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(Article begins on next page)

# 1 Extended dynamic mode decomposition for model reduction in fluid dynamics

- 2 simulations
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High computational cost and storage/memory requirements of fluid dynamics simulations constrain their usefulness as a predictive tool. Reduced-order models (ROMs) provide a viable solution to this challenge by extracting the key underlying dynamics of a complex system directly from data. We investigate the efficacy and robustness of an extended dynamic mode decomposition (xDMD) algorithm in constructing ROMs of three-dimensional cardiovascular computations. Focusing on the ROMs' accuracy in representation and interpolation, we relate these metrics to the truncation rank of singular value decomposition, which underpins xDMD and other approaches to ROM construction. Our key innovation is to relate the truncation rank to the singular values of the original flow problem. This result establishes a priori guidelines for the xDMD deployment and its likely success as a means of data compression and reconstruction of the system's dynamics from dominant spatiotemporal structures present in the data.

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### 23 I. INTRODUCTION

High computational burden of fluid dynamics simulations has propelled the develop-24 ment of model reduction techniques for problems dealing with complex flow and transport 25 rocesses in fields as diverse as geosciences and biomedicine<sup>1–5</sup>. A reduced-order model 26 (ROM) is a computationally efficient and reasonably accurate representation of the underly-27 ng dynamics of a state variable or a quantity of interest, derived from observations and/or 28 omputer-generated data. The efficiency of a model reduction technique manifests itself in 29 oth the amount of data required for the ROM construction and the ROM approximation 30 ccuracy in the interpolation and extrapolation regimes<sup>6</sup>. 31

Dynamic mode decomposition (DMD) is a data-driven technique that constructs ROMs of 32 complex dynamical systems by employing the singular value decomposition (SVD)<sup>7,8</sup>. DMD 33 aims to identify spatiotemporal structures that are dominant in the data and to reconstruct 34 an optimal linear model from these structures. A DMD variant xDMD<sup>9</sup> combines salient 35 atures of the residual learning<sup>10</sup> and the generalized DMD with a bias term<sup>11</sup>. This DMD 36 lgorithm has the ability to handle dynamical systems described by inhomogeneous partial 37 ifferential equations, which proved to be problematic for standard DMD. Numerical studies, 38 ealing with problems as diverse as the Navier-Stokes equations<sup>9</sup> and multiphase transport in 39 prous media<sup>12</sup>, suggest that the xDMD is more accurate than the standard DMD algorithm 40 (hereinafter, sDMD<sup>13</sup>). Since xDMD has more parameters than sDMD (the bias term), it is 41 otentially more sensitive to noise than. However, the numerical experiments<sup>9</sup> indicate that 42 the correction effects from the bias term may dominate the effects of over-fitting the noise. 43

These and other methods for ROM construction rely on the truncation rank of SVD 44 to control the degree of order reduction and representation accuracy. The choice of how 45 any singular values to keep depends on such factors as the quality and origin of the data 46 and the dynamic importance of low-energy modes<sup>13</sup>. The rank selection is typically done 47 ia experimentation, rendering the method's implementation subjective. A more principled 48 pproach is to balance order reduction and approximation accuracy by utilizing a general 49 eriteria<sup>12</sup>. The rank choice is also linked to xDMD's data compression ability, which is given 50 by the capability of the algorithm to preserve high accuracy for low values of the truncation 51 rank<sup>12,13</sup>. By identifying dominant coherent structures from data, the method effectively 52 reduces the dimensionality of high-dimensional datasets, thereby achieving compression-like 53

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effects. That is relevant in fluid dynamics, where DMD operates by reducing the dimension-54 ality of the flow field data while preserving its essential characteristics. Another application 55 climate science, where DMD can be used to compress large-scale climate datasets into a 56 is reduced set of dominant modes, facilitating the analysis and visualization of long-term cli-57 nate trends and variability<sup>13</sup>. In yet another setting of multiphase flow in porous media<sup>12</sup>, 58 DMD demonstrated high prediction accuracy (relative interpolation error on the order of 59  $^{-9}$ ) with a truncation rank of up to 35% of the dataset dimension. By way of a disclaimer, 60 we note that, like other SVD-based techniques, DMD often struggles to honor translational 61 and rotational invariances of low-rank objects embedded in the data<sup>13</sup>. 62

Our study has three intertwined goals. The first is to analyze how the representation error of xDMD is affected by the truncation rank in SVD, which, in turn, is linked to singular values of the problem. The second is to test the xDMD-based ROM in terms of its interpolation error, for different truncation ranks. The third goal is to explore the effect of neglecting possible irrelevant/overfit-inducing information (noise) on the accuracy of the approximation. We pursue these goals in the context of three-dimensional (3D) cardiovascular simulations of blood flow in a complex geometry of a patient-specific aorta.

