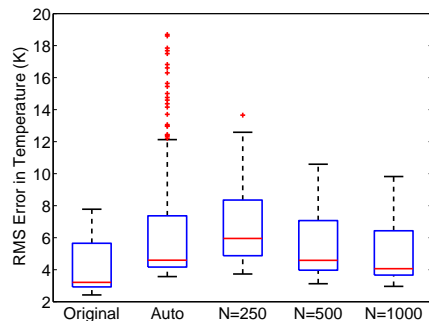
(b) AKO estimate



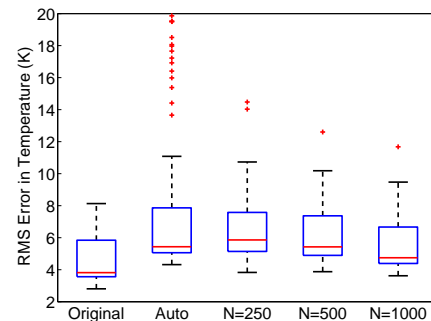
(c) Error-day (time-series)



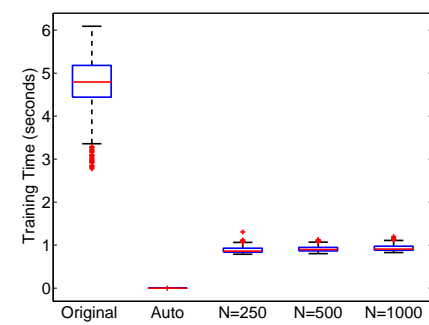
(d) Error-night (time-series)



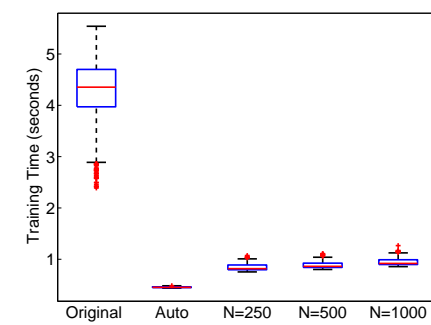
(e) Error-day (box-plot)



(f) Error-night (box-plot)



(g) Estimation time (day)



(h) Estimation time (night)

Figure 13: Performance of the kernel observer over AVVHR satellite 2012 data with different numbers of observation locations.

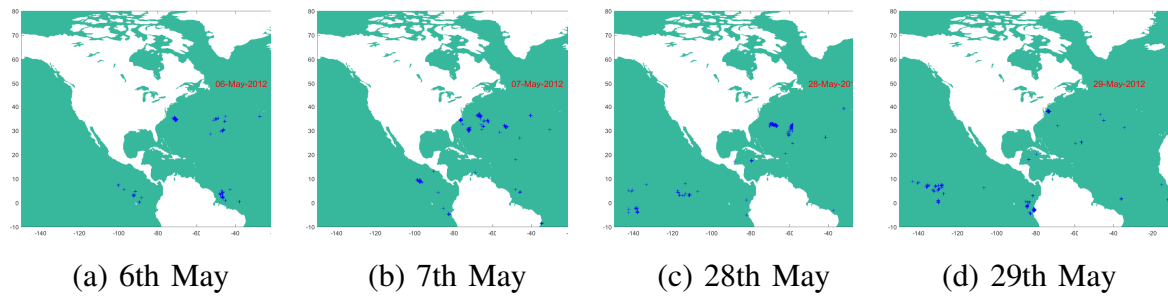


Figure 14: Locations of weather anomaly obtained based on the error between the acutual temperature and the prediction of autonomous kernel observer. Landmass is shown in white and the ocean is in green. Locations marked have error greater than two standard deviations above the mean error.

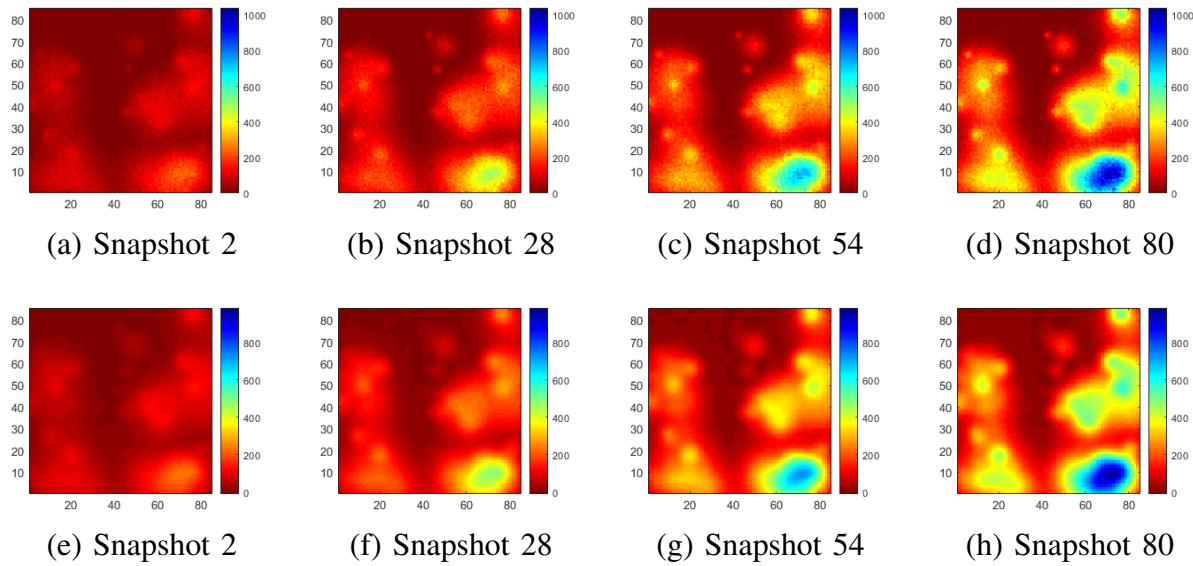
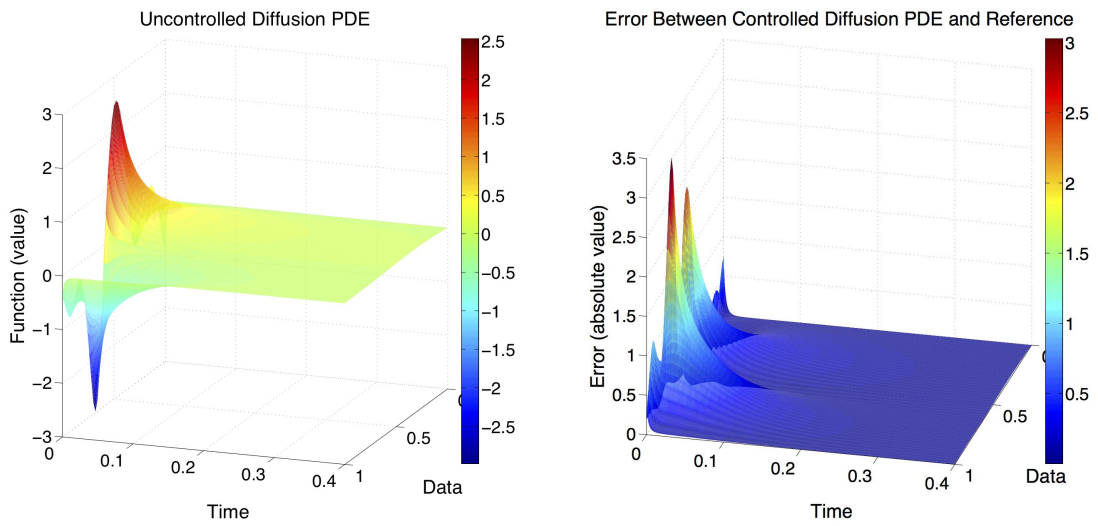


