


Model Order Reduction for Gas and Energy Networks

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Abstract:

To counter the volatile nature of renewable energy sources, gas networks take a vital role. But, to ensure fulfillment of contracts under these new circumstances, a vast number of possible scenarios, incorporating uncertain supply and demand, has to be simulated ahead of time. This many-query task can be accelerated by model order reduction, yet, large-scale, nonlinear, parametric, hyperbolic partial differential(-algebraic) equation systems, modeling gas transport, are a challenging application for model reduction algorithms.

For this industrial application, we bring together the scientific computing topics of: mathematical modeling of gas transport networks, numerical simulation of hyperbolic partial differential equation, and model order reduction for nonlinear parametric systems. This research resulted in the `morgen` (Model Order Reduction for Gas and Energy Networks) software platform, which enables modular testing of various combinations of models, solvers, and model reduction methods. In this work we present the theoretical background on systemic modeling and structured, data-driven, system-theoretic model reduction for gas networks, as well as the implementation of `morgen` and associated numerical experiments testing model reduction adapted to gas network models.

Keywords: Gas Network, Model Reduction, Empirical Gramians, Digital Twin

p	Pressure	N_s	Number of supply nodes
q	Mass-flux	N_0	Number of internal nodes
\bar{p}	Steady-state pressure	N_d	Number of demand nodes
\bar{q}	Steady-state mass-flux	N_c	Number of compressors
s_p	Supply node pressure	N_p	Dimension of pressure space
d_q	Demand node mass-flux	N_q	Dimension of mass-flux space
s_q	Supply node mass-flux	n_p	Dimension of reduced pressure space
d_p	Demand node pressure	n_q	Dimension of reduced mass-flux space

Table 1: List of recurring symbols.

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1 Introduction

Rapid transient simulations of gas flow in pipeline networks are essential for safe operations of gas networks as well as reliable delivery of denominations. Yet, in a volatile supply and demand environment, due to increasing renewable energy sources, the time horizon for planning dispatch and load forecasting shortens while more sources of accountable uncertainties, such as effects of weather on energy consumption and production are introduced; to a lesser degree this is a long standing challenge [5]. An example is the interconnection of gas and power grids through gas-fired power plants [21, 130, 82]. Thus, more simulations of dynamic gas network behavior need to be completed in less time by the gas grid operators. However, available compute power is (and was [124]) never sufficient. To this end we evaluate customized model reduction techniques for an established class of gas network models.

This work and the associated software platform is an effort to determine which model reduction methods are suitable for enabling *digital twins* [110, 48] of gas networks. Depending on the mathematical model and quantities of interest, the twin may contain redundant or superfluous information with respect to the simulations. Therefore, model reduction compresses the twin to a matched surrogate model, which is sufficiently accurate in the chosen operating region. The swift numerical simulation of gas network twins by reduced order modeling is highly relevant, not only due to the transition towards renewables at the time of writing, which is underlined by the research projects *MathEnergy*¹ (Mathematical Key Technologies for Evolving Energy Grids) [23] that the authors are part of, and *TRR154*² (Mathematical Modelling, Simulation and Optimization using the Example of Gas Networks) [75], but also because of the intriguing numerical problem of model reduction for hyperbolic, nonlinear, coupled, parametric, multiscale partial differential-algebraic equation systems.

If relevant intraday demand changes occur, established steady / stationary / static simulations may not be sufficient anymore [34]. The basic model for the simulation of unsteady / dynamic / transient flow processes in gas pipelines is based on the one-dimensional (isothermal) Euler equations, originally introduced in [45], and “rediscovered” in [94] as well as in [70, 71] around the same time. A practical extension in the context of gas networks is the repetitive modeling approach [30], which enables a modular construction. For extensive details on gas network modeling, see the works [119, 95, 36, 112, 85, 27, 14, 116], and for a concise summary of the overall approach we recommend [6]. Furthermore, a system-theoretic approach to gas networks is discussed in [29], and results on boundary reachability (controllability) and observability for this class of models have been derived in [7, 8].

In terms of complexity reduction for gas network models, earlier works applied techniques such as combining parallel pipelines [117], singular perturbation [115] and symbolic simplifications [86]. Younger works introduced projection-based model reduction methods from fluid dynamics, proper orthogonal decomposition (POD) [41, 44, 42], and system-theoretic methods [78] (matrix interpolation), [79] (balanced truncation), or exploiting a port-Hamiltonian formulation [77, 31] (Padé-type approximation).

¹<https://mathenergy.de>

²<https://trr154.fau.de>

In this work, we conceptually combine these previous approaches, by using system-theoretic but data-driven methods that are structure-preserving. The utilized data-driven assembly of the system-theoretic operators, central to the employed methods, is also a partial answer to the challenges posed in [47, Remark 5.10]. Furthermore, we note that from this work’s point of view, [130, 131, 118] are concerned rather with (valuable) model simplifications and index reduction than (projection-based) model reduction.

To avoid the analytically most complex aspects of the gas network model – the nonlinearities – one could linearize the model around an operating point. Yet, the different nonlinearities (i.e. friction, compressibility and compressors) are unlikely to have a compatible operating point for a wide range of scenarios. Furthermore, linearized and simplified models of gas flow have limitations with simulations of real scenarios [51], [121, Ch. 7], hence, we use a nonlinear model. Since there is no general theory for model reduction of nonlinear systems, and a high degree of modularity in the gas network modeling process, model reduction algorithms have to be compared heuristically to determine their applicability.

As a result of this reasoning and a demand for gas network simulation software tools [26, 59], a platform named **morgen** (**m**odel **o**rd(er) **r**eduction for **g**as and **e**nergy **n**etworks) was designed with the goal to compare different models, solvers and reducers. The **morgen**³ platform is an open-source project, and designed in a configurable and modular manner, so that specialists from the fields of modeling, discretization or model reduction can compare their methods.

In summary, this work contributes a full, but also fully modular, modeling, model reduction and simulation open-source software stack for gas networks, and potentially other energy network systems (i.e. district heating networks, water networks), which brings together research results from various disciplines. Overall, this work is composed as follows: In Section 2 the gas network model, simplifications with respect to multiscale property and algebraic constraints, the utilized approach for obtaining a steady-state as initial condition, and non-pipe elements are described. Section 3 and Section 4 outline the general model reduction idea and propose five reduction method classes. The design and features of the **morgen** platform are summarized in Section 5, followed by three sets of numerical experiments in Section 6. We conclude by an outlook (Section 7) and an evaluation of our findings in Section 8.

2 Transient Gas Network Model

The goal of this section is to describe the partial differential-algebraic equation model of a gas network as an input-output system that maps boundary values to quantities of interest. First, the model for a single pipeline is summarized, which is based on the isothermal Euler equations of gas dynamics [94, 96]. Then, it is generalized to a network of pipes, and simplified compressors are added. Finally, input and output operators are determined. Even though further non-pipe elements are common in gas networks, such as resistors, coolers, heaters, valves and control valves [36, 87], we prioritized compressors to focus on the model reduction aspect on a macro scale. Moreover, the practical numerical problems of scale homogenization, spatial discretization, index reduction and steady-state approximation are discussed in this section.

³“Morgen” is also the German language word for “tomorrow”.

2.1 Gas Pipeline Model

The principal building blocks of gas transport networks are pipelines or ducts. Since the length of pipes exceeds their diameter by far ($L > 500d$, [71]), a spatial one-dimensional model suffices. We model gas flow in a (cylindrical) pipe of length L connecting two junctions by the isothermal Euler equations:

$$\begin{aligned} \frac{1}{\gamma_0 z_0} \partial_t p &= -\frac{1}{S} \partial_x q, \\ \partial_t q &= -S \partial_x p - \left(\frac{S g \partial_x h}{\gamma_0 z_0} p + \frac{\gamma_0 z_0 \lambda_0}{2 d S} \frac{q |q|}{p} \right), \end{aligned} \quad (1)$$

which determine the evolution of the coupled pressure $p(x, t)$ and mass-flux $q(x, t)$ variables. The physical dimension of the pipe enters as its diameter d and the derived cross-section area $S = \frac{\pi}{4} d^2$, which is assumed constant, ignoring the influence of temperature and pressure on the pipe walls. These coupled partial differential equations (PDE) can also be characterized as a nonlinear, two-dimensional, first-order hyperbolic system of conservation laws: the pressure p preserves continuity, while the mass-flux q conserves momentum.

Following [69, 54, 98] and [14, Sec. 2.1], the inertia term has been neglected, which leads to the *ISO2* model in the *TRR154* classification [27], also known as *friction-dominated model* [18, Sec. 3.2.1]. Furthermore, we assume a turbulent flow with a Reynolds number exceeding $\text{Re} \gg 10^5$ [45, 32], neither line breaks or valve closings happen intraday (to preclude associated shocks [28]), and low-frequency boundary values [98], which in this work are the supply pressure and demand mass-flux, due to frequent use in literature, and use-cases such as guaranteed demand pressures [53].

In (1), the linear reaction term describes the effect of gravity (with standard gravity $g \equiv 9.80665 \frac{\text{m}}{\text{s}^2}$) due to the pipe height h , while the nonlinear reaction term models loss of momentum due to friction at the pipe walls, specified by the (Darcy-Weisbach) friction factor $\lambda_0 := \lambda(d, k, \text{Re}_0)$, given a pipe roughness k , and an estimated mean Reynolds number Re_0 , see [14, Sec. 2.2]⁴. This friction term is principal to the accuracy of the gas pipeline model [97, 84, 27, 51].

In this model variant, a (globally) constant mean compressibility factor $z_0 := z(p_0, T_0) \in \mathbb{R}$ is assumed [96, 29, 52, 101], as well as a constant gas state $\gamma_0 := R_S T_0$, whereas the temperature T_0 and the specific gas constant R_S are treated as parameters (see Section 2.6). To this end, the steady-state pressure $\bar{p} =: p_0$ is used to compute z_0 , via heuristic formulas based on the Virial expansion [19], [14, Sec. 2.3]⁵.

2.2 Homogenizing Scales

The SI-based units for pressure and mass-flux are [Pa] and [kg/s], respectively. This introduces a difference in scales of five orders of magnitude between the variables p and q , and hence induces numerical problems. To counter this multiscale structure, we simply rescale the pressure from [Pa] to [bar] which conveniently comprises a factor of 10^5 . Nonetheless, the model still consists of two interacting physical variables, hence the model still has to be treated as a coupled system, however, without numerical multiscale issues.

⁴Additionally to [14], the IGT formula [27], [87, Sec. 15.2.3] is implemented in `morgen`.

⁵Additionally to [14], the DVGW-G-2000 equation [85, Ch. 9] is implemented in `morgen`.

2.3 Gas Network Model

Given the model for a single pipe from the previous section, a (gas) network of pipes can be encoded by a finite directed graph, which is a tuple $\mathcal{G} = (\mathcal{N}, \mathcal{E})$ of finite sets symbolizing nodes \mathcal{N} , and oriented edges \mathcal{E} . The edges correspond to pipes, while the nodes represent the junctions connecting pipes. The connectivity of the network is the relationship between edges and junctions, given by the incidence matrix $\mathcal{A} \in \{-1, 0, 1\}^{|\mathcal{N}| \times |\mathcal{E}|}$, a map from edges to nodes, such that:

$$\mathcal{A}_{ij} = \begin{cases} -1 & \mathcal{E}_j \text{ connects **from** } \mathcal{N}_i, \\ 0 & \mathcal{E}_j \text{ connects **not** } \mathcal{N}_i, \\ 1 & \mathcal{E}_j \text{ connects **to** } \mathcal{N}_i. \end{cases}$$

Note, that the orientation of the edges is not enforcing the dynamic flow direction of the gas, but is necessary to determine the complexity and boundary of the overall networked model [44, 14].

We introduce the notation $|\mathcal{A}|$ for the component-wise absolute value of a matrix. Using this absolute value, the following partial incidence matrices associating edges entering and leaving nodes respectively are defined:

$$\mathcal{A}_R := \frac{1}{2}(\mathcal{A} + |\mathcal{A}|), \quad \mathcal{A}_L := \frac{1}{2}(\mathcal{A} - |\mathcal{A}|).$$

Next, based on this connectivity, certain conservation properties are enforced to maintain a network balance, and thus ensure physical relevance of the gas network model. Specifically, the Kirchhoff laws are applied to the network in vectorized (matricized) form [120, 13]:

1. The net flow (mass-flux) at every node (junction) is zero. This means that no gas gets lost in transport from one pipe to the next, with the exception of boundary nodes. Hence, a vector of flows $q \in \mathbb{R}^{|\mathcal{E}|}$ applied to the incidence matrix equals the (out-)flow at the boundary (discharge) nodes $d_q : \mathbb{R} \rightarrow \mathbb{R}^{|\mathcal{N}_D|}$, which are mapped into the network via $\mathcal{B}_d \in \{0, 1\}^{|\mathcal{N}| \times |\mathcal{N}_D|}$:

$$\mathcal{A}q(t) = \mathcal{B}_d d_q(t),$$

with $\mathcal{N}_D \subset \mathcal{N}$ denoting the subset of boundary nodes, which only connect **from one** node respectively, but **not to any** node.

2. The sum of directed pressure drops in every fundamental loop is zero. Fortunately, an equivalent representation [129, Ch. 7.3] can be used, which resolves implicitly. It remains to ensure that the nodal pressures at the in-flow boundary (supply) nodes are associated to the boundary function $s_p : \mathbb{R} \rightarrow \mathbb{R}^{|\mathcal{N}_S|}$, which are mapped to the network via $\mathcal{B}_s \in \{0, 1\}^{|\mathcal{N}| \times |\mathcal{N}_S|}$:

$$\mathcal{A}_0^\top p(t) + \mathcal{B}_s^\top s_p(t) = A_{0,R}^\top p(t) - A_{0,L}^\top p(t),$$

with $\mathcal{N}_S \subset \mathcal{N}$ denoting the subset of boundary nodes, which only connect **to one** node respectively, but **not from** any node, and the reduced incidence matrix \mathcal{A}_0 with all rows associated to supply nodes removed.

