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# The Neural Network shifted-Proper Orthogonal Decomposition: a Machine Learning Approach for Non-linear Reduction of Hyperbolic Equations

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## Abstract

Models with dominant advection always posed a difficult challenge for projection-based reduced order modelling. Many methodologies that have recently been proposed are based on the pre-processing of the full-order solutions to accelerate the Kolmogorov  $N$ -width decay thereby obtaining smaller linear subspaces with improved accuracy. These methods however must rely on the knowledge of the characteristic speeds in phase space of the solution, limiting their range of applicability to problems with explicit functional form for the advection field. In this work we approach the problem of automatically detecting the correct pre-processing transformation in a statistical learning framework by implementing a deep-learning architecture. The purely data-driven method allowed us to generalise the existing approaches of linear subspace manipulation to non-linear hyperbolic problems with unknown advection fields. The proposed algorithm has been validated against simple test cases to benchmark its performances and later successfully applied to a multiphase simulation.

*Keywords:* Deep Neural Networks (DNNs), shifted-POD (sPOD), Non-linear hyperbolic equations, Reduced Order Modelling (ROM), Multiphase simulation.

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## 1. Introduction

Reduced Order Modelling (ROM) is a well-established set of different numerical techniques whose objective is that of retrieving a low-rank representation of parametric differential models, s.a. Ordinary Differential Equations (ODEs) and Partial Differential Equations (PDEs), describing a relevant majority of models in physics and engineering [1, 2]. Lowering the computational cost of numerical simulations is a fundamental aspect of both industrial and academic research activities and therefore ROM techniques have always been, since their introduction, an important part of the modelling process. In its most general formulation, ROM deals with Initial Boundary Value Problems (IBVPs) in which

a PDE is parametrised by a finite set of  $P \in \mathbb{N}$  parameters. One such example is the following scalar, linear, first-order in space IBVP with non-parametric, steady Dirichlet's Boundary Conditions (BCs) and Initial Condition (IC)

$$\begin{cases} \partial_t u(\mathbf{x}, t, \boldsymbol{\mu}) + \mathcal{L}(u, \boldsymbol{\mu}) = 0, & \mathbf{x} \in \Omega \subset \mathbb{R}^d, t \in [0, T > 0], \boldsymbol{\mu} \in \mathcal{P}, \\ u(\mathbf{x}, t, \boldsymbol{\mu}) = g(\mathbf{x}), & \forall \mathbf{x} \in \partial\Omega, \\ u(\mathbf{x}, t = 0, \boldsymbol{\mu}) = u_0(\mathbf{x}), & \forall \mathbf{x} \in \Omega, \end{cases} \quad (1)$$

where  $d = 1, 2, 3$  represents the number of spatial dimensions of the model,  $T > 0$  is a final instant that identifies the interval in which the time evolution of the model is evaluated and  $\mathcal{P} \subset \mathbb{R}^P$  is the parameter domain of the PDE s.t.  $\dim(\mathcal{P}) = P$ . Following a domain discretisation of  $\Omega$  using e.g. Finite Volumes, Finite Elements or Finite Differences, one obtains a numerical approximation  $\mathbf{u}_h(t, \boldsymbol{\mu}) \in \mathcal{V}_h$  for the (discrete) time evolution of  $u(\mathbf{x}, t, \boldsymbol{\mu})$  as evaluated at timesteps  $t_k$ ,  $k = 0, 1, \dots$ , and where  $N_h := \dim(\mathcal{V}_h)$  represents the Degrees of Freedom (DOFs) of the problem. By defining the discrete solution manifold  $\mathcal{M}_h := \{\mathbf{u}_h(t_k, \boldsymbol{\mu}) \in \mathcal{V}_h, k \in \mathbb{N}_0, \text{ s.t. } \boldsymbol{\mu} \in \mathcal{P}\}$  one immediately recognises that a high-dimensional approximation space  $\mathcal{V}_h$  leads to an expensive computational cost whenever one wants to evaluate the time evolution of  $\mathbf{u}_h(t, \boldsymbol{\mu})$  for multiple instances of  $\boldsymbol{\mu}$  in the parameter space  $\mathcal{P}$ . Formally stated, ROM aims at reducing the computational cost of a (numerical) parametric simulation by constructing the best low-rank approximation  $\mathcal{R}$  of manifold  $\mathcal{M}$  s.t. it encodes the parameters variation of the Full Order Model (FOM) but with a reduced number of DOFs. One such example is that of evaluating the pressure field solution in response to several geometrical configurations of an airfoil parametrised by e.g. different angle of attack, chord line and mean thickness. Many different classes of ROM techniques have been developed for the low-rank reduction of several phenomena [3, 4, 5] and in particular projection based ROM have found widespread application in reducing different parametric differential problems modelling several phenomena in fluid dynamics. This class is based on the identification of a reduced set of basis functions, or modes,  $\{\phi_j\}_{j=1, \dots, R}$  s.t. their superposition spans the best possible low-rank approximation of the solution manifold i.e.  $\text{span}(\phi_1, \dots, \phi_R) = \mathcal{R} \approx \mathcal{M}_h$ . The Proper Orthogonal Decomposition (POD), on which this work is based upon, is one of those techniques in which the basis extraction is performed using Singular Value Decomposition (SVD) of a so-called snapshot matrix  $\mathcal{X}$  in which the FOM solutions  $\mathbf{u}_h(t, \boldsymbol{\mu})$  are stored as column vectors. When paired with a projection technique  $\Pi_{\mathcal{R}} \mathbf{u}_h(t, \boldsymbol{\mu}) \in \mathcal{R}$  s.a. the Galerkin projection the resulting POD-Galerkin method allows for fast computation of several simulations with different instances of  $\boldsymbol{\mu}$  in the domain  $\mathcal{P}$ . The efficiency of any ROM technique, including POD-Galerkin methods, stems directly from the number of basis functions needed to encode the parametric essential dynamics of a model at reduced order. The reduction accuracy of a model has been traditionally quantified by the Kolmogorov  $N$ -width of

the problem which is defined, in  $L_2$ -norm, as

$$\text{dist}_R(\mathcal{M}_h) := \inf_{\mathcal{R} \in \mathcal{R}_R} \sqrt{\frac{\sum_{\mathbf{u}_h \in \mathcal{M}_h} \|\mathbf{u}_h - \Pi_{\mathcal{R}} \mathbf{u}_h\|^2}{\sum_{\mathbf{u}_h \in \mathcal{M}_h} \|\mathbf{u}_h\|^2}}. \quad (2)$$

Problems that are characterised by a fast Kolmogorov  $N$ -width decay (here we used  $N \equiv R$  to be in line with the notation commonly found in the literature) are easily restricted by a low-rank linear subspace approximation; on the other hand a slow decay indicates that a linear approximation assumption might be inappropriate and inaccurate for that problem. For this reason, despite the popularity that POD-Galerkin methods gained in various areas of computational science and engineering [6, 7, 8, 9], they have been severely limited by the fact that the low-rank approximation of  $\mathcal{M}_h$  they build is sought within a sequence  $\{\mathcal{R}_R\}_{R=1, \dots, \infty}$  of  $R$ -dimensional linear subspaces. The assumption that a FOM can be accurately approximated by restricting its dynamics on a low-rank linear subspace proved in fact to be suitable for elliptic and parabolic problems (where the Kolmogorov  $N$ -width decays exponentially with  $R$  [10]) while it is not trivially extendable to problems with dominant advection such as (s.a.) those modelled by hyperbolic PDEs (for which the Kolmogorov  $N$ -width decay is much less steep [11]). It results that the reduction of advection-dominated PDEs with POD-Galerkin methods requires a large number  $R$  of basis functions to be accurately depicted as they struggle to restrict the FOM to a linear subspace approximation of  $\mathcal{M}_h$ , essentially nullifying the reduction in computational cost. Since models with dominant advection are particularly recurrent in fluid dynamics, being associated to the conservation laws describing a large multitude of phenomena (most notably Euler's equations and Riemann's problems, shallow water equations, multiphase models), a growing number of endeavours in recent years have proposed alternative modifications for the improvement in performances for the approximation of such manifolds [12, 13, 14]. The work that has been done is very heterogeneous in terms of the methodologies adopted, with many efforts that brought in the latest data-driven trends in machine learning and features extraction. Nevertheless the state-of-the-art in improving the accuracy of ROM of hyperbolic equations can be distinct into two major approaches: those based on ad-hoc transformations of the linear subspace as a support for better mode extraction of projection-based ROM [15, 16, 17, 18, 19] and those based on a construction of non-linear manifolds [20, 21, 22, 23]. The former class has been the first to be approached while the latter strategy is only most recently being explored to overcome the limitations of the earlier approaches. Among the various reasons there is the fact that the underlying methodology of pre-processing the linear subspace has been that of embedding the dominant advection of the solution at offline stage using a deterministic transformation. This requires some sort of problem-specific prior knowledge of the physical or mathematical properties of the FOM e.g. the characteristic speeds in phase space of a conservation law [24, 25]. This obviously limits the range of applicability of pre-processing based model reduction to deal with hyperbolic PDEs that feature simple advection

