Leveraging the structure of dynamical systems for data-driven modeling

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Abstract

The reliable prediction of the temporal behavior of complex systems is required in numerous scientific fields. This strong interest is however hindered by modeling issues: often, the governing equations describing the physics of the system under consideration are not accessible or, when known, their solution might require a computational time incompatible with the prediction time constraints. Nowadays, approximating complex systems at hand in a generic functional format and informing it ex–nihilo from available observations has become a common practice, as illustrated by the enormous amount of scientific work appeared in the last years. Numerous successful examples based on deep neural networks are already available, although generalizability of the models and margins of guarantee are often overlooked. Here, we consider Long-Short Term Memory neural networks and thoroughly investigate the impact of the training set and its structure on the quality of the long-term prediction. Leveraging insights from ergodic theory, we perform a thorough computational analysis to assess the amount of data sufficient for a priori guaranteeing a faithful model of the physical system. We show how an informed design of the training set, based on invariants of the system and the structure of the underlying attractor, significantly improves the resulting models, opening up avenues for research within the context of active learning. Further, the non-trivial effects of the memory initializations when relying on memory-capable models will be illustrated. Our findings provide evidence-based good-practice on the amount and the choice of data required for an effective data-driven modeling of any complex dynamical system.

I. INTRODUCTION

In the present era of mathematization of Nature and essentially every aspects of our lives, the need for modeling is ubiquitous. Focusing on engineering as an example, models are invaluable for predicting quantities of interest, designing and optimizing physical systems, understanding their behavior in an environment, allowing to derive control strategies to achieve some goals, etc. Physical models have historically been developed from a combination of observational data and first principles, in varying parts depending on the amount and type of prior knowledge. This approach is still followed today where the mathematical structure of a model is typically provided and observational data are used to inform it, i.e., learn the coefficients associated with the different terms of the retained structure. Countless
variants exist but they generally follow this philosophy. Nonetheless, in many relevant cases, the physics of the phenomena is very complex and the interplay of observation with physical intuition is not sufficient to sort out a formalized model, even roughly accurate, let alone more rigorous general laws. In those cases, it is tempting to resort to a pure data-driven approach, in which some model is built up entirely from data. In fact, this approach has been always tried and already criticized in the XIX-th century by Maxwell [9]. The idea has however received some rigorous methodology and approach, even recently, for instance after the work of Takens [7, 29, 30]. Thanks to these efforts, it is possible to derive predictive models from time-series [19], but only for low-dimensional systems.

The data-driven approach has found new motivation following the significant progress in the field of Machine Learning (ML), fostered by the massive augmentation of computer-based activities [15]. Even if a clear framework is still missing, the use of such algorithms to build models from data has been recently attempted in a variety of different physical phenomena, e.g., [3, 6]. However, this recent blooming of deep learning approaches has not necessarily changed the global picture of modeling. Neural networks can provide a highly expressive and generic model structure suitable for a wide class of applications. The potential versatility of the resulting model is however associated with a usually large amount of necessary observational data to suitably inform the associated coefficients of the model, here being the weights and biases of the network nodes. Yet, not every situations enjoy massive amount of observational data and can afford to inform a widely generic model. In practice, it is more likely that data are scarce due the cost and/or technical difficulty in obtaining them and complex high-dimensional systems may need a quantity of data significantly exceeding the observing capability, [1]. In that sense, the lack of data for inferring physical models might have direct consequence on their relevance.

A strategy to alleviate this limitation is to simplify the postulated structure of the model so that it is less data hungry, with simpler models here understood as involving fewer parameters to inform from the available data, hence a weaker constraint on the amount of training samples. Another strategy is to leverage prior knowledge one has on the physical system under consideration to guide the structure and the learning process, hence saving on data. For instance, known symmetries can be enforced by design in the structure or in the representation format of the input data, effectively reducing the need for data. A similar idea applies to invariances one may know the system obeys (conserved quantities such as energy,
translational or rotational invariances, etc.) or properties such as stability in a given sense. This physics-aware machine learning approach has now become trendy and is the motivation of numerous recent efforts, see [20, 28, 33, 36] among many more. In that sense, modeling of chaotic dynamical systems is a natural play-field [2, 26]. Remarkably, previous works seem to indicate an intriguing capability to give accurate forecasting of chaotic systems, even relatively high-dimensional [25], apparently improving over deterministic embedding techniques [19]. These results therefore deserve to be thoroughly analyzed.