Given a connected and directed graph representing a gas network topology, with the dynamic flow in the pipe edges modeled by the PDE (1), then yields a partial differential-algebraic equation (PDAE) due to the above constraints.

2.4 Discretization and Index Reduction

Next, we delineate the discretization of the spatial differential operators and reduction of the (P)DAE index in the networked system, yielding a system of ODEs (Ordinary Differential Equations). Eventually, the remaining discretization of the temporal differential operators is addressed.

We explicitly do not use the decoupling approaches from [10] or [9], as the former employs linearization and hence does not fit this setting, while compared to the latter, our equivalent analytic index reduction is more convenient here.

The partial differential(-algebraic) equation is discretized using the method of lines: First in space, then in time, yielding a (nonlinear) dynamic system. For the spatial discretization a first-order upwind finite difference scheme is utilized. We select (only) two points for each of the k pipes with length L_k , namely the start (\cdot^R) and end point (\cdot^L):

$$\begin{aligned}\partial_x p_k &\approx \frac{p_k^R - p_k^L}{L_k}, & k \in \mathcal{E}, \\ \partial_x q_k &\approx \frac{q_k^R - q_k^L}{L_k}, & k \in \mathcal{E}.\end{aligned}$$

The matter of short, long and varying lengths L_k is addressed in Section 2.4.3. For each pipe, this leads to the following equations:

$$\begin{aligned}\frac{1}{\gamma_0 z_0} \partial_t p_k^* &= -\frac{1}{S_k} \frac{q_k^R - q_k^L}{L_k}, \\ \partial_t q_k^* &= -S_k \frac{p_k^R - p_k^L}{L_k} - \left(\frac{S_k g(h_k^R - h_k^L)}{\gamma_0 z_0 L_k} p_k^* + \frac{\gamma_0 z_0 \lambda_{0,k} q_k^* |q_k^*|}{2 d_k S_k p_k^*} \right).\end{aligned}$$

Now, different choices for (\cdot^*) are surmisable. Subsequently, two specific combinations of p^* and q^* will be discussed: The midpoint discretization [41, 44, 131, 42, 13], and the left-right discretization [43, 104, 39] resulting in (implicit) ODEs. For an error analysis of these two discretization variants, see [116]. In the following, we describe a unified approach of deriving these index-reducible discretizations.

For notational ease in the coming subsections, a vectorized form for the above (networked) system including its constraints is given by:

$$\begin{aligned}D_d \partial_t p^* &= D_p (q^R - q^L), \\ \partial_t q^* &= D_q (\mathcal{A}_0^T p + \mathcal{B}_s^T s_p) - \left(D_g p^* + D_f \frac{q^* |q^*|}{p^*} \right), \\ \mathcal{A}_0 q^* &= \mathcal{B}_d d_q,\end{aligned}\tag{2}$$

using, and thus resolving, the constraint $\mathcal{A}_0^T p + \mathcal{B}_s^T s_p = p^R - p^L$, and with the diagonal matrices:

$$\begin{aligned}D_{d,kk} &= \frac{1}{\gamma_0 z_0}, & D_{p,kk} &= -\frac{1}{S_k L_k}, \\ D_{q,kk} &= -\frac{S_k}{L_k}, & D_{g,kk} &= \frac{S_k g(h_k^R - h_k^L)}{\gamma_0 z_0 L_k}, & D_{f,kk} &= \frac{\gamma_0 z_0 \lambda_{0,k}}{2 d_k S_k}.\end{aligned}$$

Note, that depending on the choices for p^* and q^* , the model's analytic and numerical character will differ.

2.4.1 Midpoint Discretization

In case of the midpoint discretization, we set p_k^* and q_k^* to the mean of its associated edge's endpoints:

$$\begin{aligned} p_k^* &= \frac{p_k^R + p_k^L}{2} =: p_k^+, \\ q_k^* &= \frac{q_k^R + q_k^L}{2} =: q_k^+. \end{aligned}$$

Furthermore, we define $q^- := \frac{1}{2}(q^R - q^L)$, and note, that in vectorized form, $p^+ = \frac{1}{2}(|\mathcal{A}_0^\top|p + |\mathcal{B}_s^\top|s_p)$.

Together with the algebraic constraints from Section 2.3, a DAE system in the variables p , q^+ , and q^- arises:

$$D_d \frac{1}{2}(|\mathcal{A}_0^\top|\dot{p} + |\mathcal{B}_s^\top|\dot{s}_p) = D_p 2q^-, \quad (3a)$$

$$\begin{aligned} \dot{q}^+ &= D_q(\mathcal{A}_0^\top p + \mathcal{B}_s^\top s_p) - \left(D_g \frac{1}{2}(|\mathcal{A}_0^\top|p + |\mathcal{B}_s^\top|s_p) \right. \\ &\quad \left. + D_f \frac{q^+ |q^+|}{\frac{1}{2}(|\mathcal{A}_0^\top|p + |\mathcal{B}_s^\top|s_p)} \right), \end{aligned} \quad (3b)$$

$$0 = \mathcal{A}_0 q^+ + |\mathcal{A}_0|q^- - \mathcal{B}_d d_q. \quad (3c)$$

Since we aim to obtain an ODE, we need to transform this DAE system. The complexity of deriving this transformation is quantified by the DAE's index. From the various DAE index concepts, we use the tractability index τ [83], for which the midpoint discretization guarantees $\tau \leq 2$ [44].

This DAE can be decoupled into an ODE by rewriting it in the variables p and q^+ . To this end,

1. the pressure boundary condition implicitly resolves (2).
2. By multiplying the differential equation (3a) by $(|\mathcal{A}_0|D_p^{-1})$ from the left, the remaining algebraic constraint (3c) is removed by replacing $\mathcal{A}_0 q^-$ by $(-\mathcal{A}_0 q^+ + \mathcal{B}_d d_q)$ in (3a). Since D_p is a diagonal matrix, this is also numerically feasible.

Thus, we obtain:

$$|\mathcal{A}_0|(\frac{1}{4}D_p^{-1}D_d)|\mathcal{A}_0^\top| \dot{p} = -(\mathcal{A}_0 q^+ - \mathcal{B}_d d_q) - |\mathcal{A}_0|(\frac{1}{2}D_p^{-1}D_d)|\mathcal{B}_s^\top| \dot{s}_p, \quad (4a)$$

$$\begin{aligned} \dot{q}^+ &= D_q(\mathcal{A}_0^\top p + \mathcal{B}_s^\top s_p) - \left(D_g \frac{1}{2}(|\mathcal{A}_0^\top|p + |\mathcal{B}_s^\top|s_p) \right. \\ &\quad \left. + D_f \frac{q^+ |q^+|}{\frac{1}{2}(|\mathcal{A}_0^\top|p + |\mathcal{B}_s^\top|s_p)} \right). \end{aligned} \quad (4b)$$

This system of a pressure and mass-flux variable now consists of only differential equations. Notably, the first equation of the ODE system contains a temporal derivative of the input function s_p , which practically would need to be approximated numerically, for example by finite differences. However, we will assume that all inputs are sums of step functions, so that effectively $\dot{s}_p \equiv 0$, which is reasonable as we assume exclusively low-frequency boundary values.

2.4.2 Endpoint Discretization

For the endpoint discretization, also called left-right discretization, we set p_k^* and q_k^* to the left and right endpoints, respectively:

$$\begin{aligned} p_k^* &= p_k^R, \\ q_k^* &= q_k^L. \end{aligned}$$

Since $(\mathcal{B}_s^\top + |\mathcal{B}_s^\top|)s_p = 0$, we can write $p^R = \mathcal{A}_{0,R}^\top p$. With the algebraic constraints from Section 2.3, a DAE system in the variables p , q^R , and q^L results:

$$\mathcal{A}_{0,R}^\top \dot{p} = D_p(q^R - q^L), \quad (5a)$$

$$\dot{q}^L = D_q(\mathcal{A}_0^\top p + \mathcal{B}_s^\top p_s) - \left(D_g \mathcal{A}_{0,R}^\top p + D_f \frac{q^L |q^L|}{\mathcal{A}_{0,R}^\top p} \right), \quad (5b)$$

$$0 = \mathcal{A}_{0,R} q^R + \mathcal{A}_{0,L} q^L - \mathcal{B}_d d_q. \quad (5c)$$

As for the midpoint discretization, we want to derive an ODE. For the endpoint discretization, it is shown in [43, 104], that the tractability index is $\tau = 1$, if all edges connecting supply nodes are directed from the supply, and each component of the graph is connected to at least one supply. This implies, no two supplies are to be directly connected. Similar to Section 2.4.1, this DAE can be decoupled into an ODE by rewriting it in the variables p and q^L . Applying equivalent steps to (5) as for the midpoint decoupling (3) yields:

$$(\mathcal{A}_{0,R} D_p^{-1} D_d \mathcal{A}_{0,R}^\top) \dot{p} = -(\mathcal{A}_0 q^L - \mathcal{B}_d d_q), \quad (6a)$$

$$\dot{q}^L = D_q(\mathcal{A}_0^\top p + \mathcal{B}_s^\top p_s) - \left(D_g \mathcal{A}_{0,R}^\top p + D_f \frac{q^L |q^L|}{\mathcal{A}_{0,R}^\top p} \right), \quad (6b)$$

using $\mathcal{A}_{0,R} + \mathcal{A}_{0,L} = \mathcal{A}_0$ in (6a).

An advantage of this endpoint discretization, in addition to the absence of a derivative of an input function, is the input-free friction term in (6b).

2.4.3 Temporal Discretization

After spatial discretization and index reduction of the gas network model, a system of stiff nonlinear ODEs (in time) remains. The remaining temporal differential operator(s) are discretized using time-stepping schemes.

Due to the hyperbolicity of the pipeline model, the temporal resolution Δt used for the discrete integration interdepends on the spatial resolution Δx employed for the discretization of the spatial differential operators. Formally, this is expressed by the Courant-Friedrichs-Levy (CFL) condition [49, 11, 126], which states that the propagation of information cannot be faster than its conveyor:

$$\lambda_{\text{CFL}} := v_{\max} \frac{\Delta t}{\Delta x} < 1,$$

with the (dimensionless) CFL constant λ_{CFL} , symbolizing the ratio of temporal and spatial discretization step-width scaled by the peak velocity v_{\max} . Since the flow is subsonic, v_{\max} could be estimated from the (linearized) characteristics [49], or via the boundary values⁶. However, we fix the peak velocity to $v_{\max} = 20 \frac{\text{m}}{\text{s}}$ (practically it is configurable).

⁶e.g. https://petrowiki.org/Pipeline_design_consideration_and_standards#Gas_line_sizing

Due to this relation of the space and time discretization and a pre-selected application-specific sampling frequency of the output trajectory (i.e., every 60s), the spatial discretization has to be adapted accordingly. The spatial discretization by finite differences in the previous sections ignores pipeline length, as each pipe is assigned only one (finite) difference. This means, pipes are potentially too long or too short with respect to a nominal length Δx , determined by the CFL condition $\Delta x = (1 - \varepsilon) v_{\max} \Delta t$, ($0 < \varepsilon \ll 1$). Thus, too long pipes are subdivided into virtual pipes of nominal length, while too short pipes, including a potential remainder of too long pipes, are “rounded” to a full nominal-length pipe, yet with a friction term scaled by the fraction of the short pipe’s length compared to the nominal length,

$$\tilde{D}_{f,kk} = \begin{cases} D_{f,kk} & L_k = \Delta x, \\ \frac{L_k}{\Delta x} D_{f,kk} & L_k < \Delta x. \end{cases}$$

This approach assumes that delays due to the forced virtual length of an actually short pipe is insignificant, hence this simple homogenization of pipe lengths may be improved by replacing short pipes by friction-less shortcuts with a static pressure-drop, as used in the quasi-static model [54, 41] and similar to the subsequent compressor model in Section 2.5.

Overall, we refine each pipe into a sequence of pipes of a selected nominal length – a graph level refinement – which is determined using the CFL condition. We note here, that this methodology is aimed at ensuring a certain minimum length for each pipe, as the shortest pipe may dictate an unnecessarily finely resolved time discretization. In terms of a pipe’s maximum length, suggestions are for example: 5km ([41]), or 10km ([130]).

As the model is composed of a stiff, linear (hyperbolic) and a nonlinear component, an implicit solution of the linear part using a DIRK (Diagonally Implicit Runge-Kutta) method, and an explicit solution of the nonlinear part via an SSP (Strong Stability Preserving) method, by an IMEX (IMplicit-EXplicit) solver, as proposed in [11, Sec. 3.2.2], is targeted. The actual quadrature rules used to compute the transient solutions are detailed in Section 5.3.

2.5 Simplified Compressors

Beyond pipes, gas networks comprise a variety of non-pipe elements, of which the most important are compressors. Compressors increase gas pressure to counteract cumulative effects of retarding forces (friction, gravity, inertia etc.), and are grouped into stations with many possible configurations. For our purposes, we just allow fixed configurations on a macro scale [81] per scenario, which leads to a compressor being modeled as a special kind of edge that boosts the pressure from its suction inlet to its discharge outlet.

Compressors are typically modeled “ideally”, based on their power consumption, for example as a special node type; in-depth discussions can be found in [53, 112]. Due to this consumption model, such ideal compressor units are useful for energy utilization optimization tasks, yet, for a simplified transient simulation aspect already too complicated. A more practical approach is taken in [130, 131, 82], where a compression ratio $\alpha_i \geq 1$ is used to scale the pressure in each node, or pipe [118], which means $\alpha_i > 1$ indicates compression / a compressor, otherwise ($\alpha_i = 1$) a pipe.