fields. To overcome such shortcomings the second class of methods based on deriving non-linear manifold reduction directly has been recently favoured with important results being obtained with the adoption of convolutional autoencoders [26] and Arbitrary Lagrangian-Eulerian (ALEs) frameworks [27, 28]. In this work we introduce a novel statistical learning framework for the generalisation of the earliest pre-processing techniques to those hyperbolic problems that feature non-linear and unknown advection fields. The resulted Neural Network shifted-Proper Orthogonal Decomposition (NNsPOD) algorithm is a fully non-intrusive, purely data-driven machine learning **offline** method that seeks for an optimal mapping of the various snapshots in the low-rank linear subspace to a reference configuration via an automatic detection that does not depend on the physical properties of the model. By construction our algorithm does not belong exclusively to either of the aforementioned classes of manifold reduction. While NNsPOD starts with the goal of building a pre-processing transformation that accelerates the Kolmogorov  $R$ -width decay for advection-dominated problems, it also achieves a virtually unlimited range of applicability to non-linear manifold reduction since it requires no prior knowledge on the properties nor complexity of the FOM. **In this work we propose NNsPOD as an efficient, pre-processing method that can be deployed for the offline of a projection-based ROM; a full ROM NNsPOD-Galerkin method will be discussed in future works where we address the practical advantages and computational issues associated to project the governing non-linear hyperbolic equations onto a low-rank manifold constructed independently from the physics of the problem.** During the development of our algorithm we became aware of a project being proposed in [29] which achieves the similar result of non-linear manifold reconstruction of transport-dominated problems via deep-learning models. Despite being both projection-based and physics-independent, our methodology substantially differs from the one proposed in [29]. **The approach undertaken in [29] is based on deriving a low-rank manifold generated via adaptive reduced basis; as remarked by the authors, those basis have the unique feature of carrying the parametric and time dependence of the PDE, which NNsPOD does not. The introduction of the two deep learning models is used to output two separate low-rank vectors whose inner product represents the non-linear approximation of the FOM  $u_h(\mathbf{x}, \boldsymbol{\mu}, t) \approx u_{NN}(\mathbf{x}, \boldsymbol{\mu}, t; \boldsymbol{\theta}) = \boldsymbol{\alpha}(\boldsymbol{\mu}, t; \boldsymbol{\theta}) \cdot \boldsymbol{\varphi}(\mathbf{x}, \boldsymbol{\mu}, t; \boldsymbol{\theta})$ . The training of the two networks will optimise the parameters  $\boldsymbol{\theta}$  in such a way that the retrieved set of adaptive reduced basis generators spans a low-rank manifold characterised by a faster Kolmogorov  $R$ -width decay than that of a traditional POD-Galerkin method. Our methodology is instead focused in automating a well-established technique of backward shifting the FOM solutions to a reference configuration by means of two independent networks that, although trained separately, are nevertheless joined together in the construction of a non-linear pre-processing operator. We argue that the methodology that we propose generalises better to those advection-dominated problems that feature unknown non-linear transports, s.a. the case for multiphase models. The underlying reason for our statement is that NNsPOD does not explicitly embed any parametric dependence on the reduced basis generators, which are therefore unaware of the paths taken to be mapped**

onto the reference configuration. This non-intrusive approach entails a stronger validity on the fundamental ansatz of ROMs  $\mathbf{u}_h(\mathbf{x}, \boldsymbol{\mu}, t) \approx \sum_{j=1}^R u_j(\boldsymbol{\mu}, t) \boldsymbol{\varphi}_j(\mathbf{x})$  in which the low-rank manifold generators  $\boldsymbol{\varphi}(\mathbf{x})$  only carry spatial dependence. The generalisation at online stage provided by NNsPOD will therefore fully encode the information regarding the parametric dependence of the transport field, with the coefficients  $u_j(\boldsymbol{\mu}, t)$  being determined by the Galerkin projection of the governing equations onto the low-rank manifold. We also emphasize that all the benchmarks that follow in the present work are 2-dimensional problems as opposed to the 1-dimensional problems presented in [29]. The present work is structured as follows: in Section 2 we contextualise the state-of-the-art of linear subspaces transformations while also showing the limitations of traditional POD-Galerkin reduction for a canonical problem with dominant advection; in Section 3 we present the methodologies adopted by NNsPOD as a generalisation of such techniques to unknown and non-linear transport fields; in Section 4 we demonstrate the performances of NNsPOD as a result of its application to a reduction of a multiphase model; in Section 5 we conclude by highlighting the outstanding challenges of our approach while also providing some insights on practical limitations we encountered during the development and test stage of our algorithm.

## 2. Pre-processing transformation of POD linear subspaces

In order to introduce the new methodology proposed in this work we shall contextualise the common approach shared by all the previous works that have been done in linear subspace pre-processing of advection-dominated models. To that end we start by formally introducing the standard procedure of a traditional POD-Galerkin technique and highlight its under-performance with a simple hyperbolic equation.

### 2.1. Full-order scalar advection equation

Let us consider a simple test case of the IBVP in (1) for which  $\mathcal{L} = \nabla \cdot \mathbf{F}$  i.e.

$$\begin{cases} \partial_t u(\mathbf{x}, t) + \nabla \cdot \mathbf{F} = 0, & \mathbf{x} \in [0, 1] \times [0, 1] \equiv \Omega, t \in [0, T = 1], \\ u(\mathbf{x}, t) = 0, & \forall \mathbf{x} \in \partial\Omega, \\ u(\mathbf{x}, 0) = \exp(-\frac{1}{2} \mathbf{x}^T \mathbf{x}), & \forall \mathbf{x} \in \Gamma \subset \Omega, \end{cases} \quad (3)$$

which is a linear first-order advection equation in a scalar unknown  $u(\mathbf{x}, t)$  that is null everywhere over a unitary square domain except for a bounded region  $\Gamma$  in which varies as a multivariate Gaussian pulse. For the sake of simplicity we set (3) to have only one parameter which is the time variable itself; this entails that  $P = \dim(\mathcal{M}_h) = 1$  which in turn means that we expect to retrieve a sufficiently accurate linear subspace approximation of the FOM with only  $R = 1$  reduced basis vector.

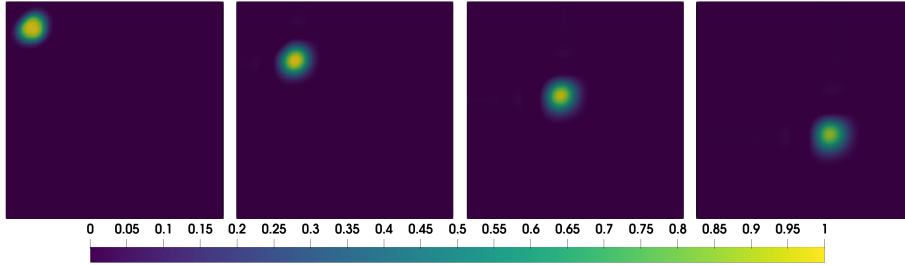


Figure 1: Different snapshots of the FOM simulation in (3) sorted left to right from the IC to increasing timesteps.

We know however that this is not the case since (3) is an hyperbolic PDE. To test the inability of traditional projection-based ROM at restricting the FOM to a linear subspace let us set a computational grid of  $50 \times 50$  points in  $\Omega$  while we discretise the time interval into 100 uniform steps to comply with the Courant–Friedrichs–Lewy (CFL) condition. We also set the vector field  $\mathbf{F} = \mathbf{f} u$  to be a linear flux function that is uniform in space and constant in time with the advection field being  $\mathbf{f} = (1, -1)$ . We obtain a FOM numerical solution  $\mathbf{u}_h(t) \in \mathcal{V}_h$ ,  $N_h = 2500$  using a Finite Volume (FV) spatial discretisation scheme, while the time evolution over the discrete timesteps is obtained via an unconditionally stable, first-order accurate, implicit Euler method. For the sake of simplicity we will not discuss the theoretical background of both numerical methods as it is out of the scope of the present work; the reader may refer to [30, 31, 32] for a comprehensive reading on standard algorithms in introductory and advanced numerical analysis. In order to present a particularly challenging advection model to our POD-Galerkin reduction method, we reduce as much as possible the numerical diffusion given by low-order cell face interpolation; as such a third-order accurate QUICK scheme [33] is adopted with the numerical stability being assured by the combined implicit Euler time advancing method and CFL-complying timesteps. Different FOM snapshots of this simulation are depicted in Figure 1.

## 2.2. The POD method for reduced basis extraction

The standard procedure of POD reduction is hereby outlined. The  $N_s = 100$  snapshots, sampled in the 1–dimensional FOM manifold during the simulation, are sorted by timestep in increasing order and stored as column vectors of a  $2500 \times 100$  snapshot matrix. We can find the best low-rank approximation of  $\mathcal{X}$  via SVD as per the Eckart–Young theorem [34]

$$\mathcal{X} = (\mathbf{u}_h(t_1 = 0), \dots, \mathbf{u}_h(t_{100} = T)) \approx \mathbf{L}\mathbf{\Sigma}\mathbf{R}, \quad \mathbf{L} \in \mathbb{R}^{N_h \times R}, R = 1, \dots, N_s. \quad (4)$$

The left singular vectors in  $\mathbf{L}$  are those that are associated with any subspace approximation  $\mathcal{R}_R$  of  $\mathcal{M}_h$ ; in Figure 2 we depict the first 8 of those. As specified in 2.1, a linear combination of  $R = P = 1$  of those vectors should, theoretically, retrieve a reduced order model with high accuracy.

