Skipping The Jacobian

(Hyper) Reduced Order Models For Moving Meshes

E. Millán Valbuena







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by



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Enrique Millán Valbuena 463 426 8

Dr. N. Dal Santo,École Polytechnique Fédérale de LausanneDr. S. J. Hulshoff,TU DelftProf. Dr. A. Manzoni,Politecnico di MilanoProf. Dr. A. Quarteroni,Politecnico di Milano

Abstract

We present a reduced order model (ROM) for a one-dimensional nonlinear gas dynamics problem: the isentropic piston. The main body of the PDE, the geometrical definition of the mesh nodes, and the boundary conditions are parametrized. The full order model is obtained with a Galerkin finite element discretization, under the Arbitrary Lagrangian Eulerian formulation (ALE). To stabilize the system, an artificial viscosity term is included. The nonlinear convective term is linearized with a second-order extrapolation. The reduced basis to express the solution is obtained with the POD technique. To overcome the explicit use of the Jacobian transformation, typical in the context of moving meshes, a system approximation technique is introduced. The (Matrix) Discrete Empirical Interpolation Method, (M)DEIM, allows us to work with a weak form defined in the physical domain (and hence the physical weak formulation) whilst maintaining an efficient assembly of the algebraic operators, despite their change with every time step. Two alternative methods are presented to collect and compress the snapshots for the discretized solution-dependent convective term. Each method leads to a different offline stage. All in all, our approach is purely algebraic and the reduced model makes no use of full order structures, thus achieving a perfect *offline-online* split. A concise description of the reduction procedure is provided. The reduced model is certified with a posteriori error estimations obtained via mode truncation.

Index Terms

finite elements, Galerkin, reduced order models, moving mesh, ALE, (M)DEIM, POD, SVD, mode truncation

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FOREWORD

These works typically set the ending of a cycle in life. In my case, it marks a restart.

I was very close to dropping out from my masters, contempt with having my bachelor and partially frustrated for multiple reasons not worth developing here. Taking into account the fact that my professional career seemed to have drifted away from Aerospace Engineering, I struggled to see the point at completing it, let alone finding the time such task required.

However, two ideas made me get back to work. First, it is easier to explain an elongated but completed academic course, rather than an uncompleted one. Second, and most important, *I do actually like the subject*. In fact, during the last year I have realized how much I enjoy *Applied Mathematics*, a field by which Aerospace Engineering is widely nurtered.

In this regard I would like to thank professors A. Quarteroni and A. Manzoni for giving me the opportunity to work with them in their research group in Milan. Apart from learning mathematics, I met great colleagues and picked up one of Europe's most beautiful languages. However, working with the FEM code available there, LifeV, was tough, and probably its complexity had to do with the growing frustration I was already experiencing in my academic life: too much time spent debugging, rather than understanding the problem with pen and paper. Nevertheless, with them I discovered a way of using mathematics that I had not learnt before, one that suits my mind and approach to scientific modelling.

I am grateful for the warmth I received from the PhD students and postdocs at the office where we worked together everyday: Federica (are you ever going to quite smoking?), Abele (namaste), Dani (south), Dani (north), Ludovica, Stefano, and a countable infinite more. I keep good memories playing volley and climbing with you all. The same goes for the staff from the Politecnico: Paola, Lucas, Susanna. Last, but not least, I would like to thank Niccolò Dal Santo for his patience, time and knowledge; which he generously dedicated to me despite being based in Lausanne. You are definitely among the most clever people I have met.

At TU Delft, I would like to thank professor Steven for his joyful encouragement during round two of this thesis. When I first wrote him back after a year and a half since he last heard from me, I thought he would (righteously) no longer want to have anything to do with this work. I was gladly surprised to receive all of the contrary, a warm welcome back and a pragmatic view and feedback to make me reach the end at the fastest pace, whilst doing a good job. I would like to ackwnoledge Simone Floreani too, my fellow student colleage and great friend at TU, who, based in his previous experience, advised me to go work in Milan and learn these mathematics. Without your words, none of this would have started. I hope we get to collaborate together in a project involving mathematics in the near future.

At last, although they are completely outside of the academic scope, I would like to thank my colleagues at work and my close friends. My two supervisors, Ana and Sarah, from whom I have adquired the pragmatism industrial problems require to be completed in time and form. To Maximiliano, again a smart and kind person across my path in life, eager to teach me professional coding skills and how to structure creativity. Juan and Javier, with whom I share great adventures and conversations. To Emmanuel, my housemate, who despite being a lawyer, would kindly ask me every now and then about the ROM convergence rates. Miquel (both of them), a friend turned brother, for your unwearing support and advice. And to all of the remaining, with whom I spend great quality time. You influence my life more than you are probably aware.

As the reader will see, this is quite an elementary work. Formulating and coding it has been laborious, but the content remains simple. Yet, its simplicity has allowed me to understand the fundamentals of two versatile and powerful mathematical tools: Finite Elements and Reduced Order Models. This has motivated me to keep on working at it once this is over, so that hopefully one day I get to tackle the real-life problem I set to solve in the first place: the fluid mechanics problem of the human heart.

Madrid (España), 2021.

| t | time coordinate |
|---------------------|---|
| x | space coordinate in the physical domain |
| X | space coordinate in a fixed reference domain |
| A | Arbitrary Lagrangian Eulerian map |
| d | mesh displacement |
| Wm | mesh velocity |
| .J | Jacobian matrix |
| Ĵ, | Jacobian transformation (matrix determinant) |
| L_0 | initial piston length |
| L^{-0} | piston movement in time |
| 11 | flow velocity |
| $\hat{\hat{u}}$ | flow velocity for the homogeneous problem |
| u _b | flow velocity finite element representation |
| a | Dirichlet boundary lifting |
| p | pressure |
| ρ | density |
| γ | specific heat ratio |
| \dot{MD} | mass defect in time (numerical integration) |
| ε | artificial viscosity value |
| ε_{POD} | POD approximation error |
| u_p | piston mach number |
| a_0 | reference speed of sound |
| p_0 | reference pressure |
| ρ_0 | reference density |
| δ | piston displacement from rest (as a fraction of |
| ω | piston oscillating frequency |
| c | dimensionless convection velocity |
| b_0 | dimensionless constant convection coefficient |
| b_L | dimensionless piston boundary condition |
| ${\mathcal F}$ | Gaussian bell |
| x_c | Gaussian bell center |
| σ_c | Gaussian bell dispersion factor |
| y_c | Gaussian bell scaling factor |
| N_h | number of finite element degrees of freedom |
| N | number of reduced basis modes |

 $L_0)$

- V_h space of Lagrangian functions
- V_N space of RB solution modes
- φ_i Lagrangian finite element basis function
- φ_i Lagrangian finite elem ψ_i RB solution modes
- $A_{h,q}$ algebraic operator modes (collateral basis)

When a painter sets out to paint, she will probably use most of the available basic colours. However, if she knew beforehand that she is going to paint landscapes, she would fare well with a farsighted palette: greens, browns, blues, whites, etc. Such is the nature of reduced order models, to find a subset among the combinations of basic colours to represent the solution to the problem of interest.

In the context of this work, the basic colours are the classical mathematical Lagrangian finite element basis functions: generic, piecewise, with local support, able to represent most functions of interest. Instead, the landscape palette will be ad-hoc: problem-dependent functions with global support, good at capturing details only specific to landscapes.

Additionally, she will not need all sorts of brushes, simply the ones with the right thickness and width for mountains, trees and hills. The brushes represent the algebraic operators that arise from the finite element discretization. As with the colours, we can find a subset of combinations of brushes that suit our problem. That is, we can find a basis for each algebraic operator to build them efficiently.

Finally, since she is a vanguard painter, the domain of our problem, her canvas, will be allowed to change in time as she paints. The landscape colours and brushes we select need to take this into account.

So far with metaphors.

1.1 The Need for Reduction

Conventional PDE discretization techniques (finite elements, finite volumes, finite differences, etc.) rely on a generic representation of the PDE solution to form an algebraic system; whose unknowns allow the determination of the PDE solution at each mesh node for every time step. The discretized operators which conform these algebraic systems are often sparse¹ and high-dimensional². For reallife scenarios, the computational time required to solve them can become prohibitive for parameter sweeps or optimization problems (two tasks which are often an important step of scientific or engineering works).

This leads to the idea of building and using ad-hoc reduced basis (RB) modes to represent the solution, in substitution of the generic functions used by conventional methods. The RB solution modes encode the main dynamics of the PDE, and thus with a few number of them the solution can be accurately represented. The reduced order model (ROM) associated to the RB solution modes is obtained by projection of the discretized operators associated with the original conventional method. The resulting linear system is dense, and several orders of magnitude smaller; which makes it suitable for many-query tasks. We refer to the RB mode identification step as the *offline stage*; and to its use to solve new problems as the *online stage*.

1. Conventional discretization techniques typically use functions with local support to represent the solution.

2. The more mesh nodes are added to the mesh (and thus the more unknowns the discretized algebraic system has), the closer the discrete PDE solution will get to the continuous PDE solution.

1.1.1 Reduction in a Moving Mesh Context

In the context of a fixed mesh, the assembly and projection steps take place once and for all. On the contrary, if the mesh moves in time, these steps need to take place for every time step³. Hence, despite the reduction in size of the linear system, there is still a considerable overhead during its assembly. To overcome it, a system approximation technique is introduced, following the same idea behind RB modes: find and use a suitable ad-hoc representation basis; thus the discrete operator is expressed as a linear combination of operator modes (collateral basis). The system approximation technique must require that an efficient assembly of the projected operator can be obtained at runtime, for each time step.

Since a basis is identified for the solution and each of the algebraic operators, the combination of these two techniques defines an hyper reduced order model (HROM). The adjective *hyper*⁴ is present because a collateral basis will be created for each algebraic operator, on top of the one created for the solution space.

To put in practice the ideas previously stated, we build and certify an HROM for a one-dimensional parametrized piston problem, whose movement is prescribed. The model PDE, the boundary conditions and the mesh geometry are parametrized. The finite element method is used as the conventional discretization technique. This work extends [1], [2] (albeit for a simpler PDE and domain), which did introduce the MDEIM system approximation technique for unsteady problems with geometrically parametrized domains, although the mesh (and the domain boundary) remained fixed in time.

1.2 The Modeling Impact of a Moving Mesh

Physical problems with moving meshes (related or not to fluid dynamics), require the introduction of a convective term to account for the movement of the mesh, and potentially a transformation in the finite element weak form. This transformation translates the integrals defined in a deforming space into a fixed, numerical grid, as shown in Figure 1.



Fig. 1. Numerical fixed grid to the left, physical moving mesh to the right. The Jacobian determinant establishes the connection between the evaluation of an integral over the square (red) in each domain (blue). Figure from [3].

3. Discrete integrals, fluxes, or interpolations associated to the conventional discretization method need to be recomputed for the new mesh.

4. From Greek *hyper* (prep. and adv.) "over, beyond, overmuch, above measure".

The entries of the Jacobian matrix might be known explicitly, if the deformation is known analytically and the domain is sufficiently simple. However, if the domain takes arbitrary shapes (likely for real-life problems in higher dimensions than one), or we are dealing with an FSI problem (where the deformation is part of the solution), it is quite possible that we have to compute the Jacobian transformation numerically. This is likely to be an undesirable situation, for the physical weak form will become contaminated with additional terms, making it more cumbersome to implement and deal with⁵, as well as the inevitable overhead in computational costs.

This overhead created by the Jacobian is likely to permeate to the reduced order model (ROM), for in the context of finite elements, ROMs are often built as a system with the same algebraic structure as the departing full order model (FOM), albeit with smaller matrices and vectors. It might even be the case that we cannot completely uncouple the ROM from the FOM, if the problem is complex enough, or that we need satellite⁶ ROMs to compute the Jacobian matrix efficiently.

Hence, to avoid all of the previous, we develop a ROM formulation which allows us to remain in the physical domain, whilst maintaining a perfect *offline-online* decomposition, allowing our reduction scheme to reach its maximum efficiency.

1.3 Literature Review

We now frame our work within the current body of knowledge with a short literature review. Additional references will be cited within the body of the document, where their appearence is more accurate and helpful.

We have used mainly two types of papers: methodology and applications. The former present a numerical method or formulation which we use, the latter make use of it for specific applications. We were especially interested in the applications to see how previous works dealt with inhomogeneous boundary conditions, on which will be discussed later on in the work and the review.

1.3.1 Burgers Terms and Piston Models

Regarding the target PDE to work with, we decided upon several constraints: it had to be one-dimensional (to ease implementation), contain non-trivial terms (to make the problem interesting), and be physically sound (to validate the outcome).

Luckily, removing viscosity and body forces (which are strong assumptions), and making use of the isentropic condition, we can transform the Navier-Stokes equations into a one-dimensional equation [4], [5] in terms of velocity. At the

^{5.} The discretization will not be formulated for the original variable ϕ , but rather for its distorted counterpart $J\phi$.

^{6.} If the Jacobian is derived from a PDE, this would require an additional ROM with its corresponding reduced basis to enhance computation times.

discretization level, we can add an artificial viscosity term, to make sure the solution remains stable [6].

This result is a simple yet complete PDE to model the movement of a piston, which can be validated with derived computations such as mass conservation, and whose solution makes intuitively sense when plotted.

The equation resembles an advection equation with a nonlinear Burgers-like convective term. The Burgers equation showed up in the gas dynamics literature several decades ago [7]–[10], for which under controlled conditions a range of implicit analytical solutions exist [11], including for moving domains [12].

Compared to modern problems with moving domains, the piston is quite simple in nature; and yet it allowed many aerodynamicists to push forward the barrier of knowledge back in the days, when computational power was not so easy to access [13].

1.3.2 Deforming Mesh (ALE)

Because the piston problem is defined in a moving domain, one needs to modify the formulation to account for the movement of the mesh nodes, with the introduction of a convective term governed by the mesh velocity vector. In fact, this needs to be done even for domains with fixed boundaries, where the interior mesh nodes move in time (front-tracking or shock-capturing schemes could be an example of such situations). This leads to the Arbitrary Lagrangian Eulerian (ALE) formulation.

An introduction to the details of the ALE formulation in a simple setting can be found in [14]–[16]. In these works, stability arguments and implementation details for finite elements and simple PDE models are provided. Work [14] contains lengthy and easy-to-read derivations which explain neatly the differences between conservative and non-conservative weak forms. For reasons that will become aparent later on, in this thesis we need to solve the non-conservative weak form, at least to use the current formulation of the system aproximation technique which we intend to use. For a complete and generic development of the subject, in higher dimensions and for complex problems, we refer the reader to [17], [18].

Regarding the stability of the integration scheme, the concept of a (*Discrete*) Geometric Conservation Law (D-GCL) is often mentioned [19]–[22]. Briefly, how the domain deforms and how this deformation is accounted for *in the discretiza-tion* of the continuous problem, could lead or not to instabilities in the solution; for the movement of the mesh could introduce artificial fluxes in the discretization. As a general rule of thumb, to guarantee some notion of stability, the scheme should be able to reproduce the constant solution (under the appropriate boundary conditions). In [14] they prove how the Implicit Euler integration scheme becomes conditionally stable for a linear advection-diffusion problem if the non-conservative weak formulation is solved.

As a final note, we would like to point out that a problem with a deforming domain could also be tackled with space-time finite elements [23]. In fact, as it is the case for us, if the boundary movement is prescribed, the domain in a space-time context will be a fixed one. However, we disregarded this line of work because it could make the implementation much more complicated. This ends the literature review regarding the FOM model. We now present the literature oriented towards the construction of the ROM.

1.3.3 Reduced Basis

We do not aim here at providing a comprehensive review of the whole field (for that could be a complete work by itself), but rather to present a good starting point from which the interested reader could start, and of course, the framing of this thesis.

A problem's complexity and its computational cost are typically something that scale together. Hence, the idea of finding a smaller subspace to represent the solution and reduce calculation times is justified.

This idea of using a problem-dependent basis with global support to solve numerically discretized PDEs is well known. The first references in this line date back to the 80s, with pioneering works in structural analysis [24]. Since then, this idea has become increasingly popular, with many papers and books explaining methods and applications for steady and unsteady problems [25]–[31], including the Navier-Stokes equations [32]. In fact, reduced basis for the Burgers' model have already been examined for a fixed domain [33].

In the following, we present a narrative for Reduced Basis methods in the finite element context, to frame our use of it. We understand and admit that there might be other narratives that suit the field, but the following has proven helpful to understand the ingredients of the ROM.

Our narrative takes the perspective of: where does the basis come from? Or in other words, how many mathematical tools are necessary to obtain it? The construction of the reduced basis needs to take into account the following facts: there must be a sampling strategy in the parameter space, the reduced basis must converge to the span of the solutions, and it must be computationally efficient.

The most plain version of reduced basis is a collection of solutions for several parametrizations. However, the modes of this basis are likely to be almost linearly dependent⁷, and no approximation arguments have been used to obtain it.

A step one can take to improve such basis is to use a greedy procedure [34], [35]. That is, the modes of the basis are still solutions of the PDE, but they are combined iteratively, by choosing the next mode which minizimes the error made by the current basis within a randomly selected parameter space (hence the name greedy, in terms of approximation accuracy). This procedure only requires the finite element discretization, and one can prove it will converge to the whole span. The difficulty in this procedure is the efficient estimation of the error of the basis at each iteration. However, it has become the established method to approach steady models [36].

An alternate approach is to rely on an external methodology to construct the basis from a collection of solution snapshots. An example is the Singular Value Decomposition (SVD) [37], which allows us to compress the span of the solution space efficiently with optimal convergence

^{7.} A strong assumption underlying reduced basis methods in this context is that the solutions of the parametrized PDE change smoothly when the parametrization varies.

properties. This is also known as the Proper⁸ Orthogonal Decomposition (POD) [38]. It has been widely used in many contexts to obtain a basis from a collection of solutions automatically, or to analyze the underlying dynamics of a flow field [39].

The POD has also wider application than the greedy method, since we could use experimental data, to obtain a reduced basis which we then use to solve a numerical model efficiently. Of particular interest is the application made in [40], where they used the POD of analytical solutions with and without a deforming grid to split effects and analyze convergence rates.

Finally, for unsteady problems, one may have the combination of both; the POD-Greedy method [36], [41]. This method uses the automatic compression feature provided by the POD in the time dimension, and the greedy approach to parameter selection in the parameter space. For our work, we will use a physics-driven approach for the sampling strategy in the parameter space, and a nested POD strategy for the time and parameter spaces [1].

Finally, some words need to be said about the handling of the inhomogeneous boundary conditions that we will encounter. According to their functional expression, parametrized essential boundary conditions (Dirichlet) can sometimes be applied directly into the algebraic system without losing reduction capacities, as it is the case with scalar-multiplicative conditions, [42], [43]. However, for arbitrary parametrized essential boundary conditions, setting them directly into the algebraic operators is not a possibility.

Neumann boundary conditions are naturally encoded in the weak form, so they do not pose a problem. Therefore, a suitable approach with essential boundary conditions is to transfer them to the weak form too, with a lifting technique [44]. Hence, the target model problem that we reduce becomes one with homogeneous boundary conditions, for which the results from most references apply.