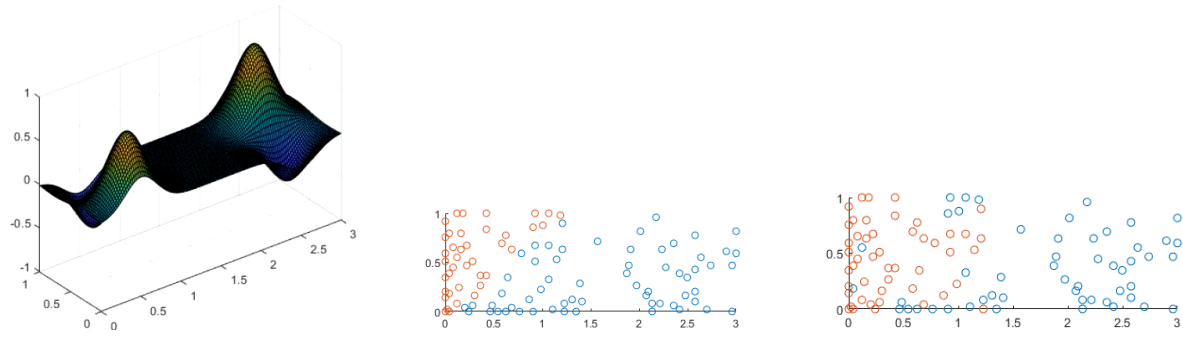
Figure 15: Visualization of Weed Density Growth over 20 days, original (a-d), E-GP (e-h)



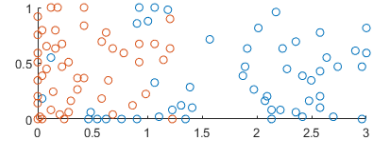
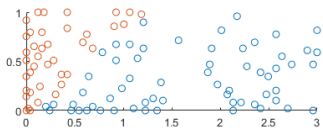
(a) Evolution of initial function  $f_{\text{init}}$ .

(b) Error in absolute value between controlled pde and  $f_{\text{ref}}$ .

Figure 16: Demonstration of the control of a linear diffusion equation.

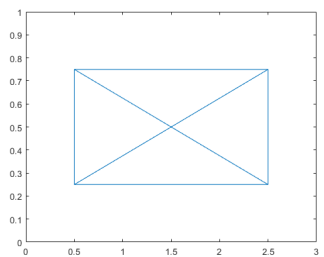


(a) Synthetic system with 2 invariant subspaces

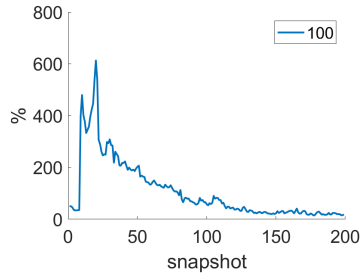


(c) Final clustering of centers

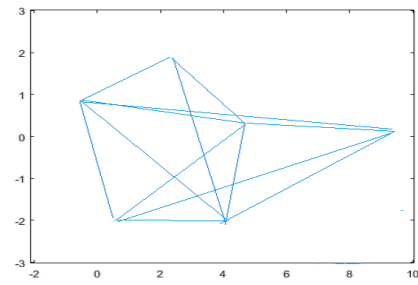
Figure 17: Synthetic System Invariant Subspaces



(a) Best Path Chosen for Synthetic Data Set



(b) Convergence of State Prediction for Synthetic



(c) Best Path Chosen for CFD

Figure 18: More Results

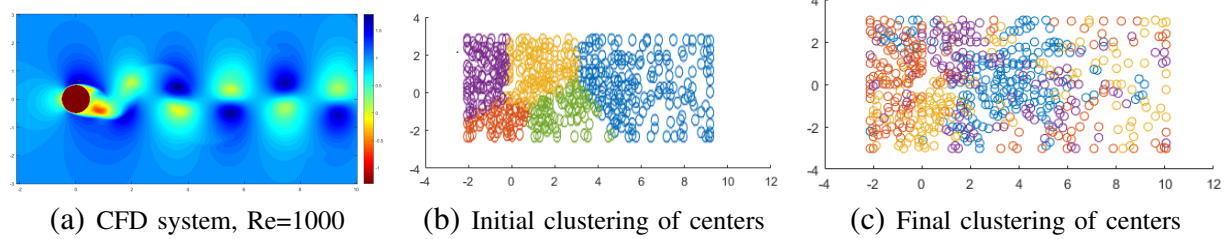
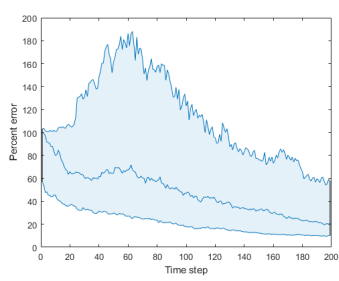
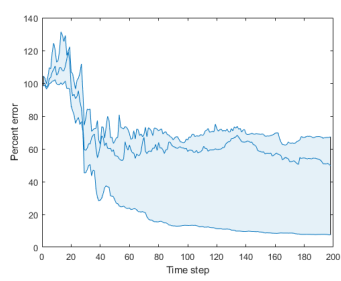


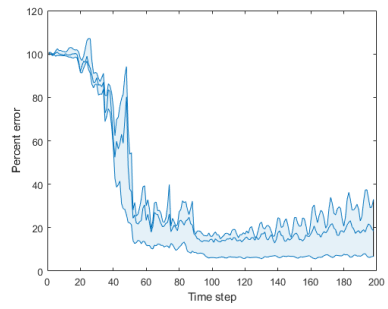
Figure 19: CFD System Invariant Subspaces



(a) Convergence of purely random paths



(b) Convergence of best scoring paths selected from random set



(c) Convergence of Best Scoring Path using Waypoints

Figure 20: Convergence results for CFD, showing mean, 15th, and 85th percentile path performance

---

**Algorithm 7** Measurement Map  $\tilde{K}$ 

---

**Input:**  $\hat{A} \in \mathbb{R}^{M \times M}$

Compute Frobenius canonical form, such that  $C = Q^{-1}\hat{A}^TQ$ . Set  $C_0 := C$ , and  $M_0 := M$ .

**for**  $i = 1$  **to**  $\ell$  **do**

    Obtain MP  $\alpha_i(\lambda)$  of  $C_{i-1}$ . This returns associated indices  $\mathcal{J}^{(i)} \subset \{1, 2, \dots, M_{i-1}\}$ .

    Construct vector  $v_i \in \mathbb{R}^M$  such that  $\xi_{v_i}(\lambda) = \alpha_i(\lambda)$ .

    Use indices  $\{1, 2, \dots, M_{i-1}\} \setminus \mathcal{J}^{(i)}$  to select matrix  $C_i$ . Set  $M_i := |\{1, 2, \dots, M_{i-1}\} \setminus \mathcal{J}^{(i)}|$

**end for**

Compute  $\mathring{K} = [v_1^T, v_2^T, \dots, v_\ell^T]^T$

**Output:**  $\tilde{K} = KQ^{-1}$

---

---

**Algorithm 8** Sampling locations set  $\mathcal{X}$ 

---

**Input:**  $\widehat{A} = C$ , lower bound  $\ell$

Decompose  $C$  to generate invariant subspaces  $\widehat{\mathcal{H}}_j$ ,  $j \in \{1, 2, \dots, \ell\}$  (see section )

**for**  $j = 1$  **to**  $\ell$  **do**

    Obtain centers  $\mathcal{C}^{(j)}$  w.r.t subspace  $\widehat{\mathcal{H}}_j$ ,

    Generate samples  $x_i^{(j)}$  to create a kernel matrix  $K^{(j)}$  that is shaded only with respect to centers  $\mathcal{C}^{(j)}$

**end for**

**Output:** Sampling locations set  $\mathcal{X} = \{x^{(1)}, x^{(2)} \dots, x^{(\ell)}\}$ .

---

---

**Algorithm 9** Kernel Observer (Transition Learning)

---

**Input:** Kernel  $k$ , basis centers  $\mathcal{C}$ , final time step  $T$ .  
**while**  $\tau \leq T$  **do**

- 1) Sample data  $\{y_\tau^i\}_{i=1}^M$  from  $f_\tau$ .
- 2) Estimate  $\hat{w}_\tau$  via standard kernel inference procedure.
- 3) Store weights  $\hat{w}_\tau$  in matrix  $\mathcal{W} \in \mathbb{R}^{M \times T}$ .