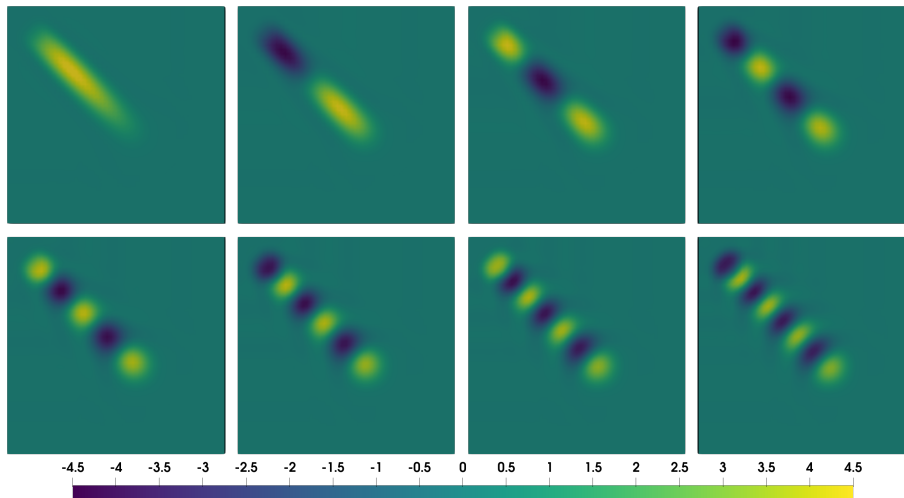


Figure 2: First 8 modes extracted via POD reduction from the FOM in (3) sorted in ascending order from top to bottom and from left to right.

However if we set such threshold to be e.g.  $10^{-3}$  we immediately realise that such degree of accuracy is obtained with a reduced basis of 14 left singular vectors. This model clearly cannot be constrained easily in a linear subspace as produced by a traditional POD method.

### 2.3. Linear subspace shift-based transformation

In order to improve the approximation of  $\mathcal{M}_h$  via linear subspaces, different methods have been proposed in the most recent years; the majority of the them started to focus in deriving an ad-hoc transformation of the linear subspace in a frame of reference that facilitates the decay of the Kolmogorov width. This shift-based pre-processing is in fact consistent with the hyperbolic character of models with dominant advection which are solved at full-order, both analytically and numerically, by exploiting the characteristics curves in phase space of the solution. Therefore, as long as the characteristic speeds of an hyperbolic PDE are known, one can pre-process the snapshots of the FOM in  $\mathcal{X}$  by e.g. mapping them to the IC following backward those curves in phase space. This approach therefore aims at finding a better frame of reference for the SVD of the POD by transporting the FOM snapshots by means of: backward-shift transformation [24, 25] and successive interpolation; displacement interpolation technique [15] or shock-fitting methods [17]; locally adapted bases [35]. In the following we will focus on the first of those linear subspace pre-processing methods. As explained in [24] one builds a discrete shift operator  $T_{\mathbf{b}}$  that acts on a space-time dependent function  $u(\mathbf{x}, t)$  transporting its frame of reference by an amount that is proportional to the transport field  $\mathbf{b}$ . The  $k$ -th FOM snapshot in (4) is an  $N_h$ -dimensional vector; each of its component store the field value of the numerical solution associated to the  $x$  and  $y$  coordinates of a centroid  $\mathbf{x}$



on the computational grid following the FV discretisation. Applying the shift operator to such snapshot amounts to deriving a new spatial distribution of those centroids for the same field values i.e.

$$T_{\mathbf{b}} \mathbf{u}_h(t_k) = \tilde{\mathbf{u}}_h = (u_h(\mathbf{x}_j - \mathbf{b}t_k, 0))_{j=1, \dots, N_h}, \quad \forall k = 1, \dots, N_s. \quad (5)$$

Furthermore an interpolation of  $\tilde{\mathbf{u}}_h$  is necessary in order to reconstructs the FOM field solution starting from the shifted points distribution on the manifold; the transformed snapshot matrix is thus decomposed in the new frame of reference by SVD. As depicted in Figure 3 in fact, a shift transformation on the snapshots allows for a better approximation with just 4 modes needed to span a linear subspace that is within the same  $10^{-3}$  accuracy threshold set in 2.2 for  $\mathcal{M}_h$ .

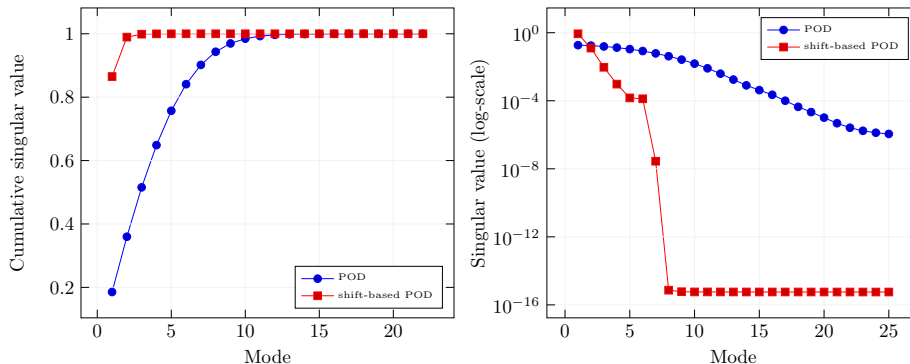


Figure 3: Comparative accuracy of traditional POD and shift-based pre-processing POD for the same advection model in (3).

#### 2.4. Limitations in deducing the shift operator

The stated claim in [24] is to build a data-driven, shift-based, pre-processing transformation that identifies the dominant structures of a hyperbolic model regardless of the characteristic velocities, with the goal of extending the methodology to non-linear advection fields. The authors also add that in order to identify those *ansatz* modes, whenever the phase-space velocities are unknown explicitly, two different approaches might be considered. One approach is based on the tracking of the peaks in the field values of the numerical solution across different snapshots; this method is however unfeasible for hyperbolic models that do not preserve the time or parameter dependent shape of the IC across their time evolution (e.g. multiphase simulations as we address in Section 4). The other suggested methodology, that is actually discussed in [24], is to perform multiple SVDs with sampled values of the shift velocities; in a data-driven fashion one should thus be able to detect the correct backward shift transformation by examining the singular values spectrum and isolate those for which the decay is maximised. This approach is indeed feasible to deal with hyperbolic PDEs that feature unknown transport velocities. However we identify two major limitations:

- The examination of the singular value spectrum as a function of trial shift operators entails an unnecessary increase in computational complexity during the offline stage that scales polynomial with  $d = 1, 2, 3$ . This is due to the fact that multiple snapshot matrices, and subsequent SVD, are required in order to detect the appropriate spatial transformation that maximise the Kolmogorov width decay since, in order to achieve a truly data-driven degree of flexibility, no prior information on the physical behaviour shall be used to sample the velocities for the trial shift operators.
- The various guesses for candidate shift operators are all, as stated by the authors in [24], limited to uniform and constant advection fields, although the shift itself can easily be computed pointwise in the grid and at different timesteps. This introduces a further degree of approximation and loss in accuracy for the case of non-linear hyperbolic PDEs; in fact guessing linear, constant and uniform shift velocities essentially amounts in linearising those characteristic curves in phase-space that may present non-trivially integrable irregularities.

Our aim is that of generalising the shift-based pre-processing linear subspace approximation by overcoming these aforementioned limitations in performance, computational scalability and models of advection. In our approach we in fact retrieve a non-linear transformation that achieves the same results in [24] but with an automatic detection that is both consistent with the method of characteristics and also robust to be applied to hyperbolic PDEs with unknown and non-linear advection.

### 3. Automatic detection of bijective mapping in non-linear manifolds

The traditional fields of application for statistical learning models, since their renewed development, have been data-analysis and data-mining, signal processing and computer vision. In the last years however they found widespread implementation in different subsets of scientific computing as well [36]. Industrial-driven requirements in improving the computational cost of numerical simulations in engineering, natural and life sciences resulted in an ever increasing integration of machine-learning algorithms within the more "traditional" numerical methods. The introduction of Artificial Neural Networks (ANNs), and specifically Deep Neural Networks (DNNs), further encouraged this coupling by providing purely data-driven non-linear mappings. ROM has been no exception to this merge of disciplines and the similarities between POD and Principal Component Analysis (PCA) is one example that justifies the natural blending of one set of techniques into the other. Given the extensive presence of complex non-linear PDEs, s.a. those modelling turbulent regimes [37], Computational Fluid Dynamics (CFD) represented the quintessential field of experimentation for ANNs integration in scientific computing with the first endeavours [38] being introduced as early as 2003. The DNN-based algorithm we hereby propose has been developed along those directions outlined above, that is by integrating, in an efficient but mathematically consistent process, a statistical learning

paradigm into a traditional ROM technique s.a. POD. The main limitation in detecting the appropriate backward transformation for shift-based POD reduction of models like (3) is that prior knowledge of the characteristic velocities is necessary in order to quantify the pre-processing map. A data-driven approach based on singular values spectrum analysis does not automate the process as, in order to be efficient, a proper sampling of the velocity space has to be performed. Furthermore if there is a parametric dependence of such transport field, i.e.  $\mathbf{b} = \mathbf{b}(\boldsymbol{\mu}, t)$ , then the Galerkin projection will not generalise well during the online phase since the pre-processed manifold does not contain the information for the evolution of the numerical solution along different characteristic curves in phase-space. The approach taken in previous works s.a. [24, 16] can thus be modelled as follows

$$T_{\mathbf{b}} : \mathcal{M}_h \mapsto \tilde{\mathcal{M}}_h, \tilde{\mathcal{M}}_h := \{\mathbf{u}_h(\mathbf{x}_j - \mathbf{b}(t_k, \boldsymbol{\mu}_p), t_k), \forall k = 1, \dots, N_s, p = 1, \dots, N_p\},$$