1.3.4 System Approximation

We reach now the final block of the literature review. In this section we review the methodology used to efficiently approximate the algebraic operators that arise from the discretization. Using an algebraic approach in the reduction scheme is of great advantage, since then most results can generalize to other discretization schemes.

We start by reviewing the approximation methodology for functions and functionals (vectors). The one for matrices is its natural extension.

The seeds of the methodology lie in what is called the Empirical Interpolation Method (EIM) [45]–[47]. It generates an ad-hoc affine decomposition of a parametrized function, by splitting the dependency into some realvalued parameter-dependent functions and a parameterindependent collateral basis. The values of the functions are obtained by enforcing that certain entries of the vector are exactly matched by the ad-hoc decomposition (hence the name interpolation). The entries at which the interpolation should be enforced are computed during the basis creation, and they represent those locations where the approximation behaves worse. The collection of the entries is referred to

8. (adj.) Fitting or right for a particular situation; suitable.

as the *reduced mesh*. The collateral basis is generated with function evaluations following a greedy procedure [48].

As with the RB scenario, the generation of the basis can be delegated to a POD procedure, leading to the Discrete Empirical Interpolation Method (DEIM) [49], [50]. Finally, if the columns of a matrix are stacked vertically to *vectorize* it, a matrix-DEIM method can be used (MDEIM) [2], [51]–[53].

These approximation methods are convenient in the finite element context. The calculation of the reduced mesh entries is the sum of evaluations of the weak form for a restricted subset of mesh elements. This operation can be done efficiently in parallel and is much cheaper than assembling the whole operator [1]. Additionally, the collateral basis can be projected in the reduced space, so that the reduced operator is approximated right away.

When nonlinearities in the solution are present, two options are available to solve the problem. Either the Newton-Rapshon (NR) method is used, and the MDEIM technique is used to approximate the NR Jacobian matrix [53]; or the nonlinear term is discretized in such a way that it is modeled by a trilinear form, whose additional argument is an approximation of the effects that using the actual solution would produce, and the MDEIM is used to approximate the trilinear form [1]. In our work, we follow the second approach.

In all of the above, time can be easily included by treating it as an additional parameter, although the implementation is not so straightforward.

This concludes our literature review.

1.4 Research Questions

In this work we answer the following specific questions to drive our research:

- How can the expense of the online application be minimised? This minimization will go through using as little information from the FOM as possible (including using no information at all).
- How does the type of mesh movement, (e.g. uniform or non-uniform stretching) affect the reducibility of the operator?
- How does the nonlinear term affect the hyperreduction? Is there more than one way to reduce a nonlinear term?
- How do the approximation errors of the solution and operators RB basis interact?
- How should a moving-domain problem be implemented to ensure a general but relatively compact HROM?

1.5 Document Layout

In Section 2 we derive the full order model (FOM), that is, the model PDE and its boundary conditions, mesh deformation and the associated Arbitrary Lagrangian Eulerian (ALE) formulation in the physical domain, both at the continuous and discrete levels. In Section 3 we explain the details of the reduction scheme, along with the system approximation technique (M)DEIM.

In Section 4 we examine the FOM parametrization and response, check the discretization convergence rates, and show GCL results. Finally, in Section 5 we present reduction results and certify the reduction scheme with an efficient a posteriori error estimators obtained via mode truncation.

2 ONE-DIMENSIONAL GAS DYNAMICS

The full order model for the parametrized one-dimensional piston problem is derived here. We depart from the onedimensional, compressible, and isentropic Navier-Stokes equations to end up with a nonlinear Burgers-like equation. Therefore, this problem contains all the necessary ingredients to show how a ROM behaves in the presence of a nonlinear term within a moving domain.

The model will be derived in the continuous, semi-discrete and fully discrete contexts for a generic parametrization and forcing term. We shall use the Galerkin projection principle to find a weak form, which we later discretize using the finite element method (FEM).

We define the vector $\vec{\mu} \in \mathcal{P}$ to collect all the parameters present in the formulation. Parameters will be present in the PDE, in the boundary conditions, or in the geometrical definition of the domain.

We consider a problem with a deforming domain in time, whose movement is known and is thus not part of the solution:

$$\Omega(t, \vec{\mu}) := \{ x \in \mathbb{R} : x \in [0, L(t, \vec{\mu})] \}.$$

From now on, we drop the dependency on time and the parameters unless it is strictly necessary.

2.1 Physical Derivation

The piston movement L(t) is a real-valued smooth sinusoidal function,

$$L(t) = L_0 [1 - \delta (1 - \cos (\omega t))],$$
 (1)

where ω is the frequency at which it oscillates, and $\delta \ll 1$ is a scale variable to adjust how much it is displaced from its original position. A sketch is given in Figure 2. The length L_0 is defined to keep physical dimensions sound, but it will be fixed to $L_0 = 1$ for the remainder of the work. See Appendix A for the derivation of this movement law.



Fig. 2. Piston sketch. The flow departs from rest to the left of the piston. Outflow (inflow) is represented by a negative (positive) velocity value.

We depart from the conservation of mass and momentum, and the isentropic relation between pressure and density,

$$\frac{\partial \rho}{\partial t} + u \frac{\partial \rho}{\partial x} + \rho \frac{\partial u}{\partial x} = 0,$$
(2a)

$$\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} + \frac{1}{\rho} \frac{\partial p}{\partial x} = 0,$$
 (2b)

$$p = k\rho^{\gamma}.$$
 (2c)

Body forces and viscosity have been neglected for the sake of simplicity⁹. A difficulty we find in this system for the

9. Viscosity terms will be introduced later on for numerical stability.

piston application is the determination of the boundary condition for density at the piston location. Ideally, we only want to solve an equation for the velocity, for which boundary conditions are easy to set.

To do so, we follow the steps given in [5]:

- 1) Remove the pressure gradient through the isentropic relation.
- 2) Relate velocity u to density ρ explicitly.
- 3) Collect the results into one equation expressed in terms of the velocity *u*.

Pressure Gradient

To remove the pressure gradient, we start by taking derivatives in the isentropic relation (2c):

$$\frac{\partial p}{\partial x} = k\gamma \rho^{\gamma - 1} \frac{\partial \rho}{\partial x} \tag{3}$$

Then, we recognize that the coefficients $k\gamma\rho^{\gamma-1}$ multiplying the spatial derivative of density are in fact the squared speed of sound. From thermodynamics, the speed of sound *a* squared is the derivative of pressure with respect to density at constant entropy,

$$a^{2} = \left. \frac{\partial p}{\partial \rho} \right|_{S} = k\gamma \rho^{\gamma - 1}. \tag{4}$$

Hence, the expression for the pressure gradients becomes

$$\frac{\partial p}{\partial x} = a^2 \frac{\partial \rho}{\partial x},\tag{5}$$

which can be plugged directly into the momentum equation. The system becomes

$$\frac{\partial \rho}{\partial t} + u \frac{\partial \rho}{\partial x} + \rho \frac{\partial u}{\partial x} = 0, \tag{6a}$$

$$\frac{\partial u}{\partial t} + u \frac{\partial u}{\partial x} + \frac{a^2}{\rho} \frac{\partial \rho}{\partial x} = 0, \tag{6b}$$

$$a = \sqrt{k\gamma}\rho^{\frac{\gamma-1}{2}}.$$
 (6c)

Compatibility Condition between u and ρ

Our next step is to find a compatibility condition between the mass and momentum equations. We define $u := V(\rho)$, which, by application of the chain rule, leads to the following equalities between the derivatives of u and ρ :

$$\frac{\partial u}{\partial t} = \mathbf{V}' \frac{\partial \rho}{\partial t},$$
 (7a)

$$\frac{\partial u}{\partial x} = \mathbf{V}' \frac{\partial \rho}{\partial x},\tag{7b}$$

where V' represents differentiation with respect to density. Introducing these relations into the mass and the momentum equations to remove u, we obtain

$$\frac{\partial \rho}{\partial t} + \frac{\partial \rho}{\partial x} \left(\rho \mathbf{V}' + \mathbf{V} \right) = 0, \tag{8a}$$

$$I'\frac{\partial\rho}{\partial t} + \frac{\partial\rho}{\partial x}\left(VV' + \frac{a^2}{\rho}\right) = 0.$$
 (8b)

Since we have an homogeneous system, it can only have a non-trivial solution if the determinant is zero.

$$\begin{vmatrix} 1 & \mathbf{V} + \rho \mathbf{V}' \\ \mathbf{V}' & \mathbf{V}\mathbf{V}' + \frac{a^2}{\rho} \end{vmatrix} = 0$$
(9)

This determinant implies that

$$\mathbf{V}'\left(\mathbf{V} + \frac{a^2}{\rho\mathbf{V}'} - \mathbf{V} - \rho\mathbf{V}'\right) = 0.$$
(10)

The above equation has two possible solutions, either

$$\mathbf{V}' = \mathbf{0} \to \mathbf{V} = C,\tag{11}$$

which is the constant solution, valid but uninteresting, or

$$\frac{a^2}{\rho \mathbf{V}'} - \rho \mathbf{V}' = 0 \to \mathbf{V}' = \pm \frac{a}{\rho}.$$
 (12)

The \pm signs come up because there are two travelling waves, each directed to one side of the domain. Since our piston is located at the right boundary of the domain, we choose the - sign so that waves propagate left. We thus imply an anechoic left boundary. We can integrate V' with respect to ρ to obtain a relation between the flow velocity u and the speed of sound a:

$$\mathbf{V} = -\int_{\rho_0}^{\rho} \frac{a}{\rho} d\rho \tag{13}$$

where ρ_0 is some reference density. At this point it is convenient to use this reference density ρ_0 to remove the constant *k* from the expression of the speed of sound:

$$a = a(\rho) \to a_0 = a(\rho_0),$$
 (14a)

$$a = \sqrt{k\gamma}\rho^{\frac{\gamma-1}{2}} \to a = a_0 \left(\frac{\rho}{\rho_0}\right)^{\frac{\gamma}{2}}$$
. (14b)

Plugging this expression into the integral, we get

$$u = \mathbf{V} = \frac{2a_0}{\gamma - 1} \left(1 - \frac{a}{a_0} \right), \tag{15a}$$

$$a = a_0 - \frac{\gamma - 1}{2}u. \tag{15b}$$

2.1.1 Burgers-like Equation

With all the above, we are now ready to obtain *one* equation which contains the three departing ones. In the momentum equation we first substitute the space derivative of density,

$$\frac{\partial u}{\partial x} = \mathbf{V}' \frac{\partial \rho}{\partial x} \to \frac{\partial \rho}{\partial x} = -\frac{\rho}{a} \frac{\partial u}{\partial x}, \tag{16a}$$

$$\frac{\partial u}{\partial t} + (u-a)\frac{\partial u}{\partial x} = 0.$$
(16b)

Then, we express the speed of sound in terms of velocity (15b), which leads to a PDE containing a Burgers-like nonlinear term and forced convection driven by the static speed of sound,

$$\frac{\partial u}{\partial t} + \frac{\gamma + 1}{2}u\frac{\partial u}{\partial x} - a_0\frac{\partial u}{\partial x} = 0.$$
 (17)

This PDE, together with its boundary conditions and the boundary's prescribed movement, represents the mathematical model of interest for our work.

2.1.2 Complete Determination of Flow Variables

Although we now have *one* equation which accounts for mass conservation and the isentropic relation between pressure and density, we still have three variables. Computing density, pressure and the speed of sound in space and time still remains useful: verification of mass conservation, computation of the force exerted by the fluid at the piston, and other secondary derivations.

Since the speed of sound *a* is defined as a function of *u* in Equation (15b), once the latter is given, we can obtain density ρ and pressure *p* as a function of flow velocity *u*:

$$\rho = \rho_0 \left(\frac{a}{a_0}\right)^{\frac{2}{\gamma-1}},\tag{18a}$$

$$p = p_0 \left(\frac{\rho}{\rho_0}\right)^{\prime}, \tag{18b}$$

$$\left(\frac{a}{a_0}\right) = 1 - \frac{\gamma - 1}{2} \left(\frac{u}{a_0}\right), \tag{18c}$$

$$\left(\frac{\rho}{\rho_0}\right) = \left(1 - \frac{\gamma - 1}{2}\left(\frac{u}{a_0}\right)\right)^{\frac{2}{\gamma - 1}},$$
 (18d)

$$\left(\frac{p}{p_0}\right) = \left(1 - \frac{\gamma - 1}{2} \left(\frac{u}{a_0}\right)\right)^{\frac{2\gamma}{(\gamma - 1)}}.$$
 (18e)

2.1.3 Mass Conservation

Before we present the complete problem and its numerical solution, we comment on the integral equation for mass conservation. This equation will not be explicitly used to integrate the system (as some methods do), but we will use it as a numerical check to assess the correcteness of the results (we expect to satisfy it up to a given accuracy).

For a control volume whose boundary moves with the piston, the integral expression for mass conservation is

$$\frac{d}{dt} \int_{\Omega_t} \rho d\Omega + \int_{\partial \Omega_t} \rho \left(\vec{u} - \vec{u}_c \right) \cdot \vec{n} dS = 0$$
(19)

At the piston location the flow moves with the piston, $\vec{u} = \vec{u}_c$, so the boundary integral vanishes. There is no flux through the walls either, so the only contribution left is the outlet. At this location, the scalar product of the velocity and normal vectors is negative¹⁰, which implies

$$\frac{d}{dt} \int_0^{L(t)} \rho(x,t) \mathrm{d}x - \rho(0,t) u(0,t) = 0.$$
 (20)

If we introduce the expression for density in terms of velocity from Equation (18d), we get

$$\frac{d}{dt} \int_{0}^{L(t)} \left(1 - \frac{\gamma - 1}{2} \left(\frac{u(x, t)}{a_0} \right) \right)^{\frac{2}{\gamma - 1}} dx - u(0, t) \left(1 - \frac{\gamma - 1}{2} \left(\frac{u(0, t)}{a_0} \right) \right)^{\frac{2}{\gamma - 1}} = 0,$$
(21)

where the constant ρ_0 has cancelled out for being a common factor to both summands, and the whole equation an equality with zero.

10. Although we expect the flow to leave the domain during compression, which means a negative magnitude, this products needs to be done taking the variables positive in the direction of their axes. To ease our task at verifying this equation, we rather calculate the integral term for each time step

$$I(t) = \int_0^{L(t)} \left(1 - \frac{\gamma - 1}{2} \left(\frac{u(x, t)}{a_0}\right)\right)^{\frac{2}{\gamma - 1}} \mathrm{d}x, \qquad (22)$$

and only then compute time derivatives, which can be done with a second order finite difference scheme. Then we can compute the mass defect MD(t) between the time derivative of volume variation and mass outflow at the open boundary,

$$MD(t) = I'(t) - u(0,t) \left(1 - \frac{\gamma - 1}{2} \left(\frac{u(0,t)}{a_0}\right)\right)^{\frac{2}{\gamma - 1}}.$$
 (23)

We shall verify to which degree of accuracy this relation is honoured by our numerical scheme.

2.2 Continuous Formulation

We continue with the definition of the problem in the continuous setting: a differential model given by a PDE and its boundary condition, referred to as strong formulation; and its weak formulation derived with the Galerkin principle.

We introduce the ALE formulation, to account for the movement of the mesh. Then, we will introduce a scaling to work with non-dimensional variables. This will complete the strong formulation.

Then, once we have obtained a weak formulation via the Galerkin projection principle, we will introduce a Dirichlet lifting of the boundary conditions. This is not a mandatory step to solve the FOM, since the boundary conditions can be directly set into the right-hand side of the algebraic vectors. However, it is a paramount step for the construction of our reduced space, to ensure the basis serves all boundary parametrizations.

2.2.1 Strong Formulation

At this point of the derivation, we allow ourselves to introduce an artificial viscosity term which was not present in the physical derivation of the model. The viscosity value ε will be small enough so that it does not remove significant amount of energy from the system, while keeping it sufficiently stable. More on this in Section 4.3.

Thus, the differential model in the physical space is given by the following PDE, boundary and initial conditions,

$$\frac{\partial u}{\partial t}\Big|_{x} + \frac{\partial u}{\partial x}\left(\frac{\gamma+1}{2}u - a_{0}\right) - \varepsilon\frac{\partial^{2}u}{\partial x^{2}} = 0, \qquad (24a)$$

$$u(L,t) = L'(t), \qquad (24b)$$

$$\lim_{t \to \infty} u(x,t) = 0, \qquad (24c)$$

$$u(x,0) = 0.$$
 (24d)

The notation for the time derivative,

$$\left. \frac{\partial u}{\partial t} \right|_x,$$

indicates that the derivative takes place in the physical domain. This is relevant for the ALE formulation, which we explain in the following section.

2.2.2 Boundary conditions

At the right boundary (moving piston) a Dirichlet condition sets the flow velocity equal to the one of the moving wall,

$$L'(t) = -\delta L_0 \omega \sin(\omega t). \tag{25}$$

At the left boundary, we have to find a procedure to model the condition that our piston tube is infinite, and hence that the far region of the flow is not aware of the piston's motion. This is an artificial boundary condition, which we choose to deal with by setting the normal derivate in space to zero

$$\frac{\partial u}{\partial x}(0,t) = 0. \tag{26}$$

This simplified model reflects that, due to the absence of incoming waves from the infinite tube, the solution does not change in space once it is far away from the piston motion.

2.2.3 ALE Formulation

Despite the fact that we will be solving the problem in the physical moving domain, we still need two basic ingredients stemming at the root of the ALE method:

- a smooth mapping between domains;
- a mesh velocity vector.

We introduce the ALE mapping $\mathcal{A} : \Omega_0 \to \Omega_t$ that connects a point in the fixed reference domain \mathcal{X} with a point in the physical domain x,

$$x = \mathcal{A}(\mathcal{X}, t), \tag{27a}$$

$$\mathcal{X} = \mathcal{A}^{-1}(x, t). \tag{27b}$$

We assume this map to be regular enough for all the operations that we carry with it. Then, we define the mesh velocity $w_m(x,t)$ function as the time derivative of the spatial coordinate, which will coincide with the time derivative of the ALE map:

$$w_m(x,t) = \frac{\partial x}{\partial t} = \frac{\partial \mathcal{A}}{\partial t} (x,t) = \frac{\partial \mathcal{A}}{\partial t} (\mathcal{A}(\mathcal{X},t),t).$$
(28)

2.2.4 ALE: Uniform Stretching

For the moving piston, a simple mapping can be derived, rescaling the spatial variable with the piston length in time,

$$\mathcal{X} = \frac{x}{L(t)} \to \mathcal{A}(\mathcal{X}, t) = \mathcal{X}L(t), \ \mathcal{X} \in [0, 1].$$
(29)

Then, we have an analytical expression for the mesh velocity too,

$$w_m(x,t) = \frac{\partial \mathcal{A}}{\partial t} = \mathcal{X}L'(t) = L'(t)\left(\frac{x}{L(t)}\right).$$
 (30)

The mesh velocity results in a linear interpolation of the moving boundary velocity.