In the present work, we aim to understand the possible limitations of the pure data-driven approach and to explore the idea of leveraging some of the known organization of the dynamics to inform the process of collecting observations on the system. In particular, we explore different strategies for collecting observations from a dynamical system in the aim of learning and modeling its behavior and predict its state over time. These strategies only differ by the region in the phase space where observational samples are taken from, every other aspects being identical (sampling frequency, number of samples, observed quantity, etc.). These different configurations are chosen to be typical of situations ranging from no prior information besides an estimated upper bound of the dimension of the dynamical system up to approximate knowledge of some invariants of the system. More specifically, dynamical systems are known to be organized around critical features such as fixed points which can be seen as constitutive of a backbone of the whole system. While fixed points do not drive the whole dynamics and only provide information on the local behavior of the flow, they are nonetheless a crucial feature. To obtain a more complete picture of the dynamical structure, higher dimensional invariant sets are also required, [13, 14].

To illustrate the role of the sampling strategy, we consider the celebrated Lorenz’63 system. While deterministic and low dimensional, the Lorenz system features many challenges encountered in more complex configurations, such as sensitivity to initial conditions and an attractor dimension $d_A$ significantly lower than the ambient dimension $d$, here $d_A \simeq 2.06 < d = 3$. It is thus somehow a minimally complex unit and justifies its relevance for illustrating some challenges in learning dynamical systems. We carry out a thorough computational analysis in order to systematically test the impact that different sampling strategies have on the quality of the resulting models in terms of prediction abilities and generalizability. From an algorithmic point of view, recurrent structures such as auto-regressive models or recurrent neural networks are popular choices in such situations.
In the present work, we focus on the LSTM (Long-Short Term Memory) architecture [16]. We have chosen this architecture for the present study both for its widespread use, including applications in prediction of dynamical systems behavior [10]. Moreover, it has been recently shown that the performance given by different architectures including reservoir computing is similar [32], although for low dimensional systems like Lorenz’63, LSTMs appear to give the best results. Given this Neural Network approach, we have also investigated the possible impact of initial conditions and memory effects on the results of supervised learning models. This is relevant in the sense of providing best-practice guidance.

The paper is organized as follows. General principles in learning a system from data are briefly discussed in Section II to set up the stage and define notations. The general class of models we consider in this work is presented in Section III. The specific configuration we retain for the present discussion is the Lorenz’63 introduced in Section IV, together with the different sampling strategies. The results are gathered and presented in Section V before the discussion in Section VI closes the paper.

II. THEORETICAL BACKGROUND

A. Learning dynamical systems

The present work lies in the framework of learning a model to approximate the time evolution of a state of the system at hand from observations $s(t) \in \mathbb{R}^{n_s}$. To do so, one needs a representation for the state and a dynamical model for describing its evolution in time. A representation of the state is given by a function $g$ mapping observations over a measurement time horizon $\Gamma$, $s_\Gamma(t)$, $t \in \Gamma$, to a set of coordinates $v \in \mathbb{R}^{d_v}$. Several mappings have been proposed in the literature, [5, 12], including delays by time intervals $\Delta t$, $v_i = s(t - i\Delta t)$ for $i = 0, 1, \ldots$, and time derivatives of varying orders $i$, $v_i = s^{(i)}$.

To obtain the size of the coordinate space, we assume that the state of the system $u$ lies on an attractor $A$ of dimension $d_A < d$ that is bounded boundaryless and a smooth submanifold of $\mathbb{R}^d$. In this case, the Whitney embedding theorem [34] shows that if the number of coordinates $d_v$ is such that $d_v > 2d_A$ then the dynamics of the state can be entirely captured by the new coordinate system. The Takens embedding theorem [29, 30] provides a way to make this result practical by asserting that one can take this number of
time delays. Recent results have extended this theorem by providing guaranteed recovery properties based on the measurement operator, type of attractor, and other parameters of the problem, [8].

We assume no prior knowledge of the governing equations of the model and rely on a purely data-driven approach. The dynamical model for describing the time evolution of \( v(t) \in \mathbb{R}^d \) can be formulated in various ways, including feed-forward neural networks, owing to their high expressiveness.

As an alternative, for instance in case an upper bound of the attractor dimension is unknown due to process noise or measurement noise, a recurrent model \( F_\theta \) can be employed for describing the dynamics of \( v \). The model is parameterized by \( \theta \in \mathbb{R}^{n_\theta} \) and writes

\[
\dot{v}(t) = F_\theta(v),
\]

where a dotted quantity denotes its time-derivative.

