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Computational Science Laboratory
Computer Science Department
Virginia Polytechnic Institute and State University
Blacksburg, VA 24060
Phone: (540)-231-2193
Fax: (540)-231-6075
Email: sandu@cs.vt.edu
Web: http://csl.cs.vt.edu
Comparison of POD reduced order strategies for the nonlinear 2D Shallow Water Equations

Răzvan Ștefănescu *, Adrian Sandu †, and Ionel M. Navon ‡

1 Computational Science Laboratory, Department of Computer Science, Virginia Polytechnic Institute and State University, Blacksburg, Virginia, USA, 24060
2 Department of Scientific Computing, The Florida State University, Tallahassee, Florida, USA, 32306

Abstract

This paper introduces tensoria calculus techniques in the framework of Proper Orthogonal Decomposition (POD) to reduce the computational complexity of the reduced nonlinear terms. The resulting method, named tensorial POD, can be applied to polynomial nonlinearities of any degree \( p \). Such nonlinear terms have an on-line complexity of \( O(k^{p+1}) \), where \( k \) is the dimension of POD basis, and therefore is independent of full space dimension. However it is efficient only for quadratic nonlinear terms since for higher nonlinearities standard POD proves to be less time consuming once the POD basis dimension \( k \) is increased. Numerical experiments are carried out with a two dimensional shallow water equation (SWE) test problem to compare the performance of tensorial POD, standard POD, and POD/Discrete Empirical Interpolation Method (DEIM). Numerical results show that tensorial POD decreases by 76× times the computational cost of the on-line stage of standard POD for configurations using more than 300,000 model variables. The tensorial POD SWE model was only 2−8× slower than the POD/DEIM SWE model but the implementation effort is considerably increased. Tensorial calculus was again employed to construct a new algorithm allowing POD/DEIM shallow water equation model to compute its off-line stage faster than the standard and tensorial POD approaches.

Keywords — tensorial proper orthogonal decomposition; discrete empirical interpolation method; reduced-order modeling; shallow water equations; finite difference methods; Galerkin projections

1 Introduction

Modeling and simulation of multi-scale complex physical phenomena leads to large-scale systems of coupled partial differential equations, ordinary differential equations, and differential algebraic equations. The high dimensionality of these models poses important mathematical and computational challenges. A computationally feasible approach to simulate, control, and optimize such systems is to simplify the models by retaining only those state variables that are consistent with a particular phenomena of interest.

Reduced order modeling refers to the development of low-dimensional models that represent important characteristics of a high-dimensional or infinite dimensional dynamical system. The reduced order methods can be cast into three broad categories: Singular Values Decomposition (SVD) based methods, Krylov based methods and iterative methods combining aspects of both the SVD and Krylov methods (see e.g. Antoulas [3]).

* rstefane@vt.edu
† sandu@cs.vt.edu
‡ inavon@fsu.edu
For linear models, methods like balanced truncation (Antoulas [4], Moore [57], Mullis and Roberts [55], Sorensen and Antoulas [73]) and moment matching (Feldmann and Freund [27], Freund [28], Grimme [35]) have been proving successful in developing reduced order models. However, balanced truncation doesn’t extend easily for high-order systems, and several grammians approximations were proposed leading to methods such as approximate subspace iteration (Baker et al. [6]), least squares approximation (Hodel [31]), Krylov subspace methods (Jaimoukha and Kasenally [32] and Gudmundsson and Laub [36]) and balanced Proper Orthogonal Decomposition (Willcox and Peraire [55]). Among moment matching methods we mention partial realization (Benner and Sokolov [12], Gragg and Lindquist [31]), Padé approximation (Gallivan et al. [29], Gragg [30], Gutknecht [35], Van Dooren [84]) and rational approximation (Bultheel and Moor [16]).

While for linear models we are able to produce input-independent highly accurate reduced models, in the case of general nonlinear systems, the transfer function approach is not yet applicable and input-specific semi-empirical methods are usually employed. Recently some encouraging research results using generalized transfer functions and generalized moment matching have been obtained by Benner and Breiten [11] for nonlinear model order reduction but future investigations are required.

Proper Orthogonal Decomposition and its variants are also known as Karhunen-Loève expansions [45, 52], principal component analysis Hotelling [42], and empirical orthogonal functions Lorenz [54] among others. It is the most prevalent basis selection method for nonlinear problems and, among other requirements, relies on the fact that the desired simulation is well simulated in the input collection. Data analysis using POD is conducted to extract basis functions, from experimental data or detailed simulations of high-dimensional systems (method of snapshots introduced by Sirovich [75, 76, 77]), for subsequent use in Galerkin projections that yield low dimensional dynamical models. Unfortunately the standard POD approach displays a major disadvantage since its nonlinear reduced terms still have to be evaluated on the original state space making the simulation of the reduced-order system too expensive. There exist several ways to avoid this problem such as the empirical interpolation method (EIM) Barrault et al. [7] and its discrete variant DEIM Chaturantabut [20], Chaturantabut and Sorensen [22, 23], best points interpolation method Nguyen et al. [60], Missing point estimation Astrid et al. [5] and Gauss-Newton with approximated tensors Amsallem et al. [2], Carlberg and Farhat [17], Carlberg et al. [18, 19, 19] methods are relying upon the gappy POD technique Everson and Sirovich [25] and were developed for the same reason. Reduced basis methods have been recently developed and utilize on greedy algorithms to efficiently compute numerical solutions for parametrized applications Barrault et al. [7], Dihlmann and Haasdonk [24], Grepl and Patera [33], Patera and Rozza [63], Rozza et al. [70].

Dynamic mode decomposition is a relatively recent development in the field of modal decomposition (Rowley et al. [69], Schmid [74], Tissot et al. [82]) and in comparison with POD approximates the temporal dynamics by a high-degree polynomial. Trajectory piecewise linear method proposed by Rewienski and White [68] follows a different strategy where the nonlinear system is represented by a piecewise-linear system which then can be efficiently approached by the standard linear reduction method. Parameter model reduction has emerged recently as an important research direction and Benner et al. [11] highlights the major contribution in the field.