Additionally, we introduce the Jacobian of the ALE mapping. The Jacobian is defined as the determinant of the ALE mapping gradient,

$$J_{\mathcal{A}} = \left| \frac{\partial x}{\partial \mathcal{X}} \right|. \tag{31}$$

The Jacobian is used when we want to map an integral in the physical domain into an integral in the reference, fixed domain,

$$\int_{\Omega_t} \phi \mathrm{d}\Omega = \int_{\Omega_0} \phi J_{\mathcal{A}} \mathrm{d}\Omega.$$
 (32)

For our piston-like one-dimensional geometry, we have a very simple transformation,

$$J_{\mathcal{A}} = L(t), \tag{33}$$

which spreads out the deformation of the domain uniformly across all the mesh elements. Since the Jacobian does not depend on space, we could pull it out of the integral,

$$\int_{\Omega_t} \phi \mathrm{d}\Omega = \int_{\Omega_0} \phi L(t) \mathrm{d}\Omega = L(t) \int_{\Omega_0} \phi \mathrm{d}\Omega.$$
(34)

That is, the Jacobian transformation for a piston-like motion does not introduce any nonlinearity in the space dimension. We will come back to this fact further down in the document, when we comment upon the reduction of the algebraic operators.

At last, we can carry out a *correctness* check in our calculations by attempting to verify a well known relation between the mesh velocity vector and the Jacobian [54],

$$\frac{\partial J_{\mathcal{A}}}{\partial t} = J_{\mathcal{A}} \nabla \cdot w_m, \qquad (35a)$$

$$\frac{\partial J_{\mathcal{A}}}{\partial t} = L'(t), \tag{35b}$$

$$J_{\mathcal{A}}\nabla \cdot w_m = L(t)\frac{L'(t)}{L(t)} = L'(t), \qquad (35c)$$

which proves the identity. Equation (35a) could be part of our problem if we did not have access to an explicit expression for the Jacobian, and we wanted to solve our problem in a fixed reference domain. If the reference domain coincided with the initial configuration of the transient domain, this would be an ODE whose initial condition is $J_A(0) = 1$.

2.2.5 ALE: Non-Uniform Mesh Displacement

We also introduce a non-uniform distortion to the nodes, to mimick the effects of mesh adapting techniques. As a proxy, we take a generic density function with a Gaussian bell,

$$\mathcal{F}(\mathcal{X}) = y_c \cdot \exp\left(-\left(\frac{\mathcal{X} - x_c}{\sigma_c}\right)^2\right),\tag{36}$$

where three geometrical parameters are defined:

- x_c : location;
- σ_c : span,
- *y_c*: magnitude.

This will concentrate nodes on a specific region of the domain. Then, for $\mathcal{X} \in \Omega_0$, where $\Omega_0 = [0, L_0]$ is the fixed reference domain, we have the following transformation in terms of a mesh displacement,

$$x = \mathcal{X} + d(\mathcal{X}, t), \tag{37a}$$

$$d(\mathcal{X}, t) = \mathcal{X} \cdot [1 + \mathcal{F}(\mathcal{X})] \Big(\hat{L}(t) - 1 \Big),$$
(37b)

$$\hat{L}(t) = 1 - \delta \cdot (1 - \cos(\omega t)).$$
(37c)

This displacement contains the piston oscillation and the non-uniform concentration of the nodes.

Effects on the Mesh Size

We demonstrate the effects of the non-uniform displacement in the mesh step size. By evaluating Equation (37a) at two consecutive nodes and computing their difference,

$$\Delta x_i = x_i - x_{i-1},\tag{38a}$$

$$\Delta x_i = \mathcal{X}_i - \mathcal{X}_{i-1} + (d(\mathcal{X}_i, t) - d(\mathcal{X}_{i-1}, t)),$$
(38b)

$$\Delta x_i = \Delta \mathcal{X}_i + (d(\mathcal{X}_i, t) - d(\mathcal{X}_{i-1}, t)), \tag{38c}$$

$$\Delta x_{i} = \underbrace{\Delta \mathcal{X}_{i} \cdot \hat{L}(t)}_{\text{Uniform stretching}} + [\mathcal{X}_{i} \cdot \mathcal{F}(\mathcal{X}_{i}) - \mathcal{X}_{i-1} \cdot \mathcal{F}(\mathcal{X}_{i-1})] \cdot (\hat{L}(t) - 1); \quad (38d)$$

we arrive to an expression for the mesh size Δx_i as a function of the reference mesh size and nodes position. If the function \mathcal{F} was set to zero, we recover the uniform stretching derived in Section 2.2.4.

2.2.6 Time Derivative in the Reference Domain

Since the equations are to be solved in the physical domain, we need to adjust the time derivative in the physical domain $\left(\frac{\partial u}{\partial t}\Big|_{x}\right)$, so that it takes into account the movement of the mesh nodes. By application of the chain rule we get

$$\left. \frac{\partial u}{\partial t} \right|_{\mathcal{X}} = \left. \frac{\partial u}{\partial t} \right|_x + w_m \frac{\partial u}{\partial x},\tag{39}$$

from where we get the necessary modification to the strong form (24a) of the PDE,

$$\frac{\partial u}{\partial t}\Big|_{\mathcal{X}} + \left(\frac{\gamma+1}{2}\right)u\frac{\partial u}{\partial x} - (a_0 + w_m)\frac{\partial u}{\partial x} - \varepsilon\frac{\partial^2 u}{\partial x^2} = 0.$$
(40)

An additional convective term shows up, to take into account the movement of the nodes. If we neglect this term, mass is no longer conserved.

2.2.7 Dimensionless Equations

Finally, we carry out a non-dimensionalization of the velocity and the spatial coordinate,

$$\tilde{x} = \frac{x}{L_0}, \qquad \tilde{u} = \frac{u}{a_0}, \tag{41a}$$

with respect to the static speed of sound a_0 and the piston's initial length L_0 . This leads to the PDE and boundary condition

$$\frac{\partial \tilde{u}}{\partial t} + \left(\frac{\gamma+1}{2}\right) \left(\frac{a_0}{L_0}\right) \tilde{u} \frac{\partial \tilde{u}}{\partial \tilde{x}} - \left(\frac{\varepsilon}{L_0^2}\right) \frac{\partial^2 \tilde{u}}{\partial \tilde{x}^2} - \left(\frac{a_0}{L_0}\right) \left(1 + \frac{w_m}{a_0}\right) \frac{\partial \tilde{u}}{\partial \tilde{x}} = 0, \quad (42a)$$

$$\tilde{u}(L(t),t) = \frac{L'(t)}{a_0} = b_L(t).$$
 (42b)

For the sake of clear notation, we condense the coefficients

$$b_0 = \frac{a_0}{L_0} \left(\frac{\gamma + 1}{2}\right),\tag{43a}$$

$$b_L(t) = -\frac{\delta L_0 \omega}{a_0} \sin(\omega t), \qquad (43b)$$

$$c(x,t) = \left(\frac{a_0}{L_0}\right) \left(1 - \frac{w_m}{a_0}\right),\tag{43c}$$

and drop the \tilde{u} notation to avoid an overloaded notation. The coefficient

$$u_p = \left(\frac{\delta L_0 \omega}{a_0}\right) \tag{44}$$

multiplying the boundary condition is the ratio between the piston motion and the static speed of sound¹¹. It is the maximum piston mach number, in terms of the static speed of sound. This coefficient will drive the response of the fluid to the motion of the piston. The larger it is, the stronger the nonlinear response will become (steepening shock wave).

With all of this done, we obtain a familiar PDE structure with diffusion, linear and nonlinear convection terms:

$$\frac{\partial u}{\partial t}\Big|_{\mathcal{X}} + b_0 u \frac{\partial u}{\partial x} - c(x, t) \frac{\partial u}{\partial x} - \varepsilon \frac{\partial^2 u}{\partial x^2} = 0.$$
(45)

2.2.8 Dimensionless Mass Conservation

If the former rescaling is applied, Equation (23) for mass conservation needs to be updated, to take into account the fact that the velocity variable is now scaled with the speed of sound.

$$I(t) = \int_{0}^{\frac{L(t)}{L_{0}}} \left(1 - \frac{\gamma - 1}{2}u(x, t)\right)^{\frac{2}{\gamma - 1}} dx,$$
(46a)
$$\frac{MD(t)}{\rho_{0}a_{0}} = \frac{1}{a_{0}}I'(t) - u(0, t)\left(1 - \frac{\gamma - 1}{2}u(0, t)\right)^{\frac{2}{\gamma - 1}}.$$
(46b)

2.2.9 Weak Formulation

Since we will be working with the Galerkin procedure to solve PDEs, we define the $L^2(\Omega)$ inner product to transform the strong formulation into a weak, variational one,

$$\langle u, v \rangle = \int_0^L uv \, \mathrm{d}\Omega.$$
 (47)

This inner product induces the so called *eyeball* norm, if two functions *look* alike ||f - g|| will return a small value,

$$||f - g|| = \sqrt{\int_0^L (f - g)^2 \, \mathrm{d}\Omega}.$$
 (48)

With this inner product, we project the residual of the strong formulation onto a given fuction space $u, v \in V$, where V is a suitable Hilbert space,

$$\left\langle \frac{\partial u}{\partial t}, v \right\rangle + \left\langle b_0 u \nabla u, v \right\rangle - \left\langle c \nabla u, v \right\rangle + \left\langle \varepsilon \nabla u, \nabla v \right\rangle + \varepsilon \left. \frac{\partial u}{\partial x} \right|_{x=0} v = 0,$$
 (49a)

$$u(x,0) = 0,$$
 (49b)

$$\frac{\partial u}{\partial t}(0,t) = 0, \qquad (49c)$$

$$u(L,t) = b_L(t); \tag{49d}$$

$$b_L(t) = -\frac{\delta L_0 \omega}{a_0} \sin \omega t. \quad (49e)$$

11. A proxy for the average speed at which information propagates.

2.2.10 Dirichlet Lifting

For reasons related to the fact that we have a parametrized boundary condition, which will become apparent when the reduction scheme is presented, it is preferable to work with a homogeneous problem in the Dirichlet boundary conditions.

To obtain so, we introduce a *lifting* g(x, t) of the Dirichlet boundary conditions. We express the solution of our problem like the linear combination of the solution of the homogeneous problem and the lifting function:

$$u(x,t) = \hat{u}(x,t) + g(x,t).$$
(50)

The lifting of the boundary conditions has to meet two restrictions:

- reach the prescribed values at the boundary nodes;
- be sufficiently smooth within the domain.

In a one-dimensional setting, the definition of a lifting function g(x,t) is straigtforward, and can be done using a linear interpolation of the boundary values,

$$g(x,t) = b_L(t)\left(\frac{x}{L}\right).$$
(51)

In higher dimensional settings a similar procedure is used. However, due to the arbitrary shape the domain can take, the construction of the lifting function can be more laborious.

Introducing the lifting breakdown (50) into the weak formulation (49a), we find additional forcing terms and PDE terms, due to the cross-product between the function and its derivative,

$$u\frac{\partial u}{\partial x} = (\hat{u} + g)\left(\frac{\partial \hat{u}}{\partial x} + \frac{\partial g}{\partial x}\right).$$
 (52)

Taking this into account, we find the following *lifted* weak formulation,

$$\left\langle \frac{\partial \hat{u}}{\partial t}, v \right\rangle - \langle c \nabla \hat{u}, v \rangle + \langle \varepsilon \nabla \hat{u}, \nabla v \rangle + \langle b_0 g \nabla \hat{u}, v \rangle + \langle b_0 \hat{u} \nabla g, v \rangle + \langle b_0 \hat{u} \nabla \hat{u}, v \rangle = - \left\langle \frac{\partial g}{\partial t}, v \right\rangle + \langle c \nabla g, v \rangle - \langle \varepsilon \nabla g, \nabla v \rangle - \langle b_0 g \nabla g, v \rangle, \qquad (53a)$$

where we have reorganized the terms to show in order: linear dependency with the solution, terms due to crossproduct effects, and finally the actual nonlinearity.

Thanks to the lifting, we find the homogenization of the left boundary condition for all time t,

$$\hat{u}(L,t) = 0, \tag{54a}$$

and recall that the initial condition should be modified accordingly to the homogeneous problem definition,

$$u(x,0) = \hat{u}(x,0) + g(x,0),$$
 (55a)

$$\hat{u}_0(x) := \hat{u}(x,0) = u(x,0) - g(x,0).$$
 (55b)

At this point, we have defined the continuous problem for the one-dimensional piston in a moving domain.

2.3 Semi-Discrete Formulation: Time Discretization

The continuous solution changes in two dimensions: space and time. Eventually, we need to discretize them both, but we can do so separately. We choose to first discretize in time.

For this task, we need to build an approximation for the time derivative. A polynomial interpolation $p_n(x,t)$ of the function $\hat{u}(x,t^{n+1})$ is built, using the solution at previous time steps, $\{\hat{u}(x,t^n), \hat{u}(x,t^{n-1}), \ldots\}$. Then, the interpolant's derivative at time t^{n+1} is used as an approximation of the actual derivative,

$$\frac{\partial \hat{u}}{\partial t}(x, t^{n+1}) \simeq \frac{\partial p_n}{\partial t}(x, t^{n+1}).$$
(56)

The accuracy of the scheme (and also its complexity) is determined by the number of previous time steps used in the interpolation.

In the coming sections, we define two time-marching schemes of order one and two, BDF-1¹², and BDF-2, respectively. Throughout our simulations we will use BDF-2, but since it requires two points from the past, it cannot be used at the beginning of the simulation. Thus, the first step is done with BDF-1, to obtain $\hat{u}(x, t^1)$.

From now on we use the abbreviated notation

$$\hat{u}^n := \hat{u}(x, t^n) \tag{57}$$

to define a function in space evaluated at time t^n . Additionally, to simplify our derivations, we will use a toy weak form, with only two spatial differential operators and one forcing term, as placeholders for the actual linear and nonlinear terms of a complete model,

$$\left\langle \frac{\partial \hat{u}}{\partial t}, v \right\rangle + \left\langle u, v \right\rangle + \left\langle l(u), v \right\rangle = \left\langle f, v \right\rangle.$$
(58)

The functional l(u) stands for a nonlinear transformation on u and its derivatives in space.

2.3.1 BDF-1

To derive the both schemes, we start by evaluating our weak formulation at time t^{n+1} , where the solution is unknown¹³,

$$\left\langle \frac{\partial \hat{u}}{\partial t}, v \right\rangle^{n+1} + \left\langle \hat{u}, v \right\rangle^{n+1} + \left\langle l(\hat{u}), v \right\rangle^{n+1} = \left\langle f, v \right\rangle^{n+1}$$
(59)

Then, we construct our interpolation polynomial using only one previous time step from the past,

$$p_1(t) := \hat{u}^{n+1} \left(\frac{t - t^n}{\Delta t} \right) - \hat{u}^n \left(\frac{t - t^{n+1}}{\Delta t} \right); \tag{60a}$$

$$\frac{\partial p_1}{\partial t} = \frac{u^{n+1} - u^n}{\Delta t}.$$
(60b)

With this approximation of the time derivative, we get the following semi-discrete weak formulation,

$$\hat{u}^{n+1} + \Delta t \left\langle \hat{u}, v \right\rangle^{n+1} + \Delta t \left\langle l(\hat{u}), v \right\rangle^{n+1} = \\ \left\langle \hat{u}^n, v \right\rangle^{n+1} + \Delta t \left\langle f, v \right\rangle^{n+1}$$
(61)

Note that the problem is still continuous in space, but no longer in time.

12. Also known as Backwards Euler.

13. For the initial time t^0 the solution is known, from the initial condition u(x, 0).

2.3.2 BDF-2

For the BDF-2 scheme, our interpolation takes two previous time steps from the past,

$$p_{2}(t) := \hat{u}^{n+1} \left(\frac{t-t^{n}}{\Delta t} \right) \left(\frac{t-t^{n-1}}{2\Delta t} \right) - \hat{u}^{n} \left(\frac{t-t^{n+1}}{\Delta t} \right) \left(\frac{t-t^{n-1}}{\Delta t} \right) + \hat{u}^{n-1} \left(\frac{t-t^{n+1}}{2\Delta t} \right) \left(\frac{t-t^{n}}{\Delta t} \right);$$
(62a)
$$\frac{\partial p_{2}}{\partial t} = \frac{3\hat{u}^{n+1} - 4\hat{u}^{n} + \hat{u}^{n-1}}{2\Delta t} = \frac{1}{\Delta t} \left(\frac{3}{2} \hat{u}^{n+1} - 2\hat{u}^{n} + \frac{1}{2} \hat{u}^{n-1} \right).$$
(62b)

With this alternative approximation of the time derivate, we get a very similar semi-discrete weak formulation as the one from Equation (61),

$$\frac{3}{2} \left\langle \hat{u}^{n+1}, v \right\rangle^{n+1} + \Delta t \left\langle \hat{u}, v \right\rangle^{n+1} + \Delta t \left\langle l(\hat{u}), v \right\rangle^{n+1} = 2 \left\langle \hat{u}^{n}, v \right\rangle^{n+1} - \frac{1}{2} \left\langle \hat{u}^{n-1}, v \right\rangle^{n+1} + \Delta t \left\langle f, v \right\rangle^{n+1}.$$
 (63)

In fact, both formulations can be synthezised into one,

$$m_{\text{BDF}} \left\langle \hat{u}^{n+1}, v \right\rangle^{n+1} + \Delta t \left\langle \hat{u}, v \right\rangle^{n+1} + \Delta t \left\langle l(\hat{u}), v \right\rangle^{n+1} = \left\langle \hat{u}_{\text{BDF}}, v \right\rangle^{n+1} + \Delta t \left\langle f, v \right\rangle^{n+1},$$
(64)

provided that the forcing term due to the time integration scheme is generalized,

$$\hat{u}_{\text{BDF}} = \begin{cases} \hat{u}^n, & \text{BDF-1,} \\ 2\hat{u}^n - \frac{1}{2}\hat{u}^{n-1}, & \text{BDF-2;} \end{cases}$$
(65)

and the parameter m_{BDF} is modified accordingly for each time integration scheme,

$$m_{\rm BDF} = \begin{cases} 1, & {\rm BDF-1}, \\ \frac{3}{2}, & {\rm BDF-2}. \end{cases}$$
 (66)

The generalized semi-discrete weak formulation from Equation (64) is much more comfortable from an implementation-wise perspective, since it has isolated the effects of the discretization scheme.

2.3.3 Nonlinear Convective Term Linearization

Now, Equation (64) is a nonlinear one, due to the presence of the $l(\hat{u})$ functional, which represents the nonlinear convection term from our original model.