Here, we utilize a likewise simple compressor model similar to [125, 37], for which we assume it is propelled by an external energy source, for example, given a compressor electrification, by excess renewable power [33], or that the off-take in gas is insignificant. But instead of using compression ratios, a constant (or parametric) target pressure is prescribed, modeling *discharge pressure control* [117].

The following affine compressor model is a variant of the compressor presented in [114] and used in [6]. We model a simplified compressor by a level, short pipe which increases the pressure at its outflow to a specified value \bar{p}_c (and without friction, $\lambda_{ij} \equiv 0$). Given the pipe from nodes i to j is treated as such a “compressor pipe”, with target pressure of \bar{p}_c , the variables p_{ij} and q_{ij} are given by the differential equations:

$$\begin{aligned} \dot{p}_{ij}(t) &= q_j - q_i, \\ \dot{q}_{ij}(t) &= \bar{p}_c. \end{aligned} \tag{7}$$

The target pressure \bar{p}_c could be a step function $\bar{p}_c(t)$, and hence a control input [118], which could be accompanied by a (nonlinear) output quantity of interest yielding the energy consumption of the compressor via [32].

A compressor could also be interpreted as an actual pipe with “negative friction”, and we considered using such nonphysical pipes as compressor model, but a difficult transformation between friction and pressure increase would have to be calibrated for every model variant (including friction factor formulas) and updated with every change in any model.

2.6 Parametrization

For the considered pipeline model (1), two scalar parameters are of practical interest: The temperature of the gas T_0 (in [K]), which is assumed to be constant throughout the network, and the global specific gas constant R_S (in [J/(kg K)]). Due to mainly underground on-shore pipelines [85, Ch. 45], and coolers in compressor stations [81], using an isothermal model is a reasonable simplification. However, temperature is relevant as a parameter, since the use of an isothermal model “freezes” the dynamic energy (temperature) component in the original Euler equation (in time), and while intraday ambient temperature variation can be neglected for simulations with a 24-hour time horizon, the temperature difference of a hot summer day and a cold winter day should be taken into account by a parameter representing an average temperature.

The specific gas constant, on the other hand, is determined by the gas composition, which may also vary. Again, local variations during an intraday simulation are neglected in this work, yet the average gas mixture of natural gas with, for example, hydrogen or bio-gas is relevant, so a parameter for the average specific gas constant is introduced.

Hence, the parameter-space Θ is given by:

$$\theta := \begin{pmatrix} T_0 \\ R_S \end{pmatrix} \in \Theta \subset \mathbb{R}^2.$$

Applied to the respective components of the input-output model of Section 2.7, this leads to parameter-dependent quantities, which need to be regarded accordingly by the model reduction as discussed in Section 3.3.

2.7 Input-Output Model

After spatial discretization and index reduction, we end up with a square input-output system, a system with the same number of inputs and outputs, consisting of an ordinary differential equation, an output function and an initial value:

$$\begin{aligned} \overbrace{\begin{pmatrix} E_p(\theta) & 0 \\ 0 & I_{N_q} \end{pmatrix}}^E \begin{pmatrix} \dot{p} \\ \dot{q} \end{pmatrix} &= \overbrace{\begin{pmatrix} 0 & A_{pq} \\ A_{qp} & 0 \end{pmatrix}}^A \begin{pmatrix} p \\ q \end{pmatrix} + \overbrace{\begin{pmatrix} 0 & B_{pd} \\ B_{qs} & 0 \end{pmatrix}}^B \begin{pmatrix} s_p \\ d_q \end{pmatrix} + \overbrace{\begin{pmatrix} 0 \\ F_c \end{pmatrix}}^f + \overbrace{\begin{pmatrix} 0 \\ f_q(p, q, s_p, \theta) \end{pmatrix}}^f, \\ \begin{pmatrix} s_q \\ d_p \end{pmatrix} &= \overbrace{\begin{pmatrix} 0 & C_{sq} \\ C_{dp} & 0 \end{pmatrix}}^C \begin{pmatrix} p \\ q \end{pmatrix}, \\ \begin{pmatrix} p_0 \\ q_0 \end{pmatrix} &= \begin{pmatrix} \bar{p}(\bar{s}_p, \bar{d}_q) \\ \bar{q}(\bar{s}_p, \bar{d}_q) \end{pmatrix}, \end{aligned} \quad (8)$$

with parameter independent linear vector field components A and B , parametric pressure mass matrix $E_p(\theta)$, and nonlinear friction and gravity retarding term $f_q(p, q, s_p, \theta)$. The actual composition of the dynamical system components depends on the discretization and index reduction, cf. Section 2.4, while the linear output function consists of $C_{sq} = |\mathcal{B}_s|$ and $C_{dp} = |\mathcal{B}_d^\top|$. The load vector $F_c \in \mathbb{R}^{N_q}$ accumulates the respective discharge pressures \bar{p}_c , as described in Section 2.5, for all compressors, and the initial state is given by a steady state, whose computation is detailed in Section 2.8, depending on given steady state boundary values \bar{s}_p, \bar{d}_q . Altogether, the gas network model is a generalized linear system (E, A, B, C) , together with a nonlinear part f .

A control system formulation of transient gas network simulation was already formulated in [71], and recently in [126]. Also in [29], a so-called ‘‘systemic interpretation’’ is discussed; inspired by [29, Figure 2], we schematically illustrate (8) in Fig. 1.

The specific structure and grouping for the single pipeline model (1), the index-reduced spatially discrete network models (4), (6), and the input-output model (8), is justified by the numerical processing: only two model components depend on the parameters, temperature T_0 and specific gas constant R_S , as well as on the compressibility factor z_0 , namely the mass matrix E_p and the jointly treated retarding forces gravity and friction f_q . Hence, the linear part of the right-hand side vector field is non-parametric and compressibility-agnostic. Overall, this system maps input boundary values, in the scope of this work, pressure at the inlets and mass-flux at the outlets, via the internal state, to output quantities of interest, here, mass-flux at the inlets and pressure at the outlets:

$$\begin{array}{ccc} \begin{pmatrix} s_p \\ d_q \end{pmatrix} & \xrightarrow{S} & \begin{pmatrix} s_q \\ d_p \end{pmatrix} \\ & \searrow & \nearrow \\ & \begin{pmatrix} p \\ q \end{pmatrix} & \end{array}$$

To this type of input-output system we can now apply (data-driven) system-theoretic model reduction methods, which preserve the input-to-output mapping S , but explicitly not the internal state $(p \ q)^\top$. Lastly, we note that based on [65], we added a model fact sheet in the Appendix.

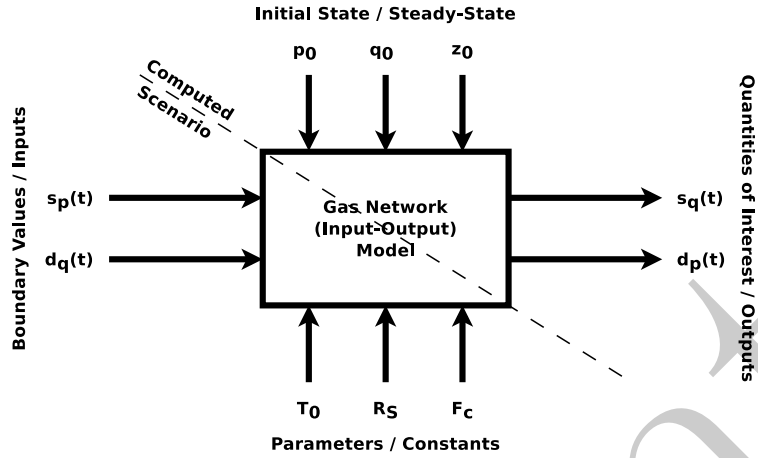


Figure 1: Schematic illustration of gas network input-output model.

2.8 Steady-State Computation

After spatial discretization, the dynamic simulation becomes an initial value problem. Yet, only the boundary values of the network model are known a-priori. This means the internal state at time $t = 0$ is unknown. We assume simulations always start at a steady-state \bar{p} , \bar{q} for which $\partial_t p = \partial_t q = 0$, given some (initial) boundary values \bar{s}_p , \bar{d}_q . The internal state is then computable as a steady-state problem. Since the employed model is nonlinear, we approximate the steady-state by a two-step procedure:

- 1a. Linear mass-flux steady-state: $A_{pq} \bar{q} = -B_{pd} \bar{d}_q$.
- 1b. Linear pressure steady-state: $A_{qp} \hat{p} = -(B_{qs} \bar{s}_p + F_c)$.
2. Corrected pressure steady-state: $A_{qp} \bar{p} = -(B_{qs} \bar{s}_p + F_c + f_q(\hat{p}, \bar{q}, \bar{s}_p, \theta))$.

Step 2 can be repeated until an error threshold is met by using the previously approximated pressure steady-state. Practically, the linear problems in Step 1 and Step 2 are solved by a QR-based least-norm method [17]. Note, that Step 1a and Step 1b can be solved in parallel and that the QR decomposition of Step 1b can be recycled in Step 2 because of the chosen model structure.

While this method works well for rooted-tree pipe-networks, it is not sufficient for cyclic networks with multiple supply nodes and non-pipe elements such as compressors. In this case, the resulting state after a limited number of the above algorithm's iterations is used as an initial value for the first order IMEX integrator detailed in Section 5.3.3, which time-steps until a steady-state is sufficiently approximated. This approximate steady-state, associated to a fixed set of boundary values and parameters, is used as initial value for the simulations:

$$\begin{pmatrix} p_0 \\ q_0 \end{pmatrix} = \begin{pmatrix} \bar{p}(\bar{s}_p, \bar{d}_q) \\ \bar{q}(\bar{s}_p, \bar{d}_q) \end{pmatrix}.$$

While other time steppers are applicable, the first order IMEX solver is related to the initial algebraic approximation, due to the synthesis of the source/input and reaction/nonlinear terms.

3 Model Order Reduction

In this section, we summarize the principal approach behind all presented model reduction methods that are extended and tested in this work. The structure of the model laid out in Section 2 is given by (8). For large networks, the differential equations in p and q become high dimensional, which impedes their solution and hence repeated simulations of scenarios. The aim of model reduction is to reduce the dimensionality of the differential equations, by computing subspaces of the phase space on which the trajectories evolve suitably similar (with regard to the quantities of interest). Furthermore, the reduced order model shall have the same form as the original model, and since two physical quantities are (bi-directionally) coupled in this system, the model reduction for interconnected systems [109] approach is used, yielding reduced operators for each subsystem:

$$\begin{aligned} \begin{pmatrix} \tilde{E}_p(\theta) & 0 \\ 0 & I_{n_q} \end{pmatrix} \begin{pmatrix} \dot{\tilde{p}} \\ \dot{\tilde{q}} \end{pmatrix} &= \begin{pmatrix} 0 & \tilde{A}_{pq} \\ \tilde{A}_{qp} & 0 \end{pmatrix} \begin{pmatrix} \tilde{p} + \tilde{p} \\ \tilde{q} + \tilde{q} \end{pmatrix} + \begin{pmatrix} 0 & \tilde{B}_{pd} \\ \tilde{B}_{qs} & 0 \end{pmatrix} \begin{pmatrix} s_p \\ d_q \end{pmatrix} + \begin{pmatrix} 0 \\ \tilde{F}_c \end{pmatrix} + \begin{pmatrix} 0 \\ \tilde{f}_q(\tilde{p} + \tilde{p}, \tilde{q} + \tilde{q}, s_p, \theta) \end{pmatrix}, \\ \begin{pmatrix} s_q \\ d_p \end{pmatrix} &\approx \begin{pmatrix} \tilde{s}_q \\ \tilde{d}_p \end{pmatrix} = \begin{pmatrix} 0 & \tilde{C}_{sq} \\ \tilde{C}_{dp} & 0 \end{pmatrix} \begin{pmatrix} \tilde{p} + \tilde{p} \\ \tilde{q} + \tilde{q} \end{pmatrix}, \\ \begin{pmatrix} \tilde{p}_0 \\ \tilde{q}_0 \end{pmatrix} &= \begin{pmatrix} 0 \\ 0 \end{pmatrix}, \end{aligned}$$

centered around the steady-state $(\bar{p} \ \bar{q})^\top$. This structure-preserving model order reduction was already used in [44, 13] in the context of model reduction for gas networks, while the centering has been used in [6] for gas network simulation and in [55] for nonlinear model order reduction. In the following, the general ansatz to obtain these reduced quantities (denoted by $\tilde{\cdot}$) is summarized.