The reference aorta geometry is selected from the Vascular Model Repository (www. 70 vascularmodel.com), a library of patient-specific cardiovascular models developed on volu-71 hetric image data sets and relevant physiologic data<sup>14</sup>. Fluid dynamics data are generated 72 with SimVascular (http://simvascular.github.io/). The latter is an open-source software 73 that provides a complete pipeline, from medical image data segmentation to patient-specific 74 lood flow simulations based on the 3D incompressible Navier-Stokes equations<sup>15</sup>. We use 75 data set consisting of  $\approx 2\cdot 10^3$  time frames of the velocity distribution (on a mesh with 76  $10^5$  elements) in a selected aorta. 77

Our research provides practical guidelines for the selection of low-rank truncation options 78 for optimal order-reduction (data compression). Our findings suggest that excluding low-79 nergy modes, which do not contribute to the elucidation of system dynamics, is beneficial 80 to ROM accuracy. We also found the ROM accuracy to be robust to both the size of time 81 intervals between the snapshots and low-rank truncations. This conclusion requires a flow 82 map of the system dynamics to be sufficiently smooth in space. An optimal rank selection 83 needs to consider the ROM's prediction reliability not only in reproducing the training data 84 (representation error) but also in making predictions at space-time locations where the data 85

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$$\| \mathbf{\Phi}_{\Delta t} \left( \mathbf{y}; \mathbf{s} \right) - \mathbf{\Phi}_{\Delta t} \left( \tilde{\mathbf{y}}; \mathbf{s} \right) \| \le e^{L\tau} \| \mathbf{y} - \tilde{\mathbf{y}} \|, \forall \tau \in [t, t + \Delta t].$$

re not available (interpolation error). Once optimized, the ROM can be used to replicate

cardiac function in a low-dimensional space, reducing the simulation cost and facilitating

the optimization and design of patient-specific interventions. At the same time, the DMD-

based modal decomposition allows for the identification of physically interpretable patterns

and possibly employed to detect pathologies<sup>17–21</sup>.

PROBLEM FORMULATION

 $\mathbf{u}(t + \Delta t)$  at any time t and time step  $\Delta t$ .

 $\mathcal{H}_{\Delta t}$ . Specifically, for any  $\mathbf{y}$  and  $\tilde{\mathbf{y}} \in \mathcal{H}_{\Delta t}$ ,

h the temporal and spatial evolution of the observed cardiovascular phenomena<sup>16</sup>. Coherent cructures and dominant flow features can be analyzed to discover the underlying physics

The paper is organized as follows: Section II is devoted to the formulation of the problem;

Section III the xDMD algorithm is described; while in Section IV its application to the

Once discretized on a numerical mesh, system states are arranged into a state vector

 $\mathbf{u}(0) = \mathbf{u}_0,$ 

(1)

 $\mathbf{u}(t) \in \mathbb{R}^N$  of length N. The temporal evolution of this discretized system is described by a

where  $\mathbf{f}(\mathbf{u}, \cdot)$  decribes the nonlinear dynamics,  $\mathbf{s} \in \mathbb{R}^N$  represents the source/sinks term and

Let  $\Phi_{\Delta t} : \mathbb{R}^N \to \mathbb{R}^N$  be a flow map, which relates the discretized system state  $\mathbf{u}(t)$  to

**Jemma 1** Assume **f** is Lipschitz continuous with Lipschitz constant L on a set  $\mathcal{H} \subseteq \mathbb{R}^N$ .

Define  $\mathcal{H}_{\Delta t} = \{ \mathbf{y} \in \mathcal{H} : \mathbf{\Phi}_{\Delta t} (\mathbf{y}) \in \mathcal{H} \}$ . Then, the flow map  $\mathbf{\Phi}_{\Delta t}$  is Lipschitz continuous on

boundary conditions, and  $\mathbf{u}_0 \in \mathbb{R}^N$  denotes the discretized initial state of the system.

system of N (nonlinear, homogeneous) ordinary differential equations,

 $\frac{\mathrm{d}\mathbf{u}}{\mathrm{d}t} = \mathbf{f}(\mathbf{u}, \mathbf{s}),$ 

test case is presented and discussed; a set of final remarks in Section V closes the paper.

The proof follows from the classical result on the continuity of a dynamical system (p. 109 in Ref. 22). The local Lipschitz continuity of the flow map ensures that nearby trajectories evolve smoothly and predictably, which is critical for the validity of the DMD approximation<sup>11</sup>. Moreover, if the flow map is locally Lipschitz continuous, the system's behavior can be accurately represented by a finite number of modes that evolve smoothly

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onto observables<sup>13</sup>.

precise representation of the system's behavior.

induced by the flow map  $\Phi_{\Delta t}$  and its DMD approximation are

reported in Section IV serve to investigate this error in detail.

algorithm describes the temporal evolution of  $\mathbf{u}(t)$  with a linear model

respectively. The error of a DMD model at time  $t_k$  is

THE XDMD ALGORITHM

(2)

(3)

(4)