In the presence of nonlinearities, one can opt to evaluate certain terms with known information from the past. This trick can be used in problems such as Burgers's equation, (or the Navier-Stokes equations) where going fully implicit leads to a nonlinear algebraic system. If the convective velocity term is extrapolated, one recovers a linear system, and yet reaches a satisfactory numerical solution [1]. Thus, we opt for the approximation of its effect by considering a second order extrapolation in time of the convection velocity,

$$\langle b_0 \hat{u} \boldsymbol{\nabla} \hat{u}, v \rangle^{n+1} \simeq \langle b_0 \hat{u}^* \boldsymbol{\nabla} \hat{u}^{n+1}, v \rangle,$$
 (67a)

$$\hat{u}^* = 2\hat{u}^n - \hat{u}^{n-1} + \mathcal{O}(\Delta t^2).$$
 (67b)

The extrapolation can be obtained by expanding \hat{u}^{n+1} and \hat{u}^{n-1} with a Taylor polynomial centered around \hat{u}^n ,

$$\hat{u}^{n+1} = \hat{u}^n + \frac{\partial \hat{u}^n}{\partial t} \Delta t + \frac{1}{2} \frac{\partial^2 \hat{u}^n}{\partial t^2} \Delta t^2 + \mathcal{O}(\Delta t^3), \quad (68a)$$

$$\hat{u}^{n-1} = \hat{u}^n - \frac{\partial \hat{u}^n}{\partial t} \Delta t + \frac{1}{2} \frac{\partial^2 \hat{u}^n}{\partial t^2} \Delta t^2 + \mathcal{O}(\Delta t^3).$$
(68b)

Combining both expansions, Δt -order terms cancel out and one gets

$$\hat{u}^{n+1} = 2\hat{u}^n - \hat{u}^{n-1} + \frac{\partial^2 \hat{u}^n}{\partial t^2} \Delta t^2 + \mathcal{O}(\Delta t^3).$$
 (69)

If the Taylor expansion terms are truncated, one gets the expression for the velocity extrapolation in time,

$$\hat{u}^* = 2\hat{u}^n - \hat{u}^{n-1} + \mathcal{O}(\Delta t^2).$$
 (70)

With this strategy we obtain a linear system. It is paramount to use a second order approximation for the convective velocity, otherwise the accuracy benefits of the BDF-2 scheme are lost; falling back to order one convergence rates, despite the second order approximation of the time derivative.

At the continuous level, this term is nonlinear, but at the discrete level, it is trilinear in all its arguments.

2.3.4 Semi-Discrete Weak Formulation

All in all, the complete semi-discretized weak formulation for the original problem can be found by replacing the placeholders for the actual spatial operators from Equation (53) into Equation (64); the generalized forcing term \hat{u}_{BDF} due to time-discretization, Equation (65); and the second-order approximation \hat{u}^* of the convection velocity in the nonlinear operator, Equation (67b),

$$m_{\text{BDF}} \langle \hat{u}^{n+1}, v \rangle - \Delta t \langle c \nabla \hat{u}^{n+1}, v \rangle + \Delta t \langle \varepsilon \nabla \hat{u}^{n+1}, \nabla v \rangle + \Delta t \left(\langle b_0 \hat{u}^{n+1} \nabla g^{n+1}, v \rangle + \langle b_0 g^{n+1} \nabla \hat{u}^{n+1}, v \rangle \right) + \langle b_0 \hat{u}^* \nabla \hat{u}^{n+1}, v \rangle = \langle \hat{u}_{\text{BDF}}, v \rangle - \Delta t \left\langle \frac{\partial g^{n+1}}{\partial t}, v \right\rangle - \Delta t \langle b_0 g^{n+1} \nabla g^{n+1}, v \rangle - \Delta t \langle \varepsilon \nabla g^{n+1}, \nabla v \rangle + \Delta t \langle c^{n+1} \nabla g^{n+1}, v \rangle.$$
(71a)

2.4 Discrete Problem: Space Discretization

To complete our discretization, we define a finite functional space $V_h \subset V$, where we can represent the solution as the linear combination of a set of finite elements (FE) basis functions $\varphi_i(x)$ with local support,

$$\hat{u}^n(x) \simeq \hat{u}^n_h(x) = \sum_j^{N_h} \hat{u}^n_{h_j} \varphi_j(x), \tag{72a}$$

$$\hat{\mathbf{u}}_{h}^{n} = \begin{bmatrix} \hat{u}_{h_{j}}^{n} \end{bmatrix}.$$
(72b)

We define the FE vector $\hat{\mathbf{u}}_h^n$ to be the collection of coefficients $\hat{u}_{h_j}^n$ which multiply the basis functions. In the FE context with Lagrangian functions, these coincide with the values of the function at each node.

Applying the Galerkin principle to solve PDEs, we enforce the orthogonality of the residual to the functional space V_h . Because the domain changes with time, both the matrices and the vectors change for each time step,

$$\left[\mathbf{M}_{h}^{n+1}\right]_{ij} = m_{\text{BDF}} \left\langle \varphi_{j}, \varphi_{i} \right\rangle^{n+1}, \tag{73a}$$

$$\left[\mathbf{A}_{h}^{n+1}\right]_{ij} = \left\langle \varepsilon \nabla \varphi_{j}, \nabla \varphi_{i} \right\rangle^{n+1}, \tag{73b}$$

$$\left[\mathbf{C}_{h}^{n+1}\right]_{ij} = -\left\langle c\nabla\varphi_{j},\varphi_{i}\right\rangle^{n+1},\tag{73c}$$

$$\begin{bmatrix} \mathbf{N}_{h}^{n+1} \end{bmatrix}_{ij} = b_0 \langle u^{*,n} \nabla \varphi_j, \varphi_i \rangle^{n+1}, \qquad (73d)$$
$$\begin{bmatrix} \mathbf{\hat{N}}_{h}^{n+1} \end{bmatrix}_{ij} = b_0 \left(\langle g \nabla \varphi_j, \varphi_i \rangle^{n+1} + \langle \varphi_j \nabla g, \varphi_i \rangle^{n+1} \right), \quad (73e)$$

$$\begin{bmatrix} \mathbf{F}_{g,h}^{n+1} \end{bmatrix}_{i} = -\left\langle \frac{\partial g}{\partial t} + b_{0}g\boldsymbol{\nabla}g - c\boldsymbol{\nabla}g, \varphi_{i} \right\rangle^{n+1} - \left\langle \varepsilon\boldsymbol{\nabla}g, \boldsymbol{\nabla}\varphi_{i} \right\rangle^{n+1},$$
(73f)

$$\left(\left\langle \hat{u}_{h}^{n},\varphi_{i}\right\rangle ^{n+1},$$
 BDF-1

$$\begin{bmatrix} \mathbf{F}_{\hat{\mathbf{u}}_{h}}^{n} \end{bmatrix}_{i} = \begin{cases} \mathbf{F}_{\hat{\mathbf{u}}_{h}}^{n} \\ 2\left\langle \hat{u}_{h}^{n}, \varphi_{i} \right\rangle^{n+1} - \frac{1}{2}\left\langle \hat{u}_{h}^{n-1}, \varphi_{i} \right\rangle^{n+1}, & \text{BDF-2.} \end{cases}$$
(73g)

This leads to the following algebraic system:

$$m_{\text{BDF}} \mathbf{M}_{h}^{n+1} \hat{\mathbf{u}}_{h}^{n+1} + \Delta t \mathbf{C}_{h}^{n+1} \hat{\mathbf{u}}_{h}^{n+1} + \Delta t \mathbf{A}_{h}^{n+1} \hat{\mathbf{u}}_{h}^{n+1}$$
$$+ \Delta t \hat{\mathbf{N}}_{h}^{n+1} \hat{\mathbf{u}}_{h}^{n+1} + \Delta t \left[\mathbf{N}_{h}^{n+1} \left(\hat{\mathbf{u}}_{h}^{*} \right) \right] \hat{\mathbf{u}}_{h}^{n+1}$$
$$= \mathbf{F}_{\hat{\mathbf{u}}_{h}}^{n} + \Delta t \mathbf{F}_{g,h}^{n+1}, \quad (74a)$$
$$\hat{\mathbf{u}}_{h}^{0} = \hat{\mathbf{u}}_{h,0}. \quad (74b)$$

The spatial boundary conditions are encoded within the matrices and the vectors. The initial condition is obtain via interpolation or projection. The assembly of the forcing term in Equation (73f) has to be done with the FE vector representation of the functional f_g (which can be obtained by projection or interpolation).

Regarding the forcing due to previous time steps, $(\mathbf{F}_{\hat{\mathbf{u}}_{h}}^{n})$, although for the FOM model we could compute the inner products at each time step, for the Reduced Order Model we will exploit an algebraic expression of these expressions. It can be expressed as the product between the mass matrix and the FE representation of the previous solution(s),

$$\mathbf{F}_{\hat{\mathbf{u}}_{h}}^{n} = \begin{cases} \mathbf{M}_{h}^{n+1} \hat{\mathbf{u}}_{h}^{n}, & \text{BDF-1}, \\ 2\mathbf{M}_{h}^{n+1} \hat{\mathbf{u}}_{h}^{n} - \frac{1}{2} \mathbf{M}_{h}^{n+1} \hat{\mathbf{u}}_{h}^{n-1}, & \text{BDF-2}. \end{cases}$$
(75)

We point out how the time step Δt has been intentionally left out of the discrete operators definition. There are two reasons to back this decision:

• Conceptually, each discrete operator encodes a spatial model, in terms of a differential operator or the presence of a forcing term. The time step Δt shows up because we first discretized the continuous problem in time. Had we gone the other way around (discretizing first in space), we would have found a system of ODEs with the previously defined spatial operators.

• When we leverage the system approximation reduction technique, we will not want to have inside the operator the presence of the time step. The reduced model could use a different time step, or the snapshots for different parameters could be gathered for different time step values.

If we collect terms and factor out the unknowns, we get a compact linear system to be solved at each time step to advance the solution,

$$\mathbf{K}_{h}^{n+1}\hat{\mathbf{u}}_{h}^{n+1} = \mathbf{b}_{h}^{n+1},\tag{76a}$$

$$\hat{\mathbf{u}}_{h}^{0} = \hat{\mathbf{u}}_{h,0}; \tag{76b}$$

$$\mathbf{K}_{h}^{n+1} = \mathbf{M}_{h}^{n+1} + \Delta t \left[\mathbf{A}_{h}^{n+1} + \mathbf{C}_{h}^{n+1} + \hat{\mathbf{N}}_{h}^{n+1} + \mathbf{N}_{h}^{n+1} + \hat{\mathbf{C}}_{h}^{n+1} \right]$$
(74)

$$+\mathbf{\hat{N}}_{h}^{n+1}+\mathbf{N}_{h}^{n+1}\left(\mathbf{\hat{u}}_{h}^{*}\right)\right],$$
(76c)

$$\mathbf{b}_{h}^{n+1} = \mathbf{F}_{\hat{\mathbf{u}}_{h}}^{n} + \Delta t \mathbf{F}_{g,h}^{n+1}.$$
 (76d)

2.4.1 Summary

The problem has been explained within three levels of abstraction: continuous with strong and weak formulations, semi-discrete in time with the approximation of the time derivative, and fully discretized with the additional FEM representation of the solution. To discretize the nonlinear term, the convective velocity has been extrapolated with a second order scheme.

A lifting of the Dirichlet boundary conditions has been introduced, which leads to an additional forcing term and the homogenization of the boundary conditions. Hence, we will focus in the solution of an homogeneous boundary value problem. This fact will prove useful when we get into the implementation details of the reduction procedure.

3 (HYPER) REDUCED ORDER MODEL

For any unseen parameter value, our aim is to be able to assemble and solve a smaller algebraic problem, and yet to obtain a solution close enough to that of the FOM. In order to do so, first we need to obtain certain algebraic structures which capture the essence of the problem at hand.

We do so by sampling the original problem at certain parameter values and processing snaphsots of the solution and the operators, obtained from the solution of the FOM problem. This is called the *offline phase*. Later on, we exploit these static structures to build reduced operators which capture the dynamics of the problem sufficiently well, solve a smaller algebraic system and then recover the solution in the original mesh variables, in order to postprocess it. This is called the *online phase*.

We will use the Reduced Basis Method (RB) to construct an ad-hoc problem-based basis to represent our ROM; the Discrete Empirical Interpolation Method (DEIM) and its matrix version (MDEIM) to build suitable approximations of the algebraic operators involved.

The continuous reduced problem formulation is skipped since it will not be used. Therefore, we jump directly into the discrete problem. We recall that we are focused on reducing the problem for the homogeneous component $\hat{u}(x)$ of our solution, that is, for the weak formulation given by the system of equations (53).

3.1 A Naive Approach

We have included in the title the word *naive* because we will define the reduction problem in a very blunt way, where many ineffiencies will show up. We do so to motivate the operator reduction procedures we will include later on.

In the reduced context, we use a finite space $V_N \subset V_h$, where we can represent the solution as the linear combination of a set of orthonormal¹⁴ problem-based basis functions $\psi_i(x)$,

$$\hat{u}_N(x) = \sum_j^N \hat{u}_{N_j} \psi_j(x).$$
 (77)

Their goal is to capture the problem dynamics, so these RB solution modes $\psi_i(x)$ have global support. Since we want to reduce the number of modes we need to represent our solution, we usually expect or desire to have $N \ll N_h$.

To maintain focus and in favor of generality, let us assume at this point that the global basis functions are given, and that they are zero at the boundary. We will give details on how to obtain them later on.

3.1.1 Reduced Space Projection

Since we can represent any function in terms of our FE basis functions $\varphi_i(x)$, we have a linear mapping between the problem-based functions $\psi_j(x)$ and the nodal basis functions. This allows us to establish the following relation between the problem solution in the reduced and original spaces,

$$\hat{\mathbf{u}}_h = \mathbb{V}\hat{\mathbf{u}}_N. \tag{78}$$

14. If they were not, we can always make them so via Gram-Schmidt.

The entries of the V matrix represent the coefficients of the global basis representation in the FE basis,

$$\psi_i(x) = \sum_j \left[\mathbb{V}\right]_{ji} \varphi_j(x). \tag{79}$$

3.1.2 Discrete Reduced Problem Assembly

In theory, to assemble the reduced problem operators, one could actually compute the inner products defined in Equations (73) with these new basis functions $\psi_i(x)$. In practice, it is more convenient to project the algebraic FOM operators with matrix-matrix and matrix-vector products into the reduced space,

$$\mathbf{X}_{N}^{n+1} = \mathbb{V}^{T} \mathbf{X}_{h}^{n+1} \mathbb{V}, \tag{80a}$$

$$\mathbf{F}_{N}^{n+1} = \mathbb{V}^{T} \mathbf{F}_{h}^{n+1}, \tag{80b}$$

where \mathbf{X}_h and \mathbf{F}_h stand for each of the FOM operators (matrices and vectors respectively). The assembly of matrices based on a FE basis can be easily done in parallel due to their local support, whereas the integration of functions with global support is not necessarily computationally efficient.

By doing so, we find the following pure ROM problem for the time evolution problem,

n

$$n_{\text{BDF}} \mathbf{M}_{N}^{n+1} \hat{\mathbf{u}}_{N}^{n+1} + \Delta t \mathbf{C}_{N}^{n+1} \hat{\mathbf{u}}_{N}^{n+1} + \Delta t \mathbf{A}_{N}^{n+1} \hat{\mathbf{u}}_{N}^{n+1}$$
$$+ \Delta t \hat{\mathbf{N}}_{N}^{n+1} \hat{\mathbf{u}}_{N}^{n+1} + \Delta t \left[\mathbf{N}_{N}^{n+1} \left(\hat{\mathbf{u}}_{N}^{*} \right) \right] \hat{\mathbf{u}}_{N}^{n+1}$$
$$= \mathbf{F}_{\hat{\mathbf{u}}_{N}}^{n} + \Delta t \mathbf{F}_{g,N}^{n+1}, \quad (81a)$$

 $\hat{\mathbf{u}}_{N}^{0} = \hat{\mathbf{u}}_{N.0}.$ (81b)

If we collect terms and factor out the unknowns we get a linear system, this time in the reduced space, to be solved for each time step to advance the solution,

$$\mathbf{K}_{N}^{n+1}\hat{\mathbf{u}}_{N}^{n+1} = \mathbf{b}_{N}^{n+1},\tag{82a}$$

$$\hat{\mathbf{u}}_N^0 = \hat{\mathbf{u}}_{N,0}; \tag{82b}$$

$$\mathbf{K}_{N}^{n+1} = m_{\text{BDF}} \mathbf{M}_{N}^{n+1} + \Delta t \left[\mathbf{A}_{N}^{n+1} + \mathbf{C}_{N}^{n+1} + \hat{\mathbf{N}}_{N}^{n+1} + \mathbf{N}_{N}^{n+1} \left(\hat{\mathbf{u}}_{N}^{*} \right) \right], \qquad (82c)$$

$$\mathbf{b}_{N}^{n+1} = \mathbf{F}_{\hat{\mathbf{u}}_{N}}^{n} + \Delta t \mathbf{F}_{g,N}^{n+1}.$$
(82d)

All of the previous has the same algebraic pattern as the FOM problem. The only difference is the size of the operators, much smaller due to the small size of N.

3.1.3 ROM Time Discretization Forcing Term

The forcing term $\mathbf{F}_{\hat{\mathbf{u}}_h}^n$ due to the time discretization has been intentionally left out in the previous section. We could naively project it too,

$$\mathbf{F}_{\hat{\mathbf{u}}_N}^N = \mathbb{V}^T \mathbf{F}_{\hat{\mathbf{u}}_h}^n, \tag{83}$$

but this would force us to reconstruct the FE vector of the ROM at each time-step, making the integration cumbersome. To get around this issue, we can exploit the algebraic representation of $\mathbf{F}_{\hat{\mathbf{u}}_h}^n$, given in Equation (75) in terms of the mass matrix. The forcing term due to the time discretization takes the following form in the ROM:

$$\mathbf{F}_{\hat{\mathbf{u}}_{N}}^{n} = \begin{cases} \mathbf{M}_{N}^{n+1} \hat{\mathbf{u}}_{N}^{n}, & \text{BDF-1,} \\ 2\mathbf{M}_{N}^{n+1} \hat{\mathbf{u}}_{N}^{n} - \frac{1}{2}\mathbf{M}_{N}^{n+1} \hat{\mathbf{u}}_{N}^{n-1}, & \text{BDF-2.} \end{cases}$$
(84)

Again, these mimic the algebraic pattern obtained in the FOM.

3.1.4 Boundary and Initial Conditions

The computation of the initial condition $\hat{\mathbf{u}}_{N,0}$ is to be done in two steps. First, the initial condition $\hat{\mathbf{u}}_{h,0}$ in the FOM space needs to be computed as a FE vector via interpolation or projection. Then, this FE representation $\hat{\mathbf{u}}_{h,0}$ needs to be projected unto the reduced space. Since we are dealing with an orthonormal basis, we can use the matrix \mathbb{V} to project the FE vector departing from the FOM-ROM relation given in Equation (78),

$$\mathbb{V}^T \hat{\mathbf{u}}_{h,0} = \underbrace{\mathbb{V}^T \mathbb{V}}_{\mathbb{T}} \hat{\mathbf{u}}_{N,0} \to \hat{\mathbf{u}}_{N,0} = \mathbb{V}^T \hat{\mathbf{u}}_{h,0}.$$
(85)

Regarding the spatial boundary conditions, at this point of the naive reduction scheme, two facts come into play, which we first present and then put together:

- The weak form has homogeneous boundary conditions.
- 2) The ad-hoc RB solution modes $\psi_i(x)$ are zero at the boundaries,

$$\psi_i(x) = 0 \quad \forall x \in \partial\Omega.$$

These two facts imply that the projection of the operators, Equation (80), does not break the original constraint of homogeneous boundary conditions; and that the linear expansion of the solution $\hat{u}_N(x)$ in the span of V_N , Equation (77), is always true.