i.e. finding a backward map  $T_{\mathbf{b}}$  in which the  $N_p$  parametric instances of the transport field at offline stage is embedded. This method changes the frame of reference of the solution manifold making difficult its generalisation at online stage. A deep-learning framework on the other hand can indeed derive a bijective transformation between the shifted manifold (of which the linear subspace approximation is build) and the original one by ignoring such parametric dependence for the advection field  $\mathbf{b}$ . This motivates the needs for the adoption of a non-intrusive technique for the computation of the backward map based on automatic shift-detection i.e. a machine learning algorithm that does not quantify the shift operator neither based on the pointwise value of the transport field nor a linear approximation of it. [Here the bijectivity property of the transformation detected by NNsPOD refers to the fact that each FOM snapshot  \$\mathbf{u}\(\mathbf{x}, t\) \in \mathcal{M}\_h\$  is associated to a unique shifted approximation of the reference configuration  \$\mathbf{u}\_{\text{ref}} \in \tilde{\mathcal{M}}\_h\$ . Our methodology in fact derives a transformation  \$\mathcal{T}\$  that is therefore more general w.r.t.  \$T\_{\mathbf{b}}\$  constructed in \[24\] which instead maps every FOM solution in the snapshot matrix to the IC. The data-driven algorithm that will be introduced in the following section can be thought of a transport-field independent transformation mapping the arbitrary points in the solution manifold to a significantly smaller region of the output shifted manifold consisting of an arbitrary small neighborhood centered in  \$\mathbf{u}\_{\text{ref}}\$ . The optimised, automatically detected, non-intrusive transformation is therefore a bijective map](#)

$$\mathcal{C}^1 \ni \mathcal{T} : \mathcal{M}_h \mapsto \tilde{\mathcal{M}}_h,$$

which does not linearise the characteristic velocities in phase-space. In the development of NNsPOD we focused in assuming that no prior information regarding the functional dependence of  $\mathbf{b}$  is known as we seek to derive a bijection that traces any snapshot in  $\mathcal{X}$  to an arbitrary [small neighborhood of the reference configuration  \$\mathbf{u}\_{\text{ref}} \in \tilde{\mathcal{M}}\_h\$](#) . It might seem trivial to choose the IC as reference configuration; however, as we will explain further on in the section, given the general framework in which NNsPOD was conceived, we will consider also other snapshots as reference configuration for improving the bijective

map. NNsPOD being a data-driven, pre-processed ROM it should also provide a consistently fast online phase for parametric formulation of time-dependent hyperbolic PDEs. We postpone the results of such models, as well as the integration of NNsPOD with a Galerkin projection, to our future works; in the present work we only highlight the generalisation of the effectiveness of shift-based linear approximations in POD reduction methods for particularly difficult advection-dominated models with unknown and non-linear transport fields (as presented in Section 4).

### 3.1. Statistical learning formulation and reference configuration

Our framework is naturally implemented as a statistical learning technique in which the objective is to detect automatically an optimal transformation of the solution manifold  $\mathcal{M}_h$  that increases the Kolmogorov width decay associated to its model. If a DNN is deployed to build such bijective map while it merely relies on the snapshots collected from FOM data, then the desired requirements are:

- It must preserve the hyperbolic nature of the PDE consistently with the dominant advection model: NNsPOD has to be flexible enough to output transformations that have to be, in principle, rigid transports of any collected snapshot to a reference configuration of choice without adding numerical diffusion while, at the same time, being able to also change the shape of those snapshots if it is required (e.g. multiphase simulation we will discuss in Section 4).
- The data-flow for the two phases of shift-detection and field-reconstruction (interpolation), as outlined in [24] has to be continuous: in order for the backward map to be generalised for the Galerkin projection the non-linear transformation provided by NNsPOD has to be invertible for any input in the training set and as such a continuity constraint in the architecture of the DNN is imposed. **Furthermore, we want the activation function of the output layer of the DNN to be itself bijective to be able to uniquely associate any FOM solution to a ROM in the low-rank manifold.**

In order to encode the automatic shift-detection in the backward map itself we must convert the variational form of singular values spectrum analysis and velocity sampling into a statistical learning theoretical framework. First and foremost we interpret the FOM snapshots as data-points of the training set for the network

$$\mathbf{M} := (\mathbf{u}_h(\boldsymbol{\mu}_j))_{j=1,\dots,P}, \quad \mathbf{X} := \boldsymbol{\mathcal{X}}^T \in \mathbb{R}^{N_s \times N_h}. \quad (6)$$

We observe that the training set is itself a subspace of the full-order solution manifold; in statistical learning theory we will thus have a  $N_h$ -dimensional features space of cardinality  $N_s$ . Secondly we assign a semi-supervised learning paradigm to our model by choosing arbitrarily, among the samples in  $\mathbf{M}$ , a reference configuration  $\mathbf{u}_{\text{ref}}$  for the field values that acts as unique label for the

training of the network on the rest of the collected data-points. It is important to clarify this aspect as it is one of the features of NNsPOD that can be exploited in order to build more refined and efficient machine learning algorithms in ROM of hyperbolic equations. For the sake of simplicity let us consider a 1–dimensional formulation of the IBVP in (3) in which the advection-dominated model features a regular, constant and uniform transport field  $b = 1$ . Given a stable full-order numerical simulation (i.e. with very low numerical diffusivity) it is intuitive that the reference configuration and any given data-point in  $\mathbf{M}$  only differ for the centroids  $x$  at which a certain value of field solution is associated. For instance, the peak of the Gaussian pulse will be located, e.g. at  $x = 0$  for a reference configuration  $\mathbf{u}_h(x, t_0)$  and we can easily predict that, after  $\Delta t = 1$ , it will be located at  $x = 1$  i.e.  $\mathbf{u}_h(x, t_1 = t_0 + 1)$ . It is trivial to see that, in this instance, choosing the IC as reference configuration is no different then choosing any other snapshots in  $\mathbf{M}$  when building a backward map either by direct calculation or automatic detection. This arbitrariness is itself an advantageous property of the hyperbolicity of the equation that is easily exploited whenever there are simple transport fields as stated above. However, if one is dealing with the more concrete case of irregular transport fields, then the IC itself does not have to be necessarily the one that fully reconstructs the shifted manifold since the stationary frame of reference can be associated to any of the snapshots collected at FOM. To take full advantage of faster Kolmogorov width decay, NNsPOD does not restrict itself in sampling backward transformations to the IC but to any reference configuration that optimises the reduction; the non-linear projection operator associated to the automatically detected map will thus be able to generalise from any frame of reference thanks to the bijective constraint discussed above. For this reason in the following we will refer to the unique label, chosen for the training of the neural network part of NNsPOD, in more general terms as the reference configuration  $\mathbf{u}_{\text{ref}} \in \mathbf{M}$  which, we reiterate, does not necessarily have to coincide with the IC of the IBVP. The setting is completed by choosing one particular metric for computing the loss function between the output associated to any data-point in  $\mathbf{M}$  and the reference configuration. While there is substantial space for testing and experimenting with this choice, in the development of the present work we restrict the development to the  $L_2$ –norm which is consistent with the Kolmogorov width decay of the finite-dimensional functional space  $\mathcal{V}_h$  for the FOM snapshots

$$J(\mathbf{x}_{\text{ref}}, N_s) := \frac{1}{N_s} \sum_{j=1}^{N_s-1} \|\tilde{\mathbf{u}}_h(\boldsymbol{\mu}_j) - \mathbf{u}_{\text{ref}}\|_2, \quad \tilde{\mathbf{u}}_h := \mathbf{u}_h(\mathbf{x}, \boldsymbol{\mu}) \circ \mathcal{T} \quad (7)$$

Future works might be able to derive a rigorous mathematical structure for the selection of appropriate metrics for the training of shift-detecting networks in projection-based ROM pre-processing.

### 3.2. Architectures for continuous data-flow: the shift-detection and field-reconstruction split

Aside from the shift detection itself, one important procedure for a pre-processing transformation of the FOM manifold is the ability of reconstructing the field values of the solution following the backward map to the reference configuration. This can be achieved via e.g. interpolation [24] of the field values from the shifted points to the nearest centroids on the grid. In developing an automatic shift-detection algorithm we therefore must include such field-reconstruction part within the algorithm itself. Being purely data-driven, it is also desirable that the interpolation is also based on the FOM snapshots and does not rely on the prior physical knowledge of the problem (e.g. upwind interpolation methods). One of the major advantages of NNsPOD is that the choice of the reference configuration for the training of the network is arbitrary. We exploit such potential by splitting the workload between two separate networks:

- *ShiftNet* is the neural network that has the duty of quantifying the optimal shift for pre-processing transformation of the full-order manifold and that maximises the Kolmogorov width decay.
- *InterpNet* is the neural network that must “learn” the reference configuration in the best possible way w.r.t. its grid point distribution s.t. it will be able to reconstruct its “shape” for every shifted centroid distribution.

The introduction of a neural network with the specific task of interpolating the field values on the nearest centroids of the grid following the shift means that, once fully trained, NNsPOD is not only capable of reconstructing the values of the solution field continuously across the computational domain but, depending on how the training is performed, it will also allow for the bijective mapping to be performed for virtually unlimited non-linear advection models and in particular with those that feature high degree of variability in the “shape” of the snapshots collected, at full-order, in  $\mathbf{M}$ . Attention must be paid in constructing such split architecture since the objective is that NNsPOD’s output  $\mathcal{T}$  is a sufficiently regular bijective map between the manifold and its shifted counterpart. As such, continuity in the data pipeline must be preserved from ShiftNet and InterpNet and viceversa. At the same time it is not desirable that the hyperparametric optimisation of the loss function of those networks has an effect on each other; a lack of separation of weights and biases updates lacks in fact the possibility of generalising NNsPOD transformation during online stage for those advection-dominated models that have parametric dependence in the transport field itself. As such we devised the DNN-based algorithm, reported in **Algorithm 1** to achieve such counteracting properties. **It is important to stress the adoption of two independent neural networks in order to derive a real physics-independent non-linear transformation of the manifold in a pure data-driven fashion. In our framework the shifted field interpolation has to be embedded in the derived shift operator, however, at the same time, it is desirable that once the reference configuration has been *learned* one could somehow retain the neural configuration as-**

sociated to it while training separately for the derivation of the actual shift map.