In a typical application,  $M \ll N$  so that the rank of **A** is at most *M*. Even though, 138 computing  $\mathbf{A}$  (or its spectral decomposition) is generally onerous. Instead, the truncated 139 SVD of  $\mathbf{X} = \mathbf{U} \mathbf{\Sigma} \mathbf{V}^{\mathsf{T}}$ , with rank r < M, is used<sup>13</sup>: 140

in time, thus aiding interpretation and forecasting. Various DMD studies<sup>23</sup> indicate that a

linear operator might not be a good approximation of the general flow map, particularly for

highly nonlinear problems. In such cases, it might be necessary to map the state variables

The DMD approach approximates the nonlinear dynamical system, i.e.,  $f(\mathbf{u}, \cdot)$ , with

linear model constructed from M temporal snapshots of the discretized state variable,

 $= \mathbf{u}(t_k)$  with  $k = 0, \dots, M - 1$ . In general, numerical simulations involve discretizing

continuous processes into time steps. The continuous nature of the flow map enables in-

terpolation between simulation time steps or extrapolation beyond them, providing a more

Let  $\mathcal{L}$  be a DMD-based ROM of the dynamical system (1). At time  $t_k$ , the true solution

 $\mathbf{x}_k = \mathbf{\Phi}_{\Delta t}(\mathbf{x}_{k-1}) \text{ and } \mathbf{x}_k^{\mathcal{L}} = \mathcal{L}(\mathbf{x}_{k-1}^{\mathcal{L}}),$ 

 $\delta_{k}^{\mathcal{L}} = \|\mathbf{x}_{k}^{\mathcal{L}} - \mathbf{x}_{k}\|,$ 

where  $\|\cdot\|$  denotes vector 2-norm. The error bounds for xDMD and sDMD, reported in

Appendix A, provide a general indicator<sup>9</sup> for the growth of  $\delta_{k}^{\mathcal{L}}$ . The numerical experiments

Consider a set of (M+1) snapshots of the vector of state variables,  $\mathbf{x}_k$  with  $t_{k+1} = t_k + \Delta t$  and

 $= 0, \ldots, M$ . Let  $\mathbf{X} \in \mathbb{R}^{N \times M}$  denote a matrix whose columns are the vectors  $\mathbf{x}_0, \ldots, \mathbf{x}_{M-1}$ .

Let  $\mathbf{X}' \in \mathbb{R}^{N \times M}$  denote a matrix whose columns are the vectors  $\mathbf{x}_1, \ldots, \mathbf{x}_M$ . The sDMD

 $\mathbf{x}_{k+1} \approx \mathbf{A}\mathbf{x}_k, \qquad \mathbf{A} = \mathbf{X}'\mathbf{X}^{\dagger} \in \mathbb{R}^{N \times N}.$ 

$$\mathbf{A} \approx \mathbf{X}' \mathbf{V} \mathbf{\Sigma}^{-1} \mathbf{U}^{\mathsf{T}},\tag{5}$$

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where 
$$\mathbf{U} \in \mathbb{R}^{N \times r}$$
,  $\mathbf{\Sigma} \in \mathbb{R}^{r \times r}$ ,  $\mathbf{V} \in \mathbb{R}^{M \times r}$ . If  $r$  is smaller than the number of nonzero singular  
values (i.e., the rank of  $\mathbf{X}$ ), then the truncated SVD is a proxy of  $\mathbf{X}$ .

To allow for a problem's inhomogeneity, the generalized DMD algorithm adds a bias term 144  $\mathbf{b}_{\varrho} \in \mathbb{R}^{N}$  to the standard formulation, 145

$$\mathbf{x}_{k+1} \approx \mathbf{A}_g \mathbf{x}_k + \mathbf{b}_g. \tag{6}$$

Here,  $[\mathbf{A}_g \ \mathbf{b}_g] = \mathbf{X}' \tilde{\mathbf{X}}^{\dagger} \in \mathbb{R}^{N \times N+1}$ , where  $\tilde{\mathbf{X}}^{\top} = [\mathbf{X} \ \mathbf{1}]$  and  $\tilde{\mathbf{X}} \in \mathbb{R}^{N+1 \times M}$ . The computational 147 cost is reduced by obtaining the best-fit linear operator through the SVD of the matrix 148  $\tilde{\mathbf{X}} \approx \mathbf{U}_{g} \boldsymbol{\Sigma}_{g} \mathbf{V}_{g}^{\mathsf{T}}$ , such that 149

$$[\mathbf{A}_g \ \mathbf{b}_g] \approx \mathbf{X}' \mathbf{V}_g \mathbf{\Sigma}_g^{-1} \mathbf{U}_g^{\mathsf{T}},\tag{7}$$

where  $\mathbf{U}_g \in \mathbb{R}^{N+1 \times r}$ ,  $\boldsymbol{\Sigma}_g \in \mathbb{R}^{r \times r}$ , and  $\mathbf{V}_g \in \mathbb{R}^{M \times r}$ . By construction, the error of this gDMD 151 method is equal to or smaller than that of sDMD (Appendix A). 152