**end while**

To infer  $\hat{A}$ , define matrix  $\Phi = \mathcal{W}^T \mathcal{W}$ . Then:

**for**  $i = 1$  **to**  $M$  **do**

At step  $i$ , solve system

$$\hat{A}^{(i)} = ((\Phi + \lambda I)^{-1} (\mathcal{W}^T \mathcal{W}^{(i)}))^T, \quad (17)$$

where  $\hat{A}^{(i)}$ , and  $\mathcal{W}^{(i)}$  are the  $i$ th columns of  $\hat{A}$  and  $\mathcal{W}^{(i)}$  respectively.

**end for**

Compute the covariance matrix  $\hat{B}$  of the observed weights  $\mathcal{W}$ .

**Output:** estimated transition matrix  $\hat{A}$ , predictive covariance matrix  $\hat{B}$ .

---



---

**Algorithm 10** Kernel Observer (Monitoring and Prediction)

---

**Input:** Kernel  $k$ , basis centers  $\mathcal{C}$ , estimated system matrix  $\hat{A}$ , estimated covariance matrix  $\hat{B}$ .

**Compute Observation Matrix:** Compute the cyclic index  $\ell$  of  $\hat{A}$ , and compute  $K$ .

**Initialize Observer:** Use  $\hat{A}$ ,  $\hat{B}$ , and  $K$  to initialize a state-observer (e.g. Kalman filter (KF)) on  $\hat{\mathcal{H}}$ .

**while** measurements available **do**

- 1) Sample data  $\{y_\tau^i\}_{i=1}^N$  from  $f_\tau$ .
- 2) Propagate KF estimate  $\hat{w}_\tau$  forward to time  $\tau + 1$ , correct using measurement feedback with  $\{y_{\tau+1}^i\}_{i=1}^N$ .
- 3) Output predicted function  $\hat{f}_{\tau+1}$  of KF.

**end while**

---

---

**Algorithm 11** Kernel Controller

---

**Input:** Kernel  $k$ , basis points  $\mathcal{C}$ , estimated system matrix  $\hat{A}$ , estimated covariance matrix  $\hat{B}$ , and function  $f_{\text{ref}}$  to drive initial function to.

**Initialize Observer:** (see Algorithm 10).

**Initialize Controller:** Use Jordan decomposition of  $\hat{A}$  to obtain no. of control locations  $\mathcal{D}$ , compute kernel matrix  $K_{CD} \in \mathbb{R}^{\ell \times M}$  between  $\mathcal{D}$  and  $\mathcal{C}$ , and initialize controller (e.g. LQR) utilizing  $(\hat{A}, \hat{B})$ .

**while** measurements available **do**

- 1) Sample data  $\{y_k^i\}_{i=1}^N$  from  $f(x, \tau)$ .
- 2) Utilize observer to estimate  $\hat{w}_{\tau+1}$ .
- 3) Use  $\hat{w}_{\tau+1}$  and  $f_{\text{ref}}$  as input to controller to get feedback.

**end while**

---

## Sidebar: Key control problems in agriculture

Shortage of qualified human labor is a key challenge facing farmers [74], [75], leading to smaller profit margins, and preventing the adoption of truly sustainable agricultural practices. Lack of timely available labor was a principle reason behind the tens of millions of dollars of unharvested fruits and vegetables that rotted in California farms in 2017 [76], [77]. Labor shortage can be a major barrier to more sustainable agricultural practices that are more labor intensive. For example, more sustainable alternatives to prevalent methods of agriculture that would not need large amounts of chemicals and other inputs, such as perennial polycultures (mixed species of fruit- and nut-producing trees and shrubs [78], see Figure S2), are currently impractical at scale with current agricultural equipment. Polyculture systems can leverage co-habitation of mutually beneficial plants (and animals, insects, or microbiomes) to create a more sustainable engineered ecosystem. Labor shortage has become a primary barrier to adoption of this sustainable agricultural alternative [79], [80].

One way to address the challenges of labor shortage in agriculture is by creating new robotic technology that can work in harsh, uncertain, and dynamically changing field environments. Here we outline some of the fundamental challenges in autonomy, estimation, and control that the controls community can help overcome to enable the future of agricultural robotics, and relate it with the problem of spatiotemporal function estimation studied in this paper:

**Persistent multi-agent autonomy under partial observability:** The digital farm of the future will employ teams of distributed heterogeneous agents to autonomously manage, optimize, and harvest large acres of diverse crops across the entire season without encumbering humans. This level of autonomy in unstructured field environments is out of reach of the current state-of-the-art which requires constant human monitoring and oversight, especially in the presence of change or unforeseen events. Efficient and reliable control will be central to the success of these robots. Small below-canopy robots (e.g. Figure S1 [81]) will need to provide precision care including pruning, weeding, and re-seeding without damaging plants or causing soil compaction. Deployed at scale, these robots can not only make large scale organic farming practical, but also enable enhanced breeding through field-scale phenotyping [81]–[83]. *The big controls challenge here is in making decisions over large spatiotemporal scales with information obtained from a few stationary and mobile sensors which can only partially observe the environment at any given time.* The work pursued in this paper lays a foundation towards this problem by enabling a team of agents to estimate the varying state of the environment.

**Dexterous and ubiquitous robotics for precise care:** The digital farm of the future will strive to eliminate costly inputs (chemicals, labor, energy, and knowledge) with low-cost,

dexterous, and highly autonomous agricultural equipment [84]. Advances in *soft* arms and grippers can enable robots that can have far better reach and dexterity around plants than robots equipped with traditional *hard* industrial robotic arms. Soft arms, which are often actuated with pressurized tubes, can be far less expensive to manufacture and significantly lighter than their hard counterparts. On the other hand, soft arms can be slow to actuate and have limited payloads. To make soft robots practical, optimal feedback control techniques are necessary that work with conformal objects with very large degrees of freedom. In particular, soft arms tend to significantly deform under weight and behave quite differently when loaded with different payloads. Unlike hard arms, encoders are not sufficient to estimate the pose of the arm or the manipulator. Strain and angle sensors need to be positioned judiciously to keep costs down, and image based feedback control will be necessary. The evolving Gaussian Process technique and the kernel observer techniques described in this paper could be utilized to create distributed observers for such soft systems.

**Control of complex designed ecosystems:** Agricultural ecosystems designed with synergistic biodiversity – known as *polycultures* – have long been proposed as alternatives to traditional monoculture-based agriculture [85]–[88]. Based on studies of a range of natural ecosystems, polycultures are expected to be sustainable, productive, resilient and ecologically beneficial, requiring far less chemical inputs for food production. However, compared to their widely practiced counterparts (traditional monocultures such as corn and soybean), polyculture production systems have not yet been demonstrated to be profitable at large scale. This is primarily due to their labor- and knowledge-intensiveness, stemming from their inherent complexity. Indeed, despite their clear benefits and the potential to be profitable, optimizing and managing such complex systems can be difficult, even for those with training and experience in polycultures.

The complex interaction between closely-spaced diverse plant species in a polyculture results in both spatial and temporal dynamics as the plants grow and interact with each other. Plant growth often exhibits hybrid dynamical systems behavior with rapid thresholded growth bursts followed by slow progression. The triggers for growth bursts are dependent on environmental factors such as temperature, soil moisture, and sunlight reaching individual and cumulative thresholds, and complex interrelations between neighboring plants, soil chemistry, insects, and soil microbes. Simulating individual plant growth is an active area of research with many open questions in modeling and plant biology [89]. Arguably, very high resolution plant growth models may not even be necessary for effective control of polycultures. Yet on the other hand, the existing models of plants and interactions with ecosystems [90]–[95] are not well suited for designing and managing polycultures because of the simplifying assumptions that are often made. A good balance could be struck with data driven machine learning models that have sufficient resolution

for aggregate prediction over multiple spatiotemporal scales and are lightweight enough for control and decision making. With these models, predictive control strategies can be created that 5 task teams of robots for management tasks. Furthermore, these predictive models can enable quantified mechanisms of design and planning of efficient agroecosystems. Obtaining the required data to train these models and using them to create effective control techniques for managing profitable polycultures remains an exciting direction of future work where the control community can help.