This paper combines standard POD and tensor calculus techniques to reduce the on-line computational complexity of the reduced nonlinear terms for a shallow water equations model. Tensor based calculus was already applied by Kunisch and Volkwein [48], Kunisch et al. [50] to represent quadratic nonlinearities of reduced order POD models. We show that the tensorial POD (TPOD) approach can be applied to polynomial nonlinearities of any degree $p$, and the its representation has a complexity of $O(k^{p+1})$, where $k$ is the dimension of POD subspace. This complexity is independent of the full space dimension. For $k$ between 10 and 50 and $p = 2$ the number of floating-point operations required to calculate the tensorial POD quadratic terms is $10^{-40}$ times lower than in the case of standard POD, and $10^{-20}$ times higher than for the POD/DEIM. However, CPU time for solving the TPOD SWE model (on-line stage) is only $2-8$ times slower than POD/DEIM SWE model for $10^3-10^5$ grid points, $k \leq 50$, and number of DEIM interpolation points $m < 180$. For example, for an integration interval of 3h, $10^5$ mesh points, $k = 50$, and $m = 70$, tensorial POD and POD/DEIM are $76 \times$ and $450 \times$ faster than standard POD, but the implementation effort of POD/DEIM is considerably increased. Many useful models are characterized by quadratic nonlinearities in both fluid dynamics and geophysical fluid flows including SWE model. In the
case of cubic or higher polynomial nonlinearities the advantage of tensorial POD is lost and its nonlinear computational complexity is similar or larger than the computational complexity of the standard POD approach. This proves that for models depending only on quadratic nonlinearities, the tensorial POD represents a solid alternative to POD/DEIM where the implementation effort is considerably larger. We also propose a fast algorithm to pre-compute the reduced order coefficients for polynomial nonlinearities of order \( p \) which allows the POD/DEIM SWE model to compute its off-line stage faster than the standard and tensorial POD approaches despite additional SVD calculations and reduced coefficients computations.

The paper is organized as follows. Section 2 reviews the reduced order modeling methodologies used in this work: standard, tensorial, and DEIM POD. Section 3 analyses the computational complexity of the reduced order polynomial nonlinearities for all three methods, and introduces a new DEIM based algorithm to efficiently compute the coefficients needed for reduced Jacobians. Section 4 discusses the shallow water equations model and its full implementation, and Section 5 describes the construction of reduced models. Results of extensive numerical experiments are discussed in Section 6 while conclusions are drawn in Section 7.

2 Reduced Order Modeling

For highly efficient flows simulations, reduced order modeling is a powerful tool for representing the dynamics of large-scale dynamical systems using only a smaller number of variables and reduced order basis functions. Three approaches will be considered in this study: standard Proper Orthogonal Decomposition (POD), tensorial POD (TPOD), and POD/Discrete Empirical Interpolation Method (POD/DEIM). They are discussed below. The tensorial POD approach proposed herein is different than the method of Belzen and Weiland [10] which makes use of tensor decompositions for generating POD bases.

2.1 Standard Proper Orthogonal Decomposition

Proper Orthogonal Decompositions has been used successfully in numerous applications such as compressible flow Rowley et al. [67], computational fluid dynamics Kunisch and Volkwein [49], Rowley [68], Willcox and Peraire [85], aerodynamics [15]. It can be thought of as a Galerkin approximation in the spatial variable built from functions corresponding to the solution of the physical system at specified time instances. Noack et al. [62] proposed a system reduction strategy for Galerkin models of fluid flows leading to dynamical models of lower order based on a partition in slow, dominant and fast modes. San and Iliescu [73] investigate several closure models for POD reduced order modeling of fluids flows and benchmarked against the fine resolution numerical simulation.

In what follows, we will only work with discrete inner products (Euclidian dot product) though continuous products may be employed too. Generally, an atmospheric or oceanic model is usually governed by the following semi–discrete dynamical system

\[
\frac{d\mathbf{x}(t)}{dt} = \mathbf{F}(\mathbf{x}, t), \quad \mathbf{x}(0) = \mathbf{x}_0 \in \mathbb{R}^n. \tag{1}
\]

From the temporal-spatial flow \( \mathbf{x}(t) \in \mathbb{R}^n \), we select an ensemble of \( N_t \) time instances \( \mathbf{x}_1, ..., \mathbf{x}_{N_t} \in \mathbb{R}^n \), \( n \) being the total number of discrete model variables per time step and \( N_t \in \mathbb{N}, N_t > 0 \). Let us define the centering trajectory, shift mode, or mean field correction (Noack et al. [61]) \( \bar{x} = \frac{1}{N_t} \sum_{i=1}^{N_t} \mathbf{x}_i \). The method of POD consists in choosing a complete orthonormal basis \( U = \{ \mathbf{u}_i \}, \quad i = 1, ..., k; \quad k > 0; \quad \mathbf{u}_i \in \mathbb{R}^n; \quad U \in \mathbb{R}^{n \times k} \) such that the mean square error between \( \mathbf{x}(t) \) and POD expansion \( \mathbf{x}^{POD}(t) = \bar{x} + U\mathbf{r}(t) \), \( \mathbf{r}(t) \in \mathbb{R}^k \) is minimized on average. The POD dimension \( k \ll n \) is appropriately chosen to capture the dynamics of the flow as follows:

To obtain the reduced model of (1), we first employ a numerical scheme to solve the full model for a set of snapshots and follow the above procedure, then use a Petrov–Galerkin (PG) projection of the full model equations onto the space \( \mathcal{X}^k \) spanned by the POD basis elements

\[
\frac{d\mathbf{\bar{x}}(t)}{dt} = W^T\mathbf{F}\left(\mathbf{\bar{x}} + U\mathbf{\bar{x}}(t), t\right), \quad \mathbf{\bar{x}}(0) = W^T\left(\mathbf{x}(0) - \bar{x}\right), \tag{2}
\]
These are also used for reduced Jacobian computation in the online stage. The projected spatial variables are stored into tensors and calculated off-line.

It can be successfully used in a POD framework for finite difference (FD), finite element (FE) and finite volume (FV) discretization methods and all other type of discretization methods that engage in spectral expansions. Tensorial POD separates the full spatial variables from reduced variables allowing fast nonlinear terms computations in the on-line stage. For time dependent nonlinearities this implies separation of spatial variables from reduced time variables. Thus, the reduced nonlinear term evaluation requires a tensorial Frobenius dot-product computation between rank \( p \) tensors, where \( p \) is the order of polynomial nonlinearity. The projected spatial variables are stored into tensors and calculated off-line. These are also used for reduced Jacobian computation in the on-line stage.

The tensorial POD technique employs the simple structure of the polynomial nonlinearities to remove the dependence on the dimension of the original discretized system by manipulating the order of computing. It can be successfully used in a POD framework for finite difference (FD), finite element (FE) and finite volume (FV) discretization methods and all other type of discretization methods that engage in spectral expansions. Tensorial POD separates the full spatial variables from reduced variables allowing fast nonlinear terms computations in the on-line stage. For time dependent nonlinearities this implies separation of spatial variables from reduced time variables. Thus, the reduced nonlinear term evaluation requires a tensorial Frobenius dot-product computation between rank \( p \) tensors, where \( p \) is the order of polynomial nonlinearity. The projected spatial variables are stored into tensors and calculated off-line. These are also used for reduced Jacobian computation in the on-line stage.