The extended DMD (xDMD) approach<sup>9</sup> endows gDMD with a residual-learning idea. It 153 approximates the relationship between  $\mathbf{Y} = \mathbf{X'} - \mathbf{X}$  and  $\mathbf{X}$ , 154

$$\mathbf{y}_{k+1} = \mathbf{B}_x \mathbf{x}_k + \mathbf{b}_x. \tag{8}$$

Here,  $[\mathbf{B}_x \mathbf{b}_x] = \mathbf{Y} \tilde{\mathbf{X}}^{\dagger} \in \mathbb{R}^{N \times N+1}$ , and  $\tilde{\mathbf{X}}^{\top} \in \mathbb{R}^{N+1 \times M}$  is defined as before. For computational 156 saving, the best-fit linear operator is obtained through the SVD of the matrix  $\tilde{\mathbf{X}}$  as 157

$$[\mathbf{B}_{x} \mathbf{b}_{x}] \approx \mathbf{Y} \mathbf{V}_{g} \boldsymbol{\Sigma}_{g}^{-1} \mathbf{U}_{g}^{\top}.$$
(9)

The error of xDMD equals to or is smaller than that of the residual DMD without bias 159 (Appendix A). The impact of the bias term and residual learning on the accuracy of the 160 DMD method is studied in Ref. 9. An efficient computational strategy to derive prediction 161 in Eq. (8) is presented in Appendix B. 162

DMD can be used as a ROM of a nonlinear PDE, whose solution is confined in  $\mathcal{H} \subseteq \mathbb{R}^N$ 163 (to satisfy the assumptions in Lemma 1). We assess the performance of xDMD, both in 164 representation and interpolation, in terms of the relative error<sup>9,12,13</sup> 165

$$\boldsymbol{\varepsilon}_{\mathcal{L}}^{k} = \frac{\|\mathbf{x}_{k}^{\mathcal{L}} - \mathbf{x}_{k}\|^{2}}{\|\mathbf{x}_{k}\|^{2}},\tag{10}$$

where  $\|\cdot\|$  denotes vector 2-norm. 167

Several criteria can be used to select the truncation rank of a ROM<sup>1,12,13</sup>. One is to use 168 the rank of the data matrix,  $r = \operatorname{rank}(\tilde{\mathbf{X}})$ , i.e., to incorporate all the information contained 169

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in the data, including the noise. Another criterion is based on the cumulative energy in the 170 SVD of  $\tilde{\mathbf{X}}$ ; for example, one could set  $r = r_{90}$ , where 171

$$r_{90} = \min(n) : \frac{\sum_{k=0}^{n} \sigma_k}{\sum_{k=0}^{M-1} \sigma_k} \ge 0.9$$
(11)

is the number of diagonal elements of  $\Sigma$  that accounts for 90% of the energy. Yet another 173 criterion defines  $r = r^*$  as the number of diagonal elements of  $\Sigma$  associated with the first 174 singular value satisfying the inequality 175

$$r^* = \min(n) : \sigma_n \le 10^{-5} \sum_{k=0}^{M-1} \sigma_k.$$
(12)

The latter two criteria allow one to ascertain the effect of truncation of low-energy modes, as we do below.

### APPLICATION

### 3D Cardiovascular Model

We deploy the SimVascular software<sup>15</sup> to solve 3D incompressible Navier-Stokes equations describing blood flow in a patient-specific aorta. A cardiovascular model and the flow-domain geometry are selected, at random, from the Vascular Model Repository<sup>14</sup>; the homogeneous Dirichlet boundary conditions imposed at the aorta walls imply no-slip velocity at the rigid wall<sup>24</sup>. SimVascular relies on the 3D Delaunay triangulation to discretize 185 the flow domain with a triangular mesh of N = 343352 elements. (The flow-domain geometry 186 and the corresponding mesh are available for download from the Vascular Model Reposi-187 tory.) A typical 3D finite-element simulations of the unsteady Navier–Stokes of two cardiac 188 vcles for this type of geometry takes a few hours<sup>24</sup>. The quantity of interest, arranged in 189 the vector  $\mathbf{u} \in \mathbb{R}^N$  (see Section II), is the velocity magnitude of which M = 1868 snapshots, 190  $(t_k)$ , are collected over 7.7 s, which covers about 12 pulsations. Columns of matrices **X** and 191  $\mathbf{X}'$  are given by the snapshots of the velocity magnitude computed by SimVascular at a 192 constant time interval (see Section III). We chose the number of snapshots to be sufficiently 193 large to perform interpolation tests for different time steps. 194

The SimVascular predictions are used to perform multiple tests, both in representation 195 and interpolation regimes, with datasets of reduced (in space and/or time) size to verify the 196