10 The discussion in this sidebar was informed by many conversations with crop scientists at UIUC. In particular, *the authors wish to thank Prof. Stephen Long and Prof. Carl Bernacchi for insights on the phenotyping bottleneck and simulation of plants (crops-in-silico), Prof. Adam Davis on inputs on the herbicide resistant weed crisis, and Prof. Sarah Lovell on sustainable agricultural production systems and perennial polycultures, as well as the members of the UIUC Center for Digital Agriculture*

## Sidebar: Feature Spaces in Machine Learning

Suppose we have data  $\mathcal{D} = \{(x_1, y_1), \dots, (x_N, y_N)\}$ , where  $x_i \in \Omega_I$  and  $y_i \in \Omega_O$ . Here,  $\Omega_I$  is the input domain, and is generally a subset of  $\mathbb{R}^D$ , although more general sets such as discrete spaces, graphs, or text documents can be considered. Similarly,  $\Omega_O$ , the output domain, can be just as general as  $\Omega_I$ . We wish to solve for functions  $f$  in some space of functions  $\mathcal{J}$  such that  $f(x_i) = y_i \forall i$ . Generally, to restrict the complexity of the space  $\mathcal{J}$ , a *loss function*  $L(f, \mathcal{D}) \mapsto \mathbb{R}$  is chosen, which measures the error between a prediction  $f(x_i)$  given a datapoint  $x_i$ , and  $y_i$ , averaged over the entire dataset  $\mathcal{D}$ , and the optimization problem becomes

$$f^* = \arg \min_{\mathcal{J}} L(f, \mathcal{D}) + \lambda g(f), \quad (\text{S1})$$

where  $\lambda \in \mathbb{R}$ , and  $g(f)$  represent some constraints on the function  $f$ , such as smoothness.  
<sup>5</sup> Control theorists are most likely familiar with input-output pairs where  $x_i \in \mathbb{R}^N$  and  $y_i$  is either in  $\mathbb{R}$  or  $\mathbb{R}^M$  (*regression*). In machine learning, the most common task is when the  $y_i$  are discrete (*classification*). Different combinations of task, loss functions, and spaces  $\mathcal{J}$  result in different algorithms to solve these problems, which can sometimes form entire subfields of machine learning.

The choice of the function space  $\mathcal{J}$  can be critical for the task we want to perform, similar to how the choice of the state space is in control theory. Let's consider a simple example. Suppose we have data from two *classes*  $\mathcal{D}_A = \{(x_1^A, y_1^A), \dots, (x_N^A, y_N^A)\}$  and  $\mathcal{D}_B = \{(x_1^B, y_1^B), \dots, (x_N^B, y_N^B)\}$ , where  $x_i^{\{A,B\}} \in \mathbb{R}^D$ , and  $y_i^{\{A,B\}} \in \{-1, +1\}$ , shown in Figure S3a. Let  $f$  be chosen from the class of linear algorithms, i.e.  $f = w^T x + b$ , where  $w \in \mathbb{R}^D, b \in \mathbb{R}$ . We pick a loss  $L$  that returns a loss of zero when the prediction is the correct class, and if the prediction is the incorrect class, returns a higher value for misclassifications that are closer to the boundary. A classical example of such a loss is that used by the *perceptron algorithm*, which can be written as

$$L(f, \mathcal{D}) = \frac{1}{N} \sum_{i=1}^N \max(0, -y_i w^T x). \quad (\text{S2})$$

<sup>10</sup> This loss measures how accurate the prediction of the perceptron is on average. The general algorithm is as follows:

- 1) Initialize  $w \in \mathbb{R}^D$  to all zeros.
- 2) For a fixed number of iterations, or until some stopping criterion is met:
  - a) For each training example  $(x_i, y_i)$ ,
    - i) Let  $\hat{y}_i = \text{sgn}(w^T x_i)$ .
    - ii) If  $y_i \neq \hat{y}_i$ , update  $w \leftarrow w + y_i x_i$ .

The perceptron was one of the first machine learning models, and the genesis of modern neural networks [96]. Figure S3 shows a visual representation of where the perceptron algorithm can solve for the decision boundary with zero error. However, if the structure of the data has some nonlinearities, no solution will be found, as seen in Figure S4. In this case, the original space the data resides in is, in some sense, not a rich enough representation. If we could construct a mapping of the data to a different space which gives a learning algorithm more degrees of freedom to work with, linear algorithms can still be deployed. If we map the same data using a nonlinear map  $\phi(x, y) := (x^2, y^2, 2xy)$ , the perceptron now finds a solution in 3 dimensions, as seen in Figure S5. This example shows why so much of the work in machine learning focuses on learning the right representation for the data, for the right representation makes the classification task easy. Two major threads of research in the arena of feature maps over the last 40 years are kernel methods, and neural networks, the latter of which has gained remarkable notoriety in the last 10 years. These lines of research represent distinctly different strategies for generating feature maps from data.

In Figure S4, the data was mapped using an explicit feature map. Kernel methods, of which Gaussian Processes are a great example, utilize an elegant strategy for generating feature maps from data, using a remarkably simple trick called *the kernel trick*. Given a positive-definite kernel function  $k(x, y) : \Omega \times \Omega \rightarrow \mathbb{R}$ , Mercer's theorem guarantees the existence of a feature map  $\psi : \Omega \rightarrow \mathcal{H}$ , where  $\mathcal{H}$  is a *reproducing kernel Hilbert space (RKHS)*, and the map  $\psi$  obeys the property

$$k(x, y) = \langle \psi(x), \psi(y) \rangle_{\mathcal{H}}. \quad (\text{S3})$$

Recall that since  $\mathcal{H}$  is an RKHS, given  $c \in \Omega$ ,  $k(x, c) = \langle \psi(x), \psi(c) \rangle_{\mathcal{H}}$ , and  $k(x, c) := \psi_c \in \mathcal{H}$ . Furthermore,  $\text{span}\{\psi_x\}_{x \in \Omega}$  is dense in  $\mathcal{H}$ . There exist kernels that generate  $\mathcal{H}$ s that are extremely high dimensional: for example, the RBF kernel  $k(x, y) = e^{-\gamma \|x-y\|^2}$  is infinite-dimensional. This high degree of freedom enables the design of powerful learning algorithms that are linear in  $\mathcal{H}$ , but nonlinear in the input domain  $\Omega$ . The canonical example of this is the support vector machine (SVM) [97], but a more instructive example for us is the perceptron algorithm.

Suppose we trained a perceptron using  $\mathcal{X} = \{x_1, \dots, x_N\}$ ,  $x_i \in \mathbb{R}^D$  for some  $D$ ,  $y_i \in \{-1, 1\}$ . The prediction of the perceptron is  $\hat{y} = \text{sgn}(w^T x)$ , where  $w \in \mathbb{R}^D$ . It can be shown that  $w = \sum_{i=1}^n \alpha_i y_i x_i$ , where  $\alpha_i$  is the number of times  $x_i$  was misclassified. This allows us to derive the dual version of this algorithm, because

$$\hat{y} = \text{sgn}(w^T x) = \text{sgn} \sum_{i=1}^n \alpha_i y_i \langle x_i, x \rangle_{\mathbb{R}^D}.$$

The dot product  $\langle x_i, x \rangle_{\mathbb{R}^D}$  can be replaced with the kernel, leading to the *kernel perceptron algorithm*:

- 1) Initialize  $\alpha \in \mathbb{R}^N$  to all zeros.
- 2) For a fixed number of iterations, or until some stopping criterion is met:
  - a) For each training example  $(x_j, y_j)$ ,
    - i) Let  $\hat{y}_i = \text{sgn} \sum_{i=1}^n \alpha_i y_i k(x_i, x_j)$ .
    - ii) If  $y_i \neq \hat{y}_i$ , update  $\alpha_j \leftarrow \alpha_j + 1$ .