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Discrete Empirical Interpolation Method. The former approach is able to calculate the reduced polynomial nonlinearities independent of \( n \), while the latter method can handle efficiently all type of nonlinearities.

\( \gamma = 1 - \frac{\text{percentage of total informations captured by the reduced space \( span\{u_1, u_2, ..., u_k\} \)}}{\text{number of variables of the full model}} \)

\( \gamma \) is taken 0.99.

To mitigate this inefficiency we propose two approaches: (1) Tensorial POD and (2) Discrete Empirical Interpolation Method. The former approach is able to calculate the reduced polynomial nonlinearities independent of \( n \), while the latter method can handle efficiently all type of nonlinearities.

\[ \tilde{N}(\bar{x}) = W^T (U\bar{x}(t))^p \]

where vector powers are taken component-wise.

To be precise, consider a steady polynomial nonlinearity \( x^p \). A POD expansion involving mean \( \bar{x} \) will unnecessarily complicate the description of tensorial POD representation of a \( p \)th order polynomial nonlinearity. Moreover the terms depending on \( \bar{x} \) are just a particular case of the term depending only on \( U\bar{x} \) since vector componentwise multiplication is distributive over vector addition. Consequently the expansion \( \tilde{x} \approx U\bar{x} \) will not decrease the generality of the reduced nonlinear term. In the finite difference case, the standard POD projection is described as follows

\[ \tilde{N}(\bar{x}) = W^T (U\bar{x})^p \]

Algorithm 1 POD basis construction

1. Calculate the mean \( \bar{x} = \frac{1}{N} \sum_{i=1}^{N} x_i \).
2. Set up the correlation matrix \( K = [k_{ij}]_{i,j=1,...,n} \) where \( k_{ij} = \langle x_i - \bar{x}, x_j - \bar{x} \rangle \), and \( \langle \cdot, \cdot \rangle \) being the Euclidean dot product.
3. Compute the eigenvalues \( \lambda_1 \geq \lambda_2 \geq ... \lambda_n \geq 0 \) and the corresponding orthogonal eigenvectors \( v^1, v^2, ..., v^n \in \mathbb{R}^n \) of \( K \).
4. Define \( I(m) = \frac{\sum_{i=1}^{m} \lambda_i}{\sum_{i=1}^{n} \lambda_i} \) and choose \( k \) such that \( k = \min\{I(m) : I(m) \geq \gamma \} \) where \( 0 \leq \gamma \leq 1 \) is the percentage of total informations captured by the reduced space \( \text{span}\{u_1, u_2, ..., u_k\} \). Usually \( \gamma \) is taken 0.99.

where \( W \in \mathbb{R}^{n \times k} \) contains the discrete test functions from the PG projection, i.e. \( W^T U = I \in \mathbb{R}^k \). The Galerkin projection may be also a choice being just a particular case of PG \( (W = U) \).

The efficiency of the POD-Galerkin techniques is limited to linear or bilinear terms, since the projected nonlinear term at every discrete time step still depends on the number of variables of the full model:

\[ \tilde{N}(\bar{x}) = W^T U \bar{x}(t). \]
where $p$-order tensors $\tilde{X}$ and $M^i$, $i = 1, 2, \ldots, k$ are defined as

$$\tilde{X} = \left[ X_{i_1i_2\ldots i_p} \right]_{i_1i_2\ldots i_p=1\ldots k} \in \mathbb{R}^{k \times \ldots \times k} ; \quad \tilde{X}_{i_1i_2\ldots i_p} = \tilde{x}_{i_1}\tilde{x}_{i_2}\ldots\tilde{x}_{i_p} \in \mathbb{R} ;$$

$$M^i = \left[ M^i_{i_1i_2\ldots i_p} \right]_{i_1i_2\ldots i_p=1,2\ldots k} \in \mathbb{R}^{k \times \ldots \times k} ;$$

$$M^i_{i_1i_2\ldots i_p} = \sum_{l=1}^{n} W_l U_{i_1l} U_{i_2l} \ldots U_{i_pl} \in \mathbb{R} ,$$

and $\tilde{x}_{i_j}, W_{li}, U_{ij}$ are just entries of POD reduced order solution $\tilde{x}$, POD test functions basis $W$ and POD trial functions basis $U$. The tensorial Frobenius dot product is defined as

$$\langle \cdot , \cdot \rangle_{\text{Frobenius}} : \mathbb{R}^{k \times \ldots \times k} \times \mathbb{R}^{k \times \ldots \times k} \rightarrow \mathbb{R} ;$$

$$\langle A , B \rangle_{\text{Frobenius}} = \sum_{i_1,i_2\ldots,i_p=1}^{k} A_{i_1i_2\ldots i_p} B_{i_1i_2\ldots i_p} \in \mathbb{R} .$$

We note that $M^i$, $i = 1, 2, \ldots, k$ are $p^{th}$ order tensors computed in the off-line stage and their dimensions do not depend on the full space dimension. For finite element and finite volume the tensorial POD representations are the same except for the type of products used in computation of $M^i_{i_1i_2\ldots i_p}$ in [5] which now are continuous and replace the sum of products used in the finite difference case.

Reduced nonlinearities depending on space derivatives are treated similarly as in equation $[3]-[5]$ since POD expansion of $x_x$ (space derivative of $x$) is $U_x \tilde{x}$, where $U_x \in \mathbb{R}^{n \times k}$ contains the space derivatives of POD basis functions $u_i$, $i = 1, 2, \ldots, k$, $u_i \in \mathbb{R}^n$.

### 2.3 Standard POD and Discrete Empirical Interpolation Method

The empirical interpolation method (EIM) and its discrete version (DEIM), were developed to approximate the nonlinear term allowing an effectively affine offline-online computational decomposition. Both interpolation methods provide an efficient way to approximate nonlinear functions. They were successfully used in a standard POD framework for finite difference (FD), finite element (FE) and finite volume (FV) discretization methods. A description of EIM in connection with the reduced basis framework and a posteriori error bounds can be found in Grepl et al. [34], Maday et al. [50].

The DEIM implementation is based on a POD approach combined with a greedy algorithm while the EIM implementation relies on a greedy algorithm Lass and Volkwein [51].

For $m \ll n$ the finite difference POD/DEIM nonlinear term approximation is

$$\hat{N}(\tilde{x}) \approx W^T V (P^T V)^{-1} F (P^T (\tilde{x} + U \tilde{x})) ,$$

where $V \in \mathbb{R}^{n \times m}$ gathers the first $m$ POD basis modes of nonlinear function $F$ while $P \in \mathbb{R}^{n \times m}$ is the DEIM interpolation selection matrix.