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**Algorithm 1:** The NNsPOD algorithm

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**Result:** Optimal shift-detected transformation  $\mathcal{T}$   
 Construction of snapshot matrix  $\mathcal{X}$ ;  
 Computing the feature matrix  $\mathbf{X} = \mathcal{X}^T$ ;  
 Setting the accuracy thresholds  $\varepsilon_{\text{SVD}}$ ,  $\varepsilon_{\text{shift}}$ ,  $\varepsilon_{\text{interp}}$ ;  
**while**  $\varepsilon > \varepsilon_{\text{SVD}}$  **do**  
   Selection of reference configuration  $\mathbf{u}_{\text{ref}}$ ;  
   **while**  $\varepsilon' > \varepsilon_{\text{interp}}$  **do**  
     **InterpNet**.forward;  
     compute  $\varepsilon'$ ;  
     **InterpNet**.backward;  
   **end**  
    $\mathcal{T}_{\text{interp}} = \mathbf{InterpNet}$ .forward;  
   **while**  $\varepsilon'' > \varepsilon_{\text{shift}}$  **do**  
      $\tilde{\mathbf{x}} = \mathbf{ShiftNet}$ .forward;  
      $\tilde{\mathbf{u}}_h = \mathcal{T}_{\text{interp}} \circ \tilde{\mathbf{x}}$ ;  
     compute  $\varepsilon''$ ;  
     **ShiftNet**.backward;  
   **end**  
    $\mathcal{T} = \mathbf{ShiftNet}$ .forward;  
   **for**  $\mathbf{u}_h \in \mathbf{X}$  **do**  
      $\tilde{\mathbf{u}}_h = \mathbf{u}_h \circ \mathcal{T}$ ;  
   **end**  
   Perform the SVD on  $\tilde{\mathbf{X}}$ ;  
   Compute  $\varepsilon_{\text{SVD}}$ ;  
**end**

---

It is therefore clear how the adoption of two independent deep learning architectures, *glued* continuously to assure bijectivity in the derived transformation, complies with the aforementioned requirement. The training stages of ShiftNet and InterpNet are thus separated with the latter being trained first. Once the network has learned the best possible reconstruction of the solution field of the reference configuration, its forward map will be used for the training of ShiftNet as well, in a cascaded fashion. For this reason we must optimise the loss of InterpNet (whose training set is composed by the field values of  $\mathbf{u}_{\text{ref}}$ ) considerably more than ShiftNet’s. A schematic of the structure of NNsPOD, as well as its input and output relation for the 1–dimensional example discussed in 3.1 is depicted in Figure 4. We highlight the simple scalability of NNsPOD architecture and algorithm to hyperbolic equations with higher dimensional space domain. As a matter of fact ShiftNet’s training set consists on the data-point’s spatial distribution meaning that its input layer will have  $d + P$  neurons while the output layer, feeding information to InterpNet, has  $d$  neurons.

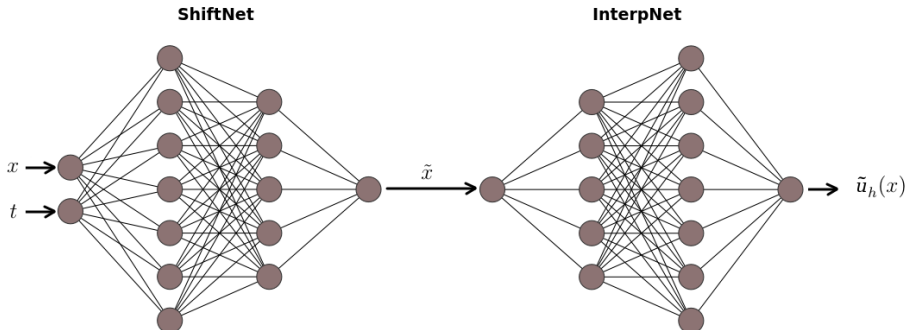


Figure 4: Deep learning architecture of NNsPOD for a 1–dimensional model showing the continuous pipeline of information passed from ShiftNet to InterpNet.

InterpNet on the other hand will have  $d$  neurons in the input layer while the output layer will always feature a single neuron, for any spatial dimension, if the numerical solution is scalar or 3 neurons if the solution is a vector field. We finally emphasise that a discussion about the fine-tuning of the networks is out of the scope of the present work. We will nonetheless specify the full details of the architecture deployed for the shift-detection of the numerical models that follow.

### 3.3. Reduction of non-uniform, non-constant linear advection equation

In order to benchmark the capability of NNsPOD automatic shift-detection the reduction of the same advection equation in (3) is hereby derived using its workflow. A non-uniform and non-constant advection field will be used to validate the performance of the proposed algorithm; such model cannot be pre-processed by traditional shift-based methods if prior knowledge of the equation is used. As outlined in 2.4 in fact, a singular values spectrum analysis on sampled constant and uniform velocities will lead to an excessive computational time during the offline phase and it will also introduce a linear approximation of the characteristic curves of the hyperbolic equation. We therefore refer to the same 2–dimensional, single parameter setting in (3) but with  $\mathbf{b} = (\frac{1}{2}y^2t, -2xt^2)$  instead; the  $N_s = 100$  collected snapshots are then pre-processed according to **Algorithm 1**. A substantial variance in shape of the snapshots is recorded, as depicted in Figure 5. This model reduction has the aim of:

- Showcasing the accurate shift-based pre-processing of linear subspace approximation for the manifold of a relatively complex advection field through automatic detection i.e. using no prior knowledge of the FOM.
- Testing the ability of reconstructing highly “diffusive” snapshots collected from an hyperbolic FOM to be later applied to the more complex multi-phase model in Section 4.

The latter is a desirable quality for our data-driven algorithm to feature; highly diffusive snapshots may arise in fact not only in models with non-uniform and



non-constant advection but also whenever the discretisation scheme of the convective term is of lower order of accuracy. The settings of NNsPOD’s neural networks for the shift-detection are the following

NNsPOD settings		
	InterpNet	ShiftNet
<b>Hidden layers</b> × <b>neurons</b>	$2 \times 40$	$3 \times 20$
<b>Activation function</b>	Sigmoid	PReLU
<b>Learning rate</b>	$10^{-3}$	$10^{-4}$
<b>Accuracy threshold</b>	$10^{-7}$	$10^{-1}$

We observed that values greater than those prescribed above for the learning rates of the two neural networks will feature either a non-convergent loss function or a resulting shift transformation that does not center around the reference configuration. The training of NNsPOD’s ShiftNet neural network lasted 6 hours on a i7 – 7700 quad processor with 16 GB of RAM; this constituted approximately 85% of the computational time of the overall offline stage (FOM snapshot collection, shift pre-processing and SVD of the shifted snapshot matrix). As for any machine learning model, and especially when treating deep learning, it is difficult to contextualise, let alone justify, the computational cost associated to the training phase. The lack of interpretability and full understanding of the networks behaviour is somewhat limiting any rigorous mathematical estimate to a more empirical assessment. In this case for instance, we observed that the training of InterpNet is significantly cheaper than the 6 hours took by ShiftNet, as a matter of fact by lasting merely 10 minutes. We argue that the difference on the datasets’ cardinality (InterpNet trains on only one snapshot field values, that of  $\mathbf{u}_{\text{ref}}$ , while ShiftNet optimises over  $N_s$  samples) and the cascaded structure of NNsPOD (ShiftNet has to optimise its parameters to minimise a loss that is in part a result of InterpNet’s behaviour, which ShiftNet cannot change directly) are the main actors for this unbalance in computational cost. The different stages of the training phase of InterpNet and ShiftNet are reported in Figure 6. We tested NNsPOD performance by selecting the 80–th snapshot as reference configuration ( $\mathbf{u}_{\text{ref}} := \mathbf{u}_h(t_{80}) \in \mathbf{M}$ ) to highlight the possibility of choosing appropriate frames of reference for the shift-based pre-processing of  $\mathcal{M}_h$ , aside for the IC’s one. The optimal transformation for the snapshots in  $\mathbf{X}$  is thus retrieved, via automatic shift-detection, for such reference configuration. During the training stage we observed that many snapshots, being different in shape among each other and specifically w.r.t. the reference configuration, were stretched differently along different directions in the spatial domain by ShiftNet in order to overlap InterpNet’s transformation.