This algorithm is nonlinear in the input domain, but linear in the feature space  $\mathcal{H}$ , which led the deep learning community to, somewhat pejoratively, label this as an example of a *shallow learning architecture*. Kernel methods can also be used in a more direct fashion: if we have a subspace  $\mathcal{H}' \subset \mathcal{H}$  with a basis generated from  $\mathcal{C} = \{c_1, \dots, c_M\}$ , i.e.  $\mathcal{H}' = \text{span}\{\psi(c_1), \dots, \psi(c_M)\}$ , a linear model in  $\mathcal{H}'$  is again given by a vector  $w \in \mathbb{R}^M$ . Suppose this weight vector represents a boundary in  $\mathcal{H}'$ : to compute which side of this boundary a point  $x$  would lie on in  $\mathcal{H}'$ , we simply compute  $\text{sgn}(\sum_{i=1}^M w_i \langle \psi(x), \psi(c_i) \rangle_{\mathcal{H}}) = \text{sgn}(\sum_{i=1}^M w_i k(x, c_i))$ .

The choice of the kernel and its parameters depends on the dataset and the loss function. The kernel and the data together form the feature space. Because kernel methods are linear in their parameters and are restricted to RKHSs, they are amenable to somewhat straightforward mathematical analysis, and are very well studied because of this. The very recent review of Gaussian processes in control that appeared in IEEE CSM contains further details, examples and pointers to software relating to GPs and their use in control [21]. We switch our attention now to a different way of obtaining the features: Deep Neural Networks. While GPs build features using positive semidefinite symmetric kernels that compare any two points, DNNs build features using nested nonlinear operations.

Deep neural networks (DNNs) are models where the representing function  $f$  has nested nonlinearities. Fix a width  $M \in \mathbb{N}$ . Deep nets are parameterized models with weight matrices  $W^l \in \mathbb{R}^{M \times M}$ , bias vectors  $b^l \in \mathbb{R}^M$ , and a pointwise nonlinearity  $\phi : \mathbb{R} \rightarrow \mathbb{R}$ , with  $l = 1, \dots, L$ . Vectors  $h^l \in \mathbb{R}^M$  are called preactivations, and  $x^l \in \mathbb{R}^M$  are called postactivations, each element of which is called a *neuron*. Let  $h^0 \in \mathbb{R}^M$  be the input: then the canonical feedforward neural network is given by

$$x^l = \phi(h^l), \quad h^l = W^l x^{l-1} + b^l.$$

Therefore, we have that  $f(h^0) = x^l$ . The individual steps  $l$  are called the *layers* of the network, and the nesting property allows these networks to learn much more complicated functions than shallow architectures given the same number of nonlinearities [98]. Different choices of nonlinearities, connections, and layer architectures lead to different types of neural networks, which are used for different applications [99]. Deep learning has had an enormous impact on both the machine learning literature and industrial applications, and that impact has

bled over rapidly to other fields. Due to the nested structure of nonlinearities in DNNs, they are more difficult to analyze using simple mathematical tools, and therefore most of the literature in  
5 the field has focused on the empirical performance of these methods, where they significantly outperform competing methods. The significance of the achievements of machine learning in general and deep learning in particular have been in its ability to simplify the implementation of complex functional representation, essentially, in its ability to make system identification more accessible for difficult problems. However, the feature spaces generated by deep networks  
10 are not as well behaved and accessible to analysis as those generated by kernel models. In particular, deep network feature spaces do not naturally have the properties of RKHSs. This presents one barrier to analyzing and understanding the nature of these models. For the reliable and verifiable inclusion of DNNs in engineering control systems, further insights are necessary into the structure of the feature spaces these models generate, to restrict certain forms of output, and to shape decision-making.



Figure S1: The TerraSentia robots developed by Chowdhary's group at UIUC and commercialized by EarthSense inc. Agricultural robots such as these present exciting possibilities for distributed agricultural management with teams of compact, ultra-light, under-canopy robots equipped with advanced autonomy and machine learning. Such robots can fill the niche between large farm equipment and manual labor, as well as enable perennial polycultures, and closer to home, they could also weed your garden! Advances in persistent multi-agent autonomy in harsh, changing, and uncertain environments driven by the controls community are driving such exciting possibilities of the future of agriculture.

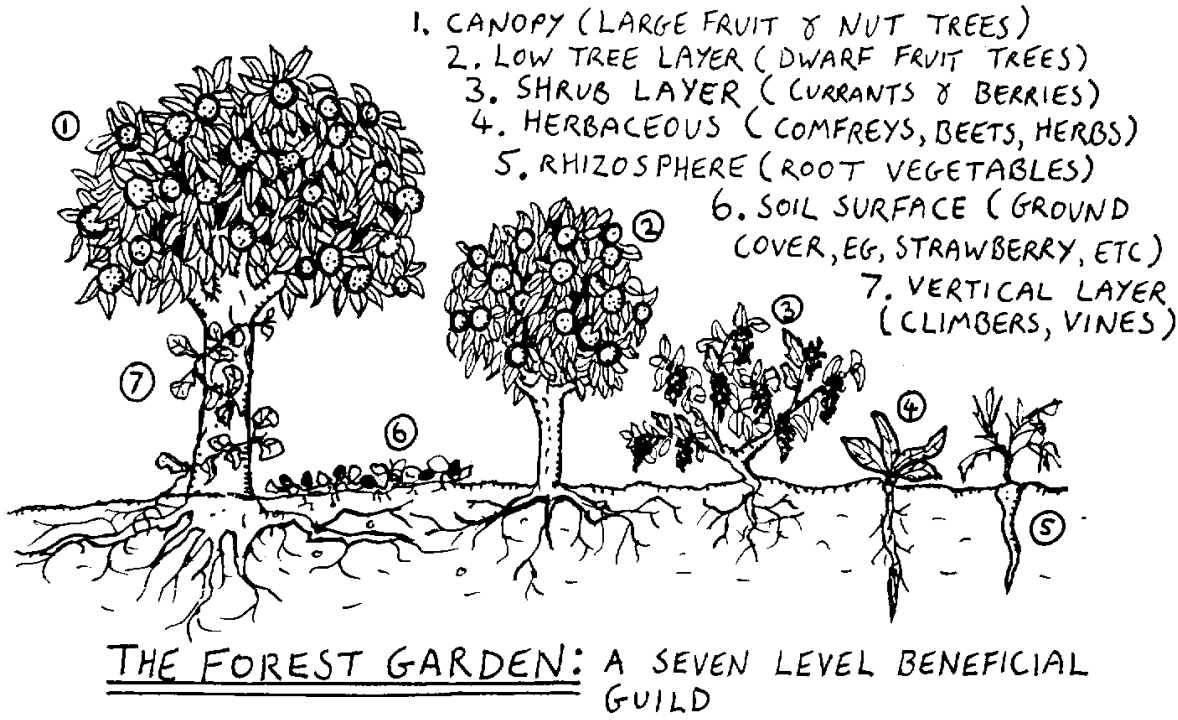
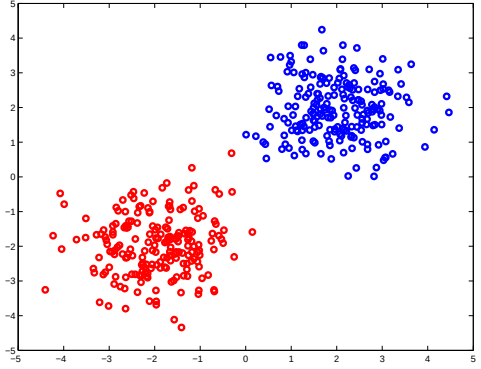
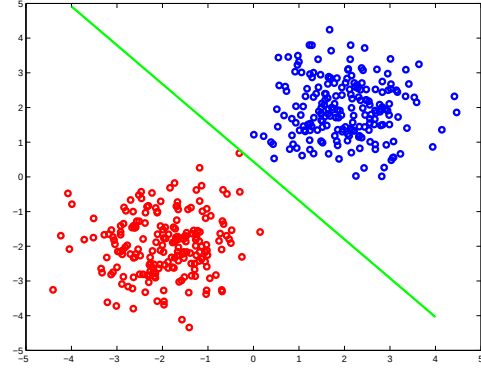


Figure S2: The seven layers of the forest garden. Polyculture agricultural production systems are designed ecosystems that can be more productive and sustainable than traditional monoculture systems. Managing such complex systems requires fundamental advances in robotics, spatiotemporal modeling, and control of complex biological processes, all areas where the control community can help. Figure source [100] see also [101].