The POD/DEIM approximation of (3) is

$$\hat{N}(\tilde{x}) \approx W^T V (P^T V)^{-1} (P^T U \tilde{x})^p$$

where vector powers are taken component-wise and $P^T V \in \mathbb{R}^{m \times m}$, $P^T U \in \mathbb{R}^{m \times k}$. $P^T U$ is also recommended for pre-computation in the off-line stage.

DEIM has developed in several research directions, e.g., rigorous state space error bounds Chatu-rantabut and Sorensen [23], a posteriori error estimation Wirtz et al. [84], 1D FitzHugh-Nagumo model...
Chaturantabut and Sorensen [22], 1D simulating neurons model Kellems et al. [46], 1D nonlinear thermal model Hochman et al. [40], 1D Burgers equation Aanonsen [1], Chaturantabut [20], 2D nonlinear miscible viscous fingering in porous medium [21], oil reservoirs models Suwartadi [81], and 2D SWE model Stefanescu and Navon [79]. We emphasize that only few POD/DEIM studies with FE and FV methods were performed, e.g., for electrical networks Hinze and Kunkel [39] and for a 2D ignition and detonation problem Bo [14]. Flow simulations past a cylinder using a hybrid reduced approach combining the quadratic expansion method and DEIM are available in Xiao et al. [87].

3 Computational Complexity of the reduced \( p^{th} \) order nonlinear representations. ROMs off-line stage discussion.

We will focus on finite difference reduced order \( p^{th} \) order polynomial nonlinearities [346]. We begin with an important observation. For POD ROMs construction the usually approach is to store each of the state variables separately and to project every model equations to a different POD basis corresponding to a state variable whose time derivative is present. This is also the procedure we employed for this study. In this context \( n \) doesn’t denote the total number of state variables but only the number of state variables of the same type which most of the time is equal with the number of mesh points. Consequently from now on we refer to \( n \) as the number of spatial points.

Standard POD representation is computed with a complexity of \( O(p \times k \times n + (p−1) \times n + k \times n) \) and the POD/DEIM term requires \( O(p \times k \times n + (p−1) \times m + k \times m) \) basic operations in the on-line stage. Tensorial POD nonlinear term has a complexity of \( O(k^{p+1}) \). While standard POD computational complexity still depends on the full space dimension the other two tensorial POD and POD/DEIM don’t. Table 1 describes the number of operations required to compute the projected \( p^{th} \) order polynomial nonlinearity for each of the three ROMs approaches and various values of \( n, k, m, p \).

<table>
<thead>
<tr>
<th>( n )</th>
<th>( k )</th>
<th>( m )</th>
<th>( p )</th>
<th>POD</th>
<th>POD/DEIM</th>
<th>Tensorial POD</th>
</tr>
</thead>
<tbody>
<tr>
<td>( 10^3 )</td>
<td>10</td>
<td>10</td>
<td>2</td>
<td>31,000</td>
<td>310</td>
<td>2,990</td>
</tr>
<tr>
<td>( 10^3 )</td>
<td>10</td>
<td>10</td>
<td>3</td>
<td>42,000</td>
<td>420</td>
<td>29,990</td>
</tr>
<tr>
<td>( 10^3 )</td>
<td>10</td>
<td>10</td>
<td>4</td>
<td>53,000</td>
<td>530</td>
<td>299,990</td>
</tr>
<tr>
<td>( 10^4 )</td>
<td>30</td>
<td>50</td>
<td>2</td>
<td>910,000</td>
<td>4550</td>
<td>80,970</td>
</tr>
<tr>
<td>( 10^4 )</td>
<td>30</td>
<td>50</td>
<td>3</td>
<td>1,220,000</td>
<td>6100</td>
<td>2,429,970</td>
</tr>
<tr>
<td>( 10^5 )</td>
<td>50</td>
<td>100</td>
<td>2</td>
<td>15,100,000</td>
<td>15,100</td>
<td>374,950</td>
</tr>
<tr>
<td>( 10^5 )</td>
<td>50</td>
<td>100</td>
<td>3</td>
<td>20,200,000</td>
<td>20,200</td>
<td>18,749,950</td>
</tr>
<tr>
<td>( 10^6 )</td>
<td>50</td>
<td>100</td>
<td>4</td>
<td>25,300,000</td>
<td>25,300</td>
<td>937,499,950</td>
</tr>
</tbody>
</table>

Table 1: Number of floating-point operations in the on–line stage for different numbers of spatial points \( n \), POD modes \( k \), DEIM points \( m \), and polynomial orders \( p \).

Clearly POD/DEIM provides the fastest nonlinear terms computations in the on-line stage. For quadratic nonlinearities, i.e. \( p = 2 \), and \( n = 10^5 \), POD/DEIM outperforms POD and POD tensorial by \( 10^3 \times \) and \( 25 \times \) times. But these performances are not necessarily translated into the same CPU time rates for solving the ROMs solutions since other more time consuming calculations may be needed (reduced Jacobians computations and their LU decompositions). It was already proven in Stefanescu and Navon [79] that for a SWE model DEIM decreases the computational complexity of the standard POD by \( 60 \times \) for full space dimensions \( n \geq 60,000 \), and leads to a CPU time reduction proportional to \( n \). CPU times and error magnitudes comparisons will be discussed in Numerical Results Section 6.

For cubic nonlinearities (\( p = 3 \)), the computational complexities are almost similar for both tensorial and standard POD while for higher nonlinearities (e.g. \( p = 4 \)) tensorial POD cost becomes prohibitive.

In the context of reduced optimization, the off–line stage computational complexity weights heavily in the final CPU time costs since several POD bases updates and DEIM interpolation points recalculations are needed during the minimization process. Since the proposed schemes are implicit in time we need to
compute the reduced Jacobians as a part of a Newton type solver. For the current study we choose to calculate derivatives exactly for all three ROMs. Consequently, some reduced coefficients such as tensors $M^i$ defined in (5) must be calculated for all three reduced approaches including POD/DEIM in the off-line stage.

A simple evaluation suggests that POD/DEIM off-line stage will be slower than the corresponding tensorial POD and POD stages since more SVD computations are required in addition to particular POD/DEIM coefficients and DEIM index points calculations (see Table 2). At a more careful examination we noticed that we can exploit the structure of POD/DEIM nonlinear term (6) like in the tensorial POD approach (4,5) and provide a fast calculation for $M^i$.

Thus, let us denote the precomputed term and $P^T U$ in (6) by

$$E = W^T V (P^T V)^{-1} \in \mathbb{R}^{k \times m}$$

and

$$U_m = P^T U \in \mathbb{R}^{m \times k},$$

where $m$ is the number of DEIM points.