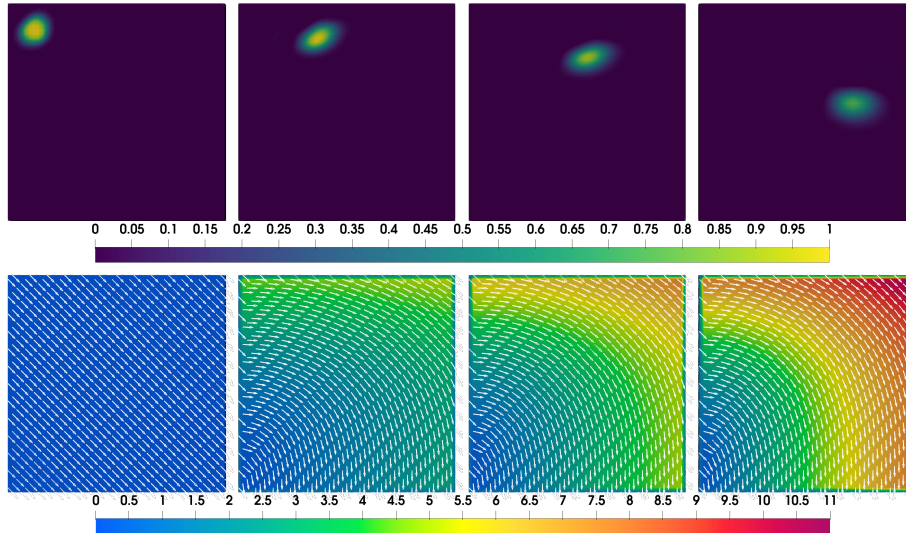


Figure 5: Different snapshots of the FOM solution (upper row) of a 2–dimensional advection equation with non-uniform and non-constant linear transport field (bottom row).

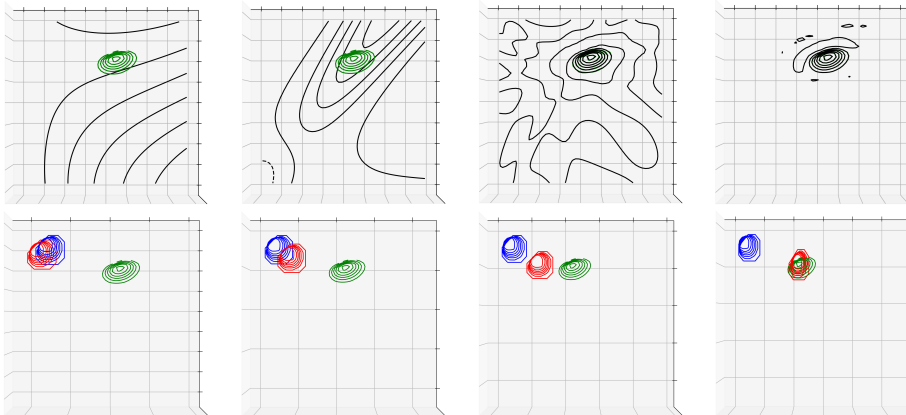


Figure 6: Output of NNsPOD's split neural networks at different epochs during their separate training stages: in the upper row IntepNet's output (black contour lines) convergence to the reference configuration (green contour lines) is shown; in the bottom row the output of ShiftNet's detected coordinates (red) for a so-called test snapshot in  $\mathbf{X}$  (blu) is depicted to converge to the reference configuration (green).

Nevertheless a pre-processing bijective map is derived by NNsPOD that uses no prior information regarding the mathematical model in (3) and its physical properties (i.e. characteristic velocities).

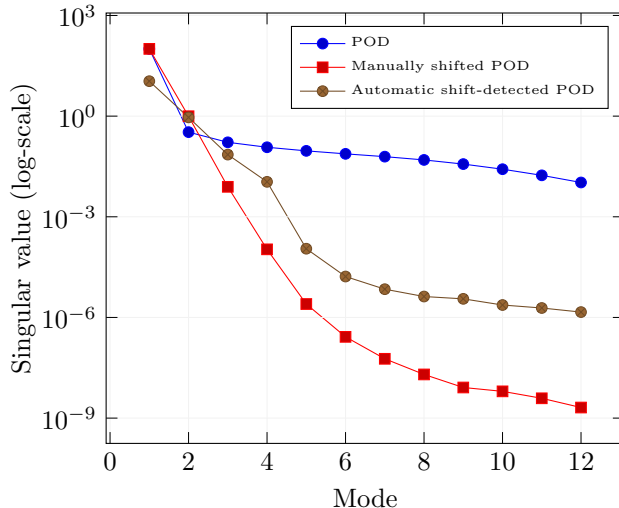


Figure 7: Accuracy comparison between traditional POD and NNsPOD for a non-uniform, non-constant, linear advection field in (3). The decay pattern follows that of an exact shift operator computed piecewise for every FOM snapshot proving the arbitrary accuracy achievable by the automatic shift-detecting bijection sought by NNsPOD.

Furthermore the non-uniform and non-constant advection field has not been linearised in its parametric dependence (in this instance only w.r.t. to the time variable  $t$ ) in order to sample the velocity space. The SVD of the reconstructed snapshots, following NNsPOD automatically detected shift-based mapping, demonstrates a faster Kolmogorov width decay (Figure 7) compared to a traditional POD algorithm; its accuracy does indeed follow the one that it would be obtained by constructing manually an exact piecewise shift operator for each of the datapoints in  $\mathbf{M}$ . To that regard NNsPOD can be interpreted as the statistical learning algorithm that constructs the best possible approximation of a shift operator via automatic, data-driven detection; as a result its precision is arbitrarily set by choosing the loss threshold for it to generate the bijective mapping and therefore  $\tilde{\mathcal{M}}_h$ . In Figure 8 the loss optimisation of InterpNet and ShiftNet are depicted for the IBVP in (3); from it we can appreciate how fundamentally different the hyper-dimensional loss functions for the two neural networks are. During testing and validation of the algorithm, we observed that the optimisation of ShiftNet during training frequently featured a steep descent followed by a slow asymptotic decay, as opposed to the more noisy but consistent minimisation of InterpNet. We believe that the reason for ShiftNet’s loss behaviour is due to the fact that in  $d > 1$ , infinitely many non-linear shifts can be traced from any given FOM snapshot to the reference configuration; once one of them is found the loss experience a steep optimisation which then quickly levels since the now shifted snapshots are centered around  $\mathbf{u}_{\text{ref}}$  but not quite overlapping its field values.

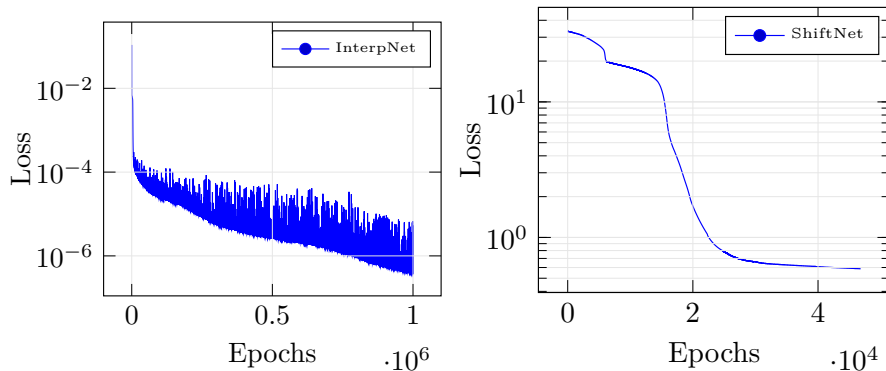


Figure 8: Loss optimisation w.r.t. the training of NNsPOD’s split neural networks for the IBVP (3) with advection field  $\mathbf{b} = (\frac{1}{2}y^2t, -2xt^2)$ .

#### 4. Reduction of a multiphase model with manifold transformation

We now present a numerical experiment to validate the fundamental property of NNsPOD being applicable to non-linear advection fields. The transport of a passive scalar field in a multiphase flows of two fluids with uniform densities  $\rho_1, \rho_2 \in \mathbb{R}$ ,  $\rho_1 \neq \rho_2$  is chosen as FOM to generalise the reduction of a 2–dimensional linear hyperbolic equation as addressed in Section 3. The reason motivating this choice lies within the mathematical model describing the multiphase flow itself being the coupling between the (incompressible) Navier-Stokes momentum balance equation and an advection equation. The coupling stems from the velocity field, which is unknown and derived numerically as a solution of the Navier-Stokes equation; the divergence of such field ultimately becomes the transport operator for a scalar field  $\alpha(\mathbf{x}, t, \boldsymbol{\mu})$  that models the pointwise fraction of volume of the two fluids within the cells of the discretised computational domain. Furthermore, being Navier-Stokes a non-linear PDE, the no-prior knowledge quality of NNsPOD is thereby assessed as its ability of deducing the proper pre-processed linear manifold approximation with a generalisation to non-linear advection fields.

##### 4.1. The Volume-of-Fluid method for Eulerian interface tracking

A multiphase flow is modelled by two fluids occupying a certain fraction of the volume domain. The time evolution of each fluid is governed by the unsteady Navier-Stokes equations

$$\begin{cases} \partial_t \rho + \nabla \cdot (\rho \mathbf{U}) = 0, \\ \partial_t(\rho \mathbf{U}) + \nabla \cdot (\rho \mathbf{U} \otimes \mathbf{U}) = -\nabla p + \nabla \cdot \boldsymbol{\tau} + \mathbf{F}, \end{cases} \quad (8)$$

in which the first equation describes the mass conservation principle (continuity equation) whereas the second models Newton’s second law for the conservation

of (linear) momentum. Assuming incompressible regime and absence of external forces ( $\mathbf{F} = \mathbf{0}$ ) acting of the fluids (8) reduces to

$$\begin{cases} \nabla \cdot \mathbf{U} = 0, \\ \partial_t \mathbf{U} + (\mathbf{U} \cdot \nabla) \mathbf{U} = -\frac{1}{\rho} \nabla p + \nu \Delta \mathbf{U}. \end{cases} \quad (9)$$