3.1 Projection-Based Model Reduction

The reduced order model is computed by projecting the high-dimensional dynamics evolving in the (coupled) pressure and mass-flux phase spaces (of dimension N_p and N_q) to low(er)-dimensional subspaces (of dimension n_p and n_q), which capture the principal components of the respective trajectories. Given suitable discrete projection mappings from the original space to the reduced space V_*^\top and mappings from the reduced space back to the original space U_* :

$$\begin{aligned} U_p : \mathbb{R}^{n_p} &\rightarrow \mathbb{R}^{N_p}, & V_p^\top : \mathbb{R}^{N_p} &\rightarrow \mathbb{R}^{n_p} : & V_p^\top \cdot U_p &= \text{id}_{n_p}, \\ U_q : \mathbb{R}^{n_q} &\rightarrow \mathbb{R}^{N_q}, & V_q^\top : \mathbb{R}^{N_q} &\rightarrow \mathbb{R}^{n_q} : & V_q^\top \cdot U_q &= \text{id}_{n_q}. \end{aligned}$$

Thus, the reduced trajectory results from applying V_* to the original trajectory's steady-state deviation, while the original trajectory is approximately recovered by applying U_* to the reduced trajectory:

$$\begin{pmatrix} \tilde{p} \\ \tilde{q} \end{pmatrix} := \begin{pmatrix} V_p^\top (p - \bar{p}) \\ V_q^\top (q - \bar{q}) \end{pmatrix} \rightarrow \begin{pmatrix} \tilde{p} + U_p \tilde{p} \\ \tilde{q} + U_q \tilde{q} \end{pmatrix} \approx \begin{pmatrix} p \\ q \end{pmatrix};$$

the initial condition is also reduced by application of V_* . Similarly, the components of the reduced system result from applying the U_* map to the argument of the respective operators, and the V_* map to the result of the operation. For

the linear operators, the matrices E_p , A_* , B_* and C_* and the vector F_c , this leads conveniently to pre-computable reduced matrices and vector respectively,

$$\begin{aligned}\tilde{A}_{pq} &:= V_p^\top \cdot A_{pq} \cdot U_q \in \mathbb{R}^{n_p \times n_q}, & \tilde{A}_{qp} &:= V_q^\top \cdot A_{qp} \cdot U_p \in \mathbb{R}^{n_q \times n_p}, \\ \tilde{B}_{pd} &:= V_p^\top \cdot B_{pd} \in \mathbb{R}^{n_p \times N_s}, & \tilde{B}_{qs} &:= V_q^\top \cdot B_{qs} \in \mathbb{R}^{n_q \times N_d}, \\ \tilde{C}_{dp} &:= C_{dp} \cdot U_p \in \mathbb{R}^{N_d \times n_p}, & \tilde{C}_{sq} &:= C_{sq} \cdot U_q \in \mathbb{R}^{N_s \times n_q}, \\ \tilde{E}_p(\theta) &:= V_p^\top \cdot E_p(\theta) \cdot U_p \in \mathbb{R}^{n_p \times n_p}, & \tilde{F}_c &:= V_q^\top \cdot F_c \in \mathbb{R}^{n_q},\end{aligned}$$

yet, the nonlinear component \tilde{f}_q remains a composition operation:

$$\tilde{f}_q := V_q^\top \cdot f_q(\bar{p} + U_p \tilde{p}, \bar{q} + U_q \tilde{q}, s_p, \theta) : \mathbb{R}^{n_p} \times \mathbb{R}^{n_q} \times \mathbb{R}^{N_s} \times \mathbb{R}^2 \rightarrow \mathbb{R}^{n_q}. \quad (9)$$

3.2 The Lifting Bottleneck and Hyper-Reduction

The gas network models considered for reduction are nonlinear (and possibly non-smooth), hence the reduced order nonlinear part involves lifting the reduced state up to the original high-dimensional space, evaluating the nonlinearity and projecting the result back down to the reduced low-dimensional space (9). As the high-dimensional space is involved, this is typically computationally demanding and may eat up the gains from the reduction of the linear part. To mitigate this so-called lifting bottleneck, hyper-reduction methods can be employed, which construct low-dimensional surrogates for nonlinearities.

In this work we discard (or rather defer) hyper-reduction due to the following reasoning: The purpose of this work is to determine which method constructs the best reduced order models, a hyper-reduction may inhibit comparability due to, for example, a dominating hyper-reduction approximation error. Second, various hyper-reduction methods for this setting are applicable (i.e. DEIM [20], DMD [128] or numerical linearization [89]), which may interact differently with the different model reduction methods. So as a first step, the bare model reduction methods are tested here (this means: which method's linear subspaces capture the nonlinear dynamics best), at a later stage the best hyper-reduction method can then be determined. Lastly we note, the nonlinear part of the vector field consists exclusively of element-wise operations, a system with repeated scalar nonlinearities (SRSN) [22], which are less difficult to handle due to "locality" of the nonlinearity.

3.3 Parametric Model Reduction

There are two common approaches for projection-based parametric model order reduction: averaging and accumulating [58]. For the selected data-driven methods, *averaging* means that for a set of parameter samples the associated trajectories or derived quantities (such as the utilized system Gramians) are averaged, while *accumulating* refers to the concatenation of trajectories or derived quantities (such as the projectors). Generally, each of the methods in Section 4 can be used with either averaging or accumulating, we opted to use the averaging ansatz for all of the following methods since their computation is without exception based on parametric empirical Gramians [62]: Structured Proper Orthogonal Decomposition, Structured Empirical Dominant Subspaces, Structured Empirical Balanced Truncation, Structured Empirical Balanced Gains, and Structured Dynamic Mode Decomposition - Galerkin.

4 Model Reduction Methods

In this section, we briefly summarize the employed model reduction methods from a practical point of view. For theoretical details and backgrounds we refer to the relevant papers, cited in the respective subsections. Due to the non-differentiable nonlinearity (friction), the sought projections U_* , V_* for all tested model reduction techniques are constructed from (transformed) time-domain trajectory data obtained from numerical simulations, which is given by discrete-time snapshots of the internal pressure nodes \widehat{X}_p and mass-flux edges \widehat{X}_q , as well as the demand node pressure and supply node mass-flux \widehat{Y}_p , \widehat{Y}_q :

$$\begin{aligned}\widehat{X}_p(t; \theta_k) &= [p^1(t; \theta_k) \quad \dots \quad p^{N_s+N_d}(t; \theta_k)], \\ \widehat{X}_q(t; \theta_k) &= [q^1(t; \theta_k) \quad \dots \quad q^{N_s+N_d}(t; \theta_k)], \\ \widehat{Y}_p(t; \theta_k) &= \begin{bmatrix} (s_q^1(t; \theta_k)) & \dots & (s_q^{N_p}(t; \theta_k)) \\ (d_p^1(t; \theta_k)) & \dots & (d_p^{N_p}(t; \theta_k)) \end{bmatrix}, \\ \widehat{Y}_q(t; \theta_k) &= \begin{bmatrix} (s_q^1(t; \theta_k)) & \dots & (s_q^{N_q}(t; \theta_k)) \\ (d_p^1(t; \theta_k)) & \dots & (d_p^{N_q}(t; \theta_k)) \end{bmatrix}.\end{aligned}$$

The state-space trajectories $\widehat{X}_*(t; \theta_k)$ are obtained for perturbations of the inputs, pressure at the N_s supply boundary nodes and mass-flux at the N_d demand boundary nodes, while the output trajectories $\widehat{Y}_*(t; \theta_k)$ are computed for perturbations in the respective N_* steady-state components. The training parameters θ_k are sampled from a sparse grid spanning the parameter space $\Theta \subset \mathbb{R}^2$.

All methods are prefixed ‘‘Structured’’, since the model structure of a pressure and mass-flux variable is preserved in the reduced order model⁷. Practically, this means while \widehat{X}_p and \widehat{X}_q are computed simultaneously due to their coupling, the individual projectors for pressure and mass-flux are constructed separately. The subsequent methods may have not been previously introduced explicitly in structured form, yet given [122, 109] introducing structured Gramians, these are trivial extensions. For ease of notation, we describe the computation of projectors $\{U_p, V_p\}$ and $\{U_q, V_q\}$ as generically for $\{U_*, V_*\}$.

We implemented a total of twelve model reduction method variants, which we compare in this work. All tested methods are data-driven and time-domain focused, as the dynamic gas network model (8) is nonlinear. Furthermore, all model reduction methods construct linear subspaces and are derived from methods for linear systems, yet differ from plain linearization: instead, the implemented methods assemble linear subspaces of the model’s phase space that, in a method specific sense, approximately enclosing the relevant nonlinear system evolution. Moreover, all methods are SVD-based [3], and their majority is originally based on (empirical) system Gramian matrices, for details see [56]. We highlight here that the time horizon for the training data is significantly shorter than for the actual simulations the reduced order model is targeted at. Furthermore, generic training inputs (transient boundary values), such as impulse, step or random signals, are utilized to avoid a *model reduction crime* [58] (comparable to an inverse crime): *Train* a reduced order model using the *test* parameters or inputs.

⁷All methods have initially also been tested in an unstructured variant, too, yet these often produced unstable and less accurate reduced models.

4.1 Empirical System Gramians

All model reduction methods currently included in `morgen` are computationally realized using empirical system Gramian matrices, which are system-theoretic operators encoding reachability and observability. From these operators, information on importance of linear combinations of states can be extracted. For linear systems, these system Gramians are typically computed via matrix equations, for general nonlinear systems there is no *feasible* closed form. However, the empirical system Gramians approximate the nonlinear Gramians locally based on state and trajectory data. The empirical Gramians used in this context are described in-depth in [56]; here only a brief summary for the gas network setting is given.

4.1.1 Empirical Reachability Gramian Matrix

Reachability quantifies how well a system can be driven by the inputs, which is encoded by the reachability Gramian. The empirical reachability Gramian is an approximation based on state trajectory data [74]:

$$\widehat{W}_{R,*} := \sum_{k=1}^K \sum_{m=1}^{N_s+N_d} \int_0^\infty \widehat{X}_*^m(t; \theta_k) \widehat{X}_*^m(t; \theta_k)^\top dt \in \mathbb{R}^{N_* \times N_*}. \quad (10)$$

Given a suitable set of input perturbations, $\widehat{W}_{R,*}$ approximates the nonlinear reachability Gramian near a steady-state [46].

4.1.2 Empirical Observability Gramian Matrix

Observability quantifies how well the state can be characterized from the outputs, which is encoded by the observability Gramian. The empirical observability Gramian is an approximation based on output trajectory data [74]:

$$\widehat{W}_{O,*} := \sum_{k=1}^K \int_0^\infty \widehat{Y}_*^m(t; \theta_k)^\top \widehat{Y}_*^m(t; \theta_k) dt \in \mathbb{R}^{N_* \times N_*}. \quad (11)$$

For a suitable set of steady-state perturbations, $\widehat{W}_{O,*}$ approximates the nonlinear observability Gramian near a steady-state [46].

4.1.3 Empirical Cross Gramian Matrix

The empirical cross Gramian concurrently encodes reachability and observability, which in conjunction quantifies redundancy also known as minimality; however, the (empirical) cross Gramian is only applicable for square systems, systems with the same number of inputs and outputs (which the gas network model (8) fortunately is), and an approximation is based on simulated state and output trajectory data [61]:

$$\widehat{W}_{X,*} := \sum_{k=1}^K \sum_{m=1}^{N_s+N_d} \int_0^\infty \widehat{X}_*^m(t; \theta_k) \widehat{Y}_*^m(t; \theta_k) dt \in \mathbb{R}^{N_* \times N_*}. \quad (12)$$

For a suitable set of input and steady-state perturbations, $\widehat{W}_{X,*}$ approximates the nonlinear cross Gramian near a steady-state [61].

4.1.4 Empirical Non-Symmetric Cross Gramian Matrix

A generalization of the empirical cross Gramian for non-square systems is the empirical non-symmetric cross Gramian, which is an approximation based on simulated state and output trajectory data [63]:

$$\widehat{W}_{Z,*} := \sum_{k=1}^K \sum_{m=1}^{N_s+N_d} \sum_{q=1}^{N_s+N_d} \int_0^\infty \widehat{X}_*^m(t; \theta_k) \widehat{Y}_*^q(t; \theta_k)^\top dt \in \mathbb{R}^{N_* \times N_*}. \quad (13)$$

For a suitable set of input and steady-state perturbations, $\widehat{W}_{Z,*}$ approximates the nonlinear cross Gramian near a steady-state. Even though the gas network model (8) is square, the empirical non-symmetric cross Gramian is included here, since, heuristically, it could provide better results than the regular cross Gramian [63].

4.2 Structured Proper Orthogonal Decomposition

Proper orthogonal decomposition (POD) is a basic data-driven method for model reduction: given a matrix of state snapshots over time, the dominant left singular vectors are computed as a basis via a singular value decomposition (SVD) [88]. The basis vectors are assigned their principality with respect to the conveyed energy by the associated (relative) singular value magnitude. In the context of this work, the POD is constructed from a system-theoretic point of view, that connects to the system property of reachability. Due to the overall structured approach to model reduction, a structured POD refers in this context to the separate PODs for pressure and mass-flux variables p and q as in [44, 42, 13].

4.2.1 Reachability-Gramian-Based

The singular vectors to the principal singular values of the empirical reachability Gramian (10) correspond to the POD modes. To obtain a reduced order model, first, a truncated SVD (tSVD) of the empirical reachability Gramian,

$$W_{R,*} \stackrel{\text{tSVD}}{=} U_{R,*} D_{R,*} U_{R,*}^\top,$$

reveals the principal subspace of the respective trajectories $\widehat{X}_*^m(t)$, with respect to the default L_2 inner product, whereas the importance of each basis vector (column) in $U_{R,*}$ is determined by (the square-root of) the associated singular value $\sigma_i := D_{R,*}^{1/2}$:

$$U_* := U_{R,*}.$$

The matrix of basis vectors (POD modes) constitutes a Galerkin projection $V_* := U_*$. Notably, (structured) POD only considers the input-to-state mapping, not the state-to-output mapping, and hence approximates the state variables, p and q , not the output quantities of interest s_q and d_p .

The POD could also be computed directly from an SVD of the trajectory data, yet the computational overhead of using the empirical reachability Gramian is small compared to the trajectory simulation runtimes, and the systematic perturbations of the empirical Gramian approach [56] are exploited.

4.3 Structured Empirical Dominant Subspaces

The previous (structured) POD method considers only the reachability information, hence the data only reflects the input-to-state mappings, and thus the POD derived ROMs (Reduced Order Model) approximate the state variables p and q . To approximate the outputs s_q and d_p , the state-to-output mappings, encoding observability information, need to be considered, too.

The (empirical) dominant subspaces method initially developed in [100], and originally named DSPMR (Dominant Subspace Projection Model Reduction), conjoins and compresses the dominant reachability and observability subspaces of an input-output system, such as the gas network model (8), obtained from (empirical) system Gramians. Heuristically, this method seems to be useful for hyperbolic input-output systems [40]. Here, we consider three variants: first, based on the empirical reachability and observability Gramians, second, based on the empirical cross Gramian and third, on the empirical non-symmetric cross Gramian.