The $p$-tensor $M^i$ can be computed as follows:

$$M^i_{i_1i_2...i_p} = \sum_{l=1}^{m} E_{il} U^m_{i_1l} U^m_{i_2l} ... U^m_{i_pl} \in \mathbb{R}, \quad i, i_1, i_2, ..., i_p = 1, 2, ..., k.$$  \hspace{1cm} (7)

Clearly, this estimation is less computationally expensive then (5) since the summation stops at $m \ll n$.

During the numerical experiments we observed that tensors $M^i$, $i = 1, 2, ..., k$ calculated in POD/DEIM off-line stage (7) are different than $M^i$, $i = 1, 2, ..., k$ obtained in the tensorial POD case (5), but $M$ the reduced nonlinear term estimations of $\tilde{N}(\tilde{x})$ are accurate for both methods. We also mention for both standard POD and POD/DEIM approaches terms as $M^i$, $i = 1, 2, ..., k$ are used only for reduced Jacobian computations. In the case of POD/DEIM method, this leads to different derivatives values than in the case of tensorial POD or standard POD but the output solution error results are accurate as will see in Section 6.

4 The Shallow Water Equations

In meteorological and oceanographic problems, one is often not interested in small time steps because the discretization error in time is small compared to the discretization error in space. SWE can be used to model Rossby and Kelvin waves in the atmosphere, rivers, lakes and oceans as well as gravity waves in a smaller domain. The alternating direction fully implicit (ADI) scheme Gustafsson [37] considered in this paper is first order in both time and space and it is stable for large CFL condition numbers (we tested the stability of the scheme for a CFL condition number equal up to 8.9301). It was also proved that the method is unconditionally stable for the linearized version of the SWE model. Other research work on this topic include efforts of Fairweather and Navon [26], Navon and Villiers [59].

We are solving the SWE model using the $\beta$-plane approximation on a rectangular domain Gustafsson [37]

$$\frac{\partial w}{\partial t} = A(w) \frac{\partial w}{\partial x} + B(w) \frac{\partial w}{\partial y} + C(y) w, \quad (x, y) \in [0, L] \times [0, D], \quad t \in (0, t_f],$$  \hspace{1cm} (8)

where $w = (u, v, \phi)^T$ is a vector function, $u, v$ are the velocity components in the $x$ and $y$ directions, respectively, $h$ is the depth of the fluid, $g$ is the acceleration due to gravity, and $\phi = 2\sqrt{gh}$.

The matrices $A, B$ and $C$ are

$$A = \begin{pmatrix} u & 0 & \phi/2 \\ 0 & u & 0 \\ \phi/2 & 0 & u \end{pmatrix}, \quad B = \begin{pmatrix} v & 0 & 0 \\ 0 & v & \phi/2 \\ 0 & \phi/2 & v \end{pmatrix}, \quad C = \begin{pmatrix} 0 & f & 0 \\ -f & 0 & 0 \\ 0 & 0 & 0 \end{pmatrix},$$

where $f$ is the Coriolis term

$$f = \dot{f} + \beta(y - D/2), \quad \beta = \frac{\partial f}{\partial y}, \quad \forall y,$$

with $\dot{f}$ and $\beta$ constants.
We assume periodic solutions in the $x$ direction for all three state variables while in the $y$ direction
\[
v(x, 0, t) = v(x, D, t) = 0, \ x \in [0, L], \ t \in (0, t^f]
\]
and Neumann boundary condition are considered for $u$ and $\phi$.

Initially $w(x, y, 0) = \psi(x, y), \ \psi : \mathbb{R} \times \mathbb{R} \rightarrow \mathbb{R}, \ (x, y) \in [0, L] \times [0, D]$. Now we introduce a mesh of $n = N_x \times N_y$ equidistant points on $[0, L] \times [0, D]$, with $\Delta x = L/(N_x - 1), \ \Delta y = D/(N_y - 1)$. We also discretize the time interval $[0, t^f]$ using $N_t$ equally distributed points and $\Delta t = t^f/(N_t - 1)$. Next we define vectors of unknown variables of dimension $n$ containing approximate solutions such as
\[
w(t_N) \approx [w(x_i, y_j, t_N)]_{i=1,2,...N_x, j=1,2,...N_y} \in \mathbb{R}^n, \ N = 1, 2, ...N_t.
\]
The semi-discrete equations of SWE \cite{3} are:
\[
\begin{align*}
    u' &= -F_{11}(u, \phi) - F_{12}(u, v) + F \odot v, \\
    v' &= -F_{21}(u, v) - F_{22}(v, \phi) - F \odot u, \\
    \phi' &= -F_{31}(u, \phi) - F_{32}(v, \phi),
\end{align*}
\]
where $\odot$ is the Matlab componentwise multiplication operator, $u', v', \phi'$ denote semi-discrete time derivatives, $F = [f, f, ..., f]$ stores Coriolis components $f = [f(y_j)]_{j=1,2,...N_y}$ while the nonlinear terms $F_{11}$ and $F_{12}, i = 1, 2, 3,$ involving derivatives in $x$ and $y$ directions, respectively, are defined as follows:
\[
\begin{align*}
    F_{11}(u, \phi) &= u \odot A_x u + \frac{1}{2} \phi \odot A_x \phi, \\
    F_{12}(u, v) &= v \odot A_y u, \\
    F_{21}(u, v) &= u \odot A_x v, \\
    F_{22}(v, \phi) &= v \odot A_y v + \frac{1}{2} \phi \odot A_y \phi, \\
    F_{31}(u, \phi) &= \frac{1}{2} \phi \odot A_x u + u \odot A_y \phi, \\
    F_{32}(v, \phi) &= \frac{1}{2} \phi \odot A_y v + v \odot A_y \phi.
\end{align*}
\]

Here $A_x, A_y \in \mathbb{R}^{n \times n}$ are constant coefficient matrices for discrete first-order and second-order differential operators which take into account the boundary conditions.

The numerical scheme was implemented in Fortran and uses a sparse matrix environment. For operations with sparse matrices we employed SPARSEKIT library Saad \cite{71} and the sparse linear systems obtained during the quasi-Newton iterations were solved using MGMRES library Barrett et al. \cite{8}, Kelley \cite{47}, Saad \cite{72}. Here we didn’t decouple the model equations like in Stefanescu and Navon \cite{79} where the Jacobian is either block cyclic tridiagonal or block tridiagonal. We followed this approach since we plan to implement a 4D-Var data assimilation system based on ADI SWE and the adjoints of the decoupled systems can’t be solved with the same implicit scheme applied for solving the forward model.

## 5 Reduced Order Shallow Water Equation Models

Here we will not describe the entire standard POD SWE, tensorial POD SWE and POD/DEIM SWE discrete models but only we introduce the projected nonlinear term $F_{11}$ for all three ROMs. ADI discrete equations were projected onto reduced POD subspaces and a detailed description of the reduced equations for standard POD and POD/DEIM is available in Stefanescu and Navon \cite{79}.