Once the reduced homogeneous solution $\hat{u}_N(x)$ is obtained, it can be brought back to the original space V_h via Equation (78), and then added to the FE representation of the Dirichlet lifting, to obtain the solution $u_h(t, \mu)$ in the physical domain,

$$u(x,t;\mu) \simeq u_h(t,\mu) = \mathbb{V}\hat{u}_N(t,\mu) + g_h(t,\mu).$$
 (86)

3.1.5 Naive Reduction Scheme: Final Remarks

At this point, the naive reduction scheme for the RB-ROM has been defined and explained. However, some aspects of it remain unattended.

The construction of the problem-based basis functions $\psi_i(x)$: there are many methods to build them, and which combination of them we choose defines which reduction method we are applying, along with their advantages and requirements.

The offline-online decomposition: that is, to uncouple the usage of FOM operators in as much as possible from the integration of the ROM. Ideally, no FOM structures should be assembled during the online phase. In this naive scheme, we are clearly not meeting such requirement, since we need to assemble all the FOM operators and then project them for each time step.

Hence, some additional sections need to be brought up, in order to complete our definition of the reduction scheme. We now treat the construction of the basis functions, Section 3.2; and the approximation of the algebraic operators, Section 3.3.

3.2 Reduced Basis Construction

Hereby we layout the details of the basis construction, that is, the definition of the $\psi_i(x)$ functions. There are many techniques to build such basis. We opt for an automatic and out-of-the-box technique: the nested POD approach [1].

On paper, the most simple basis anyone could come up with is a collection of solution snaphsots for different parameter values and time steps,

$$\Psi_{\hat{u}} := [\psi_i]
= [\hat{u}_h(t^0; \mu_0), \hat{u}_h(t^1; \mu_0), \dots, \hat{u}_h(t^j; \mu_i), \dots],
= [\Psi_{\hat{u}}(\mu_0), \Psi_{\hat{u}}(\mu_1), \dots, \Psi_{\hat{u}}(\mu_{N_\mu})]$$
(87)

Yet, this basis $\Psi_{\hat{u}}$ is unpractical from a computational point of view: we could not possibly store all the basis vectors or compute efficiently all the required algebraic operations with them. Additionally, it would probably lead to illposed linear systems, since the vectors are almost linearly dependent.

However, since all these vectors arise from the same PDE (although for different parametrizations of it), and for each time step the solution is close to the previous one, in a way, we could expect there to be a lot of repeated information inside each $\Psi_{\hat{u}}(\mu_i)$, and consequently, inside $\Psi_{\hat{u}}$. We can exploit this fact by using a compression algorithm, such as the Proper Orthogonal Decomposition (POD), to find a set of vectors which sufficiently capture the span of $\Psi_{\hat{u}}$, and yet do so with less basis vectors.

3.2.1 POD Space Reduction

We define the $POD : X \to Y$ function between two normed spaces, such that $Y \subseteq X$, which takes as inputs a collection of N_S vectors $y_k \in X$, a prescribed tolerance error ε_{POD} ,

$$[\psi_i]_{i=1}^{N_{POD}} = POD\left([y_k]_{k=1}^{N_S}, \varepsilon_{\text{POD}}\right).$$
(88)

and returns a basis of modes $\psi_i \in Y$ to approximate the span of y_k . This function returns a collection of orthonormal N_{POD} vectors, whose span is a subset of the input vector span. The modes ψ_i are obtained from an optimality problem in the L^2 norm,

$$(v_i, \psi_i)$$
 s.t. $\min \sum_k \left\| y_k(x) - \sum_i v_i \psi_i \right\|^2 \le \varepsilon_{\text{POD}},$ (89)

that is, the representation of any vector in the original space can be reconstructed by the POD basis, and the error in the L^2 norm should be less or equal to ε_{POD} .

There is a relation between the prescribed tolerance error ε_{POD} and the outcoming number of modes N_{POD} ,

$$N_{POD} = N_{POD}(\varepsilon_{\text{POD}}). \tag{90}$$

This function usually shows exponential decay, or a sharp drop beyond a given number of modes. If a problem can be reduced, beyond a given number of modes, including more in the approximation will not improve significantly the approximation error.



Fig. 3. Nested POD treewalk. For each parameter μ a simulation is run, and snaphsots are collected and compressed in time (red dots). This provides a collateral basis for each value of the parameter. These bases are finally compressed into one, which summarizes the changes in time and parameters.

3.2.2 Nested POD Basis Construction

We now describe the nested POD algorithm, sketched in Figure 3. First, for a fixed parameter value, we build a POD basis from the snapshots at different time steps,

$$\Psi_{\mu_0} = POD_{\varepsilon} \left(\left[\hat{u}_h(t^0; \mu_0), \dots, \hat{u}_h(t^T; \mu_0) \right] \right),$$

$$\Psi_{\mu_1} = POD_{\varepsilon} \left(\left[\hat{u}_h(t^0; \mu_1), \dots, \hat{u}_h(t^T; \mu_1) \right] \right),$$

$$\dots,$$

$$\Psi_{\mu_{N_{\mu}}} = POD_{\varepsilon} \left(\left[\hat{u}_h(t^0; \mu_{N_{\mu}}), \dots, \hat{u}_h(t^T; \mu_{N_{\mu}}) \right] \right).$$

Then, all the μ -fixed POD basis (which sum up the information contained in the time evolution direction), are compressed again using a POD,

$$\mathbb{V} := \Psi = POD_{\varepsilon} \left(\left[\Psi_{\mu_0}, \Psi_{\mu_1}, \dots, \Psi_{\mu_{N_{\mu}}} \right] \right).$$

In the end, we obtain a basis $\Psi = [\psi_i] = \mathbb{V}$, which contains information for parameter variations and time evolution. This leads to an efficient basis, in terms of storage and numerical stability Different error tolerances could be prescribed at the time and parameter compression stages for demanding¹⁵ problems.

We call this an automatic and out-of-the-box procedure because it does not require further developments beyond the storage of the snapshots and the implementation of the POD algorithm. It only requires the construction of a collection of parameter values to solve for. This can be done with random sampling techniques, or if some physical knowledge is available, a custom selection of parameter subsets for which the solution will present strong variations (leading to the identification of richer RB solution modes).

Since we are using an SVD-based POD, we need to set a threshold for the acceptable singular values/modes. We set this threshold at 10^{-7} , since after visual inspection of the resulting modes revealed numerical noise beyond this figure.

15. Ideally, the whole POD basis is stored and then partially loaded during the online stage, according to the desired ROM accuracy. However, the offline stage might still be a memory-consuming one, which could require triming at runtime the POD basis before storing it.

3.3 (M)DEIM: System Approximation

During the offline stage (where the FOM problem is solved), we have to assemble all the discrete operators for each time step; during the online stage we additionally have to project them onto the reduced space as well, as shown in Equations (80). This projection step can mean a ROM is still costly, if it only relies on RB solution modes to reduce the problem. This will be our main motivation to include a system approximation technique, with the goal of speeding up the construction of the operators.

An essential ingredient for our system approximation is the concept of *parameter and time separable* problems (or the existence of an *affine decomposition*). This takes place when the spatial operators (bilinear or linear forms) present the following functional separable form:

$$A_h(t,\mu) = \sum_q^{Q_a} \Theta_q^a(t,\mu) A_{h,q},$$
(91)

where the coefficient functions are real-valued, $\Theta_q^a(t,\mu) \in \mathbb{R}$; and the RB operator modes $A_{h,q}$ are parameter-independent.

This expansion can be used for both matrices or vectors provided the topology of the mesh does not change. If this is the case, the matrices can be transformed into vectors by stacking the columns¹⁶, and later on brought back to matrix form once any necessary operation has been carried out.

If we had such a decomposition, once we had computed the basis matrix \mathbb{V} , we could project each mode of the operator basis $A_{h,q}$ to obtain an expression for the reduced operator,

$$A_N(t,\mu) = \mathbb{V}^T A_h(t,\mu) \mathbb{V},$$

$$= \sum_q^{Q_a} \Theta_q^a(t,\mu) \mathbb{V}^T A_{h,q} \mathbb{V},$$

$$A_N(t,\mu) = \sum_q^{Q_a} \Theta_q^a(t,\mu) A_{N,q}.$$

(92)

Since $A_{N,q}$ is fixed, provided that we had a way to evaluate each $\Theta_q^a(t,\mu)$, we would be able to build the reduced operator for a given parameter for each time step without having to use any FOM operator.

3.3.1 Discrete Empirical Interpolation Method

Naturally, not many problems are likely to present a separable form as the one shown above. Even a simple linear heat equation problem, due to the time-deformation of the mesh, cannot be presented in such a form.

To tackle this issue, we use the Discrete Empirical Interpolation Method (DEIM). This method is a numerical extension of its analytical sibling, the Empirical Interpolation Method (EIM). Basically, it mimicks the idea of creating a basis for the solution space, but this time centered around the operator space. By means of a nested POD as we explained in Section 3.2.2, if we replace the solution snapshots with operator snapshots, we can build the static and problemdependent basis $A_{h,q}$.

16. We used a CSR storage format.

Since we will be creating the operator basis with an approximation technique, an error is expected in the reconstruction of the actual operator, and so we introduce the notation $A_h^m(t,\mu)$ to reference the approximation of the operator via the (M)DEIM algorithm,

$$A_h(t,\mu) \simeq A_h^m(t,\mu) = \sum_q^{Q_a} \Theta_q^a(t,\mu) A_{h,q}.$$
 (93)

Naturally, this idea leads to the concept of approximated reduced operators,

$$A_N(t,\mu) \simeq A_N^m(t,\mu) = \sum_q^{Q_a} \Theta_q^a(t,\mu) A_{N,q}, \qquad (94)$$

which is the approximation of the reduced operators when the collateral basis has been projected unto the reduced space.

3.3.2 Evaluation of the Coefficient Functions

To evaluate the $\Theta_q^a(t, \mu)$ functions, we set and solve an interpolation problem; that is, we enforce the approximation to actually match certain entries of the operator,

$$[A_{h}(t,\mu)]_{k} = [A_{h}^{m}(t,\mu)]_{k}$$

= $\sum_{q}^{Q_{a}} \Theta_{q}^{a}(t,\mu)[A_{h,q}]_{k},$ (95)

for certain indices $k \in \mathcal{I}_a$ such that $|\mathcal{I}_a| = Q_a$. These indices \mathcal{I}_a are known as the reduced mesh nodes, sketched in Figure 4. The notation $[A_h(t, \mu)]_k$ stands for the value of



Fig. 4. Reduced mesh nodes \mathcal{I}_a sketch for $\mathbb{P}1$ Lagrangian finite element basis functions. The weak form integral evaluation only takes place for a restricted subset of mesh elements, to assemble the entries corresponding to the selected nodes. In dashed are shown the interactions with adjacent finite elements basis functions. A sketch for a 2D domain can be found in [2].

the operator at the given mesh node k. In the FE context it can be obtained by integrating the weak form locally.

This leads to a determined system, where each $\Theta_q^a(t, \mu)$ is unknown. The indices \mathcal{I}_a are selected during the offline stage, according to error reduction arguments of the reconstruction error [49],

$$e_a(t,\mu) = \|A_h(t,\mu) - A_h^m(t,\mu)\|.$$
(96)

3.3.3 Reduction of the Trilinear Term

We recall that the trilinear operator $N_h(u^*)$ from Equation (73d) takes as input in its first argument an extrapolation of the velocity u^* . This extrapolation lives in the same function space as the solution: from Equation (70), it is a linear combination of past discrete timestamps.

Therefore, two approaches can be followed to build the the trilinear operator modes,

- (*u**-general) collecting the snapshots from the FOM simulation [1];
- (*u**-restricted) collecting the snapshots from evaluations of the operator with RB solution modes.

The first approach to simply collect and compress the operator snapshots during the offline phase of the FOM, with the general nested POD algorithm described in Section 3.2.2. This would tie the operator snapshots to the parameter space sampled to identify the RB solution modes.

The second approach is to recognise that in the execution of the ROM, the first component of the trilinear term, the extrapolated solution, will always be expressed in the finitedimensional subspace of the RB solution modes (as by Equation (77)). Thus, why not use these modes to assemble the operator snapshots? Both approaches are summarized in Table 1.

The u^* -restricted strategy requires a slight modification of the nested POD algorithm. Instead of two, we have three nested levels: parameters, modes and time, as sketched in Figure 5.



Fig. 5. Nested POD treewalk for the trilinear term, u^* -restricted strategy. The parameter space is sampled, for each parameter value and RB solution mode snaphsots are collected and compressed in time (red dots). This provides a collateral basis for each parameter. These bases are finally compressed into one, which summarizes the changes in time, parameters and RB solution modes.

We set a parameter, we pick a mode, collect as many snapshots as time steps, and compress:

$$\Psi_{\mu_i,\psi_0} = POD(\{N_h^n(\psi_0)\}^{n=T}).$$
(97)

This gives us a basis for the parameter/mode tuple Ψ_{μ_i,ψ_0} . Then we pick the next mode, and obtain a basis for it, Ψ_{μ_i,ψ_1} . So on and so forth, we get as many bases as modes for that given parameter. We then compress thoses bases altogether to obtain one final parameter basis,

$$\Psi_{\mu_i} = POD(\{\Psi_{\mu_i,\psi_j}\}^{j=N}),$$
(98)

which collects the effects of each mode and time variation (for that specific parameter). We repeat the process again for another parameter, until we end up with one operator basis, which is the result of compressing all parameter bases,

$$\Psi_{N_h} = POD(\{\Psi_{\mu_i}\}^{i=N_{\mu_i}}).$$
(99)

An advantage of this procedure is that it can be used when the FOM simulation is not available, e.g. when the RB solution modes are obtained from experimental data.

TABLE 1 Trilinear operator N-MDEIM reduction approaches. The snapshots outset ("Op. assembly"), the first argument of the weak form ("1st argument"), and the treewalk pattern are compared. The restricted strategy has an additional nesting level, but it can be run as an independent step during the offline stage (provided that the RB solution modes are given).

| | Op. assembly | 1st argument | Treewalk |
|----------------------------------|-------------------|---------------------|-----------------------|
| u^* -general u^* -restricted | FOM simulation | Extrapolation u^* | Figure <mark>3</mark> |
| | Ind. offline step | RB solution modes | Figure <mark>5</mark> |

3.4 Hyper Reduced Order Model

With an approximation of the reduced operators available, we can define yet another algebraic problem to integrate and obtain the reduced solution in time for a given parametrization,

$$m_{\text{BDF}}\mathbf{M}_{N}^{m,n+1}\hat{\mathbf{u}}_{N}^{m,n+1} + \Delta t\mathbf{C}_{N}^{m,n+1}\hat{\mathbf{u}}_{N}^{m,n+1} + \Delta t\mathbf{A}_{N}^{m,n+1}\hat{\mathbf{u}}_{N}^{m,n+1} + \Delta t\hat{\mathbf{N}}_{N}^{m,n+1}\hat{\mathbf{u}}_{N}^{m,n+1} + \Delta t\left[\mathbf{N}_{N}^{m,n+1}\left(\hat{\mathbf{u}}_{N}^{n}\right)\right]\hat{\mathbf{u}}_{N}^{m,n+1} = \mathbf{F}_{\hat{\mathbf{u}}_{N}}^{m,n} + \Delta t\mathbf{F}_{g,N}^{m,n+1};$$
(100a)
$$\hat{\mathbf{u}}_{N}^{0} = \hat{\mathbf{u}}_{N,0}.$$
(100b)

If we collect terms and factor out the unknowns we get a linear system, in the reduced space and with approximated operators, to be solved for each time step to advance the solution,

$$\mathbf{K}_{N}^{m,n+1}\hat{\mathbf{u}}_{N}^{m,n+1} = \mathbf{b}_{N}^{m,n+1},$$
(101a)

$$\hat{\mathbf{u}}_N^0 = \hat{\mathbf{u}}_{N,0}; \tag{101b}$$

$$\mathbf{K}_{N}^{m,n+1} = m_{\text{BDF}} \mathbf{M}_{N}^{m,n+1} + \Delta t \left[\mathbf{A}_{N}^{m,n+1} + \mathbf{C}_{N}^{m,n+1} + \hat{\mathbf{N}}_{N}^{m,n+1} + \mathbf{N}_{N}^{m,n+1} \left(\hat{\mathbf{u}}_{N}^{m,*} \right) \right], \qquad (101c)$$

$$\mathbf{b}_{N}^{m,n+1} = \mathbf{F}_{\hat{\mathbf{u}}_{N}}^{m,n} + \Delta t \mathbf{F}_{g,N}^{m,n+1}.$$
 (101d)

Each of the operators present in the problem will have associated an operator basis and will require the solution of the interpolation problem (95) for each time step and parameter value. Although this could still seem like a costly procedure, if the operators are actually reduceable, the number of basis functions $Q_m, Q_a, Q_f, Q_{f,g}$ should be small, and thus way simpler problems than assembling the whole operator and then carrying out the projection.

Once the reduced homogeneous solution $\hat{u}_N^m(x)$ is obtained with approximated operators, it can be brought back

to the original space V_h via Equation (78), and then added to the Dirichlet lifting, to get the solution in the physical domain,

$$u(x,t;\mu) \simeq u_h^m(t,\mu) = \mathbb{V}\hat{u}_N^m(t,\mu) + g_h(t,\mu).$$
 (102)

4 FOM CALIBRATION

This section is devoted to present the full order model discretization details. Certain parameters need to be defined, along with consistency checks to validate the simulation.

Sections 4.1 and 4.2 define the parametrization space and show some results. In Section 4.3 we show the effects of artificial viscosity, in Section 4.4 we explore the differences between BDF-1 and BDF-2 time integration schemes.

4.1 Parameter Space and Discretization

To identify the reduced and collateral bases, we randomly sample the parameter space. We set the range for each parameter in Table 2. The discretization parameters are given in Table 3.

TABLE 2 Parameter range for random sampling. Note that mesh parametrizations which produce an invalid mesh are discarded at runtime.

| Variable | Min. | Max. |
|--|--------------------|---------------------|
| $egin{array}{c} a_0 \ \omega \ \delta \end{array}$ | 18 15 0.15 | 25 30 0.3 |
| $egin{array}{c} x_c \ \sigma_c \ y_c \end{array}$ | 0.2 0.1 0.25 | 0.75 0.2 1.75 |

TABLE 3 Space-time domain definition, and discretization parameters.

| L_0 | N_x | T | N_t | Δx | Δt |
|-------|-------|-----|-------|----------------|--------------------|
| 1 | 1000 | 1.0 | 500 | $\sim 10^{-3}$ | $0.5\cdot 10^{-3}$ |

There are two relevant dimensionless groups for this problem, presented in Table 4. Dimensionless groups are useful to sample the parameter space wisely, probing different dynamical scenarios rather than different parametrizations which actually represent the same kind of response. For our sampling during the offline stage (and online testing), we determined that taking into consideration the piston mach u_p was sufficient to collect physically distinct system responses.