Learning a model for the system under consideration takes the form of a classical supervised learning problem based on the misfit between observations \( s_\Gamma \equiv s(u(t \in \Gamma)) \) collected through a measurement operator \( h : u \mapsto s \) over a time domain \( \Gamma = [t_0, t_T] \) and predictions, in the sense of some norm \( \| \cdot \|_\Gamma \). It typically results in an optimization problem in terms of the model parameters, possibly including a regularization term \( r \):

\[
\theta \in \arg \inf_{\tilde{\theta} \in \mathbb{R}^{n_\theta}} \| s(t) - h(v(t; \tilde{\theta})) \|^2_\Gamma + r(\tilde{\theta}), \quad \text{s.t.} \quad \dot{v}(t; \tilde{\theta}) = F_{\tilde{\theta}}(v). \tag{2}
\]

B. Collecting data

The main issue impacting the predictions based on observations is the quantity of data actually needed to approximate the dynamics in a given sense. The issue is related to the ergodic theory of dynamical systems, whose founding idea is that the long-time statistical properties of a deterministic system can be equivalently described in terms of the invariant (time-independent) probability, \( \mu \), such that \( \mu(S) \) is the probability of finding the system in any specified region \( S \) of its phase space. If the trajectories of a \( d \)-dimensional ergodic system evolve in a bounded phase space \( \Omega \subset \mathbb{R}^d \), the Poincaré recurrence theorem ensures that analogues exist as it proves that the trajectories exiting from a generic set \( S \subset \Omega \) will return to such set \( S \) infinitely many times. The theorem holds for any points in \( S \), almost surely. It was originally formulated for Hamiltonian systems, but it can be
straightforwardly extended to dissipative ergodic systems provided initial conditions are chosen on the attractor. In $d$-dimensional dissipative systems, the attractor $A$ typically has a dimension $d_A < d$. The dimension $d_A$ describes the small scale (that is, with resolution $\varepsilon \ll 1$) driving the probability $\mu(B^d_x(\varepsilon))$ of finding points on the attractor which are in the $d$-dimensional ball of radius $\varepsilon$ around $x \in A \subset \mathbb{R}^d$:

$$\mu(B^d_x(\varepsilon)) = \int_{B^d_x(\varepsilon)} d\mu(\tilde{x}) \sim \varepsilon^{d_A}. \quad (3)$$

When $d_A$ is not an integer, the attractor and its probability measure are termed fractal. The attractor is typically multifractal [24], but we can assume without lack of generality that it is homogeneous. Given that the Poincaré theorem proves the existence of good analogues, the issue is shifted to finding out how much time is needed for a Poincaré cycle to end.

This key problem was considered by Smolouchoski and resolved by Kac [18], who showed that for an ergodic system, given a set $S \equiv B^d_{x^*}(\varepsilon)$, the mean recurrence time $\langle \tau_S(x^*) \rangle$ relative to $S$ is inversely proportional to the measure of the set $S$, i.e.,

$$\langle \tau_S(x^*) \rangle \propto \frac{1}{\mu(S)} \sim \varepsilon^{-d_A}, \quad (4)$$

where the average $\langle \cdot \rangle$ is computed over all the points $x \in S$ according to the invariant measure. Therefore, if we require $\varepsilon$ to be small and if $d_A$ is large, the average recurrence time becomes huge, a symptom of the curse of dimensionality in data-driven methods.

III. MACHINE-LEARNING APPROACH

A. Architecture

As discussed above, a set of $d$-dimensional observations retained to express a state of the system under consideration might not be a state vector, in the sense that it does not necessarily contain enough information to uniquely define its future evolution as an autonomous deterministic dynamical system. Even though the system were Markovian, it might not be so in the retained representation and may implicitly involve latent variables. This is typically the case in situations of partial observability or when measurements are affected by noise. As discussed in Sec. II, under conditions rigorously studied in the embeddology line of research, [7, 29], latent state variables can be substituted with observations over a recent
FIG. 1: Sketch of the retained architecture. Each hidden layer is made of LSTM cells.

past, that is, temporal information is traded for unavailable high-dimensional phase space information.

As said in the Introduction, in the present work, we consider the LSTM (Long-Short Term Memory) architecture. Compared with standard multi-layer perceptrons, LSTMs additionally feature internal variables $\zeta$ carried-over through the recurrence and effectively acting as a “memory”. Training an LSTM then allows to derive a discrete-in-time model predicting the future “state” $v_{k+1} \equiv v(t_k + \Delta t)$ from the current state and memory:

$$v_{k+1} = F_\theta(v_k; \zeta_k).$$ (5)

The retained architecture comprises 2 LSTM layers of 50 neurons each and one fully connected layer, with tanh activation functions in the LSTM cells and sigmoids for the state output, see also a sketch in Fig. 1. The present architecture was deemed as a good compromise between accuracy and computational expenses, after an analysis with comparable architectures with one and three layers has been performed. A 1-step ahead model is learned based on a whole trajectory, or a set of trajectories, in the $L^2$-sense; the training is performed with ADAM optimizer and an adaptive learning rate.