The system above is coupled in the pressure  $p(\mathbf{x}, t)$  and velocity  $\mathbf{U}(\mathbf{x}, t)$  fields which are the unknowns of the problem at hand that has to be solved numerically. We observe that (9) describes the dynamics of one single fluid with (uniform) volumetric density  $\rho$  and kinematic viscosity  $\nu$ . In the multiphase flow however we have two fluids, separated by a sharp interface, that evolve in a coupled fashion with each other. By assuming that the two fluids are immiscible we can describe the physical properties of the flow as averaged across the domain; to that end the fraction of volume fields  $\alpha_1, \alpha_2 \in [0, 1]$  are introduced which are defined through a constitutive relation  $\alpha_1 \Omega + \alpha_2 \Omega = \Omega \Rightarrow \alpha_1 + \alpha_2 = 1 \Rightarrow \alpha := \alpha_1 = 1 - \alpha_2$  that acts as a constraint on the model. The new density field is thus defined as  $\alpha(\mathbf{x}, t) = \alpha(\mathbf{x}, t) \rho_1 + (1 - \alpha(\mathbf{x}, t)) \rho_2$  and its time evolution is governed by an advection equation coupled with the constitutive relation derived above

$$\begin{cases} \alpha \rho_1 + (1 - \alpha) \rho_2 = \alpha, \quad \alpha \in [0, 1] \quad \forall \mathbf{x} \in \Omega, \\ \partial_t \alpha + \nabla \cdot (\mathbf{U} \alpha) = 0, \end{cases} \quad (10)$$

with  $\mathbf{U}(\mathbf{x}, t)$  being the (non-linear) velocity field in (9) which is thus coupled with (10). Eulerian multiphase modelling tracks the instantaneous and pointwise changes in the sharp interface between the two fluids; continuous-continuous and dispersed-continuous phase interaction are the two large families of algorithms used to model multiphase flow. The latter is used in simulating dispersed particles (solid phases), droplets (liquid phases) and bubbles (gaseous phases) within a larger continuous fluid phase; the former instead is used whenever there is a presence of a continuous sharp interface between two fluids. The presence of an additional transport operator in the advection equation for field  $\alpha(\mathbf{x}, t)$  suggests that traditional bounded schemes for its numerical discretisation might not suffice. The operator becomes in fact highly diffusive in proximity of the interface in a process called excessive smearing. One possibility to mitigate the phenomena, as outlined in [39], is to introduce a numerical compression term  $\nabla \cdot (\alpha(1 - \alpha) \mathbf{U}_r)$  in the advection equation in (10) built ad-hoc s.t. it takes null values everywhere in the domain except in proximity of the interface where  $\alpha \in (0, 1)$  (being  $\mathbf{U}_r := C_\alpha \|\mathbf{U}\| \frac{\nabla \alpha}{\|\nabla \alpha\|}$  with  $C_\alpha \in [0, 1]$  as a free parameter). The resulting system from (10) thus is

$$\begin{cases} \alpha \rho_1 + (1 - \alpha) \rho_2 = \alpha, \quad \alpha \in [0, 1] \quad \forall \mathbf{x} \in \Omega, \\ \partial_t \alpha + \nabla \cdot (\alpha(\mathbf{U} + (1 - \alpha) \mathbf{U}_r)) = 0. \end{cases} \quad (11)$$

Then, a discretisation scheme is devised for the advection terms, with the compressive interface capturing (CICSAM) and piecewise linear interface construction (PLIC) being the most widespread (we refer to [40] for a detailed explanation and derivation).

#### 4.2. Full-order multiphase numerical modelling

To benchmark the performance of NNsPOD automatic shift-detection and construction of pre-processed linear manifold approximation for the non-linear advection of an hyperbolic equation, a simplified model of a multiphase simulation is solved numerically. While retaining the parametric dependence on the density and viscosity for the couple of fluids, for the sake of simplicity we restrict the parameter space, paired to the FOM, to be the exclusively the time variable as previously derived for the linear models in Section 2 and 3. To this end we consider the following IBVP for the time-evolution of a sharp interface separating water ( $\rho_1 = 10^3$ ,  $\nu_1 = 10^{-6}$ ) from air ( $\rho_2 = 1$ ,  $\nu_2 = 1.48 \cdot 10^{-5}$ )

$$\begin{cases} \nabla \cdot \mathbf{U} = 0, \\ \partial_t \mathbf{U} + (\mathbf{U} \cdot \nabla) \mathbf{U} = -\frac{1}{\rho} \nabla p + \nu \Delta \mathbf{U}, \\ \mathbf{U} = (0, 0), \quad \forall \mathbf{x} \in \Gamma_b, \quad \partial_n \mathbf{U} = 0, \quad \forall \mathbf{x} \in \partial\Omega \setminus \Gamma_b, \\ \mathbf{U}(\mathbf{x}, 0) = (0.25, 0), \quad \forall \mathbf{x} \in \Omega, \\ \alpha \rho_1 + (1 - \alpha) \rho_2 = \alpha, \quad \alpha \in [0, 1] \quad \forall \mathbf{x} \in \Omega, \\ \partial_t \alpha + \nabla \cdot (\alpha(\mathbf{U} + (1 - \alpha)\mathbf{U}_r)) = 0, \quad C_\alpha = 1, \\ \alpha = 1, \quad \forall \mathbf{x} \in \Gamma_b, \quad \nabla \alpha = 0, \quad \forall \mathbf{x} \in \partial\Omega \setminus \Gamma_b, \\ \alpha(\mathbf{x}, 0) = 1, \quad \forall \mathbf{x} \in \Omega \text{ s.t. } y < e^{-\frac{x^2}{2}}, \end{cases} \quad (12)$$

where the IC for field  $\alpha(\mathbf{x}, t)$ , which identifies the initial configuration of interface between the two fluids, features a Gaussian profile. The BCs are set to comply with the no-slip condition for the bottom partition of the boundary  $\Gamma_b$  whereas a null gradient through top, the inlet and outlet partitions is allowed (see Figure 9 for reference). The setting is intended to simulate a multiphase flow through a 2-dimensional canal with open lid at atmospheric pressure. The domain  $\Omega = [-2.5, 3.5] \times [-0.5, 1.25]$  is discretised in a homogeneous set of collocated rectangular cells with a computational grid of  $250 \times 75$  centroids ( $N_h = 18,750$ ). The time interval for the simulation is set to  $t \in [0, 5]$  which is in turn discretised into 100 steps to comply with the CFL condition thereby leading to the following manifold definition

$$\mathcal{M}_h := \{\alpha_h(t_k) \in \mathcal{V}_h, k = 1, \dots, 100\}$$

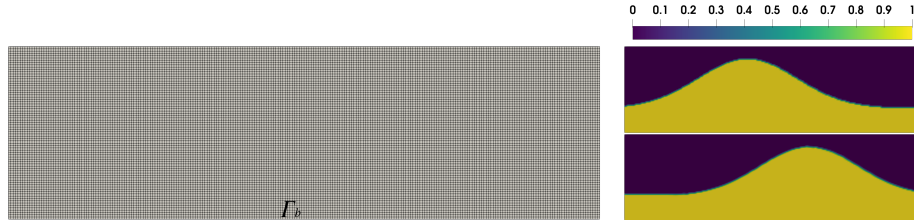


Figure 9: Illustration of the FOM for the IBVP in (12): on the left the discrete computational grid is shown with the indication for the bottom partition of the boundary where the no-slip condition is applied; on the right two snapshots of the solution field  $\alpha_h$  are reported (IC on the top, 100-th snapshot on the bottom).

In the statistical learning configuration of NNsPOD we remark that the offline snapshot collection generates a dataset  $\mathbf{M} \in \mathbb{R}^{N_s \times N_h}$ , associated to  $\mathcal{M}_h$ , of cardinality  $N_s = 100$  and dimensionality (number of features)  $N_h = 18.75 \cdot 10^3$ .

#### 4.3. Automatic shift-detection and linear manifold reconstruction

The shift-detection of the algorithm regards the correct reconstruction of the interface separating the two phases of fluids w.r.t. a reference configuration for the problem. As discussed in Section 3 the choice for an appropriate snapshot for the training of InterpNet does not affect the ability of ShiftNet in constructing a pre-processed manifold approximation. Nevertheless, even if for highly diffusive transports one can always select any configuration of the numerical field  $\alpha_h$  for the subsequent automatic shift detection, we remark that InterpNet optimises only on those field values. It is therefore to be expected that, the more complex and non-linear the advection field is, the more training for InterpNet and ShiftNet will be necessary. Here we selected the 35–th snapshot to be the reference configuration  $\alpha_{\text{ref}} := \alpha_h(t_{35}) \in \mathbf{M}$ . The split architecture of NNsPOD was set with the following parameters.

NNsPOD settings		
	InterpNet	ShiftNet
<b>Hidden layers</b> ×neurons	$4 \times 40$	$5 \times 25$
<b>Activation function</b>	HardSigmoid	PReLU
<b>Learning rate</b>	$10^{-5}$	$10^{-6}$
<b>Accuracy threshold</b>	$10^{-4}$	$10^2$

The field  $\alpha_h$  targeted by the shift-detection, and therefore involved in the minimisation of the loss function, has values in  $[0, 1]$ ; intuitively in most of the cells of the FOM its values is either one of the infimum (i.e. air only) or the supremum (i.e. water only) with the exception of the interface where the gradient has non-null values. As such, in order for NNsPOD to emulate such behaviour, the hard sigmoid activation function for InterpNet’s neurons, which is responsible for *learning* the best possible approximation for the reference configuration, has been adopted. As expected the training of NNsPOD lasted considerably longer than the previous linear simulation reported in Section 3 with the convergence time towards the threshold amounting to 4 times what was previously measured (27 hours circa), on the same hardware, for the previous case with the linear setting. Although this is partially attributed with the higher dimensional full-order discrete space  $\mathcal{V}_h$  we did observe a non-trivial behaviour for the convergence of the shift-detection towards the reference configuration. The non-linearity of the FOM in (10) forced the sampling of NNsPOD to not be constrained to simple linear backward shifts which is indeed the goal for the generalisation process itself.