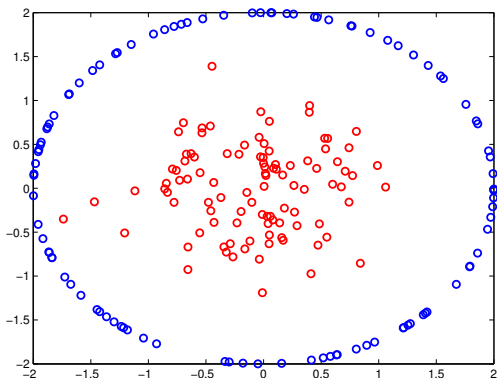


(a) Data from classes  $A$  and  $B$ .

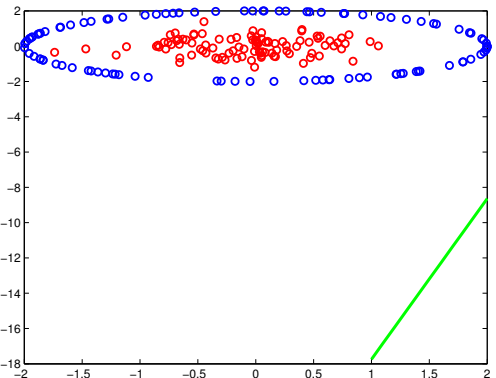


(b) Linear boundary separating data.

Figure S3: Example of linearly separable data. Any simple linear learning algorithm e.g. perceptron, finds a solution.

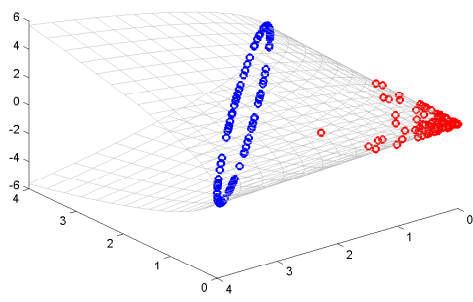


(a) Data from classes  $A$  and  $B$ .

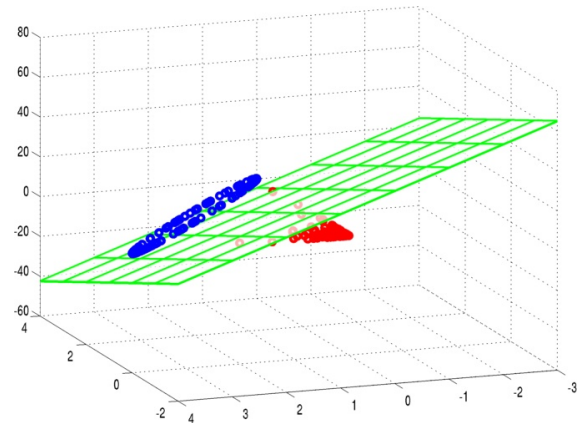


(b) Linear boundary fails to separate data.

Figure S4: Example of nonlinearly separable data. Perceptron fails to find a solution, and diverges.



(a) Nonlinear mapping of data.



(b) Linear boundary in new space.

Figure S5: If we map the same data using a nonlinear map  $\phi(x, y) := (x^2, y^2, 2xy)$ , the perceptron now finds a solution in 3 dimensions.

## Sidebar: Learning Fluid Flows with Evolving Gaussian Processes

2nd order Partial Differential Equations (PDEs) are ubiquitous in practical science and engineering, from mechanics to transport phenomena to electromagnetics. In our view, the Navier-Stokes equations governing fluid dynamics represent most, if not all of the overall complexity of modeling these as (a) it exhibits of hybrid system behavior, e.g. elliptic-hyperbolic, and (b) the nonlinearity results in the complex spatio-temporal dynamics that are prevalent in many practical situations. The form of the NSE for compressible Newtonian fluids is expressed below, where  $\mathbf{u}$  is the fluid velocity,  $p$  is the fluid pressure,  $\rho$  is the fluid density, and  $\mu$  is the fluid dynamic viscosity. [102]

$$\rho \left( \frac{\partial \mathbf{u}}{\partial t} + \mathbf{u} \cdot \nabla \mathbf{u} \right) = -\nabla p + \nabla \cdot (\mu(\nabla \mathbf{u} + (\nabla \mathbf{u})^T)) + \nabla \left( -\frac{2\mu}{3} \nabla \cdot \mathbf{u} \right) + \rho \mathbf{g}$$

We demonstrate the Evolving Gaussian Processes method on CFD data of flow over a bluff body (a cylinder) over a range of Reynolds numbers from 100 to 1000 (the Reynold number is dimensionless flow rate). This deterministic, high-dimensional spatiotemporal dynamical system is well-studied in the fluid dynamics literature, both experimentally and numerically [68]–[70]. The conventional wisdom would be to learn a separate model over each Reynolds number, but our results show that the E-GP method is capable of learning the dynamics of all the flow patterns at once. Using the learned dynamics over weights of successive kernel models, E-GP is capable of predicting the future states of functional evolution in a recursive manner. The key advantage of E-GP is that evolution of large function spaces can be transformed into learning the evolution of a relatively smaller Hilbert space which is encoded by the kernels and the associated weight vector. Furthermore, the values of the weights and the associated linear dynamical systems provide critical insights, such as spatial-correlations through the structure of the transition matrix, local modes of dynamic evolution through invariant subspaces of the transition matrix, and eigenmodes of evolution. The latter is of significant importance to the ongoing work in Koopman operator models of spatiotemporal phenomena.

In our CFD simulation, we used a 4th-order polynomial expansion with spectral element method on the incompressible Navier-Stokes equation to generate the cylinder flow data for  $Re = 100, 300, 600, 800$ , and  $1000$ . The spatial domain is  $[-2, 10] \times [-3, 3]$ , excluding the diameter-1 cylinder at the origin. Neumann boundary conditions are applied to the far-field of the cylinder in the  $y$ -direction and the outlet of the flow field; and a Dirichlet boundary condition is applied to the inlet. Each data set contains at least 200 snapshots with a uniform time step of  $\tilde{0.03}$  sec. Each snapshot contains 24,000 velocity data points for  $Re=100$  or 95,000 velocity data points for  $Re=300,600,800,1000$ . Each data set took at least 10 hours in a high performance computer

cluster to generate. Figures S6,S7(a-d) visualize the horizontal velocity for Re=100 and Re=1000, with red being the greatest negative velocity and blue the greatest positive velocity. The flow is  
5 unstable, periodic, and clearly nonlinear.

We used the Gaussian RBF kernel  $k(x, y) = e^{-\|x-y\|^2/2\sigma^2}$  in our E-GP model, with  $\sigma$  estimated to be 0.4. Using a budget of 600 kernel centers (see Figure S9a-S9b, and note how they cluster in the most dynamic regions), we find a  $600 \times 600$  matrix  $\hat{A}$  which accurately (Figure S8a) captures the dynamics of the nonlinear system. We can use this to propagate single initial  
10 condition  $w_0$  forward to make predictions, then compare the predictions to the original training data. We found total percentage errors between 3% for Re=100 and 7-8% for Re=1000, as can be seen in the solid lines in Figure S8a. We define the total percentage errors as  $E_\tau = \frac{\|\bar{y}_\tau - \bar{y}_\tau\|_2}{\|\bar{y}_\tau\|_2}$  where  $\bar{y}_\tau$  is the output vector for time  $\tau$  and  $y_\tau$  is the E-GP estimate at that time. Note that *the size of the model has been reduced by almost two orders of magnitude* from the original CFD  
15 data. This process takes about 13 minutes in MATLAB for a 200 snapshot by 95,000 point set on an ordinary Intel i7 4.00 GHz processor.