TABLE 4 Relevant dimensionless groups. Note that δ is a dimensionless scale parameter.

| Name | Expression | Min. | Max. |
|----------------------------------|---------------------------------|------|------|
| Reduced frequency $\hat{\omega}$ | $\frac{\omega L_0}{a_0}$ | 0.6 | 1.65 |
| Piston mach u_p | $\frac{\delta \omega L_0}{a_0}$ | 0.1 | 0.4 |

4.2 FOM Response to Piston Motion

We present the response of the system for a piston mach value $u_p = 0.37$. The dimensionless velocity profile is shown in Figure 6; the mass conservation check is examined in Figure 7. The PDE was derived using the mass conservation principle, but the mass conservation integral

equation has not been used in the discretization of the weak form. Hence it is a good independent quality check of our simulation results.



Fig. 6. Flow response (blue) to piston motion (dashed). (Top) Piston outlet, artificial boundary condition. (Bottom) Piston mid-length. A weak shock wave is formed due to Burgers' nonlinear convection.



Fig. 7. Mass conservation check. (Top) The two resulting summands from the mass conservation integral equation: density integral derivative in time and outlet boundary mass flow. The weak shock wave is also present in the mass conservation. (Bottom) Numerical value of the mass conservation principle, coined mass defect MD. The mean value (red line) is the corresponding y-axis value in the convergence plot in Figure 10.

4.3 Artificial Viscosity

As stated previously in the document, we included an artificial viscosity term in the final formulation to go around the need for more involved¹⁷ stabilization schemes. A value needs to be choosen for the viscosity constant ε . For con-

17. Note that the reduction approach used remains valid, since it is purely algebraic.

sistency with the PDE, this constant should scale with the square of the mesh size,

$$\varepsilon \sim k \cdot (\Delta x)^2$$
.

For a mesh size of $\Delta x \sim 10^{-3}$, the *k*-value which keeps the system stable whilst purely convective (at least for our simulation time scale) is

$$k = 10^{-4} \to \varepsilon \sim 10^{-10}$$

In Figure 8 we present a comparison for these two viscosity values. We show the motion of the fluid at the outflow and a phase plot, to ease the analysis of the correlation between the weak shock wave at the outflow and the piston motion. The smallest viscosity value, (10^{-10}) leads to a convection-like phase plot: the shape of the input is distorted but the maximum and mimimum values are honored, the same path is repeated over and over. Instead, the solution with a higher viscosity value shows a diffusion pattern, reducing the extrema and changing its path on every pass.



Fig. 8. Artificial viscosity comparison. (Top) Outer boundary velocities for two different values of the viscosity term. In dashed, the piston motion, to help in the visualization of the nonlinear distortion. (Bottom) Phase plot between the piston motion (x-axis) and the response at the outer boundary (y-axis) for the same artificial viscosity values. Due to the creation of a weak shock wave, the piston motion is distorted. The viscosity value (10^{-10}) presents a stable phase plot, going over and over the same path.

4.4 BDF Convergence Rates

To check the quality of our simulations we examine the convergence rates for the two time-integration schemes, BDF-1 and BDF-2. We analyze two convergence plots: the solution itself and mass preservation.

For the solution, since we do not have an analytical reference, we establish one numerically. It will be the one corresponding to the solution for a small time step, $\Delta t = 10^{-4}$. In Figure 9 we present the convergence rates for the solution, where the BDF-2 scheme (1.98 \sim 2) decreases twice as fast as the BDF-1 scheme (0.95 \sim 1).

For the mass defect, we do know it should tend to zero analytically. In Figure 10 we give the convergence rates for the mass defect, where the BDF-2 scheme ($2.08 \sim 2$) is seen to converge twice as fast as the BDF-1 scheme ($1.24 \sim 1$). Hence, the BDF-2 scheme is correctly implemented.



Fig. 9. Convergence rates to numerical reference solution. The reference was obtained with a small time step, $\Delta t = 10^{-4}$. Both schemes decrease at their expected rates.



Fig. 10. Convergence rates for mass defect MD. The analytical solution of this variable is zero (mass is perfectly preserved). Both schemes decrease at their expected rates.

4.5 Non-Uniform Mesh Displacement

Not all geometrical parametrizations of the mesh displacement are valid. As shown in Table 5, some parametrizations lead to non-invertible mappings. This takes place when the displacement is so large locally that the mesh nodes ordering is lost in the physical domain. When this happens, two points in the reference domain map to the same point in the physical space.

TABLE 5 Mesh deformation parametrizations.

| | δ | x_c | σ_c | y_0 | Invertible |
|------------------------|-----|-------|------------|-------------|------------|
| Figure 11 Figure 12 | 0.3 | 0.5 | 0.1 | 0.5 0.75 | Yes |
| Figure 13 | | | | 1.75 | No |

The correct procedure to deal with the mesh feasibility problem would be to compute the Jacobian analytically and derive from it upper and lower bounds for each parameter. However, for such a simple domain transformation, we opt to check numerically at runtime if the mesh is feasible or not. In fact, to prevent round-off errors, we impose a lower bound on the mesh step size. When it is breached,

$$\forall i \quad \Delta x_i < 10^{-6}, \tag{103}$$

we discard that parametrization.



Fig. 11. Feasible mesh compression. The green line shows the maximum piston compression. The nodes are locally compressed to the left of the Gaussian curve.



Fig. 12. Feasible mesh expansion. The green line shows the maximum piston expansion. The nodes are locally compressed to the right of the Gaussian curve.



Fig. 13. Unfeasible mesh expansion. Due to the large mesh distortion, the ordering is lost in the physical domain, leading to negative mesh step sizes.

4.6 Geometric Conservation Law

As mentioned in the literature review regarding deforming meshes, in Section 1.3.2, a proxy to determine how affected is a discretization by a moving mesh is to attempt the resolution of the constant solution.

This is so because if the discretization is not handling the movement of the mesh correctly, artificial fluxes of information might be introduced. Nevertheless, no strict conclusions can be reached with this numerical test. Succesfully integrating the constant solution seems to be a sufficient but not a necessary condition for stability.

Our implementation seems unable to reproduce correctly the constant solution. This happens in a fixed and moving mesh setting (see Figure 14). Therefore, we believe it could be due to the accumulation of round-off errors, as previous works have reported in the literature [55].

In a way, we are surprised that this blowing-up effect does not show up in the piston problem. We hint towards the fact that, in that context, the constantly changing boundary conditions somehow lead to balanced round-off errors, which cancel out in time. In the constant solution test, the boundary condition is constant, thus potentially preventing round-off error cancellation.

Chasing to the detail this behaviour is certainly meaningful, but since our reference solution (the piston) does not suffer from these effects, we limit ourselves to report this behaviour and skip any further investigation, as we believe it falls out of the scope of this work.



Fig. 14. Constant solution simulation for fixed (top) and moving mesh (middle, bottom). The round-off error increases as the number of DoFs increases too (N_x). The errors in the fixed mesh take more to accumulate or do not accumulate at all. For the moving mesh, for the smallest number of DoFs the solution drifts away from the constant solution. In the bottom plot, the values have been clipped to the [0.01, 10] interval.

5 **RESULTS AND CERTIFICATION**

We are now going to see the HROM in action. This section is divided in two conceptual blocks, summarized in Table 6.

The first block will show HROM results with a uniform mesh stretching. This scenario is interesting to establish certain procedures and results, but due to its simplicity, the reduction of the bilinear operators is trivial (limiting the range of the discussion). Section 5.2 shows the expected error decay of the standalone RB solution modes; Section 5.3 shows the results of the u^* -general and u^* -restricted approaches to deal with the N-MDEIM of the trilinear form. An unexpected behaviour is present in the singular value decay for the u^* -restricted approach, which we discuss and clarify in Section 5.4. Finally, in Section 5.5 we show how to certify the HROM solution by mode truncation.

To compensate for the simplicity of the bilinear operator reductions, we introduce a second block, where the HROM will be applied in the presence of a non-uniform mesh stretching (all operators then require a non-trivial number of RB operator modes). In Section 5.6 we present u^* -general reduction results. In Section 5.7 we present the limits of mode truncation for HROM error estimation in this context. Finally, in Section 5.8 we show the improvement and limits of the u^* -restricted approach.

TABLE 6 Summary table with the FOM and HROM characteristics for each section. Certain sections contain results without the (M)DEIM methodology activated, to isolate effects.

| Mesh Stretching | Section | (M)DEIM | N-MDEIM Approach |
|-----------------|--------------------------|------------------------|--|
| Uniform | 5.2 5.3 5.4 5.5 | No Yes Yes No | - general / restricted restricted - |
| Non-Uniform | 5.6 5.7 5.8 | Yes | general general restricted |

5.1 RB Solution Basis

The piston is moving according to a harmonic law, and we have seen in Figure 6 from the previous section how the flow response is periodic and nonlinear. Additionally, the solution is stretched in time with the boundary movement. We expect to find these effects in the RB solution modes.

We show the first ten RB solution modes obtained by the POD in Figure 15. They are represented in the reference domain with $L_0 = 1$. The first five modes contain some sort of oscillatory response near the right boundary, where the piston movement is located, but they are not harmonic at all. The next five modes look similar to Fourier modes, which is an expected behaviour given the harmonic movement of the piston and the fact that we sample for different ω values during training. The forcing of the piston at the right boundary can be guessed. The first five modes capture the nonlinear response, whereas higher order modes present the oscillatory nature of the solution.

We do not represent the RB operator bases because their interpretation would be too abstract, given that the operator itself does not have a simple representation in space.



Fig. 15. First ten RB solution modes (0-indexing is used to be aligned with code results). (Top) First five modes. (Bottom) Next five modes. The first modes seem to collect the effects of the weak shock wave formation. The next five modes look like Fourier modes for different frequencies (which is an expected behaviour given the harmonic movement of the piston and the fact that we sample for different ω values during training).

5.2 RB Solution Basis Error

We explore how the number of modes influences the approximation error of the FOM solution. We have trained our model with 10 random samples according the parameter ranges given in Tables 2 and 4. In Table 7 the resulting bases size are reported. Since the stretching is uniformly spread across the mesh elements, despite their change in time, all the linear operators are trivially reduced. The RHS requires two RB operator modes because it contains the lifting and its gradient. Since we are using the POD to build

TABLE 7 For each operator, basis size after the tree walk and final size after tree walk compression. The trilinear term and the RB space have the same size, since the RB solution modes were used to evaluated the trilinear term. The Nonlinear-lifting operator corresponds to the cross-term lifting matrix that arises from the nonlinear convective term.

inting matrix that anses nom the nominear convective term

| | After Treewalk | Final Size |
|-------------------|----------------|------------|
| RB | 295 | 69 |
| RHS | 20 | 2 |
| Mass | 10 | 1 |
| Stiffness | 10 | 1 |
| Trilinear | 690 | 69 |
| Convection | 20 | 2 |
| Nonlinear-lifting | 10 | 1 |
| | | |

our reduced basis, we attempt to relate POD a posteriori error estimations with the actual error between ROM and FOM.

5.2.1 POD: A Posteriori Errors

The POD returns a collection of modes and sigular values, $(\psi_i(x), \sigma_i)$, each related to the other. The magnitude of each singular value σ_i somehow encodes how much information is carried by its associated mode about the original span.

We define the energy of the basis that contains up to the *i*-th mode as

$$\mathcal{E}_i = \frac{\sum_{j=0}^i \sigma_j^2}{\sum \sigma_k^2}.$$
(104)

This magnitude is derived from the *a posteriori* error bounds for a POD reconstruction, where the following inequality holds,

$$\left\| y(x) - \sum_{j=0}^{n} w_j \psi_j(x) \right\|_2^2 \le \sum_{i=n+1}^{N} \sigma_i^2,$$
(105)

with y(x) belonging to the snapshots used to build the POD basis. This inequality is telling us that if up to *n* basis modes are used to reconstruct a function belonging to the original span, the average error in the L_2 sense will be smaller or equal to the sum of the remaining squared singular values.

We shall see if this variable has any predictive power on the ROM error.

5.2.2 Error Decay

We want to find the smallest size N^* for which the FOM is correctly reproduced. Naturally, the exact value of this variable is problem-dependent, but the way in which we approach its search would suit any RB problem.

TABLE 8 Online sampling parameters, sorted in ascending order by piston mach u_p values.

| a_0 | ω | δ | u_p |
|-------|-------|------|-------|
| 22.96 | 29.55 | 0.15 | 0.20 |
| 19.28 | 22.87 | 0.20 | 0.23 |
| 18.24 | 18.88 | 0.29 | 0.30 |
| 24.64 | 27.13 | 0.29 | 0.32 |
| 20.62 | 25.98 | 0.29 | 0.37 |

In Figure 16 we present the ROM error with respect to the number of RB solution modes and the energy of the truncated basis. At this stage, the (M)DEIM approximation is not used (the FOM operators are projected at each stage). The more RB solution modes, the smaller the error. The more



Fig. 16. Error decay vs. the number of RB solution modes and basis energy level. Expectedly, the more RB solution modes we have, the lower the error.

energy the basis contains, the better the approximation.

We present in Figures 17 and 18 the actual solution at the outflow for each model (FOM and ROM). We can see how the ROM model is not accurately resolved for N = 5, as reflected by the error, but how it improves for N = 10. Clearly, the problem is quite simple to reduce if these few modes are sufficient to reduce the problem. The flow



Fig. 17. Outflow velocities for different models, for N = 5 RB solution modes. It is inaccurately resolved at the beginning of the transient and some diffusion is introduced, reducing the extrema values.



Fig. 18. Outflow velocities for different models. The ROM model is better resolved for ${\cal N}=10~{\rm RB}$ solution modes.

departs from rest as the piston starts to move. Therefore, the first snapshots contain a nonlinearity in the sense that the flow starts to move on one side, but remains still along the rest of the tube.

However, for most of the time interval, this kind of nonlinearity is not present again, since the flow is in constant motion, flowing in and out of the system. Hence, it is an expected behaviour that for the initial interval the ROM is inaccurate, unless we include a sufficient number of RB solution modes. To fix this problem, certain techniques exist which enhance the POD basis without adding excessive complexity [56].

5.3 N-MDEIM Basis Error

In this section we explore the reduction of the trilinear operator for the mesh with uniform stretching.

In Section 5.3.1 we present the results for the basis resulting from the snapshots collected from the FOM simulation. In Section 5.3.2 we investigate the interaction between the RB and MDEIM errors. Finally, in Section 5.3.3 we present the results for the snapshots resulting from the snapshots collected from RB mode evaluation in the trilinear form.

We refer to the collateral basis of the trilinear operator as the N-MDEIM basis.

5.3.1 N-MDEIM: Hierarchical Basis

We present the results for the u^* -general strategy, for the uniform stretching mesh displacement. During the simulation of the FOM, we collect snapshots of the trilinear term and compress them with the same nested POD strategy as we did for the reduced basis, Section 3.2.2.

By doing so, we obtain a hierarchical basis, which allows us to tune the error, as proved by the singular value decay (Figure 19). The singular value decay for the trilinear operator presents the same pattern as the one for the solution. The reduction difficulty is the same as the one required for the reduced basis.



Fig. 19. Singular values decay for the reduced basis and the trilinear operator. The dashed vertical line corresponds to the number of modes, N=69. The basis for the trilinear operator presents the same pattern (and hence the same size) as the solution reduced basis. This is due to the fact that the operator is a trilinear form. Hence, due to the simplicity of the Jacobian, the reduction difficulty is similar to the one required for the reduced basis.

In Table 9 we show the parameter range for which we tested this basis against the first ten modes. In Figure 20 we present the reconstruction error decay, as the N-MDEIM basis is truncated. The basis shows the expected error decay as modes are removed, hence allowing to speed up even further computations if some compromise is made in the error.

TABLE 9 Parameter space sampling for trilinear term reconstruction.

| a_0 | ω | δ | u_p |
|--------|--------|-------|-------|
| 24.275 | 18.549 | 0.203 | 0.155 |
| 21.055 | 15.969 | 0.236 | 0.179 |
| 23.627 | 24.427 | 0.211 | 0.218 |
| 21.533 | 24.376 | 0.200 | 0.226 |
| 20.440 | 18.913 | 0.289 | 0.267 |
| 23.849 | 27.396 | 0.261 | 0.300 |
| 19.976 | 23.190 | 0.259 | 0.301 |
| 20.633 | 24.184 | 0.280 | 0.329 |
| 18.803 | 29.705 | 0.226 | 0.357 |
| 19.890 | 27.986 | 0.278 | 0.391 |
| | | | |



Fig. 20. N-MDEIM reconstruction error decay, as we truncate the modes from the MDEIM basis for some individual parametrizations. Due to the simplicity of the problem, the (implicit) Jacobian and the fact that the linearized term is trilinear in all of its arguments, there is no noticeable difference for different parametrizations.

5.3.2 RB and N-MDEIM Error Interaction

With two bases (solution and trilinear operator) which show error decay, we can truncate them simultaneously, to determine their interaction. In Table 10 we present the timewise L_2 error for different numbers of RB solution and RB trilinear modes. In Figure 21 we present the same table, albeit in a heatmap plot.

Expectedly, we reach a situation where one error dominates the other one. Increasing the size of one basis will not improve the results, unless the other one is enlarged too. If we wanted to achieve a determined error threshold, we would have to analyze and take into account both bases errors simultaneously.

5.3.3 N-MDEIM: Non-Hierarchical Basis

Now we see what happens if we build the MDEIM basis from trilinear snapshots assembled with RB solution modes $(u^*$ -restricted).

The first ten RB solution modes are used to assemble the operator for the ten parameters from Table 9. In Figure 22 we show the mean approximation error in time for the trilinear operator on these same RB solution modes. We see that unless the complete basis is used (N = 69), the approximation error is quite poor.

TABLE 10 Timewise L_2 error decay for different truncation levels of the reduced basis space (N) and the trilinear operator basis (N-MDEIM).

| N-MDEIM | 5 | 10 | 15 | 30 | 40 | 50 |
|---------|--------|--------|--------|--------|--------|--------|
| N | U | 10 | 10 | 00 | 10 | 00 |
| IN | | | | | | |
| 5 | 3 40-2 | 3 40-2 | 3 40-2 | 3 40-2 | 3 40-2 | 3 40-2 |
| 5 | 0.40-2 | 0.40-2 | 0.40-2 | 0.40-2 | 0.40-2 | 0.40-2 |
| 10 | 3.4e-3 | 7.8e-4 | 7.8e-4 | 7.8e-4 | 7.8e-4 | 7.8e-4 |
| 15 | 3.3e-3 | 2.1e-5 | 1.4e-5 | 1.4e-5 | 1.4e-5 | 1.4e-5 |
| 20 | 3.3e-3 | 1.7e-5 | 8.0e-7 | 7.8e-7 | 7.8e-7 | 7.8e-7 |
| 25 | 3.3e-3 | 1.7e-5 | 2.8e-7 | 1.5e-7 | 1.5e-7 | 1.5e-7 |
| 30 | 3.3e-3 | 1.7e-5 | 2.5e-7 | 5.1e-8 | 5.3e-8 | 5.3e-8 |
| 35 | 3.3e-3 | 1.7e-5 | 2.5e-7 | 4.5e-8 | 4.6e-8 | 4.6e-8 |
| | | | | | | |



Fig. 21. Timewise L_2 error decay for different truncation levels of the reduced basis space (N) and the trilinear operator basis (N-MDEIM). The log_{10} transformation was applied, so that the heatmap color gradient is smooth.