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generality of our results. In the representation regime, these tests start with an analysis 197 of the representation error performed on the entire dataset of M = 1868 snapshots, each 198 consisting of N = 343352 grid elements. Next, ROMs are trained on randomly selected 199 data sets in which N is reduced by a tenth and a hundredth. Finally, ROMs are trained 200 on randomly selected data sets in which M is reduced to 200 snapshots associated with 201 different time intervals. In the interpolation-error analysis, we perform several tests for 202 different interpolation rates. Results and analysis of these tests are presented in the following 203 section. 204

### 205 B. Results and Discussion

### 206 1. Representation Error and Data Compression

We use xDMD to construct ROMs from the entire collection of snapshots of the velocity magnitude and testing these ROMs' ability to reproduce these training data. This exercise quantifies the representation error of xDMD. A sequence of ROMs differ from each other in the truncation rank applied to the SVD. We explore the xDMD accuracy at low-rank truncations, which are relied upon to identify dominant spatiotemporal structures in the computer-generated data. This analysis is also relevant for the exploration of xDMD's effectiveness for data compression and storage.

Figure 1a shows the ROMs' representation error, computed with Eq. (10) for different 214 truncation ranks r and averaged over all the time steps. As expected, the representation 215 error decreases with the truncation rank r. High accuracy is reached for relatively low r: 216 when  $r = \operatorname{rank}(\tilde{X}) = 1868$ , i.e., in the absence of truncation, the representation error is 217  $3.6 \cdot 10^{-16}$ ; setting  $r = r^* = 357$  or  $r = r_{90} = 24$  leads to errors of  $2.8 \cdot 10^{-5}$  or  $1.5 \cdot 10^{-1}$ , 218 respectively. By considering only 20% of the modes, with  $r = r^*$ , the result is remarkably 219 ccurate. Additionally, the cumulative energy associated with  $r^*$  is approximately equal to 220 (Fig. 1c). That is linked to the rate at which the singular values decrease to 0 (Fig. 1b), 1 221 indicating that the limited number of modes captured by  $r^*$  are dominant in the dynamics. 222 The remaining features (n > 357) are low-energy modes that do not affect the ROM accuracy; 223 as such, they can be interpreted as noise and, for the purpose of data compression, can be 224 neglected. 225





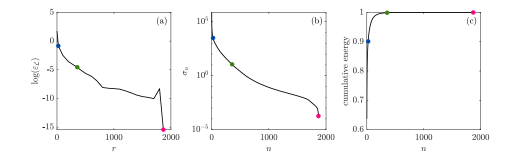


FIG. 1. (a) Representation error (averaged over the time instants) as function of the truncation rank r of the SVD of  $\tilde{\mathbf{X}}$  when all data (N = 343352, M = 1868) are used to train the ROMs. (b) Singular values and (c) cumulative energy associated with the SVD, both plotted as function of the singular values number n. In all panels, the blue, green, and red dots correspond to  $r = r_{90}$ ,  $r = r^*$ , and  $r = \operatorname{rank}(\tilde{\mathbf{X}})$ , respectively. In this example,  $r_{90} = 24$ ,  $r^* = 357$ , and  $\operatorname{rank}(\tilde{\mathbf{X}}) = M = 1868$ resulting in no truncation.

To elucidate further the effects of the truncation rank on the prediction accuracy of 226 xDMD, we compare the original data with the corresponding reconstructed snapshots pro-227 vided by the ROMs truncated at  $r_{90}$  and  $r^*$  (Fig. 2). Both ROMs reproduce the general 228 velocity patterns, although the  $r_{90}$  truncation rank returns a slightly worse approximation. 229 This comparison demonstrates the ROM ability to capture the salient features of the flow, 230 which suggests that xDMD is suitable for the interpretation and reproduction of 3D car-231 diovascular simulations. Depending on the accuracy required by the application, one can 232 select an appropriate truncation criteria and employ the xDMD-based ROM to replace the 233 onerous numerical simulations with compressed reconstructions. 234

To test the method's robustness, we train ROMs on data sets with missing spatial data. Specifically, elements of the original mesh of size N are randomly selected to obtain two reduced-size data sets of dimensions N/10 and N/100. Representation accuracy of the same values of  $r = r_{90}$ ,  $r = r^*$ , and  $r = \operatorname{rank}(\tilde{\mathbf{X}})$ . When only the dominant spatiotemporal structures of the underlying flow are considered, the accuracy close to locations where the training data are sampled is not affected by the data loss. The error increases with r,





FIG. 2. Magnitude of the flow velocity u in the aorta, as predicted by (left column) direct numerical simulations, (middle column) xDMD with truncation ranks  $r_{90}$ , and (right column) xDMD with  $r^*$ . The velocity is plotted at times k = M/3, k = 2M/3 and k = M in the first, second and third rows, respectively.

<sup>242</sup> reaching tens of orders of magnitude for  $r = \operatorname{rank}(\tilde{\mathbf{X}})$  when all the features contained in the <sup>243</sup> data are accounted for. This finding suggests that when the data are not sufficiently rich <sup>244</sup> to cover the solution space of interest, considering low-energy modes does not increase the <sup>245</sup> ROM accuracy.