Depending on the type of reduced approaches, the Petrov-Galerkin projected nonlinear term $\tilde{F}_{11}$ has the following form

### Standard POD.

\[
\tilde{F}_{11} = W^T F_{11} = \sum_{k=1}^{n} \left( W \odot (U_k u) \right) + \frac{1}{2} \sum_{k=1}^{n} \left( \Phi \odot \Phi_{x} \phi \right), \quad (9)
\]
where $U$ and $\Phi$ contains the POD bases corresponding to state variables $u$ and $\phi$ while the POD basis derivatives are included in $U_x = A_x U \in \mathbb{R}^{n \times k}$ and $\Phi_x = A_x \Phi \in \mathbb{R}^{n \times k}$.

**Tensorial POD.**

$$\tilde{F}_{11} = W^T F_{11} \in \mathbb{R}^k; \quad \tilde{F}_{11}^i = (M_1^i, \tilde{U})^{\text{Frobenius}} + (M_2^i, \tilde{\Phi})^{\text{Frobenius}}, \quad i = 1, 2, \ldots, k;$$

$$\tilde{U} = [\tilde{U}_{ij}]_{i,j=1,\ldots,k} \in \mathbb{R}^{k \times k}, \quad \tilde{U}_{ij} = \tilde{u}_i \tilde{u}_j \in \mathbb{R}, \quad \tilde{\Phi} = [\tilde{\Phi}_{ij}]_{i,j=1,\ldots,k} \in \mathbb{R}^{k \times k}, \quad \tilde{\Phi}_{ij} = \tilde{\phi}_i \tilde{\phi}_j \in \mathbb{R},$$

where $\tilde{u} \in \mathbb{R}^k$ and $\tilde{\phi} \in \mathbb{R}^k$ are reduced state variables.

$$M_1^i = [M_{1i_1i_2}]_{i_1,i_2=1,\ldots,k} \in \mathbb{R}^{k \times k}; \quad M_1^i = \sum_{l=1}^n W_{il} U_{il} U_{xli_2} \in \mathbb{R}$$

$$M_2^i = [M_{2i_1i_2}]_{i_1,i_2=1,\ldots,k} \in \mathbb{R}^{k \times k}; \quad M_2^i = \sum_{l=1}^n W_{il} \Phi_{il} \Phi_{xli_2} \in \mathbb{R},$$

and $U_x$ and $\Phi_x$ were defined above.

**POD/DEIM.**

$$\tilde{F}_{11} \approx W^T V_{F1} (P_{F1}^T V_{F1})^{-1} \left( (P_{F1}^T \tilde{U} \tilde{u}) \odot (P_{F1}^T U_x) \tilde{u} + (P_{F1}^T \tilde{\Phi} \tilde{\phi}) \odot (P_{F1}^T \Phi_x \tilde{\phi}) \right),$$

where $V_{F1} \in \mathbb{R}^{n \times m}$ collects the first $m$ POD basis modes of nonlinear function $F_{11}$ while $P_{F1} \in \mathbb{R}^{n \times m}$ is the DEIM interpolation selection matrix. Let us denote the precomputed term by $E_{11} = W^T V_{F1} (P_{F1}^T V_{F1})^{-1}$.

Tensors like $M_1^i$ and $M_2^i$ [11] must also be computed in the case of standard POD and POD/DEIM since the analytic form of reduce Jacobian was employed. This approach reduces the CPU time of standard POD since usually the reduced Jacobians are obtained by projecting the full Jacobian at every time step. A generalization of DEIM to approximate operators is not been yet developed but has the ability to decrease more the computational complexity of POD/DEIM approach. Some related work includes Tomm [33] who developed Multi-Component EIM for deriving affine approximations for continuous vector valued functions. Wirtz et al. [38] introduced the matrix-DEIM approach to approximate the Jacobian of a nonlinear function. Chaturantabut [20] proposed a sampling strategy centered on the trajectory of the nonlinear functions in order to approximate the reduced Jacobian. An extension for nonlinear problems that do not have component-wise dependence on the state has been introduced in Zhou [85].

Table 2 contains the procedure list required by all three algorithms in the off-line stage. $M_1^i$, $M_2^i$ and $E_{11}$ are POD and POD/DEIM coefficients related to nonlinear term $F_{11}$, similar coefficients being required for computation of other reduced nonlinear terms.

#### 6 Numerical Results

For all tests we derived the initial conditions from the initial height condition No. 1 of Grammeltvedt 1969 [32] i.e.

$$h(x, y, 0) = H_0 + H_1 + \tanh\left(\frac{g D/2 - y}{2D}\right) + H_2 \text{sech}^2\left(\frac{g D/2 - y}{2D}\right) \sin\left(\frac{2\pi x}{L}\right).$$

The initial velocity fields are derived from the initial height field using the geostrophic relationship

$$u = \left(\frac{-g}{f}\right) \frac{\partial h}{\partial y} \quad v = \left(\frac{g}{f}\right) \frac{\partial h}{\partial x}.$$
Table 2: ROMs off-line stage procedures - POD coefficients $M_1^i$, $M_2^i$ are required for reduced Jacobian calculation. Only tensorial POD uses them also for right-hand side terms computations during the quasi-Newton iterations required by Gustafsson’s nonlinear ADI finite difference scheme.

Figure 1: Initial condition: Geopotential height field for the Grammeltvedt initial condition and windfield (the velocity unit is 1km/s) calculated from the geopotential field using the geostrophic approximation.

We use the following constants $L = 6000km$, $D = 4400km$, $\hat{f} = 10^{-4}s^{-1}$, $\beta = 1.5\cdot10^{-11}s^{-1}m^{-1}$, $g = 10ms^{-2}$, $H_0 = 2000m$, $H_1 = 220m$, $H_2 = 133m$. Figure 1 depicts the initial geopotential isolines and the geostrophic wind field.

Most of the depicted results are obtained in the case when the domain is discretized using a mesh of $376 \times 276 = 103,776$ points, with $\Delta x = \Delta y = 16km$. We select two integration time windows of 24h and 3h and we use 91 time steps ($NT = 91$) with $\Delta t = 960s$ and $\Delta t = 120s$.

ADI FD SWE scheme proposed by Gustafsson 1971 in [37] is first employed in order to obtain the numerical solution of the SWE model. The implicit scheme allows us to integrate in time at a Courant-Friedrichs-Levy (CFL) condition of $\sqrt{gh}(\Delta t/\Delta x) < 8.9301$.

The nonlinear algebraic systems of ADI FD SWE scheme is solved using quasi - Newton method, and the LU decomposition is performed every 6 time steps.