This has to do with the fact that the RB solution modes, which form an orthogonal basis, were used to assemble the trilinear operator snapshots. Hence, all the result RB operator modes contain fundamental information, as hinted by the abrupt drop in the singular value, see Figure 23. We have obtained a basis that allows us to reduce the assembly of the operator, but it doesn't allow us to tune the error. We name such basis *non-hierarchical*, because there is no decay in its singular value spectrum. We will clarify the loss of hierarchy in Section 5.4.



Fig. 22. N-MDEIM error approximation on the first ten RB solution modes for different collateral basis size. The first ten RB solution modes were used to assemble the operator. The approximation is quite poor, unless the complete basis is used (N = 69).



Fig. 23. Singular values decay for the reduced basis and the trilinear operator. The dashed vertical line corresponds to the number of reduced modes, N = 69. The singular value decay for the trilinear operator does not present the same pattern as the one for the reduced basis. All RB trilinear operator modes seem to contain the same amount of information from the original span. This has to do with the fact that the modes, which form an orthogonal basis, were used to assemble the trilinear operator snapshots.

5.4 Non-Hierarchical POD Bases

We investigate further the non-hierarchical character of the N-MDEIM basis that showed up in Section 5.3.3. After we discretized the nonlinear term we obtained a trilinear form, whose first argument is the velocity extrapolation.

To build a basis in the u^* – *restricted* approach, we obtain the snapshots from evaluations of the trilinear operator with RB solution modes,

$$\mathbf{N}_{h}^{n+1}(\psi_{h})]_{i,j} = b_{0} \langle \psi_{h}(x) \nabla \varphi_{j}, \varphi_{i} \rangle^{n+1},$$

$$= b_{0} \int_{0}^{L(t^{n+1})} \psi_{h}(x) \nabla \varphi_{j} \varphi_{i} \mathrm{d}x.$$
(106)

Now, the RB solution modes are expressed themselves in terms of the finite element Lagrangian basis functions,

$$\psi_h(x) = \sum_k \psi_k \varphi_k(x), \tag{107}$$

which leads to the following exact expression for each of the entries of the trilinear form,

$$\left[\mathbf{N}_{h}^{n+1}\left(\psi_{h}\right)\right]_{i,j} = b_{0} \sum_{k} \psi_{k} \underbrace{\int_{0}^{L(t^{n+1})} \varphi_{k} \nabla \varphi_{j} \varphi_{i} \mathrm{d}x}_{\text{Only affected by } J(x,t)}.$$
 (108)

Each summand takes a nonzero value only when the local support of the three basis functions coincides simultaneously.

For the assembly of the trilinear operator during the FOM integration, the extrapolated velocity u^* is also expressed within the Lagrangian basis (as shown in Equation (70), it is a linear combination of the two previous time steps). And yet, the snapshots compressed from that source lead to a hierarchical basis. Therefore, the interaction between the finite element basis functions is not the effect leading to a non-hierarchical basis.

The remaining degrees of freedom are the coefficient values ψ_k . Since the solution is smooth, when the snapshots are collected during the FOM simulation, each of the u_k^* values is quite similar to the ones from the previous time step. However, when the modes are used to create the trilinear operator snapshots, for any two different basis modes $\psi_1(x)$ and $\psi_2(x)$, we know that they satisfy an orthogonality condition. To determine if this condition is playing a role, we remove the complexity of the reduced basis construction (application of the POD on the solutions from the PDE); and consider an example using functions from a naturally orthogonal basis: the Fourier basis.

Our working hypothesis is that if the functions used to evaluate the trilinear form are orthogonal, so will be the resulting POD base, and hence non-hierarchical (there is no reduction capacity, since the input is already forming a base).

5.4.1 Fourier Basis: Insights

We kick-off by studying the effects of attempting an SVD over a Fourier basis,

$$\psi_q = \{\cos(2\pi\omega_q x) : \omega_q \in \{1, \dots, 3\}\}.$$
 (109)

We carry out two experiments:

- (I) Run the Fourier basis through the SVD and analyze its spectrum decay. We would expect no decay, since the basis is orthogonal¹⁸.
- (II) Compute snapshots of the trilinear operator with the Fourier basis and run them through the SVD as well. The spectrum decay will be used to establish conclusions.

18. It the input modes are normalized with the L_2 norm, the singular values will be equal to one. Else, the singular values will be equal to the L_2 norm.

5.4.2 (I) SVD of a Fourier Basis

When the POD is applied to a basis of orthogonal modes, the resulting basis is potentially¹⁹ formed by linear combinations of the original modes.

Our implementation of the POD is based on the SVD breakdown, which splits a matrix into two rotations and one scaling,

$$A = U\Sigma V^{T}.$$
 (110)

If the matrix A is orthogonal²⁰, we would expect that the transformation is such that the modes remain unaltered,

$$U = A, \Sigma = I, \text{ and } V = I$$

(with their respective dimensions well set). However, the only guarantee we actually have is that $\Sigma = I$, the rotations U and V are not unique. And so is the case with the implementation we are using of the SVD.

Instead of returning the original Fourier modes, it is creating linear combinations of it, where the coefficients of the linear combination are collected in the columns of

$$V = \begin{bmatrix} 0.577 & 0 & 0.816\\ 0.577 & 0.707 & -0.408\\ 0.577 & -0.707 & -0.408 \end{bmatrix}.$$
 (111)

There is no spectrum decay, for all singular values are equal to 1.0, as shown in Table 11.

 TABLE 11

 Singular value spectrum for the SVD of the Fourier basis. Since the input is orthogonal, the resulting POD basis is non-hierarchical. To be aligned with the code, a 0-indexing presentation is used.

| Singular index | σ_i |
|----------------|----------------|
| 0 | 1.002 |
| 1 | 0.999 |
| 2 | 0.999 |
| 1 2 | 0.999 0.999 |

POD Basis Fourier Basis 0.08 First POD mode 0.06 0.04 0.02 Э 0.00 -0.02 -0.04 Linear comb. of Fourier modes 0.0 0.2 0.4 0.6 0.8 1.0 0.0 0.2 0.4 0.6 0.8 1.0

Fig. 24. Fourier basis and its SVD transformation. Since the Fourier basis is orthogonal, the resulting SVD transformation is a linear combination of Fourier modes.

19. This is an implementation-wise feature. 20. $AA^T = A^T A = I$

5.4.3 (II) N-MDEIM with Fourier Basis

To confirm our working hypothesis, we build the trilinear form (Equation (108)) using the first five modes of the Fourier basis,

$$\psi_q(x) = \{\cos(2\pi\omega_q x) : \omega_q \in \{1, \dots, 5\}\}.$$
 (112)

Since the Jacobian transformation is a linear function of time, despite the change in time of the domain, the trilinear form remains linear. If there were no additional argument ψ_k , we would only require one mode to represent the whole operator. Therefore, we predict that we will only find 5 orthogonal resulting modes, that is, a non-hierarchical basis. This is confirmed by the singular value decay, shown in Table 12 (to be aligned with the code, a 0-indexing presentation is used).

TABLE 12

Singular value spectrum for the N-MDEIM reduction with a Fourier basis. We have built the snapshots in time for two parameters and five Fourier modes. Since we know the resulting bases will be of the same size as the original input, we expect to have only 5 non-zero singular values, which is the case. To be aligned with the code, a 0-indexing presentation is used.

| Singular index | σ_i |
|----------------|------------|
| 0 | 1.414 |
| 1 | 1.414 |
| 2 | 1.414 |
| 3 | 1.414 |
| 4 | 1.414 |
| 5 | 0.000 |
| 6 | 0.000 |
| 7 | 0.000 |
| 8 | 0.000 |
| 9 | 0.000 |

Again, as depicted by Figure 25, we obtain the same poor results in the approximation error when we attempt to reconstruct the operator with a truncated basis.



Fig. 25. N-MDEIM reconstruction error. The basis for this N-MDEIM has been built using the first five Fourier modes to evaluate the trilinear form. According to the singular value decay, the basis is not hierarchical, that is, all the operator modes (continuous) need to be used to recover an accurate representation of the space. Hence, when one operator mode is removed (dashed), the approximation error increases to unaffordable values for a simulation.

Now that we have been able to reproduce the same behaviour, and to validate our working hypothesis, we go back to the use of RB solution modes.

5.4.4 N-MDEIM with RB Solution Modes

What would happen if we only used one mode from the RB solution modes to reduce the operator? Could we use that basis to approximate the trilinear operator evaluated with other functions? We take the first three RB solution modes and create a linear combination f(x) with them,

$$f(x) = 2\psi_0(x) + \psi_1(x) + 3\psi_2(x).$$
(113)

We now reduce the trilinear form with two strategies:

- (a) Subspace: using only the first RB solution mode.
- (b) Complete space: using the three RB solution modes.

With each approach, we obtain a non-hierarchical orthogonal basis, which we then use to reconstruct $N_h(f(x))$. Since f(x) is made out of the three orthogonal modes, we could expect approach (a) to fail. However, it succeeds as well as approach (b). This is proved by Figures 26 and 27. Additionally in Figure 27 we have included the reconstruction error for the truncated base (we have removed operator mode).

Using *one* RB solution mode is sufficient to construct a perfect collateral basis due to the linearity of the integrand within the trilinear form, and the linearity of the Jacobian transformation. Despite the fact that the function f(x) contains other modes, the collateral basis built with one mode is able to reproduce its effect on the operator perfectly. In



Fig. 26. Reconstruction error for the trilinear operator evaluated with linear combination f(x). The basis has been obtained only using the first RB solution mode.

the next section we explore what happens if a nonlinear function is present in the integrand.



Fig. 27. Reconstruction error for the trilinear operator evaluated with linear combination f(x). The basis has been obtained using the three RB solution modes. In dashed is the approximation error using a truncated basis (N = 2 operator modes). The truncated basis shows a poor approximation error because the original basis is non-hierarchical, hence all the operator modes are required to reconstruct the operator.

5.4.5 N-MDEIM with a Nonlinearity in Space

We have shown in the previous section how one can achieve a perfect collateral basis for the trilinear using exclusively one RB solution mode.

We now show results for a form with a nonlinear function inside the weak form, e.g., a non-uniform mesh displacement. Hence, we take on the reduction of

$$\begin{bmatrix} \mathbf{W}_{h}^{n+1}(\psi_{h}) \end{bmatrix}_{i,j} = b_{0} \left\langle \psi_{h}(x) \cos(1+x) \nabla \varphi_{j}, \varphi_{i} \right\rangle^{n+1},$$
$$= b_{0} \int_{0}^{L(t^{n+1})} \psi_{h}(x) \cos(1+x) \nabla \varphi_{j} \varphi_{i} dx.$$
(114)

The results are shown in Figures 28 and 29. Both figures include the reconstruction error for a truncated base.

We find out that in the presence of a nonlinearity, it is unsufficient to use only one RB solution mode. When the basis built with approach (a) is truncated, the approximation error goes up to values near unity, leading to poor reconstruction results.

Instead, when the first three RB solution modes are used to build the collateral basis, truncating the outcome leads to a feasible basis. The truncation leads to a higher approximation error, but still sufficiently small to produce meaningful results if it were used in a simulation.



Fig. 28. Reconstruction error for the nonlinear operator evaluated with function f(x). The basis has been obtained only using the first RB solution mode ($N_{\text{full}} = 7$). In dashed is the approximation error using a truncated basis, from which only one operator mode has been removed ($N_{\text{truncated}} = 6$). The truncated basis fails, due to the presence of the nonlinearity, it is now unsufficient to use only one RB solution mode space.



Fig. 29. Reconstruction error for the nonlinear operator evaluated with function f(x). The basis has been obtained using the three RB solution modes ($N_{\rm full} = 19$). In dashed is the approximation error using a truncated basis, from which several operator modes have been removed ($N_{\rm truncated} = 10$). The truncated basis does not fail as drastically as the previous ones did, it simply presents a higher error, but still within an acceptable order of magnitude.

5.4.6 Wrap-Up

We have clarified the behaviour of the POD in the presence of a trilinear form and a linear Jacobian transformation.

Our first conclusion was that the POD of an orthogonal basis will produce another orthogonal basis, as a linear combination of the input vectors. This orthogonality is preserved under the linear transformation of the trilinear form. As a consequence, the construction of the collateral basis can be achieved collecting snapshots with the evaluation of the trilinear form with only one RB mode. If more RB solution modes are used to build the snapshots, the POD will return a larger collateral basis, but one that will not have the hierarchical property. That is, truncating that basis will consistently lead to poor approximation results. The full basis will have to be used. This is so because the SVD has created a linear combination of the orthogonal input snapshots.

In the presence of a non-uniform mesh movement, the reduction trick of using one RB solution mode is unsufficient. More RB solution modes have to be used to obtain a collateral basis that appropriately spans the whole domain. Then, the basis is hierarchical, and allows for error control through basis truncation.

If we collect snapshots of the operator evaluated with the PDE solutions, we couple the reduction of the RB space and the operator space. Hence, it is best to reduce trilinear forms with RB solution modes. If the trilinear form does not contain any spatial or time nonlinearity, one RB solution mode will be enough. Instead, if nonlinearities are present within the integrand, more RB solution modes will be necessary to build a satisfactory basis. One way or the other, this approach will reduce the number of snapshots to collect and compress, leading to a lighter offline stage.

5.5 Certification by Truncation

To compute the HROM error in the preceding sections we required the assembly and solution of the FOM model. This is an undesirable fact, since if to validate our ROM solution we need to compute its FOM counterpart, we are better off computing the FOM model straightaway (which will be more accurate by definition). Hence, we need an efficient a posteriori error estimator.

5.5.1 Sacrificial ROM

We use the mode truncation technique. That is, we carry with us *two* ROMs, with different basis sizes. The additional ROM is called *sacrificial* ROM (SROM). It will be used to compute the actual error for the smaller ROM, without requiring the calculation of the FOM. By doing so, we bound the SROM error. We assume it is smaller than the one of base ROM, since it contains more modes, but we cannot know how much smaller it is. Hence, we would bound the SROM errors by using a smaller ROM, but would use the SROM to calculate and present results.

For this error estimation procedure, the question we seek to answer is: how many more modes does the SROM need to carry to estimate accurately the base ROM error? Again, the answer to this question is problem-dependent, but our approach to find it could be used for any problem.

5.5.2 Error Estimation

We depart from the error with respect to the FOM, which we seek to bound. Then, we proceed to add and subtract the solution from the sacrificial ROM,

$$\|u_h - \mathbb{V}_N u_N\| = \|u_h \pm \mathbb{V}_{\hat{N}} u_{\hat{N}} - \mathbb{V}_N u_N\|, \qquad (115)$$

where $\hat{N} > N$. By the triangle inequality, we get

$$\|u_{h} - \mathbb{V}_{N}u_{N}\| \leq \|u_{h} - \mathbb{V}_{\hat{N}}u_{\hat{N}}\| + \|\mathbb{V}_{\hat{N}}u_{\hat{N}} - \mathbb{V}_{N}u_{N}\|.$$
(116)

If we define the error function $e_N = ||u_h - \mathbb{V}_N u_N||$, we get an upper bound of the desired error in terms of the sacrificial ROM error and the error between ROMs,

$$e_N \le e_{\hat{N}} + \left\| \mathbb{V}_{\hat{N}} u_{\hat{N}} - \mathbb{V}_N u_N \right\|. \tag{117}$$

With sufficient RB solution modes, it should be safe to assert that the error made with additional modes should be smaller or equal to the one made without it,

$$e_{\hat{N}} \le e_N. \tag{118}$$

Thus, we get the following error bound,

$$e_N \le \left\| \mathbb{V}_{\hat{N}} u_{\hat{N}} - \mathbb{V}_N u_N \right\|,\tag{119}$$

It remains to determine how sharp this error estimate is.

5.5.3 Error Between Two ROMs

The error estimator

$$\tilde{e}_N(\tilde{N}) = \left\| \mathbb{V}_{\hat{N}} u_{\hat{N}} - \mathbb{V}_N u_N \right\| \tag{120}$$

can be expressed as a sum in terms of the RB solution modes. Due to the hierarchical character of the RB basis,

 $\mathbb{V}_{\hat{N}}$ contains the same columns up to N as \mathbb{V}_N . Thus, the error between ROMs can be expressed as

$$\|\mathbb{V}_{\hat{N}}u_{\hat{N}} - \mathbb{V}_{N}u_{N}\| = \\ \left\|\sum_{i=N+1}^{\hat{N}} u_{i}^{\hat{N}}\psi_{i} + \sum_{j}^{N} (u_{j}^{\hat{N}} - u_{j}^{N})\psi_{j}\right\|.$$
(121)

The difference $(u_j^{\hat{N}} - u_j^N)$ between ROM coefficients is relatively small, since it represents the difference between the two coefficients associated to the same mode. If they were notably different, it would mean that the dynamics between the original ROM and the sacrificial one are different. Since the RB solution basis is hierarchical, this will not be the case for sufficiently well resolved ROMs: adding an additional mode should only refine the solution, not change drastically how the previous modes are scaled. They are not strictly the same because the ROM matrix changes if more modes are added to the basis, but again, it does so in a way that dynamics are preserved.

In Figure 30 we show in the behaviour of the error estimator for each time step. In Figure 31 we present how close the estimator is to the actual error, where both are aggregated in the time direction. We have computed the estimator for $\Delta N = 1,5$ and 10 additional modes. Then, we compute *estimator accuracy*, how far it is from the actual ROM error, aggregated for all time steps,

Estimator Accuracy =
$$||e_t - \tilde{e}_t||$$
 (122)

We conclude that it seems to be better to carry more than one mode. Nevertheless, if to compute all these additional modes becomes too costly, one extra mode will produce a valid estimator, provided that the base ROM is well resolved. Later on in Section 5.7 we explore the limits of this certification technique.



Fig. 30. Time-wise a posteriori error estimator accuracy for $N_{\rm ROM}=10,$ $N_{\rm SROM}=11.$ We observe how accurate the estimator is carrying only one additional mode.



Fig. 31. A posteriori error estimator accuracy. We see how it can become more effective to carry more additional modes than rather just one.

5.6 Results for Non-Uniform Mesh Stretching

We now present results for a non-uniform mesh displacement. In Table 13 we give the resulting basis size for each operator at each step of the nested POD. The final size ("Param. space" column) is sufficient to accurately reconstruct the operators. All operators require more than a trivial number of elements, but the bases size differ. The trilinear term snapshots have been obtained from the FOM simulation (u^* -general). Again, the reduction of this operator requires as many terms as the solution space.

We have purposefully used a different parameter sampling size for the FOM simulation and the collection of operator snapshots; to show a weakness of this methodology. Assembling operators is a time-consuming operation, but it is cheaper than carrying out a full FOM simulation. Hence, it is encouraged to have a methodology which splits the collection of operator snapshots and solution snapshots. The former are cheaper to obtain than the latter, and hence a larger parameter space can be sampled to produce a richer collateral basis. By richer basis we intend one that is likely to work better during the online stage, for it has potentially a higher generalization degree.