B. Strategies for sampling

Learning a model here consists in estimating the best set of parameters $\theta$ in the sense of the optimization problem defined in Eq. (2). In the context of neural networks, this acts as the loss function used for training. The regularization term may promote several properties of the optimal solution $\theta$, such as the popular choice of low magnitude (ridge regression), $r(\theta) \sim \|\theta\|_2$, or sparsity, $r(\theta) \sim \|\theta\|_1$, possibly mimicked with a suitable dropout strategy.
We adopt a discrete-in-time viewpoint, in a formulation consistent with the discrete model (5). The loss function $\mathcal{L}$ associated with the training problem then writes

$$
\mathcal{L} (\theta) = \sum_{k=1}^{K} \| s_k - h (v_k (\theta)) \|_2^2 + r (\theta),
$$

with $K$ such that $T = K \Delta t$ and the 1-step ahead dynamical model $v_{k+1} = F_{\theta} (v_k; \zeta_k)$.

The loss is here simply defined in terms of the Euclidean distance but many alternative definitions could be employed. For instance, optimal transport-based metrics, such as Dynamic Time Warping (DTW), can prove useful when observations come at a poorly known or controlled time pace. While the way the prediction from the model is compared with observational data is pivotal, the focus in this work is instead on the relevance of the training data. The present findings are believed to hold disregarding of the definition of the loss. A standard $L^2$ distance, implicitly assuming regularly sampled measurements, is hence retained.

The measurement operator is the identity, $h \equiv I$, so that the measurements are associated with the state of the system, $s_k \equiv u_k$, $\forall k$. No noise affects the observations, so that one is in a full observability situation. In this context, the observed system is Markovian in the representation space of the observations. However, a recurrent architecture is here considered as a general structure for learning, relevant for a wide class of situations. It further has the benefit of improving upon the robustness of the learning by virtue of some degree of redundant representation.

Without loss of generality, we disregard the regularization term $r (\theta)$ in the definition of the loss function which finally writes

$$
\mathcal{L} (\theta) = \sum_{k=1}^{K} \| s_k - v_k (\theta) \|_2^2, \quad \text{s.t.} \quad v_{k+1} = F_{\theta} (v_k), \quad v_0 = s_0.
$$

While this context is favorable for learning (low-dimensional deterministic system, noiseless observations, full observability, etc.), it will be seen below that the measurement strategy still significantly affects the learning performance.
IV. LEARNING THE LORENZ’63 SYSTEM

A. Presentation

To illustrate the different learning strategies, we consider the celebrated Lorenz’63 system [23], originally introduced to mimic the thermal convection in the atmosphere. It consists of three coupled nonlinear ordinary differential equations (ODEs):

\[
\begin{align*}
\dot{x} &= \sigma(y - x), \\
\dot{y} &= \rho x - y - xz, \\
\dot{z} &= xy - \beta z,
\end{align*}
\]

The system of ODEs is integrated in time with a RK4 scheme and a time step \( \Delta t = 0.01 \). Constants \( \sigma, \rho, \beta \) are positive parameters, related to dimensionless scales such as the Prandtl and the Rayleigh numbers. For \( \rho > \sigma (\sigma + \beta + 3) / (\sigma - \beta - 1) \) and \( \sigma > (\beta + 1) \), the solution \( u := (x, y, z) \) exhibits a chaotic behavior. It is important for the following to note that the system features three unstable fixed points \( u_{FP} \), associated with a vanishing time-derivative:

\[
\begin{align*}
u_{FP_+} &= \left( \sqrt{\beta(\rho - 1)}, \sqrt{\beta(\rho - 1)}, \rho - 1 \right), \\
u_{FP_-} &= \left( -\sqrt{\beta(\rho - 1)}, -\sqrt{\beta(\rho - 1)}, \rho - 1 \right), \\
u_{FP_0} &= (0, 0, 0).
\end{align*}
\]

In the following we shall consider the parameters fixed to \( \beta = 8/3, \sigma = 10, \rho = 28 \) as in the original work, such that the solution is in the chaotic regime. In this regime, the Kaplan-Yorke dimension of the attractor is \( d_A \approx 2.06 \).