## One Transition Matrix for Everything

In order to approach the challenge of generalizing across similar spatiotemporally evolving systems, the first question we had to answer is whether we can find an  $\hat{A}$  matrix that accurately  
20 captures the dynamics of multiple similar flows. The answer to that question is *yes*, using the trajectory concatenation method. Amazingly, a single model generated this way works almost as well on all five data sets as do five individual models trained on each data set separately. This is confirmed by both the total error plots (Figure S8a), which show only slight increases in each of the total percentage error plots, and visual inspection of the dynamic modes displayed.  
25 This result is even more surprising in light of the fact that the rate of vortex shedding for each Reynolds number is different. By taking a Fourier transform of the time evolution of a data point located at (0.5,8), we find that for the original data sets the vortex shedding frequency is 0.448 Hz, 1.260 Hz, 1.380 Hz, 1.388, and 1.401 Hz for Re=100, 300, 600, 800, and 1000 respectively, and for the E-GP models the frequencies are 0.452 Hz, 1.21 Hz, 1.36 Hz, 1.36 Hz, and 1.36 Hz  
30 respectively.

## Generalizing from Learned Dynamics to Unknown Dynamics

Having seen that it is possible to find a single transition in the weight space that models the dynamics systems over a range of parameters, the next challenge is to be able to model flows with parameters that the model has not been trained on. We derived an  $\hat{A}$  matrix from the Re=100, 300, 600, and 1000 data sets and tested it against the Re=800 data set. The results are below in Figure S8b. For the first 120 snapshots, the total percentage error remains under

10%, which is satisfactory. After this, however, the total percentage error curves upwards as the slight errors in the transition matrix compound. Over 800 snapshots, we found an average total  
5 percentage error of less than 25%.

## Linear Dynamical Layer Analysis & Insights

Due to the spatial encoding of the weights which the linear transition model operates on, we are able to analyze the dynamics and find physical insights into the process. We demonstrate two techniques: (1) using eigendecomposition of the transition matrix to discover the eigenfunctions  
10 and invariant subspaces of system, and (2) visualizing the most significant spatial interactions in the system.

An invariant subspace of a linear operator is a subspace of the Hilbert space such that any vector in the subspace remains in the subspace under transformation by the operator. By marking which kernel centers are associated with different subspaces, we can spatially separate  
15 the space into multiple dynamic modules. The physical insight is some areas of the space are dynamically entangled with each other, and other are independent of each other. For those interested in monitoring spatiotemporally evolving systems, the number and location of the invariant subspaces determines how many and where feedback sensors ought to be for robust prediction of the weights. Before doing the Jordan decomposition of  $\hat{A}$ , we zero any elements  
20 smaller than some small  $\varepsilon$  in order to stabilize the algorithm for matrices with many elements close to zero. Afterwards we visualize the eigenvector matrix using a *logarithmic* color chart, as seen in Figures S9a,S9b,S9c. These plots are for models trained individually on Re=100 and Re=1000 with 300 kernels, and on all five with 600 kernels, for comparison. We see three categories of eigenvector in the rows: (1) Rows at the bottom that have exactly one non-zero  
25 elements, (2) In the middle, a couple rows with a dozen significant elements, and (3) at the top a number of rows that affect the majority of the kernel centers in the space.

Each eigenvector of (1) spans its own invariant subspace, and is depicted in magenta circles in Figure S10. Category (3) is one invariant subspace, depicted with black crosses. Category (2) is subsumed in Category (3). The figures show that the dynamics near/around the cylinder and  
30 in its wake are so entangled that a single sensor measurement in that area may be sufficient to estimate over that entire subspace. On the other hand, areas far from the core of dynamic excitement are their own independent, invariant subspaces, and thus must be monitored locally.

Another way to visualize the operation of the linear transition matrix is to plot lines between kernel centers that are influencing each other strongly. That is, if we draw a line center  $c_j$  to  $c_i$  for each of the (relatively) largest elements  $a_{ij}$  of  $\hat{A}$ , one can see how the system dynamics are coupled spatially (Figures S11a,S11b,S11c). We can also plot the magnitude of  $a_{ij}$  in a third

axis for further insight into the most dominant dynamic connection in the system.

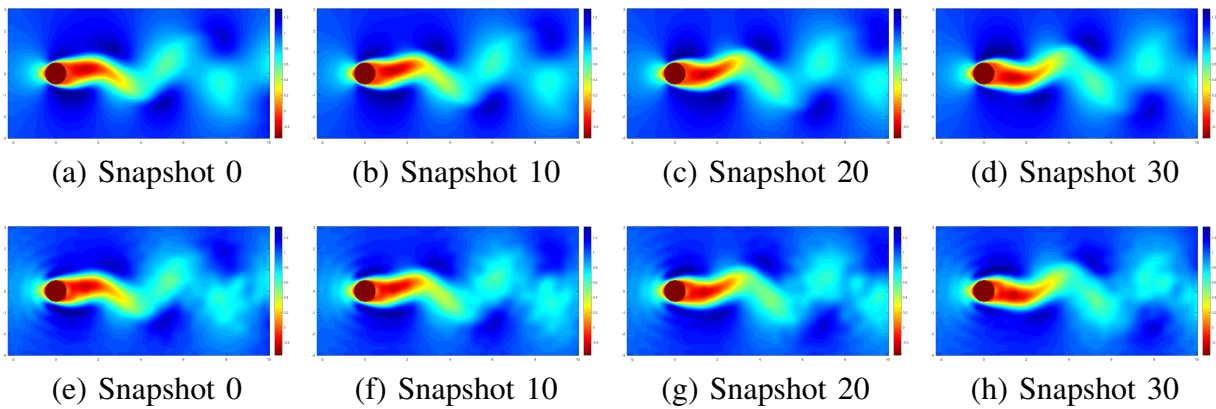


Figure S6: Visualization of Fluid Flow at  $\text{Re} = 100$ , CFD (a-d), E-GP (e-h)