4.3.1 Reachability- and Observability-Gramian-Based

The singular vectors associated to the principal singular values of the empirical reachability and observability Gramians span these dominant subspaces, which are first extracted by tSVDs, and then, after concatenation, compressed by orthogonalization via another tSVD:

$$\begin{aligned}\widehat{W}_{R,*} &\stackrel{\text{tSVD}}{=} U_{R,*} D_{R,*} V_{R,*}^\top, \\ \widehat{W}_{O,*} &\stackrel{\text{tSVD}}{=} U_{O,*} D_{O,*} V_{O,*}^\top, \\ [(\omega_R U_{R,*} D_{R,*}) \quad (\omega_O U_{O,*} D_{O,*})] &\stackrel{\text{tSVD}}{=} U_{RO,*} D_{RO,*} V_{RO,*}^\top,\end{aligned}$$

from which the singular vectors $V_* = U_* := U_{RO,*}$ make up a Galerkin projection. The weights $\omega_R := \|\widehat{W}_{R,*}\|_F^{-1}$, and $\omega_O := \|\widehat{W}_{O,*}\|_F^{-1}$ equilibrate the potentially mismatched scales of the respective Gramians, akin to the *refined DSPMR* method from [100].

4.3.2 Cross-Gramian-Based

A truncated SVD of the cross Gramian also engenders the sought dominant subspaces [15]: The (empirical) cross Gramian's left and right singular vectors (approximately) span the reachability and observability subspaces, respectively, and their orthogonalized concatenation, via a truncated SVD,

$$\begin{aligned}\widehat{W}_{X,*} &\stackrel{\text{tSVD}}{=} U_{X,*} D_{X,*} V_{X,*}^\top, \\ [(U_{X,*} D_{X,*}) \quad (V_{X,*} D_{X,*})] &\stackrel{\text{tSVD}}{=} U_{RO,*} D_{RO,*} V_{RO,*}^\top,\end{aligned}$$

yields a Galerkin projection $V_* = U_* := U_{RO,*}$. The cross-Gramian-based variant does not need to additionally weight the reachability subspace $U_{X,*}$ and observability subspace $V_{X,*}$, as they are both extracted from the same matrix. This cross-Gramian-based empirical dominant subspaces method seamlessly extends to the empirical non-symmetric cross Gramian $\widehat{W}_{Z,*}$.

4.4 Structured Empirical Balanced Truncation

The dominant subspaces method combines separately quantified input-to-state and state-to-output energies, but not the actual input-to-output energy. Such can be accomplished by balanced truncation and based on the Hankel operator, which maps past inputs to future outputs. This operator's singular values measure the sought input-to-output energy, and the singular vectors constitute a basis. To obtain the Hankel operator's truncated SVD, first, the underlying system needs to be balanced, and then singular vectors associated to small magnitude singular values are truncated.

Balanced truncation is the reference model reduction method for linear input-output systems, due to stability preservation in the ROMs and a computable error bound. For (control-affine) nonlinear input-output systems, such as the gas network model (8), balanced truncation can be generalized to *empirical balanced truncation* [74], while a structured variant is introduced as *interconnected system balanced truncation* [122], which we combine.

Again, we consider three variants: First, based on the empirical reachability and observability Gramian, second, based on the empirical cross Gramian and third, on the empirical non-symmetric cross Gramian; additionally, the reachability and observability-based balanced POD variant is included.

4.4.1 Reachability- and Observability-Gramian-Based

The original balanced truncation method is based on the reachability and observability Gramians [90]. A transformation into a balanced coordinate system in which both system Gramians are diagonal and equal is obtained via simultaneous diagonalization. Various balancing algorithms are available for this task [123], in this setting we selected the general balancing algorithm [107, 108], which utilizes the magnitude-based truncated eigenvalue-decomposition (tEVD) of the Gramians: The matrices U_* , V_* constitute a Petrov-Galerkin projection, whereas the importance of each column is determined by the approximate Hankel singular values, the (diagonal) elements of $D_{B,*}$:

$$\begin{aligned}
 \widehat{W}_{R,*} \widehat{W}_{O,*} T_{R,*} &\stackrel{\text{tEVD}}{=} \Lambda_{R,*} T_{R,*}, \\
 (\widehat{W}_{R,*} \widehat{W}_{O,*})^\top T_{O,*} &\stackrel{\text{tEVD}}{=} \Lambda_{O,*} T_{O,*}, \\
 T_{O,*}^\top T_{R,*} &\stackrel{\text{tSVD}}{=} U_{B,*} D_{B,*} V_{B,*}, \\
 U_* &:= T_{R,*} V_{B,*} D_{B,*}^{-\frac{1}{2}}, \\
 V_* &:= T_{O,*} U_{B,*} D_{B,*}^{-\frac{1}{2}}.
 \end{aligned} \tag{14}$$

4.4.2 Cross-Gramian-Based

For linear, symmetric systems, alternatively a balanced and truncated reduced order model can be computed via the cross Gramian. Yet, the gas network model (8) is neither linear nor symmetric, not even in a nonlinear sense of symmetric systems, i.e. gradient systems [66], but the considered input-output system is square. Hence, an empirical cross-Gramian-based reduced model is computable, but it will differ from the reduced model obtained by reachability- and observability-Gramian-based balanced truncation.

Given a cross Gramian with full rank, an (approximate) balancing projection is computable in a similar manner as for the (empirical) balanced truncation (14), but based on the left and right eigen spaces of the cross Gramian [68]:

$$\begin{aligned}
\widehat{W}_{X,*} T_{R,*} &\stackrel{\text{tEVD}}{=} \Lambda_* T_{R,*} \\
\widehat{W}_{X,*}^\top T_{O,*} &\stackrel{\text{tEVD}}{=} \Lambda_* T_{O,*} \\
T_{O,*}^\top T_{R,*} &\stackrel{\text{tSVD}}{=} U_{X,*} D_{X,*} V_{X,*}, \\
U_* &:= T_{R,*} V_{X,*} D_{X,*}^{-\frac{1}{2}}, \\
V_* &:= T_{O,*} U_{X,*} D_{X,*}^{-\frac{1}{2}}.
\end{aligned} \tag{15}$$

The matrices U_* , V_* again constitute a Petrov-Galerkin projection, and the importance of each column is determined by the absolute value of the (diagonal) elements of $D_{X,*}$, which are only equal to the approximate Hankel singular values for linear and symmetric systems.

As for the empirical dominant subspaces method, the cross-Gramian-based empirical balanced truncation variant directly extends to the non-symmetric cross Gramian $\widehat{W}_{Z,*}$.

One could assume that if only one system Gramian has to be computed, instead of two for dominant subspaces or balanced truncation, that the cross-Gramian-based computation is significantly faster, but the overall number of simulated trajectories is the same for both methods, which causes the dominant fraction of computational cost. Thus, the empirical cross Gramian computation is merely slightly quicker than the computation of both empirical reachability and observability Gramians.

4.4.3 Structured Balanced POD

Instead of balancing the system using the product of the (empirical) reachability and observability Gramians, the dominant subspaces of the respective Gramians can be used to approximately balance the system. This variant of reachability- and observability-Gramian-based balanced truncation is called balanced POD [127], and the approximate balancing algorithm reads:

$$\begin{aligned}
\widehat{W}_{R,*} T_{R,*} &\stackrel{\text{tEVD}}{=} \Lambda_{R,*} T_{R,*}, \\
\widehat{W}_{O,*} T_{O,*} &\stackrel{\text{tEVD}}{=} \Lambda_{O,*} T_{O,*}, \\
T_{O,*}^\top T_{R,*} &\stackrel{\text{tSVD}}{=} U_{B,*} D_{B,*} V_{B,*}, \\
U_* &:= T_{R,*} V_{B,*} D_{B,*}^{-\frac{1}{2}}, \\
V_* &:= T_{O,*} U_{B,*} D_{B,*}^{-\frac{1}{2}},
\end{aligned} \tag{16}$$

with the matrices U_* , V_* inducing a Petrov-Galerkin projection.

Here, we categorized balanced POD as a variant of balanced truncation, due the algorithmic similarity to the balancing algorithm (14). Alternatively, we could have classified balanced POD as a variant of POD, since it can be described as POD with the observability Gramian defining the POD's inner product [105].

We leave it to the reader to test other balancing algorithms, e.g. [123, 12], while we explicitly excluded the modified POD [93, 58], as it is not a Petrov-Galerkin method.

4.5 Structured Empirical Balanced Gains

The empirical balanced gains method is a variant of the empirical balanced truncation method [58]: While balanced truncation selects principality of subspaces based on the Hankel singular value magnitude, balanced gains sorts by the impulse response norm:

$$\begin{aligned} \|s_q\|_2 &\approx \sqrt{\text{tr}(C_{sq}\widehat{W}_{R,q}C_{sq}^\top)} \approx \sqrt{\text{tr}(B_{qs}^\top\widehat{W}_{O,q}B_{qs})} \\ &\approx \sqrt{|\text{tr}(C_{sq}\widehat{W}_{X,q}B_{qs})|} \approx \sqrt{|\text{tr}(C_{sq}\widehat{W}_{Z,q}B_{qs})|}, \\ \|d_p\|_2 &\approx \sqrt{\text{tr}(C_{dp}\widehat{W}_{R,p}C_{dp}^\top)} \approx \sqrt{\text{tr}(B_{pd}^\top\widehat{W}_{O,p}B_{pd})} \\ &\approx \sqrt{|\text{tr}(C_{dp}\widehat{W}_{X,p}B_{pd})|} \approx \sqrt{|\text{tr}(C_{dp}\widehat{W}_{Z,p}B_{pd})|}, \end{aligned}$$

for a system in balanced form. Hence, either method, balanced truncation and balanced gains compute the same balancing transformation, via (14) or (15), but the sequence of basis vectors differs due to the variant measures.

We note, that due to the linear input and output operators, the balanced gains method can be directly applied for gas network models (8). However, for general nonlinear control systems, linear input and operators need to be obtained via linearization [80], or system identification [76].

4.6 Structured Dynamic Mode Decomposition - Galerkin

In addition to the four energy-based method classes (Sections 4.2, 4.3, 4.4, 4.5), also an alternative method from [1], based on system identification, is investigated. However, it is utilized as a structured projection-based model reduction method with separate projections for pressure and mass-flux. After summarizing the parent system identification method, the derived model reduction technique is presented.

4.6.1 Dynamic Mode Decomposition

Dynamic mode decomposition (DMD) identifies a discrete-time operator from (discrete) trajectory time-series data, preserving certain modal behavior [106]. This system identification method is based on the Koopman operator, which is an infinite dimensional, but linear operator, mapping (a transformation, or observable of the) state x^k at discrete-time step k to $(k+1)$. DMD yields a (linear) finite-dimensional approximation of the Koopman operator preserving its dominant eigenmodes. Here, using the identity observable and given time series data $X = [x^0 \ \dots \ x^K]$, DMD identifies an operator \widehat{A} , via least-squares:

$$x^{k+1} \approx \widehat{A}x^k, \quad (17a)$$

$$\widehat{A} := [x^1 \ \dots \ x^K] [x^0 \ \dots \ x^{K-1}]^+. \quad (17b)$$

Since the underlying model is a control system, DMD with control (DMDc) [102] (and known input operator B), i.e. using $X^c := [x^0 - Bu^0 \ \dots \ x^K - Bu^K]$ instead of X for DMD, is applicable, yet, due to the perturbed steady-state training of ROMs (see Section 6.1) not beneficial.

4.6.2 Reachability-Gramian-Based DMD-Galerkin

DMD is rather a system identification than a model reduction method. To fit into the projection setting, we utilize the DMD-Galerkin method [1], which forms a Galerkin projection U from the orthogonalized dominant eigenvectors (based on the associated eigenvalue magnitudes),

$$\begin{aligned} \widehat{A}T &\stackrel{\text{EVD}}{=} T\Lambda, \\ \rightarrow T\Lambda &\stackrel{\text{SVD}}{=} UDV^\top. \end{aligned}$$

Since only state-space trajectories are utilized, as for the POD, the DMD-Galerkin method approximates the state, not the output. Curiously, we note, that this method uses discrete-time information to assemble projections for continuous-time systems.

Practically, U_* are computed as a subset of singular vectors for the largest magnitude singular values, instead of orthogonalized principal eigenvectors. In the structured setting at hand $\widehat{X}_* = [x_*^0 \ \dots \ x_*^K]$, the Galerkin projection $U_* = V_*$ is given by:

$$[x_*^1 \ \dots \ x_*^K] [x_*^0 \ \dots \ x_*^{K-1}]^+ \stackrel{\text{tSVD}}{=} U_* D_* \widetilde{V}_*^\top.$$

Effectively, the approximate Koopman operator \widehat{A} is computed via an empirical reachability Gramian, yet instead of the standard inner product, the DMD-“kernel” (17b) is used. Additionally, and in line with the original empirical Gramians [74], the utilized trajectories are centered, following [64]. This means theoretically, DMD-Galerkin is POD with a specific kernel, and practically, that by computation via an empirical Gramian the systematic perturbation properties can also be exploited for DMD-Galerkin.

5 morgen

The **morgen** (Model Order Reduction for Gas and Energy Networks) platform implements the mathematical methods presented above, in MATLAB ($\geq 2020b$) and compatible to Octave (≥ 6.1). Compared to, for example [52], **morgen** does not feature a graphical user interface or Simulink integration, since it is designed for batch testing on (headless) workstations and multi-query supercomputer use. The source code is organized into five main components:

- networks** (holds network and scenario data-sets)
- models** (discretizes networks and assembles input output systems)
- solvers** (computes solution time series for discrete models and scenarios)
- reducers** (reduces state variables of discrete models)
- tests** (defines experiments for data-model-solver-reductor combinations).