B. Set-up of the learning method

We turn our attention to the training set \( T = \{s_k\}_{k=1}^K \) for learning the Lorenz’63 system. It is a crucial ingredient of the learning process and largely conditions the quality of the resulting model. We consider four different training sets (\( T_{\text{ergo}}, T_{\text{split}}, T_{\text{ens}}, \) and \( T_{FP} \)) of identical settings (sampling frequency, number of samples, measurement operator, etc.) but following different sampling strategies.
1. **Ergodicity-compliant training set**

The $\mathcal{T}_{\text{ergo}}$ dataset is a long trajectory on the attractor thought to act as a reference, providing sufficient statistics for learning a reliable and accurate model. With $\varepsilon = 10^{-2}$ and the Kaplan-Yorke dimension $d_A \simeq 2.06$ of the Lorenz’63 for the retained settings $\{\sigma, \rho, \beta\}$, the required number of samples resulting from a constant sampling period of $\Delta t = 0.01$ is about 27,000.

Another dataset, $\mathcal{T}_{\text{split}}$, is derived from the previous dataset, and consists of the trajectory on the attractor, yet split in 9 chunks of equal size and reshuffled, such that each chunk is thus composed by 3000 samples. As it will be clear soon, this dataset will allow us to investigate some effects of memory.

2. **Fixed points, $\mathcal{T}_{\text{FP}}$**

The training set $\mathcal{T}_{\text{FP}}$ exploits prior knowledge on the structure of the system at hand, namely the fixed points. As mentioned in the Introduction, it is well known dynamical systems are organized around critical features, such as fixed points, and that invariant sets provide information on the behavior of the solution, [13]. Besides fixed points, these features include higher-dimensional invariant sets such as slow invariant manifolds. The sampling strategy here consists in acquiring observations from the system originating from its (unstable) fixed points, in contrast with other training sets obtained from the attractor. Specifically, three 3000-sample trajectories are acquired from each of the three fixed points of the Lorenz’63 system (see Sec. IV A). These trajectories originate from a small deviation from each fixed points along each of the $d = 3$ directions given by the eigenvectors of the local Jacobian matrix. Note that if the Jacobian is not available, a set of orthonormalized deviations can also be used. The resulting training set then consists of $3 \times 3 \times 3000 = 27,000$ samples, consistent in size with the other training sets illustrated in Fig. 2.

3. **Ensemble sampling, $\mathcal{T}_{\text{ens}}$**

With this strategy, the dynamical system is sampled along short trajectories on the attractor. Nine trajectories are considered, each originating from a state randomly chosen on the attractor. Each trajectory is 3000 sample long as for the other strategies introduced
FIG. 2: The figure qualitatively illustrates the different training sets used in this investigation. Each $\mathcal{T}_{\text{ergo}}$ dataset is composed by one long trajectory (a). The $\mathcal{T}_{\text{FP}}$ and $\mathcal{T}_{\text{ens}}$ datasets are composed by an ensemble of 9 trajectories, of which one example for each is shown in (b) and (c), respectively.

above so that the total size of the training set is again 27,000 samples, similar to the reference training set $\mathcal{T}_{\text{ergo}}$. This training set is representative of the widespread situation where information about a system comes in several independent glimpses, as resulting from several measurement campaigns. It is illustrated in Fig. 2.

V. RESULTS

We now assess the quality of the models learned from various observation datasets of the Lorenz’63 system.

A. How much training information?

The influence of the amount of information to train a model is first considered. Two models with the same architecture, cf. Sec. III, are trained from two different datasets. These datasets consist of observations of the solution of the Lorenz system along the same trajectory on the attractor, but of different length. One dataset, $\mathcal{T}_{\text{ergo}}$, comprises 27,000 samples while the second one $\mathcal{T}_{\text{short}}$ is its restriction to the first 3000 samples, so that $\mathcal{T}_{\text{short}} \subset \mathcal{T}_{\text{ergo}}$. Once trained, the resulting models, hereafter respectively termed $F_{\text{ergo}}$ and $F_{\text{short}}$, are evaluated in terms of their long-term prediction capability.
Results are gathered on Fig. 3. The performance of the models is here evaluated when predicting from a known (i.e., contained in their training set) and unknown (not contained) initial condition, to assess the generalization capability of the models. The model is the LSTM architecture described in Sec. III A, with a null initial memory.

A known initial condition is first considered. As can be seen from the top row (left column) where it is plotted against the truth (red), the $F_{\text{short}}$ model allows a reasonable quality of the prediction (plotted in blue) over a significant time horizon of about 6 Lyapunov times (7 time units). Since training the models is achieved via a stochastic procedure, e.g. stochastic gradient descent and random initialization of the parameters, and the landscape of the loss function is not globally convex, the underlying optimization problem ends up in a local optimum and bears an aleatoric contribution. To strengthen the above conclusions, we hence consider an ensemble of 100 identical independent models and evaluate their performance. The right column of Fig. 3 shows the mean and standard deviation of the prediction of the ensemble of models for the same conditions as in the left column. From the top row, it is clear that the prediction of the ensemble of models is very consistent and that the learning is robust.