A possible scenario where the split between the assembly and snapshot-collection steps is not feasible is one where the displacement had to track the solution of the PDE, e.g. in fluid-structure interaction. In that case we have no alternative but to collect *all* the snapshots (operators and solution) from FOM simulations.

TABLE 13

Basis size at each step of the nested POD strategy. The operators are sorted by collateral basis final size. The Nonlinear-lifting operator corresponds to the cross-term lifting matrix that arises from the nonlinear convective term.

| | Time int. ($N_t = 500$) | Param. space | N_{μ} |
|-------------------|---------------------------|--------------|-----------|
| Reduced-basis | 660 | 93 | 20 |
| Trilinear | 670 | 94 | 20 |
| Stiffness | 121 | 68 | |
| Rhs | 180 | 32 | |
| Mass | 60 | 21 | 30 |
| Convection | 60 | 20 | |
| Nonlinear-lifting | 60 | 19 | |

In Figure 32 we show the singular value decay (SV decay) for each operator. On the top plot, we present the decays for each time integration path (for a fixed parameter, they represent the branches of the treewalk). These decays correspond to the first compression of the snapshots, with which we obtain a basis Ψ_{μ_i} for a specific parametrization. On the bottom plot, we present the decay of the final POD step, which compresses (for each operator) all of the previous bases into one.

In Figure 33 we present a zoom-in for the first ten singular values. This shows how some operators are reduced to very few modes in the time direction.

Indeed, in the time direction most operators still remain simple to reduce, but when each parameter-fixed basis is compressed with the others, most information is retained. This is so because the mesh parameters have a nonlinear effect, whereas time has a linear one. That is why in time it is simple to reduce the operator, but not so much across the geometrical parameter space. When we were dealing with a uniform deformation of the mesh, since the stretching parameter δ is present linearly, the piston oscillation can be captured by sampling one parametrization, two²¹ at most.

5.6.1 Operators Error Decay

Now that we have multiple operators with a non-trivial basis, we have to tune all of their errors simultaneously. We plot in Figure 34 the error decay for each operator as a function of the basis percentile size. The offline and online sampling pools are shown in Figure 41 in the Appendix.

All the errors decay as the basis size increases, but each operator needs a different basis size to reach the same error threshold. For the offline samples, as the basis size is shrunk, the errors remain concentrated for different parametrizations. For the online samples, this behaviour is not present, with more dispersion between the different parametrizations. The errors with the full basis are smaller for the offline pool, compared to those from the online pool. This is an expected behaviour, since the basis is optimized for the offline sample.



Fig. 32. Singular value decay for the two nested POD steps: time integration (top) and parameter space (bottom). Both axes are represented in logarithmic scale. We observe how due to the presence of the non-uniform displacement the operators are no longer trivially reduced. It takes several basis terms to accurately represent the effects of the nonlinearity. The solution space and the trilinear operator have an identical decay. This is so because the trilinear form is equally affected by the values of the extrapolated velocity u^* and the effects of the non-uniform displacement.

21. The oscillating frequency ω is inside a sinusoidal function, but this was only relevant for the mesh velocity, not the overall displacement.



Fig. 33. This figure is a zoom-in of Figure 32. Only the first ten singular values (modes) are shown. We can see how for the time integration step most operators are perfectly summarized with two to six modes. This is due to the fact that the displacement is separable in time and space; and only the space function is not linear.



Fig. 34. Operator error decay as a function of basis size percentile. (Top) Offline parameter sample. (Bottom) Online parameter sample. All the errors decay as the basis size increases, but each operator needs a different basis size to reach the same error threshold. For the offline samples, the errors are concentrated. For the online samples, the errors present more dispersion. The errors with the full basis are smaller for the offline pool, compared to those from the online pool. This is an expected behaviour, since the basis is optimized for the offline sample.

5.7 Certification by Truncation: Limits

We already saw in the heatmap from Figure 21 in Section 5.3.2 how the RB and MDEIM approximation errors could set a lower bound to the HROM error. We show how this interaction can make the HROM certification by truncation fail. To do so, we set the experiments shown in Tables 15 and 16. The online parametrization is given in Table 14. We want to determine the error estimator behaviour when the RB solution approximation error is smaller or greater than the (M)DEIM error.

TABLE 14 Online parametrization. The results will only be obtained for one parameter, since the conclusions scale for many.

| Variable | Value |
|--|--------------------------------|
| $egin{array}{c} a_0 \ \omega \ \delta \ u_p \end{array}$ | 18.64 24.78 0.28 0.37 |
| $egin{array}{c} x_c \ \sigma_c \ y_c \end{array}$ | 0.32 0.14 0.26 |

The conclusion of the experiments is that if the RB solution error is smaller than the (M)DEIM error, the HROM error is saturated by the (M)DEIM errors; and hence the mode truncation technique cannot be used to certify the HROM results, because both HROMs produce the same result. When the RB solution error is greater than the (M)DEIM error, the trucation technique will produce effective error estimations.

TABLE 15 Numerical experiments configuration. Pure ROM means no (M)DEIM is used. HROM means all collateral bases are used.

| | Pure ROM | HROM |
|--|--|------------------------|
| $ \begin{aligned} \varepsilon_{\text{RB}} &> \varepsilon_{\text{MDEIM}} \\ \varepsilon_{\text{RB}} &< \varepsilon_{\text{MDEIM}} \end{aligned} $ | Figure <mark>35</mark> Figure <mark>36</mark> | Figure 37 Figure 38 |

TABLE 16 Numerical experiments summary. The bases size and the order of magnitude of the error are provided.

| Experiment | (M)DEIM | ROM N, Error | SROM N, Error | Estimator |
|--|------------------------|--|--|--|
| Figure 35 Figure 36 Figure 37 Figure 38 | 10^{-6} 10^{-6} | $ \begin{array}{r} 15,10^{-4}\\ 25,10^{-6}\\ 15,10^{-3}\\ 25,10^{-4}\\ \end{array} $ | $25, 10^{-6} \\ 30, 10^{-7} \\ 25, 10^{-4} \\ 30, 10^{-4}$ | Accurate Accurate Regular Ineffective |



Fig. 35. ROM vs. FOM approximation error, no (M)DEIM. The FOM operators are assembled and projected for each time step. The certification by truncation procedure is accurate, with the estimator matching the ROM error.



Fig. 36. ROM vs. FOM approximation error, no (M)DEIM. The FOM operators are assembled and projected for each time step. The certificationby-truncation procedure is accurate, with the estimator matching the ROM error.



Fig. 37. ROM vs. FOM approximation error, (M)DEIM is active. The approximation error for all operators is 10^{-6} . The error of the SROM has increased, but it remains smaller than the ROM error. Hence, the certification-by-truncation procedure produces a valid estimator.



Fig. 38. ROM vs. FOM approximation error, (M)DEIM is active. The approximation error for all operators is 10^{-6} . The error for the ROM and the SROM is the same, leading to an ineffective truncation estimator.

5.8 Non-Uniform Mesh: Restricted N-MDEIM

In this final section we build the N-MDEIM basis for the trilinear term from snapshots obtained from RB solution modes evaluation (u^* -restricted).

We anticipate that we obtain a non-hierarchical basis as in Section 5.4.4 (N-MDEIM with RB Solution Modes), despite the non-uniform spacing of the mesh. This is due to the following facts:

- The domain is one-dimensional.
- The FEM basis functions are ℙ1 polynomials.
- The trilinear integrand is linear in all of its arguments (including parameters).
- The trilinear integrand contains one derivative (trial function *u*).

The last fact is crucial, making the non-uniform mesh size go unnoticed. For $\mathbb{P}1$ polynomials, the derivative of the Lagrangian function is the inverse of the mesh size, which gets cancelled by the integration over the mesh element. This is a known fact of convection-like forms, for which the mesh size does not show up in the FEM matrix. We could have expected that due to the non-uniform displacement, the mesh size would have an impact in the matrix elements. However, in the specific context described in the list above, this is not the case. For our particular scenario (despite the the non-uniform of the mesh displacement), to reduce the trilinear form it is sufficient to collect snapshots for one parameter for as many RB solution modes as we will be using for the HROM simulation (taking into account the number of modes carried by the SROM).

As we saw in previous sections, when the basis is not hierarchical, the reconstruction error cannot be used as a proxy to determine the accuracy of the collateral basis. Our conclusion here is that for this trilinear form, the collateral basis error will be as good as the error associated to the RB solution basis used to assemble the snapshots.

This is shown in Figure 39. The N-(M)DEIM has been assembled with the first five RB solution modes. In comparison to the results in Figure 35, which only reflect the RB solution error²²; these HROMs underperform because the N-MDEIM captures the dynamics for the first five RB solution modes, where the ROM and the SROM have 15 and 25 respectively. Instead, when the N-MDEIM is assembled with 15 modes, the ROM recovers its original accuracy, see Figure 40.

Furthermore, since the SVD creates a linear combination of the inputs; for non-hierarchical operator bases, trucating the RB operator basis is not equivalent to collecting the snapshots without the same number of removed RB solution modes.



Fig. 39. ROM vs. FOM approximation error. The (M)DEIM bases size for the linear operators is such that the error is 10^{-6} . The trilinear N-MDEIM has been assembled with N = 5 RB solution modes. The ROM and the SROM are poorly resolved due to unsufficient representation of the excess modes they carry with respect to the N-MDEIM training modes.



Fig. 40. ROM vs. FOM approximation error. The (M)DEIM bases size for the linear operators is such that the error is 10^{-6} . The trilinear N-MDEIM has been assembled with N = 15 RB solution modes. The ROM recovers its original accuracy, since the N-MDEIM is accurately represented for that number of modes. Instead, the SROM presents a larger error than expected, due to the excess RB solution modes it carries with respect to the N-MDEIM training modes.

6 CONCLUSIONS

An hyper reduced order model has been succesfully created and certified for the one-dimensional isentropic moving piston. This simplified gas dynamics problem contains all the ingredients that conform a real life problem: a Burgerslike nonlinear term and a moving mesh. Two parametrized moving meshes have been introduced: uniform and nonuniform mesh stretching. We are not restricted to moving boundaries, the conclusions and methods are also valid for domains whose mesh nodes move inside the domain, whilst remaining fixed at the boundaries.

The reduction strategy permits us to remain in the physical domain, thus skipping the Jacobian transformation.

6.1 Research Answers

Thanks to the fact that the methodology is purely algebraic, and that we have dealt in a general way with the parametrized Dirichlet boundary conditions, (without any assumption on their functional form), our reduction strategy adquires non-intrusive and compactness properties, that should allow its implementation on existing solvers.

We have identified a reduced basis to carry out the Galerkin projection; and a collateral basis for each of the algebraic operators (vectors and matrices), to approximate their projection onto the reduced space without explicitly assembling and projecting the original FOM operator. Building the collateral basis increases the cost of the offline stage, but it allows a perfect offline-online split: no FOM operators are used during the online run. For the linear operators, the offline stage is carried out separately from the solution RB basis identification, so that a wider parameter range can be spanned, and thus a richer operator basis obtained (a simulation is more expensive than the assembly of an operator). To approximate the operators at the same level of accuracy, the presence of the non-uniform mesh movement requires more operator modes than the uniform mesh.

The interaction between the RB and (M)DEIM errors has been analyzed. The number of RB solution and RB operators modes needs to be such that the (M)DEIM error is always smaller or equal to the RB solution error. Otherwise, the certification by truncation technique will fail.

6.1.1 Discretization Flexibility

The (M)DEIM technique determines certain operator entries which have to be computed at runtime during the online stage to carry out the empirical interpolation. In the finite element context, this is quite convenient. The restricted operator entries are computed by evaluating the weak form integral for the mesh elements associated with the selected entries. However, all the content of this work could be reused in a finite difference or finite volume formulation, provided that the implementation is capable of an efficient calculation of individual operator entries.

6.1.2 Nonlinear Convection MDEIM

Due to the velocity extrapolation in Burgers' nonlinear convection term, a trilinear operator is present in the FOM discretization. We have compared two approaches to create the MDEIM basis for this operator:

- (*u**-general) collecting the snapshots from the FOM simulation [1];
- (*u**-restricted) collecting the snapshots from evaluations of the operator with RB solution modes.

The u^* -general approach has the following drawbacks:

- it limits the sampling pool of the N-MDEIM to the one used to identify the solution reduced basis;
- the FOM operators from the simulation would not be available if the reduced basis is identified from experimental data.

Our enhancement of the trilinear MDEIM offline stage $(u^*$ -restricted) tackles these two issues, uncoupling snapshot collection from the FOM simulation. Once the RB solution modes have been identified (and trimed to a given threshold error) they can be used to build the N-MDEIM collateral basis.

The u^* -restricted approach is not limited to Burgers' nonlinear convective term. It could be used with the nonlinear convective term present in the Navier-Stokes equations, and potentially with any trilinear form whose additional argument is expressed in terms of a linear combination of the solutions. In fact, if the additional argument belonged to another space (different from the solution space), the modes²³ of such function space could be used to sample the operator efficiently.

6.2 Limitations and Future Work

We are aware of how we have benefited from solving our PDE in a one-dimensional domain. Despite the presence of a nonlinear term and a moving mesh, the formulation of the problem remained benign.

Nevertheless, our whole procedure would scale switfly to higher dimensions. We have restricted ourselves to $\mathbb{P}1$ finite elements. Although for one-dimensional domains scaling to higher order polynomials does not raise dimensionality issues, for actual three-dimensional domains it would. Therefore, we saw no need for rising our degree at this point.

The oscillating cylinder problem is a good candidate to test the formulation in a more realistic setting. Most of the problem algebraic formulation would remain the same (albeit larger matrices), except for a difficulty which was not present in this work: the calculation of the Dirichlet lifting. In this work we have leveraged the fact that the lifting could be computed analytically. For higher dimensions, the boundary conditions need to be numerically extended to the domain, e.g., via harmonic extensions [14], or solving the elastodynamic equations [57]. These extension problems can be expressed as HROMs too [1].

These are not the only challenges when the problem is dealt with in higher dimensions. For the creation of the Navier-Stokes ROM, the velocity field needs to be enriched [58], so that the inf-sup condition is also satisfied at the ROM level.

^{23.} Not necessarily obtained with a POD decomposition.

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APPENDIX A DETERMINATION OF PISTON MOVEMENT LAW

We impose the movement of the piston, so we need to make sure we do so in a physically meaningful way. This appendix is born after making the mistake of believing that any sinusoidal function would do the job. Since our flow will depart from rest, we need to set the piston motion so that at the initial instant the piston also starts its movement from rest.

To derive the piston movement law, we depart from a force defined by the elemental harmonic functions,

$$A\cos(\omega t) + B\sin(\omega t) = m\ddot{x}(t), \qquad (123a)$$

$$x(0) = 0,$$
 (123b)

$$\dot{x}(0) = 0.$$
 (123c)

Integrating in time, we get

$$\frac{A}{m\omega}\sin(\omega t) - \frac{B}{m\omega}\cos(\omega t) + C_1 = \dot{x}(t).$$
(124)

If we enforce the initial condition of rest to find the value of the integration constant, we arrive to an incongruency,

$$-\frac{B}{m\omega} + C_1 = 0 \to C_1 = \frac{B}{m\omega}.$$
 (125)

If we were to integrate in time again, due to the presence of the constant C_1 , a linear term proportional to $\sim t$ would show up.

This is not the physical result we expected, given the fact that we departed from two linear harmonic functions (which introduce and remove energy with fixed frequency and amount from the system). Hence a harmonic movement was expected, not one changing linearly in time.

This conflictive result comes from the sinusoidal term in the definition of the force moving the piston. If we set B = 0, the first integration constant will become identically zero, $C_1 = 0$. When this is the case, the linear term vanishes, and we recover a harmonic piston movement,

$$\frac{A}{m\omega^2}\cos(\omega t) + C_2 = x(t).$$
(126)

By setting the initial piston location, we get the value for C_2 ,

$$x(t) = L_0 - \frac{A}{m\omega^2} (1 - \cos(\omega t)).$$
 (127)

If we now define A such that $\frac{A}{m\omega^2}$ represents a fraction of the initial piston length, δL_0 , we get a compact expression for the piston movement,

$$x(t) = L_0 \left[1 - \delta \left(1 - \cos(\omega t) \right) \right].$$
 (128)

APPENDIX B LARGE FIGURES

B.1 Non-Uniform Displacement

| | Convection | Mass | N-Lifting | RHS | Stiffness | Trilinear |
|------|------------|---------|-----------|---------|-----------|-----------|
| р | | | | | | |
| 0.05 | 7.4e-01 | 3.6e-05 | 2.9e-02 | 4.2e-03 | 4.0e-08 | 2.5e-01 |
| 0.10 | 3.5e-02 | 2.1e-05 | 2.9e-02 | 4.6e-04 | 2.2e-08 | 6.8e-04 |
| 0.20 | 2.1e-02 | 1.7e-05 | 7.7e-03 | 9.7e-05 | 4.3e-09 | 5.1e-07 |
| 0.40 | 3.4e-03 | 3.2e-06 | 1.0e-03 | 1.4e-05 | 6.8e-10 | 1.5e-09 |
| 0.60 | 2.8e-04 | 2.4e-07 | 4.9e-05 | 1.5e-07 | 2.0e-11 | 4.1e-11 |
| 0.80 | 9.4e-06 | 4.8e-09 | 2.2e-06 | 3.2e-09 | 6.7e-13 | 3.3e-13 |
| 0.85 | 2.5e-06 | 2.6e-09 | 3.6e-07 | 2.5e-09 | 3.9e-13 | - |
| 0.90 | 6.6e-07 | 1.2e-09 | 1.9e-07 | 5.6e-10 | 4.2e-13 | - |
| 0.95 | 6.1e-07 | 6.3e-10 | 1.6e-07 | 2.7e-10 | 3.5e-13 | - |
| 1.00 | 1.7e-07 | 1.7e-10 | 6.5e-08 | 1.4e-10 | 4.8e-14 | 1.3e-15 |

TABLE 17 Online operator approximation errors.

TABLE 18 Basis size according to percentile position.

| | Convection | Mass | N-Lifting | Rhs | Stiffness | Trilinear |
|------|------------|------|-----------|-----|-----------|-----------|
| р | | | | | | |
| 0.05 | 1 | 1 | 0 | 1 | 3 | 4 |
| 0.10 | 2 | 2 | 1 | 3 | 6 | 9 |
| 0.20 | 4 | 4 | 3 | 6 | 13 | 18 |
| 0.40 | 8 | 8 | 7 | 12 | 27 | 37 |
| 0.60 | 12 | 12 | 11 | 19 | 40 | 56 |
| 0.80 | 16 | 16 | 15 | 25 | 54 | 75 |
| 0.85 | 17 | 17 | 16 | 27 | 57 | 79 |
| 0.90 | 18 | 18 | 17 | 28 | 61 | 84 |
| 0.95 | 19 | 19 | 18 | 30 | 64 | 89 |
| 1.00 | 20 | 21 | 19 | 32 | 68 | 94 |



Fig. 41. Sampling space for offline and online stages for the nonlinear displacement test case.