These components are briefly described in the following. For an illustration of the internal structure of **morgen**, see Fig. 2. Further code that is used by the main function or multiple components is contained in an **utils** folder, and the stand-alone network format converters⁸ are stored in a **tools** folder.

⁸Converters for GASLIB XML and *MathEnergy* JSON data-sets are available in **morgen**.

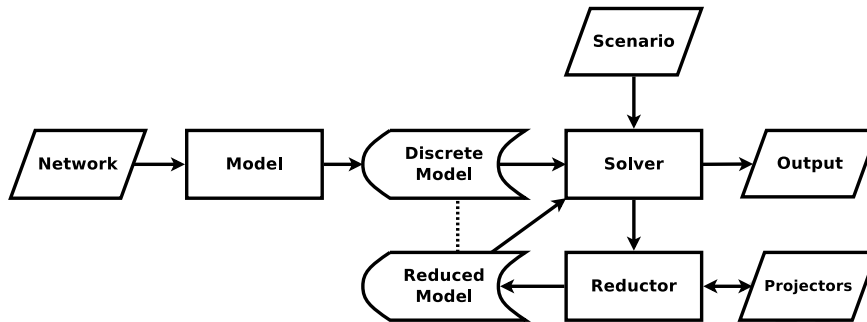


Figure 2: Internal data flow and process of morgen.

5.1 Networks

The `networks` directory stores the network `.net` files, and a folder for each network with the same name as the associated network file's base name, which hold the scenario definition files. The `.net` file, in comma-separated value (CSV) format, defines the network topology and gas network components through an edge list. Each row encodes one edge from the information: edge type (pipe, shortcut, compressor, valve), "from"-node, "to"-node, length, diameter, incline, roughness; the latter three are only relevant for pipes.

Note that boundary nodes have to be leaf nodes of the network graph in order to be identified by `morgen` as such. Furthermore, if a boundary node shall act as supply and demand, it needs to be artificially split into two leaf nodes, as the edge connecting a supply node has to be directed from the leaf, while the edge connecting a demand node has to be directed towards the leaf (Section 2.3).

5.1.1 Scenario

A scenario is defined via a set of key-value pairs in an `.ini` file, whereas sequence of pairs does not matter. Each network has at least a generic `training.ini` scenario. The following keys are mandatory for a scenario definition:

- Mean temperature `T0`.
- Mean specific gas constant `RS`.
- Time horizon `tH`.
- List of time instances `ut`.
- List of supply node pressures `up` at time instances `ut`.
- List of demand node mass-fluxes `uq` at time instances `ut`.

Depending on the network composition, the following keys may need to be provided:

- List of compressor discharge pressures `cp`.
- List of valve settings `vs`.

Note that amongst other configurations, the parameter ranges of temperature and specific gas constant for the parametric model reduction are set in the `morgen.ini` file.

5.2 Models

A model encodes a spatial discretization of the simplified Euler equations (8) on a gas network topology in a structure with the members: A , B , C , E , F , f and the Jacobian J ([116, 2]), together with the system dimensions in terms of number of pressure-at-nodes and mass-flux-on-edges states, and total number of boundary nodes. It is ensured during the assembly of the model that the sparsity of the model components is preserved.

The model interface is given by the following signature:

```
discrete = model(network,config);
```

While the linear components A , B , C , and F are provided as sparse matrices, the parameter-dependent linear component E is a closure⁹ returning a sparse matrix, and the nonlinear component f and the (nonlinear) Jacobian J are closures returning the application of a state (as well as steady-state, input, parameters, compressibility).

Two spatial discretizations are currently provided in `morgen`:

- Midpoint ODE Discretization (Section 2.4.1): `ode_mid`
- Endpoint ODE Discretization (Section 2.4.2): `ode_end`

Even though both provided models are ODEs, DAE models can also be implemented, given a solver (and reductor) is available.

5.3 Solvers

The `morgen` platform provides four solvers: An adaptive step-size method, a fixed step-size explicit method and two fixed step-size implicit-explicit methods. The solver interface is given by the following signature:

```
solution = solver(discrete,scenario,config);
```

with the return value `solution`, being a structure, and the arguments `discrete` (model), `scenario`, and `config`(uration).

All provided fixed step-size solvers cache matrix decompositions. The initial steady-state is also cached, as is the QR decomposition used to compute it.

Even though the overall model (8) has a two dimensional structure and the reductors exploit this structure, the simulations itself can be performed on a lumped model (we omit parametrization here for ease of notation):

$$\begin{aligned} E\dot{x}(t) &= Ax(t) + Bu(t) + f(x(t), u(t)), \\ y(t) &= Cx(t). \end{aligned} \tag{18}$$

Practically, the vector field is split slightly different as shown in the Appendix.

5.3.1 Second-Order Adaptive Implicit Solver: `generic`

For validation purposes, the adaptive step-size solver for stiff systems `ode23s`¹⁰ included in MATLAB (and Octave) [113, Sec. 2.3], based on a modified second order Rosenbrock formula is used and encapsulated as a generic solver. Due to preferential performance demonstrated in [42], `ode23s` is preferred over alternatives such as `ode15s` (or `ode45`).

⁹A closure is a pair of a function together with its scoped environment.

¹⁰<https://mathworks.com/help/matlab/ref/ode23s.html> (accessed: 2020-11-18)

5.3.2 Fourth-Order “Classic” Runge-Kutta Solver: rk4

Since in [94, 95], the fourth-order explicit Runge-Kutta (RK) method [73] is employed, `morgen` provides it, too. This method is strong stability preserving [38], however not SSP-optimal. Expectedly, it works only for very small time-steps.

5.3.3 First-Order Implicit-Explicit Solver: imex1

The lumped gas network model (18) can be split into a linear and a nonlinear part, of which the linear part is numerically stiff and hence should be solved with an implicit solver, while an explicit solver is preferred for the nonlinear part as to avoid solving a root-finding (optimization) problem in each time step. An IMEX method allows this separate treatment of operators and is thus suitable for this hyperbolic and nonlinear system. Combining the first-order explicit Euler’s method with the first order implicit Euler’s method yields the first order IMEX method:

$$\begin{aligned} Eh^{-1}(x_{k+1} - x_k) &= (1 - \gamma)Ax_k + \gamma Ax_{k+1} + Bu_k + f(x_k, u_k) \\ \Rightarrow Ex_{k+1} - \gamma hAx_{k+1} &= Ex_k + (1 - \gamma)hAx_k + hBu_k + hf(x_k, u_k) \\ &\Rightarrow x_{k+1} = x_k + (E - \gamma hA)^{-1}h(Ax_k + Bu_k + f(x_k, u_k)), \end{aligned}$$

with the associated Butcher tableaux Table 2. Even though this IMEX method is not a Runge-Kutta method [4], it was successfully applied to gas network models in [43] with a relaxation parameter set to $\gamma = 1$.

Explicit:	Implicit:
$\begin{array}{c c} 0 & 0 \\ \hline & 1 \end{array}$	$\begin{array}{c c} 1 & 1 \\ \hline & 1 \end{array}$

Table 2: Butcher tableaux for the 1st-order IMEX method (Section 5.3.3).

5.3.4 Second-Order Implicit-Explicit Runge-Kutta Solver: imex2

A second-order (two-stage) IMEX Runge-Kutta solver is provided, based on the combination of a second-order explicit SSP Runge-Kutta method [38], and a second-order DIRK method. Following [99], such an IMEX-SSP2(2,2,2) method with relaxation γ is given by:

$$\begin{aligned} z_1 &= (E - h\gamma\lambda A)^{-1} Ex_k, \\ z_2 &= (E - h\gamma\lambda A)^{-1}(Ex_k + hBu_k + hf(x_k, u_k) + h\gamma(1 - 2\lambda)Az_1), \\ x_{k+1} &= x_k + E^{-1}\frac{h}{2}(Bu_k + f(x_k, u_k) + \gamma Az_1 + Bu_{k+1} + f(z_1, u_{k+1}) + \gamma Az_2). \end{aligned}$$

The explicit component of this IMEX method is SSP-optimal [38], while depending on the choice for the free parameter λ , different properties of the implicit component can be achieved (see Table 3). Practically, we found $\lambda = \frac{1}{2}$, making the implicit part SDIRK (Singly DIRK) and stiffly accurate, to work best. Additionally, we would like to highlight passive Runge-Kutta methods [92], specifically PDIRK (passive DIRK), as implicit IMEX component, which have various desirable stability and conservation properties [35].

Explicit:			Implicit:		
0	0	0	λ	λ	0
1	1	0	$1 - \lambda$	$1 - 2\lambda$	λ
	$\frac{1}{2}$	$\frac{1}{2}$		$\frac{1}{2}$	$\frac{1}{2}$

Table 4: Butcher tableaux for the 2nd-order IMEX-RK method (Section 5.3.4).

Parameter	Property	Source
$\lambda = 0.24$	“Efficient”	[72]
$\lambda \geq \frac{1}{4}$	A-stable	[67]
$\lambda = \frac{2-\sqrt{2}}{2} = 1 - \frac{1}{\sqrt{2}}$	L-Stable	[99]
$\lambda = \frac{1}{2}$	Stiffly accurate	<code>morgen</code> , cf. [67, Sec. 5.1]
$\lambda = \frac{3+\sqrt{3}}{6} = \frac{1}{2} + \frac{1}{\sqrt{12}}$	Passive	[92], cf. [11, Sec. 3.2.2]

Table 3: Parameter choices for the implicit 2nd-order IMEX-RK component.

The associated Butcher tableaux are given in Table 4. In our experiments, the first-order IMEX method Section 5.3.3 allowed larger time-steps and exhibited less numerical oscillations or artifacts compared to the second-order IMEX-RK methods. The generic (adaptive) method Section 5.3.1 also solves sufficiently accurate, but takes longer to compute. Thus, by default, we recommend the first-order IMEX integrator for gas network simulations.

5.4 Reductors

The reductor module provides methods that compute (Petrov-)Galerkin projectors for a given discretization. These projectors can be stored on disk for reuse. In the current version of `morgen`, the twelve reductors, described in Section 4 are included:

- Structured Proper Orthogonal Decomposition: `pod_r`
- Structured Empirical Dominant Subspaces: `eds_ro`, `eds_wx`, `eds_wz`
- Structured Balanced Proper Orthogonal Decomposition: `bpod_ro`
- Structured Empirical Balanced Truncation: `ebt_ro`, `ebt_wx`, `ebt_wz`
- Structured Empirical Balanced Gains: `ebg_ro`, `ebg_wx`, `ebg_wz`
- Structured Dynamic Mode Decomposition Galerkin: `dmd_r`

The reductors have the interface:

```
ROM = reductor(solver, discrete, scenario, config);
```

returning a closure `ROM`, which in turn returns a reduced-order model for a given `discrete` model, projection matrices and reduced order, as well as the arguments `solver`, `discrete` model, and `configuration`; the latter containing the sampled parameters available to the reductor. The projectors specific to model and solver defining the ROM closure are stored in a `.rom` file.

5.4.1 Empirical Gramian Framework

The compute backend for the model reduction methods is `emgr` – empirical Gramian framework [56]; currently in version 5.8 [57]. This (open-source) Octave and MATLAB toolbox computes the empirical Gramians, which are essential to construct the reduced order models via the Gramian-based model reduction methods from Section 4, including the structured DMD-Galerkin method.

5.5 Tests

A test is a script defining an experiment by specifying network, scenario, model, solver and reducers. The `tests` component is a collection of test scripts probing primarily model reduction for various networks. A typical test contains two calls to the main `morgen` function. The first call computes the reduced order model (offline phase):

```
morgen(network,training_scenario,model,solver,reducers);
```

which computes a ROM from a short, generic, steady-state `training_scenario`. The projectors defining the reduced order models are then stored in `rom_files`. The second call tests the reduced order model(s) on a longer `test_scenario` (online phase):

```
morgen(network,test_scenario,model,solver,rom_files);
```

Generally, a model reduction method can also be tested in a single call, disregarding that scenario’s boundary value time series, yet for productive use, a reduced order model is constructed once (first call). and then employed for many different scenarios (second call).

Included in `morgen` are two types of tests: First, tests prefixed with “`sim_`” only simulate the test scenarios, second, tests prefixed with “`mor_`” compute the reduced order models using training scenarios, and benchmark these ROMs on the test scenarios.

6 Numerical Experiments

In the following, we present three sets of numerical experiments, with the purpose of demonstrating the reducibility of gas network models via the data-driven, parametric, system-theoretic model order reduction algorithms from Section 4, and illustrating the capabilities of the `morgen` framework summarized in Section 5. The first set uses a pipeline “`network`”, which is interesting in the context of model reduction, while the second set tests an academic toy network as a sanity check and a simple functionality test. Lastly, a realistic gas network topology is evaluated.

We note that various further networks are included for testing in `morgen`; among others: the CANVEY-LEEDS network [45, 69], the Belgium transport network [25, 86] and a part of the FERMACA network [103]. A synthetic pipeline model and associated simulation results were provided by the PSI SOFTWARE AG for validation of `morgen` against the commercial `PSIganesi`¹¹ solver.

¹¹<https://www.psigasandoil.com>

6.1 Workflow

For each of the numerical tests, the same workflow is employed, which is composed of a training phase (offline phase), in that the ROMs are computed, using a generic test scenario, with a (virtual) time horizon of 1h, and boundary value input functions typical for system identification, i.e. Dirac impulse, step signal, random-binary signal or Gauss noise [91, Ch. 16]. In the test phase (online phase), the ROMs are tested on scenarios with a (virtual) time horizon of 24h [21, 81], (starting at 6am [32]). In addition to shorter offline phases, this difference in training and test time horizons emphasizes generality of the ROMs. To verify models and solvers [97] this offline/online procedure is performed for all combinations of:

- Models: `ode_mid`, `ode_end`,
- Solvers: `imex1`, `imex2`,
- Reductors: `pod_r`,
`eds_ro`, `eds_wx`, `eds_wz`,
`bpod_ro`, `ebt_ro`, `ebt_wx`, `ebt_wz`,
`ebg_ro`, `ebg_wx`, `ebg_wz`,
`dmd_r`.