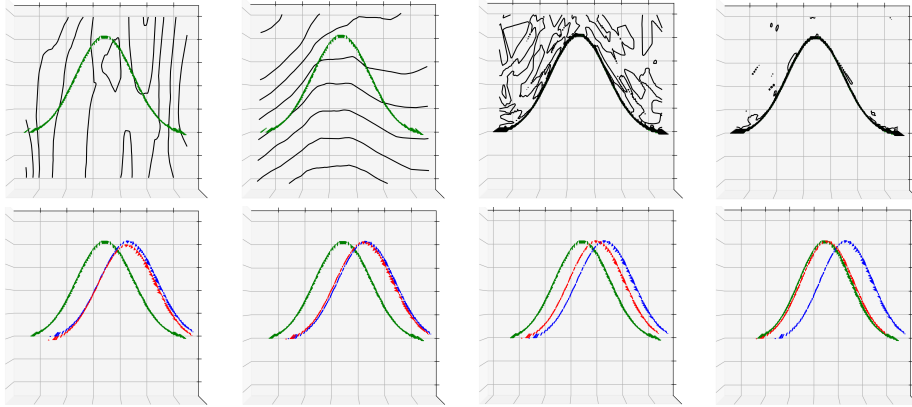


Figure 10: NNsPOD’s neural networks at different epochs during the training for the multi-phase FOM in (10): the upper row displays IntepNet’s (black) convergence to  $\alpha_{\text{ref}}$  (green) while the bottom row shows the convergence of ShiftNet’s output (red) for test snapshot in  $\mathbf{X}$  (blu).

In fact, as motivated in Section 3 and later shown in Figure 6 for the non-uniform linear case, NNsPOD’s capability is not limited to linear backward shift for a test snapshot but rather to a non-linear stretching of the shifted coordinates.

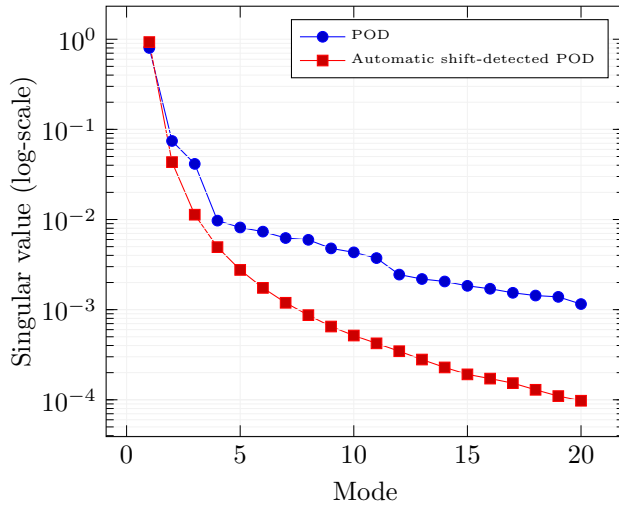


Figure 11: Comparison of singular values decay for the POD prior and following NNsPOD’s pre-processing bijective mapping. We can appreciate a substantial improvement in accuracy retained per number of modes, in line with the shift-based treatment of the snapshot. This result highlights the possibility of NNsPOD to replicate the results of sPOD (with arbitrary accuracy) to those models for which the manual construction of an exact shift operator is not possible.



The search for alternative and non-trivial backward maps, as shown by ShiftNet’s outputs in Figure 9 and 10 entails to our algorithm the property of generating automatically the best possible approximation of the backward shift in the form of a bijective map for the linear manifold reconstruction for unknown transport fields. Obviously, the non-linear transformation required by (12) requires more refined DNN architectures; this can be seen by the relatively slower singular values decay reported in Figure 11. As we explained in previous sections, we cannot derive a mathematically rigorous selection of the appropriate networks’ configuration by merely relying on the task of increasing the decay of the singular value; nevertheless the authors believe that more advanced architectures at the state-of-the-art may outperform the still significant gains shown in Figure 11 obtained by much simpler DNNs.

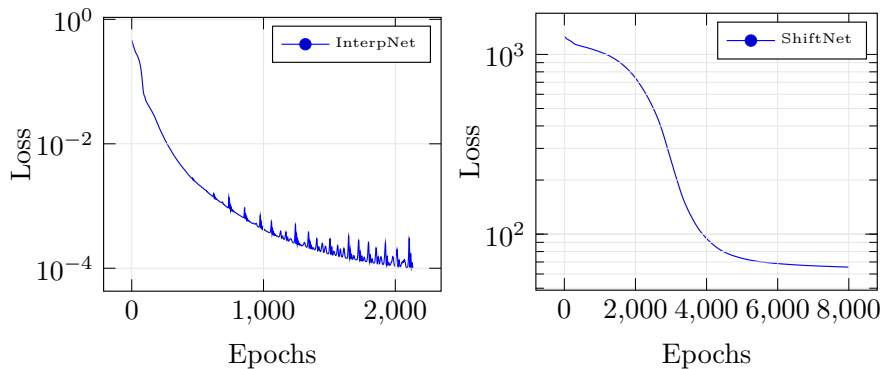


Figure 12: NNsPOD training loss optimisation for the multiphase IBVP (12).

As a final remark, we validate the slow asymptotic convergence of the loss function of ShiftNet, firstly reported in 3.3 for the linear advection equation in (3) and depicted in Figure 12 for the multiphase IBVP (12). Once again such behaviour matches the visual observations of FOM snapshots taking a variety of different paths to be centered around the reference configuration. The authors reiterate the need of setting a much tighter threshold for the optimisation of InterpNet as more complex and non-linear the advection field of the model is.

## 5. Conclusions

The purpose of this work was to derive a novel approach for the model order reduction of advection dominated problems. At this aim, the development of NNsPOD allowed us to emphasize the advantages posed by the construction of a non-linear transformation that does not rely on prior knowledge of the physical model. The automatization of the shift-detection process opposes the sampling of the phase space for the advection field to derive better linear approximations of the manifold at the current state of the art. The multiphase test showcases that the time required for the training of NNsPOD falls within

the current estimation for the offline phase with no pre-processing of the snapshots (e.g. traditional POD) while leading to more accurate low-rank subspaces. The online phase of a pre-processing based POD, which is of paramount importance for the industrial applications of any ROM, requires for the backward map to generalise for new instances of the parameter vector; in order to achieve such result the map itself has to be independent on the advection field. In the early development stage of NNsPOD, we realised that the traditional sPOD [24] method could not generalise well during the online phase due to the fact that the transformation that maps the FOM snapshots to the IC is linear and advection field dependent. Our methodology on the other hand is non-intrusive and data-driven, allowing for the parameter vector to be unaccounted in the construction of the low-rank manifold since it disregards the transport field; this entails that performing the Galerkin projection of the governing equations onto the reduced order subspace constructed by NNsPOD would result in numerical coefficients that fully and solely carry the information on the parametric dependence embedded in the transport field. We remark once again that, in order for NNsPOD’s automatically detected transformation to be used at online stage, it must be bijective; such constraint is mathematically enforced by assuring continuity in the pipeline of the two neural networks and by adopting a bijective activation function for the output layer of InterpNet. Our algorithm will therefore map every FOM snapshot to a unique shifted approximation of the reference configuration within an arbitrary small neighborhood centered in it. Future works will focus on the validation of the capability of the proposed framework on a more complex setting, e.g. higher-dimensional formulation for the multiphase model in (10) with a parametric treatment of the fluid viscosity. Moreover, since this work has the main purpose of presenting the NNsPOD methodology for improving the classical POD reduction, we have postponed to future works the development of a full NNsPOD-Galerkin ROM. In particular the authors suggest that the objective of further investigation in the creation of a full, NNsPOD-based ROM, should include:

- validation on the accuracy of the reduced order solution for parametric inputs that result in advection fields that are significantly different from those used for training the networks during the online stage. We recall that the underlying mathematical structure of the algorithm relies on a transformation mapping multiple FOM solutions  $\mathbf{u}_h \in \mathcal{M}_h$  into a relatively small neighborhood of the shifted manifold, centered in  $\mathbf{u}_{\text{ref}}$ . Therefore if the advection field at online stage is e.g. several orders of magnitude different from those used for training the networks at offline stage, perhaps larger, more refined and fine-tuned deep learning architectures will have to be deployed by NNsPOD.
- investigate a fast and efficient algorithm to compute the Jacobian of NNsPOD non-linear transformation. At online stage the linear superposition of the spatial modes of the shifted low-rank manifold, although carrying non-explicit dependence of the parametric advection field, is characterised by a dependence on the non-linear map by the parametric coefficients

$\hat{a}(\boldsymbol{\mu}, t)$  of the linear combination computed in the shifted frame of reference. When projecting the governing equation onto the reduced order manifold, the explicit form of  $J(\hat{a}(\boldsymbol{\mu}, t))$  will have to be computed which is non-trivial.

Of course, the NNSPOD effectiveness strongly depends on the problem at hand and on the machine learning architecture used for both the networks — Shiftnet and Interpnet —. It is impossible indeed, at the current state, to predict rigorously the optimal structure of the two networks with the choice of the activation function being the sole setting that can be deduced by the FOM solution field as outlined in Sections 3 and 4. Future studies will better investigate the sensitivity of the method with respect to the networks architecture, considering a set of hyper-parameters larger than the one analysed in the current article.

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