We derive the reduced order models by employing a Galerkin projection. The POD basis functions are constructed using 91 snapshots (number of snapshots equal with the number of time steps $N_t$) obtained from the numerical solution of the full - order ADI FD SWE model at equally spaced time steps for each time interval $[0, 24h]$ and $[0, 3h]$. Figures 2 show the decay around the eigenvalues of the snapshot solutions for $u$, $v$, $\phi$ and the nonlinear snapshots $F_{11}$, $F_{12}$, $F_{21}$, $F_{22}$, $F_{31}$, $F_{32}$. We notice that the singular values decay much faster when the model is integrated for 3h. Consequently this translates in a more
accurate solution representations for all three ROM methods using the same number of POD modes. For both time configurations and all tests in this study, the dimensions of the POD bases for each variable is taken to be 50, capturing more than 99% of the system energy. The largest neglected eigenvalues corresponding to state variables $u$, $v$, $\phi$ are 2.23, 1.16 and 2.39 for $t_f = 24h$ and 0.0016, 0.0063 and 0.0178 for $t_f = 3h$, respectively.

![Figure 2](image1.png)

(a) State variables $u$, $v$, $\phi$

(b) Nonlinear terms

Figure 2: The decay around the singular values of the snapshots solutions for $u$, $v$, $\phi$ and nonlinear terms for $\Delta t = 960s$ and integration time window of $24h$.

![Figure 3](image2.png)

(a) State variables $u$, $v$, $\phi$

(b) Nonlinear terms

Figure 3: The decay around the singular values of the snapshots solutions for $u$, $v$, $\phi$ and nonlinear terms for $\Delta t = 120s$ and a time integration window of $3h$.

Next we apply DEIM algorithm and calculate the interpolation points to improve the efficiency of the standard POD approximation and to achieve a complexity reduction of the nonlinear terms with a complexity proportional to the number of reduced variables, as in the case of tensorial POD. Figures illustrate the distribution of the first 100 spatial points selected by the DEIM algorithm together with the isolines of the nonlinear terms statistics. Each of these statistics contain in every space location the maximum values of the corresponding nonlinear term over time. Maximum is preferred instead of time averaging since a better correlation between location of DEIM points and physical structures was observed in the former case.
However, in most cases, the spatial positions of the interpolation points don’t follow the nonlinear statistics structures. This is more visible in Figure 5 for $t_f = 3h$. The exceptions are $F_{12}$ and $F_{21}$ (Figures 5b,c), nonlinear terms depending only on velocity components, where DEIM interpolation points target better the underlying physical structures. This proves that DEIM algorithm doesn’t particularly take into account the physical structures of the nonlinear terms but search (in a greedy manner) to minimize the error (residual) between each column of the input basis (POD basis of the nonlinear term snapshots) and its proposed low-rank approximations Stefanescu and Navon [79, p.16].

Figures 6,7 depict the grid point absolute error of the standard POD, tensorial POD and POD/DEIM solutions with respect to the full solutions. For POD/DEIM reduced order model we use 180 interpolation points. The magnitude of the errors are similar for each of the method proposed in this study. Moreover, we observed that error isolines distribution in Figure 7 is well correlated with the location of interpolation points illustrated in Figure 5 underlying the empirical characteristics of DEIM.

In addition, we propose two metrics to quantify the accuracy level of standard POD, tensorial POD and standard POD/DEIM approaches. First, we use the following norm

$$\frac{1}{N_t} \sum_{i=1}^{N_t} \frac{||w^{\text{FULL}}(:,t_i) - w^{\text{ROM}}(:,t_i)||_2}{||w^{\text{FULL}}(:,t_i)||_2}$$

$$\frac{1}{N_t} \sum_{i=1}^{N_t} \frac{||w^{\text{FULL}}(:,t_i) - w^{\text{ROM}}(:,t_i)||_2}{||w^{\text{FULL}}(:,t_i)||_2}$$

$i = 1, 2, \ldots, t_f$ and calculate the relative errors for all three variables of SWE model $w = (u, v, \phi)$. The results are presented in Table 3. We perform numerical experiments using two choices for number of DEIM points 70 and 180. For 24h tests we notice that more than 70 number of DEIM points are needed for convergence of quasi-Newton method for POD/DEIM SWE scheme explaining the absence of numerical results in Table 3 (left part).
on quadratic nonlinearities, the tensorial POD poses the appropriate characteristics of a reduced order approach. Given that the implementation effort is much reduced, in the cases of models depending only on a few nonlinear terms in comparison with the POD ADI SWE model and overall decreases the computational time with a factor of 77× for 24h time integration and 76× for a 3h time window integration. POD/DEIM outperforms standard POD being 450× and 250× time faster for 70 and 180 DEIM interpolation points and a time integration window of 3h. For \(t_f = 24h\) and \(m = 180\), POD/DEIM SWE model is 183× time faster than standard POD SWE model. In terms of CPU time, tensor POD SWE model is only 2.38× slower than POD/DEIM SWE model for \(m = 180\) and \(t_f = 24h\) while for \(t_f = 3h\) the new tensorial POD scheme is 5.9× and 3.3× less efficient than POD/DEIM SWE model for \(m = 70\) and \(m = 180\). This suggests that operations like Jacobian computations and its LU decomposition required by both reduced order approaches weight more in the overall CPU time cost since the quadratic nonlinear complexity of the tensorial POD requires 374,950 floating-point operations and POD/DEIM only 27,180 (\(m = 180\), see Section 3). Given that the implementation effort is much reduced, in the cases of models depending only on quadratic nonlinearities, the tensorial POD poses the appropriate characteristics of a reduced order approach.
method and represent a solid alternative to the POD/DEIM approach.

For cubical nonlinearities and larger, tensorial POD loses its ability to deliver fast calculations (see Table 1), thus the POD/DEIM should be employed.

In our case the Jacobians are calculated analytically and its computations depend only on the reduced space dimension $k$. However, more gain can be obtain if DEIM would be applied to approximate the reduced Jacobians but this is subject of future research.

The computational savings and accuracy levels obtained by the ROMs studied in this paper depend on the number of POD modes and number of DEIM points. These numbers may be large in practice in order to capture well the full model dynamics. For example, in the case of a time window integration of 24h, if someone would ask to increase the ROMs solutions accuracy with only one order of magnitude, the POD basis dimension must be at least larger than 100 which will drastically compromise the time performances of ROMs methods. Elegant solutions to this problem were proposed by Peherstorfer et al. [64], Rapún and Vega [65] where local POD and local DEIM versions were proposed. Machine learning techniques such as $K$-means Lloyd [62], MacQueen [55], Steinhaus [80] can be used for both time and space partitioning. A recent study investigating cluster-based reduced order modeling was proposed by Kaiser et al. [74].