The models are specialized by the *Schiffrinson* friction factor formula and the *AGA88* compressibility factor formula. We excluded the `generic` and `rk4` solvers in this comparison as they are too slow or too fragile, respectively. However, the visualizations of the test scenarios in Fig. 4a and Fig. 6a are simulated using the `generic` solver.

For the parametric model reduction, the temperature range for training and testing is set to $[0^\circ, 15^\circ]\text{C}$, while the specific gas constant range is chosen as $[500, 600] \frac{\text{J}}{\text{kg K}}$. During training, samples from the parameter space are drawn from a sparse grid, whereas for the tests, parameters are drawn from a uniform random distribution. For either test and training, five parameters are sampled. The input perturbations for the steady-state training scenario are selected to be a step function, which heuristically works well for hyperbolic systems [40]. The empirical Gramian configuration uses default values except for constraining perturbations to only positive values.

The reduced order models are compared via the approximate, discrete, parametric, $(L_2 \otimes L_2)$ -norm of the output error [58]:

$$\|y - \tilde{y}\|_{L_2 \otimes L_2} \approx \sqrt{\sum_{\theta_h \in \Theta_h} \Delta t \|\text{vec}(y_h(\theta_h) - \tilde{y}_h(\theta_h))\|_2^2},$$

for a finite sample Θ_h of the parameter space Θ and discrete output samples $y_h(\theta_h), \tilde{y}_h(\theta_h)$. This energy norm is chosen, since all methods are at least related to an energy-based method. However, `morgen` can also provide the errors in the *approximate* parametric $(L_k \otimes L_\ell)$ parameter-space-state-space norms for $k \in \{1, 2, \infty\}$, $\ell \in \{0, 1, 2, \infty\}$, cf. [41]. To enhance comparability of the results, also the reducibility measure MORSCORE [58] for each experiment is computed. The MORSCORE for a certain method and model is essentially the area above the method's error graph in the relative error plot such as Fig. 4c.

Lastly, we note that the following numerical experiments are conducted using a computer with an AMD RYZEN 4500U @ 2.3Ghz hexa-core processor and 16GiB memory running MATLAB 2020b on UBUNTU 20.04 LINUX.

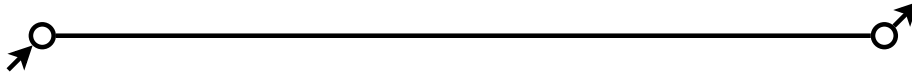


Figure 3: Pipeline topology.

6.2 Yamal-Europe Pipeline

First, a pipeline is tested, which is an interesting test case, since the trivial topology (Fig. 3) comprises little redundancy, hence pipelines are a useful benchmark for model reduction methods.

The Yamal-Europe pipeline connects gas fields in Russia with western Europe¹². A section of this pipeline was also benchmarked in [19, 97, 18, 14], from which the technical properties and test scenario are taken. The considered pipeline section is 363km long, has a diameter of 1.422m, no (reported) inclination, and a pipe roughness of 0.01mm. A steady-state, used as initial state, is set by a supply pressure of 84bar and demand mass-flux of $46.3 \frac{\text{kg}}{\text{s}}$.

In semi-discrete form, the nonlinear state-space system has two inputs and outputs as well as 908 states; and a time step width of 60s is used.

The employed test scenario is taken from [19], compressed to 24h, and shown in Fig. 4a, the associated model reduction errors are given in Fig. 4c, Fig. 4d, Fig. 4e, and Fig. 4f, for up to reduced order 150, while the resulting MORSCORES are listed in Table 5.

Generally, in this comparison the choice of solver is more relevant than the choice of model: while the MORSCORES for different models but same solver are similar, for the same model but different solver, they are significantly dissimilar. The (balancing) Petrov-Galerkin methods perform generally worse than the Galerkin methods, even when ignoring unstable ROMs, only produced by the balancing methods.

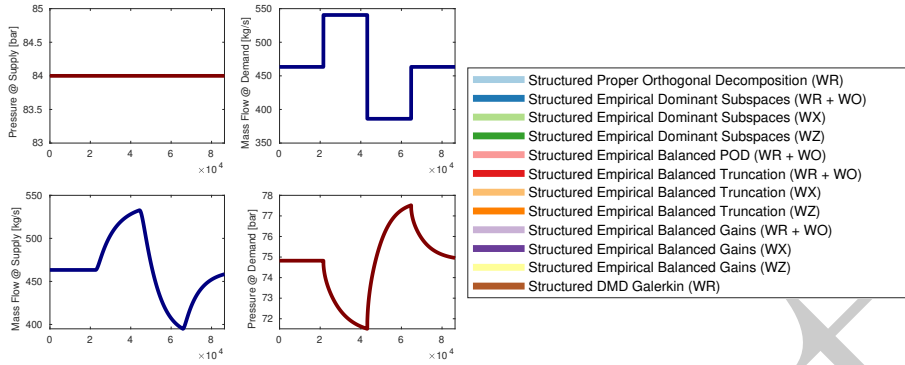
For both models, and the first-order IMEX solver, the structured empirical dominant subspaces methods perform best, followed (closely) by the DMD-Galerkin and POD methods; then, among the balancing methods, structured balanced POD. The most overall accurate method is the non-symmetric-cross-Gramian-based dominant subspaces method.

For both models, in combination with the second-order IMEX-RK solver, the structured proper orthogonal decomposition method leads, followed by the DMD-Galerkin reductor. Notably, for this model-solver setup, the balanced truncation variant `ebt_ro` and balanced gains variant `ebg_ro` result in no unstable reduced order model, while in contrast to the first-order IMEX solver, also Galerkin methods produce unstable ROMs.

For both solvers, the endpoint model performs better than the midpoint model. We note that the dominant subspaces methods produce the lowest errors. However, the purely reachability-based POD and DMD-Galerkin methods only need to perturb each input component, and hence are the more efficient, especially DMD-Galerkin is the most efficient method.

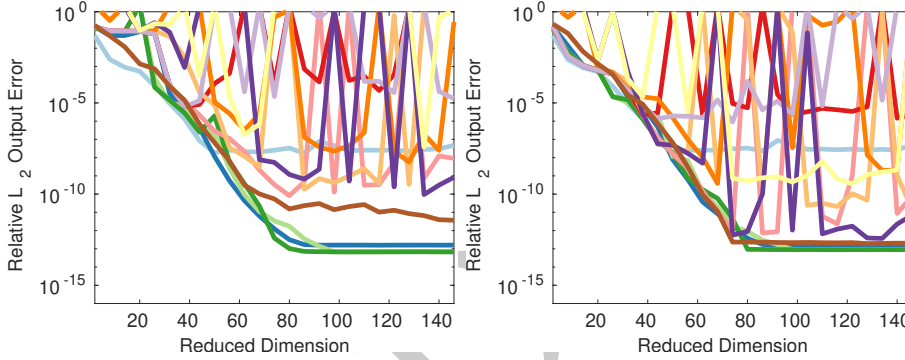
Interestingly, while the second-order IMEX-RK solver is better suited for simulations of the full order model, in terms of data-driven model reduction and/or reduced order model simulation it is significantly worse than the first-order IMEX solver. This is also demonstrated in the subsequent experiments.

¹²See also https://en.wikipedia.org/wiki/Yamal-Europe_pipeline.



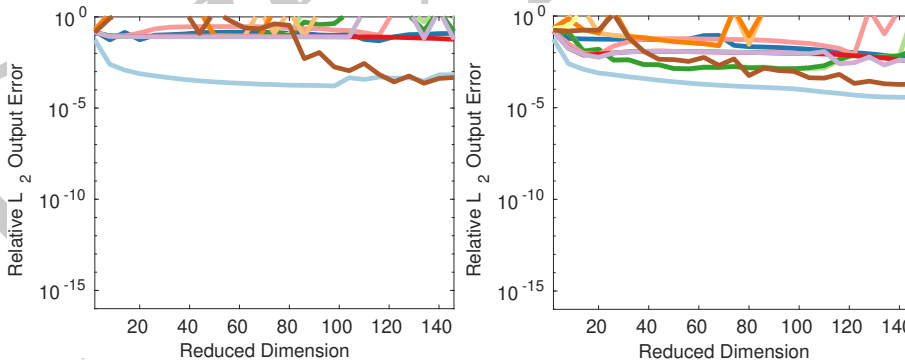
(a) The test scenario's transient boundary values and quantities of interests for the Yamal-Europe pipeline.

(b) Common legend for the model reduction error plots Fig. 4c, Fig. 4d, Fig. 4e, and Fig. 4f.



(c) L_2 output error of the ROMs using the `ode_mid` model and the `imex1` solver for increasing reduced orders.

(d) L_2 output error of the ROMs using the `ode_end` model and the `imex1` solver for increasing reduced orders.



(e) L_2 output error of the ROMs using the `ode_mid` model and the `imex2` solver for increasing reduced orders.

(f) L_2 output error of the ROMs using the `ode_end` model and the `imex2` solver for increasing reduced orders.

Figure 4: Visualization of the test scenario, and model reduction errors of the tested ROMs for the Yamal-Europe pipeline from Section 6.2.

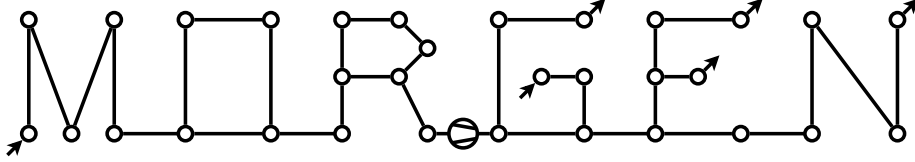


Figure 5: MORGEN network topology.

6.3 MORGEN Network

The second set of tests encompasses a synthetic network for testing `morgen`'s capabilities. This "MORGEN" network, with topology as in Fig. 5, tests the interaction of simplified compressors from Section 2.5 for various network features, such as cycles, multiple supply and demand nodes, and is in the spirit of a test network from [31].

Specifically, six sub-networks (in the shape of letters) are connected, the second and third sub-network contain a cycle, a compressor connects the third and fourth sub-network, and the fourth and fifth sub-network contain additional supply and demand nodes. The edges vary in length between 20km and 60km, while the diameter and roughness are consistently 1m and 0.01mm, respectively. A steady-state, used as initial state, is set by supply (and discharge) pressures of 50bar at both supply nodes and the compressor, and demand mass-fluxes of $30 \frac{\text{kg}}{\text{s}}$ at all demand nodes.

In semi-discrete form, the nonlinear state-space system features six inputs and outputs as well as 901 states; and a time discretization with 60s time steps.

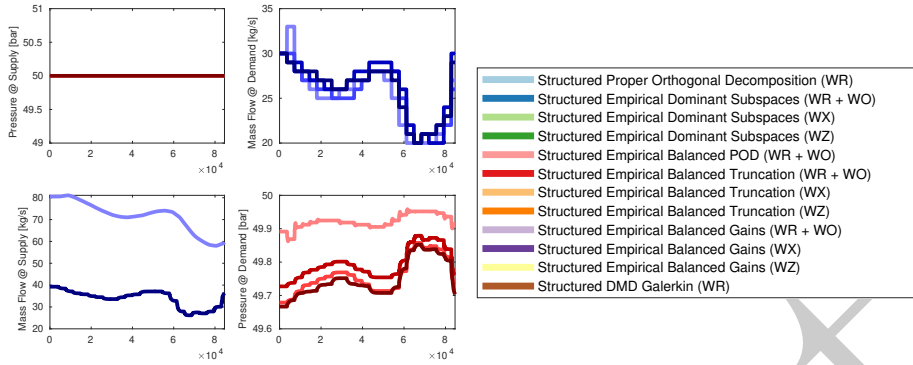
The employed 24h test scenario is made from hourly standard load profiles [50] and shown in Fig. 6a, the associated model reduction errors are given in Fig. 6c, Fig. 6d, Fig. 6e, and Fig. 6f, for up to reduced order 150, while the resulting MORSCORES are listed in Table 6.

Again for this comparison, the choice of solver is more relevant than the choice of model, yet the MORSCORES are much lower, due to the complexities (cycles, compressor, multiple demands) of the network.

Both models in conjunction with the first-order IMEX solver only produce workable results with Galerkin methods, with the exception of the balanced POD (Petrov-Galerkin) method. The endpoint ROMs are more accurate than the midpoint ROMs. Notably, the DMD-Galerkin ROMs of the endpoint model are leading the MORSCORES.

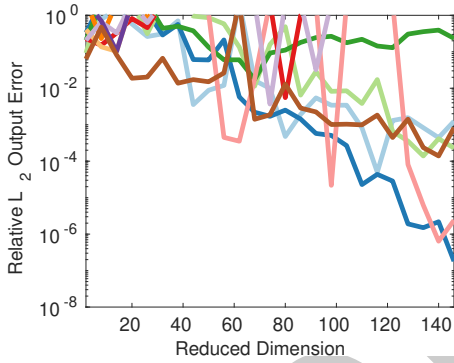
The ROMs for both models with the second-order IMEX-RK solver perform worse, with the exception of the balanced truncation `ebt_ro` and balanced gains `ebg_ro` variants. However, the second-order IMEX-RK solver related ROMs are of no practical use due to the high error.

Overall, as for the pipeline test, the dominant subspace methods produce the lowest error, while the POD and DMD-Galerkin methods are more efficient. In terms of models, the endpoint model seems better suited for model reduction, while the first-order IMEX solver results in significantly better ROMs than the second-order IMEX-RK solver.