With the same short training set $T_{\text{short}}$, the trained model is now evaluated on an initial condition not contained in the training data, see Fig. 3 middle row. Compared with the previous situation (top row), the performance is seen to deteriorate, both for a single model (left column) and in average (right). In particular, the time horizon before the prediction significantly differs from the truth is much shorter, indicating a poor generalizability properties.

When more training data are used, $T_{\text{ergo}}$, the resulting model remains accurate over a long time horizon even when predicting from an unknown initial condition, as seen from Fig. 3 (bottom row). Again, this is a reliable finding as evidenced from the ensemble performance seen on the right column.

These results illustrate the fact that the training data, relied on to learn a model, have to be informative enough of the system under consideration to allow for generalizable models to be inferred. In particular, the superior performances shown in the top row, when the initial conditions are contained in the learning dataset, appear to be due to overfitting.
FIG. 3: Left column: Time-evolution of the $y$-component of the Lorenz’63 system, both from the true (true, solid red) and the learned model (prediction, dashed blue) from different training sets and initial conditions. Right column: zoom on the $[0, 2]$ time units interval, with the mean prediction (dashed blue line) from an ensemble of 100 trained models, and the associated 1 standard deviation. Top row: The prediction of the best (out of the ensemble set) model trained from $T_{\text{short}}$ is shown. The initial condition is drawn from the training set. The prediction is seen to remain reliable up to about 4.3 Lyapunov times (5 time units) before drifting significantly from the truth. Middle row: When the same model is evaluated with an unknown initial condition (test set), the performance deteriorates and the accuracy is lost after less than 4 time units. When a unique ergodic-compliant trajectory $T_{\text{ergo}}$ is used to train a model (bottom row), the prediction from an unknown initial condition improves significantly.

B. Relevance of the data

Besides the mere size of the training set, a critical aspect is its relevance for informing a model in view of a given objective function. To shed light on this pivotal aspect, the four different training sets introduced in Sec. III B, namely $T_{\text{ergo}}$ (“Ergodic”), $T_{\text{split}}$ (“Ergodic split”), $T_{\text{ens}}$ (“Random”) and $T_{\text{FP}}$ (“Structured”), all of the same size of 27,000 samples, are now evaluated in terms of the quality of the associated resulting model. To alleviate any bias due to the initialization of the parameters, an ensemble of 100 models is trained for
FIG. 4: The $d_2$ dimension is here adopted as a metric for assessing the prediction quality of the 100 models trained with each of the 4 training strategies $\mathcal{T}_{\text{ergo}}$, $\mathcal{T}_{\text{ens}}$, $\mathcal{T}_{\text{FP}}$, and $\mathcal{T}_{\text{split}}$; the resulting $d_2$ dimension is estimated by analysing time predictions from trajectories not belonging to the training set. For each training set, the proportion of models associated with an estimated dimension differing by more than 25% from the true dimension $d_A = 2.06$ is indicated in percent.

A first metric to quantify the quality of the resulting model is the $d_2$-dimension of the predicted model. This criterion is not associated with a given initial condition and is affected by the way the trained model consistently reproduces the fractal nature of the Lorenz’63 system. Results are gathered in Fig. 4 in terms of a histogram of the estimated $d_2$ dimension for the ensemble of 100 models for each of the four training sets. For a more quantitative appreciation, a threshold is defined when the dimension resulting from the trained model differs by more than 25% from the truth $d_A = 2.06$. As expected, the ergodicity-compliant training set $\mathcal{T}_{\text{ergo}}$ leads to a good performance, with most models associated with a $d_2$ dimension close to the true value and a few degenerate models such that $d_2 \approx 0$. Only 14% of the models are beyond the threshold. Models trained from $\mathcal{T}_{\text{FP}}$ are also seen to achieve a similar level of performance. Again, almost all these models are associated with a rather good $d_2$ dimension, with very few degenerate models. Only 8% of the models are beyond the threshold, an even better performance than models trained from $\mathcal{T}_{\text{ergo}}$.

In contrast, the $\mathcal{T}_{\text{ens}}$ (random) and $\mathcal{T}_{\text{split}}$ (split) dataset leads to a rather poor performance model, with a significant part of the models associated with $d_2 \approx 0$ and respectively 34 and 26% of the models beyond the threshold. It may be not quite surprising that models.
(a) t-distributed stochastic neighbor embedding (t-SNE) of the parameters of the trained models analyzed in Fig. 4. Each dot corresponds to a different model and its size visually indicates the accuracy: the larger the dot, the larger the error in respect of $d_A$.