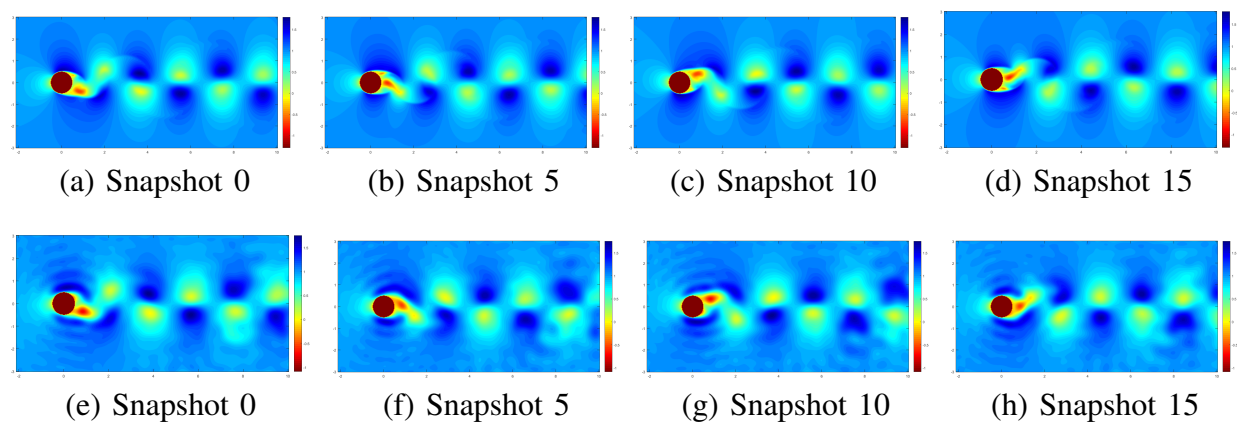
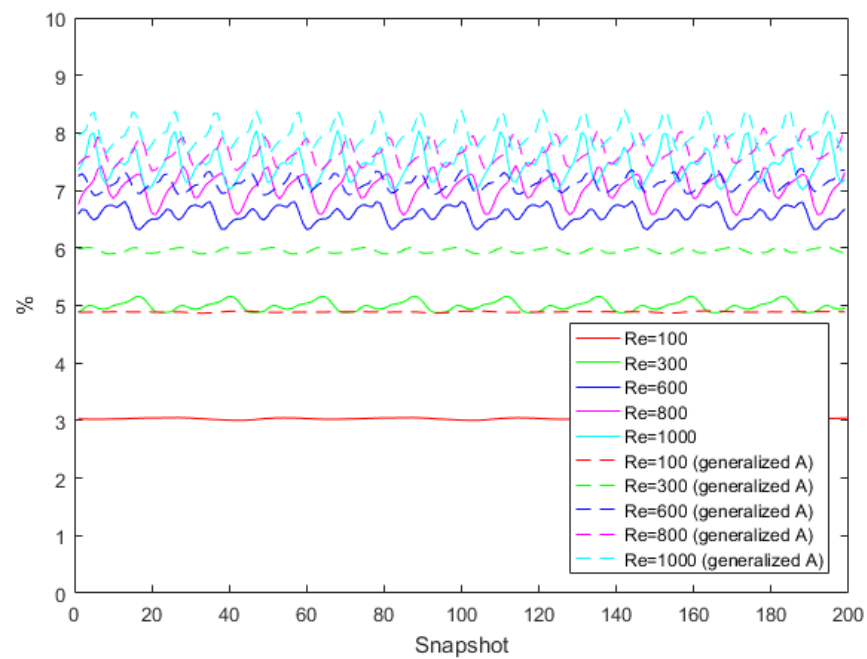
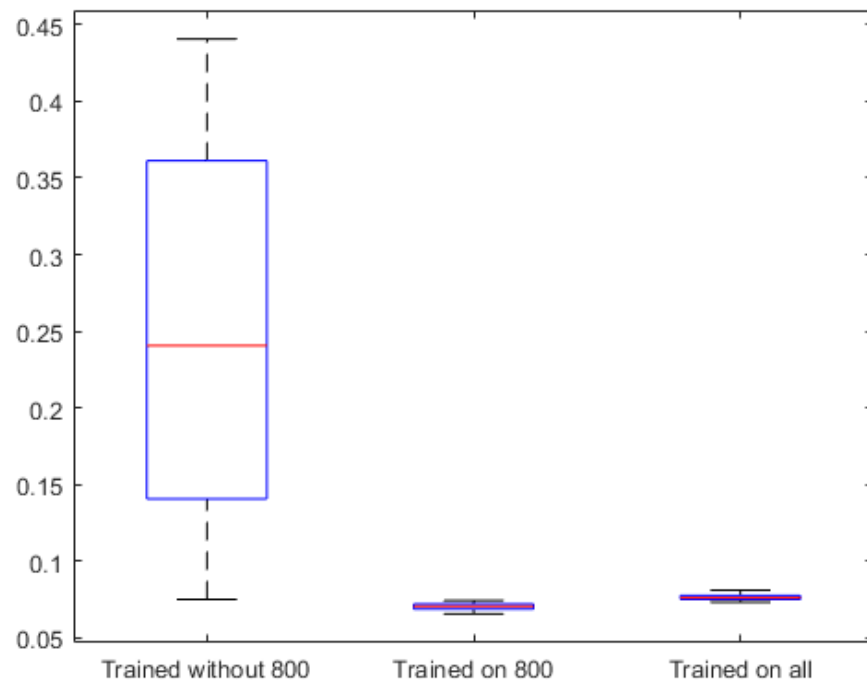


Figure S7: Visualization of Fluid Flow at  $\text{Re} = 1000$ , CFD (a-d), E-GP (e-h)



(a) Universal Generalizer vs Individual Models



(b) Different Models Tested on  $\text{Re}=800$

Figure S8: Total Percentage Errors

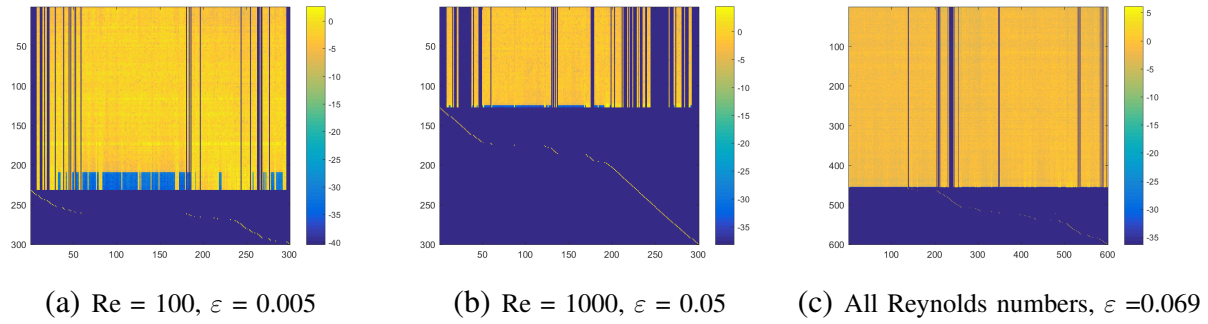


Figure S9: Eigenvector Heat Maps

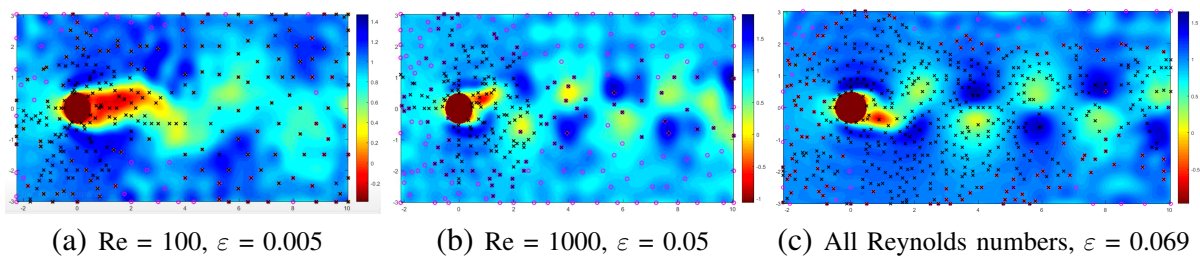


Figure S10: Invariant Subspaces

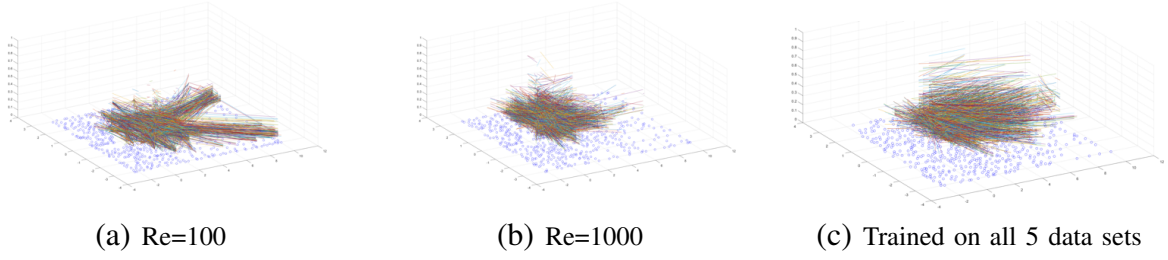


Figure S11: Visualization of Co-Relations in Transition Matrix

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**Harshal Maske** completed his PhD in 2018 from the University of Illinois in Urbana-Champaign,  
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