Figure 8 depicts the efficiency of tensorial POD and POD/DEIM SWE schemes as a function of...
Baumann [9] to solve a reduced 4D-Var data assimilation problem and good results were obtained for a different than the ones used to generate the DEIM interpolation points. DEIM was first employed by Pod/DEIM SWE scheme to have the fastest off-line stage (Figure 9b). This gives a good advantage of POD scheme is 10\times faster than the standard POD scheme. The performances of POD/DEIM depends on the number of DEIM points and the numerical results displays a 10\times time reduction of the CPU costs in comparison with the standard POD outcome when \( n \geq 61 \times 45 \) and \( n \geq 101 \times 71 \) for \( m = 70 \) and \( m = 180 \) respectively.

The new algorithm introduced in Section 3 relying on DEIM interpolation points delivers fast tensorial calculations required for computing the reduced Jacobian in the on-line stage and thus allowing POD/DEIM SWE scheme to have the fastest off-line stage (Figure 9b). This gives a good advantage of ROM optimization based on Discrete Empirical Interpolation Method supposing that quality approximations of nonlinear terms and reduced Jacobians are delivered since during optimization input data are different than the ones used to generate the DEIM interpolation points. DEIM was first employed by Baumann [9] to solve a reduced 4D-Var data assimilation problem and good results were obtained for a 1D Burgers model. Extensions to 2D models are still not available in the literature.
Table 4: CPU time gains and the root mean square errors for each of the model variables at \( t_f = 24h \). Number of POD modes was \( k = 50 \) and we choose 180 number of DEIM points.

<table>
<thead>
<tr>
<th></th>
<th>Full ADI SWE</th>
<th>Standard POD</th>
<th>Tensorial POD</th>
<th>POD/DEIM m=180</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU time</td>
<td>1813.992s</td>
<td>191.785</td>
<td>2.491</td>
<td>1.046</td>
</tr>
<tr>
<td>( u )</td>
<td>-</td>
<td>9.095e-3</td>
<td>9.095e-3</td>
<td>1.555e-2</td>
</tr>
<tr>
<td>( v )</td>
<td>-</td>
<td>8.812e-3</td>
<td>8.812e-3</td>
<td>1.348e-2</td>
</tr>
<tr>
<td>( \phi )</td>
<td>-</td>
<td>6.987e-3</td>
<td>6.987e-3</td>
<td>1.13e-2</td>
</tr>
</tbody>
</table>

Table 5: CPU time gains and the root mean square errors for each of the model variables at \( t_f = 3h \) for a 3h time integration window. Number of POD modes was \( k = 50 \) and two tests with different number of DEIM points \( m = 180, 70 \) were simulated.

<table>
<thead>
<tr>
<th></th>
<th>Full ADI SWE</th>
<th>Standard POD</th>
<th>Tensorial POD</th>
<th>POD/DEIM m=180</th>
<th>POD/DEIM m=70</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU time</td>
<td>950.0314s</td>
<td>161.907</td>
<td>2.125</td>
<td>0.642</td>
<td>0.359</td>
</tr>
<tr>
<td>( u )</td>
<td>-</td>
<td>5.358e-5</td>
<td>5.358e-5</td>
<td>5.646e-5</td>
<td>7.453e-5</td>
</tr>
<tr>
<td>( v )</td>
<td>-</td>
<td>2.728e-5</td>
<td>2.728e-5</td>
<td>3.418e-5</td>
<td>4.233e-5</td>
</tr>
<tr>
<td>( \phi )</td>
<td>-</td>
<td>8.505e-5</td>
<td>8.505e-5</td>
<td>8.762e-5</td>
<td>9.212e-5</td>
</tr>
</tbody>
</table>

7 Conclusions

It is well known that in standard POD the cost of evaluating nonlinear terms during the on-line stage depends on the full space dimension, and this constitutes a major efficiency bottleneck. The present manuscript applies tensorial calculus techniques which allows fast computations of standard POD reduced quadratic nonlinearities. We show that tensorial POD can be applied to all type of polynomial nonlinearities and the resulting nonlinear terms have a complexity of \( O(kp+1) \) operations, where \( k \) is the dimension of POD subspace and \( p \) is the polynomial degree. Consequently, this approach eliminates the dependency on the full space dimension, while yielding the same reduced solution accuracy as standard POD. Despite being independent of number of mesh points, tensorial POD is efficient only for quadratic nonlinear terms since for higher nonlinearities standard POD proves to be less time consuming once the POD basis dimension \( k \) is increased.

The efficiency of tensorial POD is compared against that of standard POD and of POD/DEIM. We theoretically analyze the number of floating-point operations required as a function of polynomial degree \( p \), the number of degrees of freedom of the high-fidelity model \( n \), of POD modes \( k \), and of DEIM interpolation points \( m \). For quadratic nonlinearities and \( k \) between 10–50 modes, the tensorial POD needs 10–40 times fewer operations than the standard POD approach and 10–20 times more operations than the POD/DEIM with \( m = 100 \). But these performances are not translated into the same CPU time rates for solving the ROMs solutions since other more time consuming calculations are needed.

Numerical experiments are carried out using a two dimensional ADI SWE finite difference model. Reduced order models were developed using each of the three ROM methods and Galerkin projection. The spectral analysis of snapshots matrices reveals that local versions of ROMs lead to more accurate results. Consequently, we focus on three hours time integration windows. The tensorial POD SWE model becomes considerably faster than standard POD when the dimension of the full model increases. For example, for 100,000 spatial points the tensorial POD SWE model yields the same solutions accuracy as standard POD but is 76 times faster. Numerical experiments of POD/DEIM SWE scheme revealed a considerable reduction of the computational complexity. For a number of 70 DEIM points, POD/DEIM SWE model is 450 times faster than standard POD, but only 6 times faster than tensorial POD.

For models depending only on quadratic nonlinearities, the tensorial POD represents a solid alternative to POD/DEIM where the implementation effort is considerably larger. However, for cubic and higher order nonlinearities, tensorial POD loses its ability to deliver fast calculations and the POD/DEIM approach should be employed.

We also propose a new DEIM-based algorithm that allows fast computations of the tensors needed by reduced Jacobians calculations in the on-line stage. The resulting off-line POD/DEIM stage is the fastest among the ones considered here, even if additional SVD decompositions and low-rank terms are
Figure 8: CPU time vs. the number of spatial discretization points for $t_f = 3h$; number of POD modes $= 50$; two different numbers of DEIM points 70 and 180 have been employed.

computed. This is an important advantage in optimization problems based on POD/DEIM surrogates where the reduced order bases need to be updated multiple times.

On-going work by the authors focuses on reduced order constrained optimization. The current research represents an important step toward developing tensorial POD and POD/DEIM four dimensional variational data assimilation systems, which are not available in the literature for complex models.

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