(b) Radial distribution of the t-SNE projected model parameters. $r$ indicates the distance from the barycenter of the cluster in Fig. 5a while the distribution of the models is reported on the vertical axis using the same number of bins. We order from light cyan to darker blue models performing progressively worse in terms of resulting $d_2$ accuracy; note that the level set of the blue gradient is normalized for each subplot, such that only a qualitative information on the distribution along $r$ is included in the graphs.

FIG. 5: Illustration of the model parameters via a t-SNE representation. The LSTM memory is initialized to zero.

trained with $\mathcal{T}_{\text{ens}}$ and $\mathcal{T}_{\text{split}}$ are associated with a similar performance. Since the only difference between the two datasets is that the samples in $\mathcal{T}_{\text{ens}}$ originate from a collection of 9 independent segments of trajectory, randomly located on the attractor, while $\mathcal{T}_{\text{split}}$ is made of 9 contiguous segments. Yet, such behavior for the $\mathcal{T}_{\text{split}}$ dataset appears to be not trivial, because it contains exactly the same information of the $\mathcal{T}_{\text{ergo}}$ one about the phase space.

Further assessment of these observations can be made using a different technical tool,
namely t-distributed stochastic neighbor embedding (t-SNE) of the parameters, shown in Fig. 5a. It is essentially a dimensionality reduction technique convenient for assessing high-dimensional quantities \[31\]. The model parameters vectors are here mapped onto a 2-dimensional space where each model can hence be represented in terms of its corresponding coefficients \(\gamma_1\) and \(\gamma_2\).

\(F_{\text{ergo}}\) and \(F_{\text{FP}}\) are mostly clustered in the center of the points cloud, indicating that, while the models are different, they are however similar to each other in terms of parameter distribution. Instead, the models obtained from the random sampling strategy (\(T_{\text{ens}}\)) and the ones obtained stacking chunks from an ergodicity-compliant trajectory (\(T_{\text{split}}\)) are distributed in the outer region.

LSTM parameters are initialized as independent realizations of a Gaussian random variable, whose t-SNE representation is typically visually close to a Gaussian as well since it penalizes deviation from a t-Student distribution. Recent works have shown that neural networks associated with parameters weakly evolving during training while yet able to significantly improve the loss function, are often associated with good generalization properties, \[17\]. In the present study, the t-SNE representation of models trained from the ergodicity-compliant and the structured datasets both appear close to a Gaussian distribution, a further indication that they might enjoy good generalization properties. In contrast, the t-SNE distribution of the models trained from \(T_{\text{ens}}\) or \(T_{\text{split}}\) appear to deviate significantly from their initial distribution, which corroborates well with their poor performance.

This distribution indicates that the best models are robust, in the sense of having similar parameters. Furthermore, the training using the information about fixed points appears equivalent to that based on a long trajectory. This observation is also quantified in terms of a radial distribution, see Fig. 5b. Both the models trained from \(T_{\text{ergo}}\) and \(T_{\text{FP}}\) present a distribution most dense around zero, while \(T_{\text{ens}}\) and \(T_{\text{split}}\) lead to a bias in the parameter distribution, hence being associated with poor models.

C. Handling the memory

The role of the memory in the recurrent model is now studied. Instead of the null memory (initialized to zero) considered so far, an initial memory chosen at random, drawn from a multivariate Gaussian distribution \(\mathcal{N}(0, \mathbf{I})\) is used. Compared to the null initial memory, the
performances are seen to significantly increase for all models, as evidenced in Fig. ???. This is consistent with the fact that a non-zero hidden state improves the numerical conditioning of the training steps and prevents degenerate gradients, [11].

The contrast in performance between models learned from $\mathcal{T}_{\text{ergo}}$ and $\mathcal{T}_{\text{split}}$, which rely on the same information, leads to identify a key point. The LSTM model trained with $\mathcal{T}_{\text{ergo}}$ learns to best fit one long trajectory. In the case of $\mathcal{T}_{\text{split}}$, it learns to best fit 9 short trajectories, which however, when stitched together, match the $\mathcal{T}_{\text{ergo}}$ long trajectory. Yet, their performances are consistently very different in the null initialization situation, emphasizing that the initial memory of the retained recurrent network (LSTM) dramatically affects the resulting performance. In the case of $\mathcal{T}_{\text{ergo}}$, the impact of an incorrect initial memory is diluted within a long trajectory and weakly affects the learning quality. When short trajectories are considered instead, the relative impact of an incorrect initialization is stronger and can introduce a significant bias, hence a poor resulting model.