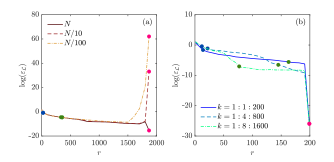


FIG. 3. Dependence of time-averaged representation error of ROMs on the SVD truncation rank r. In (a), the ROMs are alternatively trained on the data in all N pixels and on the data at randomly selected N/10 and N/100 pixels; in all three cases, using M snapshots. In (b), the ROMs are alternatively trained on the first 200 snapshots and on the 200 snapshots selected with time intervals 4 or 8; in all three cases, using N/100 pixels. The blue, green and red dots correspond to  $r = r_{90}$ ,  $r = r^*$  and  $r = \operatorname{rank}(\tilde{\mathbf{X}})$ , respectively.

Another facet of xDMD's robustness is its sensitivity to the number of temporal snapshots available for training. Figure 3b shows the representation error of the xDMD trained on N/100 velocity measurements and 200 snapshots. (These snapshots are selected from the full data set (M = 1868) using either the first 200 images or every fourth or every eighth image.) This experiment reveals that the ROM's accuracy is not affected by either the reduction of the number of snapshots or the time step between the snapshots. Hence, xDMD is robust and provides a good approximation of nonlinear flow phenomena.

### 253 2. Interpolation Error

ROMs are typically employed to make predictions at space-time points wherein the output of fluid dynamic simulations is not available. We test our ROMs' performance in the interpolation regime for several values of the interpolation rate  $\eta$ . The data-matrix dimensions and truncation ranks for all the cases considered are reported in Table I.

We start by constructing three ROMs associated with the truncation rank  $r = \operatorname{rank}(\mathbf{\tilde{X}})$ ,  $r = r^*$  and  $r = r_{90}$ , and trained on half of the snapshots, i.e.,  $\eta = 0.5$  (Case 1 in Table I).

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TABLE I. Cases considered for interpolation tests.					
Case	η	Train set	$r = r_{90}$	$r = r^*$	$\mathrm{rank}( ilde{\mathbf{X}})$
1	0.5	k = 1 : 2 : M	23	252	934
2	0.67	k=1:3:M	23	261	622
3	0.8	k = 1 : 5 : M	16	119	373
4	0.9	k = 1 : 10 : M	10	62	186

TABLE I. Cases considered for interpolation tests.

Interpolation errors of these ROMs are shown in Figure 4a; the errors are defined in Eq. (10) 260 and predictions are carried out for the missing half of time steps. The ROM truncated at 261 =  $r^*$  assures high accuracy and stability (the error varies between 10<sup>-5</sup> and 10<sup>-4</sup> at all 262 times), while the truncation at  $r = \operatorname{rank}(\tilde{\mathbf{X}})$  results in the error that increases with time; 263  $r = r_{90}$  the error is stable in time but about three orders of magnitude higher than in 264 the case of  $r = r^*$  (it varies between  $10^{-2}$  and  $10^{-0.5}$ ). Reducing the size of the training 265 et, i.e., setting  $\eta = 0.67$  (Case 2 in Table I), yields the two different ROMs truncated at  $r^*$ 266 and rank( $\tilde{\mathbf{X}}$ ) with similar interpolation errors, while  $r = r_{90}$  produces a significantly higher 267 error (Figure 4b); for all r considered, the respective ROMs' error peaks are aligned and 268 the periodicity is similar, with  $r = r^*$  providing a smaller error. In the cases of  $\eta = 0.8$  and 269 = 0.9 (Cases 3 and 4 in Table I, respectively) the interpolation errors of all the ROMs 270 increase with time and the difference when truncating at  $r^*$  and rank( $\hat{\mathbf{X}}$ ) relative to  $r = r_{90}$ 271 decreases till about one order of magnitude in the case of  $\eta = 0.9$  (Figure 4c-d). 272

To provide a local view on the ROMs' accuracy, Figure 5 compares the reference and 273 reconstructed velocity time series at two points in a cross-section of the aorta for Cases 3 274 and 4 in Table I in panels (a) and (c) and (b) and (d), respectively. As expected, the ROM 275 runcated at rank  $r = r^*$  (panels (c) and (d)) has high accuracy both in representation and 276 interpolation for all the points considered; instead, the ROM truncated at  $r = r_{90}$  (panels 277 (a) and (b)) fails to adequately reproduce the overall system state and loses accuracy when 278 or t increases. The ROM's performance is not affected by the selection of the points near 279 the wall or in the middle of the aorta. 280

These results provide actionable indicators for the rank choice and the role played by the non-dominant modes. When all the modes are included in the training phase,  $r = \operatorname{rank}(\tilde{\mathbf{X}})$ ,