This observation is further supported by the model learned from $\mathcal{T}_{\text{FP}}$. While it also relies on samples originating from 9 distinct, non sequential, segments, its performance is good, as evidenced by Fig. [4] What is unique about this particular training set is that the samples are from segments of trajectories originating from (unstable) fixed points of the Lorenz’63. These points of special interest are associated with a locally linear dynamics, and this allows the incorrect initial memory to be progressively wiped out in time without significantly affecting the predicted dynamics. In this regard, fixed points are privileged training elements which allow to reliably learn memory-based models by initializing the training in a configuration where the memory does not have a harmful impact.

VI. CONCLUDING DISCUSSION

In this work, we have performed a thorough analysis of some aspects inherent to the process of learning a complex physical system from observational data. More specifically, we have considered the following issues: (i) What is the impact of the initial state of the retained model on its resulting performance? (ii) Is there a minimum amount of data needed to obtain a robust model able to generalize? (iii) Is it possible to go beyond this limitation thanks to some knowledge about the system?

The first issue is related to the objective of discussing best-practice guidelines for prac-
FIG. 6: The figure depicts the $d_2$ dimension associated with predictions obtained for models trained by initializing the LSTM memory at random, using the four different learning strategies. For each training set, the proportion of models associated with an estimated dimension differing by more than 25% from the true dimension $d_A = 2.06$ is indicated.

tioners. The others are more fundamental. The second is related to what is known for classical embedding approaches, which is bound to fail when time-series data are not enough to reconstruct the entire phase-space. The last one explores perspectives to find a physics-informed learning approach capable of overcoming some of the intrinsic limitations of data-driven supervised learning.

We have considered the classical Lorenz’63 system as the benchmark problem, because of its simplicity and hence the possibility to carry out an exhaustive campaign of simulations. Moreover, the system is known in details and allows to investigate the importance of singular points of the dynamics. For the learning method, we have considered an LSTM, an established recurrent structure often used in such context.

In these memory-based models, flushing an initially incorrect memory is crucial for obtaining a relevant and faithful model able to generalize its prediction beyond the sole situations encountered during the training step. This issue may be particularly severe if one learns from a set of short-termed observations. Our numerical experiments show clearly that there is an impact of the initialization of the memory on the quality of the predictions provided by the LSTM models. As a matter of fact, it should be always initialized as random to
avoid bias in the learning since, in all cases, we have obtained better performance in such a configuration. In particular, when the initial condition is random, we have found little difference in the results obtained by training on a long trajectory or using exactly the same data but randomly shuffled in several short trajectories, consistently with the problem of exploring the phase-space of a dynamical system.

Generally speaking, we have found that, in absence of prior information on the system at hand besides an estimated upper bound of its dimension, the ergodic theory, and notably the Kac lemma, provides a criterion on the minimal amount of data necessary to train a model. This amount of data grows exponentially with the dimension of the attractor. Models trained on a smaller amount of training data have been shown to consistently lead to poor performance, as illustrated both via the time-series prediction and the resulting model estimated dimension assessment criteria.

Remarkably, numerical experiments have neatly shown that starting from the fixed points of the dynamical system constituted a workaround in learning the dynamics of the system from situations where the memory is essentially harmless to the learning process. Furthermore, this choice leads to consistent and accurate models even with a decreased amount of data with respect to what prescribed by ergodic theory. The information of the structure of the dynamics appears therefore key to improve the learning process.

On-going efforts further explore the idea that an additional benefit of training a model from fixed points is that the resulting dataset encompasses a wide range of solution regimes, from a linear behavior close to the fixed points where the non-linearities are comparatively weak, to a full nonlinear regime when the state reaches the attractor. This essentially allows to disentangle the contribution of the different terms of the underlying system and to sequentially focus the learning effort on some solution regimes from specific parts of the training set. In this sense, this is reminiscent of a form of curriculum learning approach.

Another relevant perspective is to put additional efforts focusing on the impact of noise in the observations onto the learning quality. On one hand, the presence of noise may be a source of regularisation \[7\], notably in the loss function, and thus might be beneficial to the learning process. On the other hand, adding noise would imply a fairly large parametric space to be analyzed, with non-trivial consequences. Moreover, the distinction between noise and chaotic behavior in real data is often problematic \[4\], and generally speaking, the inverse problem of learning an unknown dynamics is even more challenging for stochastic
systems [22]. Providing an assessment of the learning of stochastic dynamics and novel ideas for improving it is a cutting-edge subject of research [35].

In conclusion, while we here focused on the Lorenz’63 system, we believe the findings of this work to be widely applicable and to provide some evidence-based good practice for data-driven modeling of physical systems. Work in progress concerns a finer analysis of the interplay between the structure of the dynamical system and the learning process in order to explain why fixed points are so informative, in the hope to propose an active learning strategy aimed at leveraging the underlying physics of a dynamical system.

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