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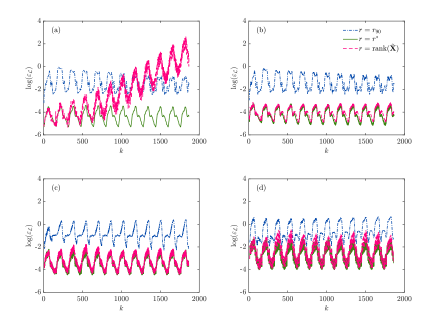


FIG. 4. Interpolation errors for (a) Case 1, (b) Case 2, (c) Case 3, and (d) Case 4 in Table I. In each plot different lines correspond to the ROMs with different truncation ranks  $r = r_{90}$ ,  $r = r^*$  and  $r = \operatorname{rank}(\tilde{\mathbf{X}})$ .

the ROM suffers from noise overfitting and loses its interpolation accuracy, especially when the training set is larger ( $\eta = 0.5$ ). The loss in accuracy is difficult to predict given the lack of a priori error estimators. Hence, the use of a low-rank truncation not only aligns with a ROM's purpose (identification of the dominant modes and data compression) but also increases the ROM's prediction reliability at space-time locations where data are not available.

### 289 V. CONCLUSION

We analyzed the performance of an extended dynamic mode decomposition (xDMD)<sup>9</sup> on the task of ROM construction to approximate the fluid dynamics simulations of 3D blood

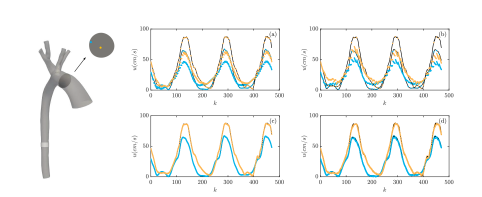


FIG. 5. Flow velocity u at kth time step, provided by SimVascular (continuous line) and estimated by the ROMs (dots) in the interpolation regime. The ROMs are trained for Case 3 in Table I in panels (a) and (c), and for Case 4 in Table I in panels (b) and (d). The data are reported for two points in one aorta's cross-section, as shown on the left. Panels (a) and (b) refer to the ROMs truncated at  $r = r_{90}$ , while (c) and (d) refer to the ROMs truncated at  $r = r^*$ .

flow in a patient-specific aorta. Our results show that xDMD is able to identify dominant 292 spatiotemporal structures in the simulated data set and to provide an accurate approxima-293 tion of numerical simulations. We explored relevant indicators of a ROM's performance in 294 both representation and interpolation. These indicators are related to the choice of the trun-295 ation rank and linked to the number of retained singular values corresponding to the most 296 relevant spatiotemporal structures. We found that a low-rank truncation, which preserves 297 almost all the cumulative energy in the data, avoids overfitting and yields high accuracy and 298 error stability. The xDMD-based ROMs demonstrate a remarkable robustness to the num-299 ber of space-time training data. Finally, we verified the local accuracy of xDMD when used 300 to predict time series at selected points in the flow domain. Overall, our study suggests that 301 the use of xDMD is beneficial for time-dependent data compression and for computational 302 saving when used in place of onerous numerical simulations. 303

The use of DMD for order reduction offers other benefits as well. By identifying the dominant spatially correlated structures (modes) in a given dataset and analyzing their temporal evolution (time dynamics), we can gain insight into the main features of the physical process, facilitating both data interpretation and reconstruction. DMD not only enables data

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compression, which is beneficial in many fields, but also allows us to reconstruct the sys-308 tem's behavior where data is unavailable (in interpolation or extrapolation regimes) with a 309 single linear model providing predictions everywhere in space at any given time. This linear 310 model is readily interpretable and is cleansed of noise, which would otherwise impede the 311 reconstruction. 312

Our study demonstrates that the identification of an optimal DMD structure requires 313 the selection of a low-rank approximation able to guarantee the ROM's accuracy in both 314 representation and interpolation. This instill trust in the ROM's predictions, paving the way 315 for their use in clinical practice. For example, DMD can be employed to predict blood flow 316 beyond the available data to study variations in the flow waveform<sup>17</sup>, to provide reliable 317 real-time forecasting of tumor ablation treatment $^{25}$ , and to facilitate spectral analysis in 318 dynamic MRI acquisitions to advance the diagnostic potential<sup>20</sup>. 319

Since DMD is formulated entirely in terms of (observational and/or simulated) data, it 320 can be readily deployed in a wide range of applications, including in real-time simulation 321 environments. In this context, newly available data can be absorbed in the training phase 322 while updating the future state prediction. 323

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## Appendix B: Strategy to increase the xDMD efficiency 348 Direct evaluation of (9) requires the computation of $[\mathbf{B}_x \mathbf{b}_x] \in \mathbb{R}^{N \times N+1}$ . Since N is large 349 in any application of practical significance, this computation decreases the efficiency and 350 accuracy of the algorithm. To avoid this bottleneck, we decompose the computation into 351 two parts. First, we multiply only the first three terms of (9) thus leading to the matrix 352 353 Second, we multiply the last term in (9) by $\tilde{\mathbf{x}}_k$ , which gives a vector 354 355

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 $\mathbf{d}_x = \mathbf{U}_g^\top \tilde{\mathbf{x}}_k \in \mathbb{R}^{r \times 1}.$ (B2)

This procedure leads to 356

Appendix A: Error bounds

defined in (3), satisfies the inequality

general guideline for the growth of errors.

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347 а

$$\mathbf{y}_{k+1} = \mathbf{C}_x \mathbf{d}_x,\tag{B3}$$

(B1)

which is equivalent to (8). 358

An overall step-by-step implementation of xDMD with the efficient computational strat-359 egy described in this Section, is illustrated in Algorithm 1. 360

In addition to the assumptions in Lemma 1, we assume that  $\|\mathcal{L} - \Phi_{\Delta t}\|_{L^{\infty}(\mathcal{H}_{\Delta t})} < +\infty$  and

that  $\mathbf{x}_k$  and  $\mathbf{x}_k^{\mathcal{L}} \in \mathcal{H}_{\Delta t}$  for  $k = 0, \dots, M - 1$ . If  $\mathcal{L}$  is sDMD, then the error  $\delta_{\mathcal{L}}^M$  at time  $t_M$ ,

 $\delta_{\mathcal{L}}^{M} \leq \mathrm{e}^{ML\Delta t} \delta_{\mathcal{L}}^{0} + \parallel \mathcal{L} - \Phi_{\Delta t} \parallel_{L^{\infty}(\mathcal{H}_{\Delta t})} \sum_{l=0}^{M-1} \mathrm{e}^{kL\Delta t}.$ 

The proof, based on the triangle inequality, follows that for Theorem 4.3 in Ref. 11. More-

 $\delta_{\mathcal{L}}^{M} \leq (1 + \mathrm{e}^{L\Delta t})^{M} \delta_{\mathcal{L}}^{0} + \parallel \mathcal{L} - \Phi_{\Delta t} \parallel_{L^{\infty}(\mathcal{H}_{\Delta t})} \sum_{k=0}^{M-1} (1 + \mathrm{e}^{L\Delta t})^{k}.$ 

The xDMD is proven to have a tighter error bound than rDMD.<sup>9</sup> The error bounds provide

 $\mathbf{C}_{x} = \mathbf{Y}\mathbf{V}_{g}\boldsymbol{\Sigma}_{o}^{-1} \in \mathbb{R}^{N \times r}.$ 

Similarly, if  $\mathcal{L}$  is the xDMD, then the error  $\delta_{\mathcal{L}}^{M}$  at time  $t_{M}$  satisfies the inequality

over, the gDMD is proven to have a tighter error bound than sDMD.<sup>9</sup>

1	Algorithm 1: xDMD implementation based on the efficient computational
	strategy.
1. (	Compute the residual matrix $\mathbf{Y}$ : $\mathbf{Y} = \mathbf{X}' - \mathbf{X}$ , where $\mathbf{Y} \in \mathbb{R}^{N \times M}$
2. 1	Introduce the matrix $\tilde{\mathbf{X}}$ : $\tilde{\mathbf{X}}^{\top} = [\mathbf{X} \ 1]$ , where $\tilde{\mathbf{X}}^{\dagger} \in \mathbb{R}^{N \times N+1}$
3. (	Compute the truncated SVD of $\tilde{\mathbf{X}}$ : $\tilde{\mathbf{X}} \approx \mathbf{U}_g \boldsymbol{\Sigma}_g \mathbf{V}_g^{T}$ , where
$\mathbf{U}_g$	$\in \mathbb{R}^{N+1  imes r}, \mathbf{\Sigma}_g \in \mathbb{R}^{r  imes r}, \mathbf{V}_g \in \mathbb{R}^{M  imes r}$
4. (	Compute the matrix $\mathbf{C}_x$ : $\mathbf{C}_x = \mathbf{Y}\mathbf{V}_g \boldsymbol{\Sigma}_g^{-1}$ , where $\mathbf{C}_x \in \mathbb{R}^{N \times r}$
5. (	Compute the vector $\mathbf{d}_x$ : $\mathbf{d}_x = \mathbf{U}_g^\top \tilde{\mathbf{x}}_k$ , where $\mathbf{d}_x \in \mathbb{R}^{r \times 1}$
6. (	Compute the residual at $k + 1$ : $\mathbf{y}_{k+1} = \mathbf{C}_x \mathbf{d}_x$ , where $\mathbf{y}_{k+1} \in \mathbb{R}^{N \times 1}$
7. (	Compute the state at $k + 1$ : $\mathbf{x}_{k+1} = \mathbf{y}_{k+1} + \mathbf{x}_k$ , where $\mathbf{x}_{k+1} \in \mathbb{R}^{N \times 1}$

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