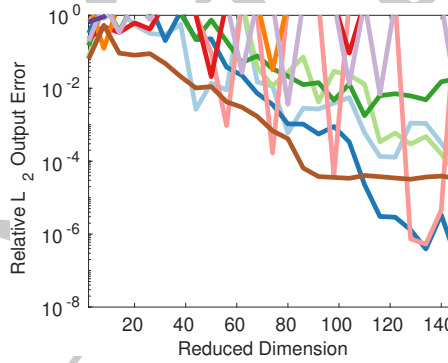


(a) A plot of the transient boundary values used as a test scenario for the ROMs of the MORGEN network.

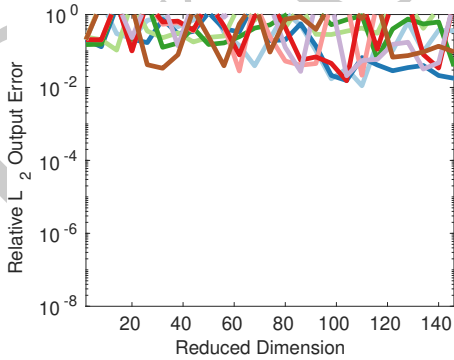
(b) Common legend for the model reduction error plots Fig. 6c, Fig. 6d, Fig. 6e, and Fig. 6f.



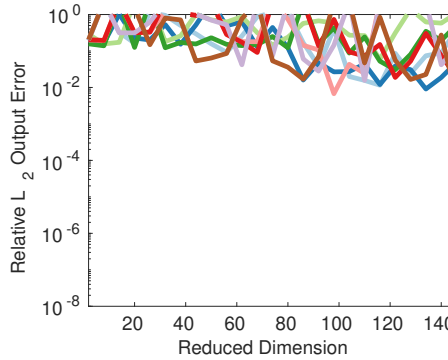
(c) L_2 output error of the ROMs using the `ode_mid` model and the `imex1` solver for increasing reduced orders.



(d) L_2 output error of the ROMs using the `ode_end` model and the `imex1` solver for increasing reduced orders.



(e) L_2 output error of the ROMs using the `ode_mid` model and the `imex2` solver for increasing reduced orders.



(f) L_2 output error of the ROMs using the `ode_end` model and the `imex2` solver for increasing reduced orders.

Figure 6: Visualization of the test scenario, and model reduction errors of the tested ROMs for the MORGEN network from Section 6.3.

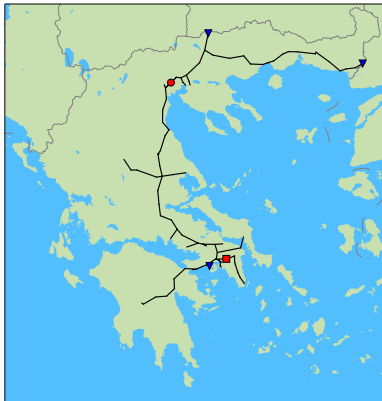


Figure 7: Topology of the GasLib134v2 network depicting a part of the Greek gas network (taken from <https://gaslib.zib.de>, licensed under CC-BY.).

6.4 GasLib Network

Lastly, a network topology derived from real-life is tested. The GasLib-134v2 network [111], modeling a part of the Greek gas transport system, is overlaid on a map of Greece in Fig. 7. The network has a total length of 1412km and features a compressor. A steady-state, used as initial state, is set by supply (and discharge) pressures of 80bar at supply nodes and the compressor, and demand mass-fluxes up to $16 \frac{\text{kg}}{\text{s}}$ at all demand nodes.

In semi-discrete form, the nonlinear state-space system has 48 inputs and outputs as well as 3886 states; and 20s time steps are employed.

For testing, a random (24h) load profile is generated, by adding samples from a scaled uniform random distribution on top of the steady-state¹³, shown in Fig. 8a, the associated model reduction errors are given in Fig. 8c, Fig. 8d, Fig. 8e, and Fig. 8f, for up to reduced order 250, while the resulting MORSCORES are listed in Table 7.

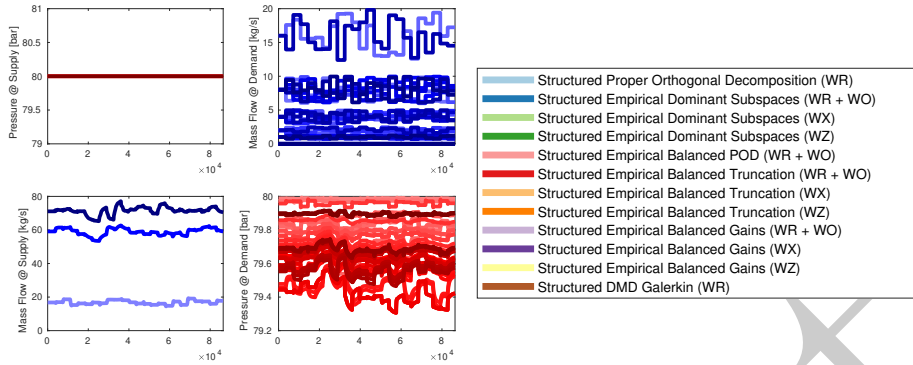
As before, the choice of solver is more relevant than the choice of model. Challenges in this network, beyond the compressor, are the high number of boundary nodes, which are predominantly demand nodes ($N_d = 45$).

First, we note that only POD and DMD-Galerkin produce consistently stable ROMs. Furthermore, in comparison with the previous experiments, the dominant subspace methods perform worse, and all variants based on reachability and observability Gramians perform relatively better.

The endpoint model seems to be slightly better suited for the tested model reduction methods than the midpoint model. And as for the other experiments, the first order IMEX solver outmatches the second order IMEX-RK solver.

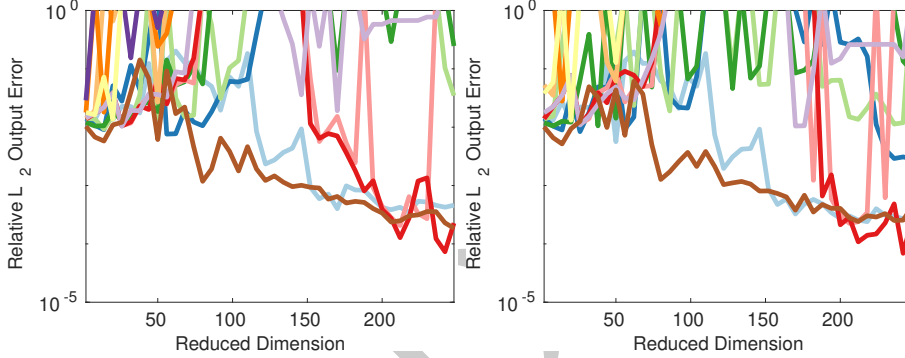
Overall, the DMD-Galerkin method performs best in terms of MORSCORE, accuracy and efficiency, followed by the POD method. As an initial evaluation in terms of the tested model reduction methods, we note, that purely reachability-based methods perform better than the reachability-observability-based methods, which need further investigation in terms of workable results and computation time. Moreover, Galerkin methods perform better than Petrov-Galerkin methods in terms of accuracy and stability.

¹³`morgen` can generate such profiles for all included networks.



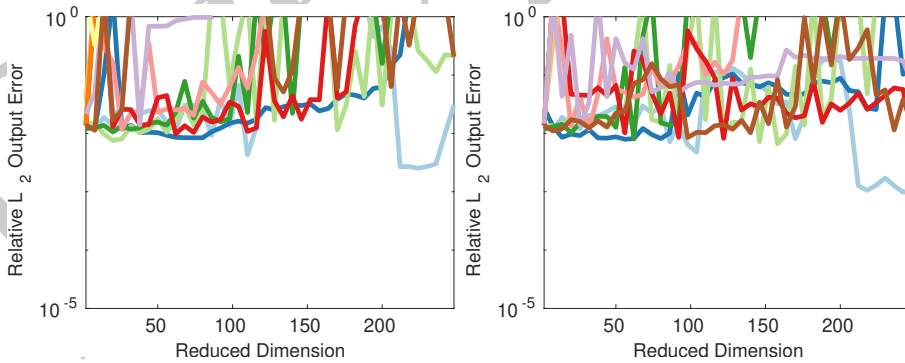
(a) A plot of the transient boundary values used as a test scenario for the ROMs of the GasLib-134v2 network.

(b) Common legend for the model reduction error plots Fig. 8c, Fig. 8d, Fig. 8e, and Fig. 8f.



(c) L_2 output error of the ROMs using the `ode_mid` model and the `imex1` solver for increasing reduced orders.

(d) L_2 output error of the ROMs using the `ode_end` model and the `imex1` solver for increasing reduced orders.



(e) L_2 output error of the ROMs using the `ode_mix` model and the `imex2` solver for increasing reduced orders.

(f) L_2 output error of the ROMs using the `ode_end` model and the `imex2` solver for increasing reduced orders.

Figure 8: Visualization of the test scenario, and model reduction errors of the tested ROMs for the GasLib-134v2 network from Section 6.4.

	ode_mid imex_1	ode_end imex_1	ode_mid imex_2	ode_end imex_2
pod_r	0.40	0.40	0.21	0.23
eds_ro	0.56	0.58	0.06	0.10
eds_wx	0.56	0.58	0.03	0.15
eds_wz	0.57	0.59	0.04	0.15
bpod_ro	0.35	0.38	0.03	0.08
ebt_ro	0.13	0.23	0.07	0.13
ebt_wx	0.26	0.32	0.00	0.02
ebt_wz	0.23	0.21	0.00	0.02
ebg_ro	0.11	0.22	0.05	0.13
ebg_wx	0.23	0.42	0.00	0.00
ebg_wz	0.07	0.31	0.00	0.00
dmd_r	0.50	0.56	0.08	0.15

Table 5: MORSCORES $\mu(150, \epsilon_{\text{mach}(16)})$ in the $(L_2 \otimes L_2)$ error norm for the “Yamal-Europe” pipeline network from Section 6.2.

	ode_mid imex_1	ode_end imex_1	ode_mid imex_2	ode_end imex_2
pod_r	0.12	0.12	0.04	0.05
eds_ro	0.16	0.16	0.05	0.06
eds_wx	0.08	0.08	0.02	0.02
eds_wz	0.03	0.07	0.02	0.04
bpod_ro	0.07	0.07	0.02	0.02
ebt_ro	0.00	0.00	0.03	0.03
ebt_wx	0.00	0.00	0.00	0.00
ebt_wz	0.00	0.00	0.00	0.00
ebg_ro	0.00	0.01	0.02	0.02
ebg_wx	0.00	0.00	0.00	0.00
ebg_wz	0.00	0.00	0.00	0.00
dmd_r	0.14	0.18	0.03	0.04

Table 6: MORSCORES $\mu(150, \epsilon_{\text{mach}(16)})$ in the $(L_2 \otimes L_2)$ error norm for the “MORGEN” test network from Section 6.3.

	ode_mid imex_1	ode_end imex_1	ode_mid imex_2	ode_end imex_2
pod_r	0.14	0.15	0.07	0.11
eds_ro	0.03	0.05	0.08	0.09
eds_wx	0.02	0.03	0.05	0.07
eds_wz	0.02	0.02	0.04	0.03
bpod_ro	0.08	0.05	0.02	0.02
ebt_ro	0.10	0.07	0.06	0.08
ebt_wx	0.00	0.00	0.00	0.00
ebt_wz	0.00	0.00	0.00	0.00
ebg_ro	0.03	0.04	0.00	0.05
ebg_wx	0.00	0.00	0.00	0.00
ebg_wz	0.00	0.00	0.00	0.00
dmd_r	0.17	0.17	0.00	0.09

Table 7: MORSCORES $\mu(250, \epsilon_{\text{mach}(16)})$ in the $(L_2 \otimes L_2)$ error norm for the “GasLib-134v2” benchmark network from Section 6.4.

7 Outlook

The next stage in the development of `morgen` involves testing larger real-life networks, such as the deliverable of the *SciGRID_gas*¹⁴ (Open Source Model of the European Gas Network) project. Yet, various further venues of linked modeling and model reduction questions are still not covered by `morgen`.

From a modeling perspective, the aspects of non-constant local compressibility, and controllable compressors are interesting, as well as testing the presented model reduction techniques on a (nonlinear) port-Hamiltonian gas network model [24, 77, 31].

In terms of model reduction, an interesting issue are intraday switchable valves, which change the topology of the gas network graph and likely require to extend the utilized model reduction methods towards these switched systems.

Another interesting question in need of further investigation is the minimal time horizon of the training phase. A lower bound is the time step times the longest path from a supply to a demand node, but this is likely not sufficient.

Also as detailed in Section 3.3, the pipe roughness is a relevant parameter for (transient) simulations [118], yet the entailing high-dimensional parameter space, due to the locally differing roughness and attrition rates, would have to be treated, too. This in turn would raise the question for combined state and parameter reduction [61, 60, 55], and is postponed to future investigations.

Finally, a tunable efficiency factor [95, 97, 101] could be added to the model's friction term, which as a free parameter could be used to tune the models to match real data.

8 Conclusions

In more than a quarter century of computational transient gas network simulation research and industrial use, `morgen` seems to be the first open-source platform (modularly) covering modeling, simulation, and model order reduction of gas (and energy) networks. The target applications for `morgen` are finding the best model reduction method or best reduced order model for a network by heuristic comparison, as well as comparing model, solver, reductor simulation configurations.

Summarizing the numerical experiments, the method yielding the lowest model reduction errors is the empirical structured dominant subspaces method. However, the most efficient reductor is the DMD-Galerkin method as it requires the least training data for the lowest error. In this regard, it is interesting to note that centering of data is essential, while for all other methods centering is not beneficial. The best model-solver combination is the endpoint model and the first-order IMEX solver, whereas the solver is significantly more decisive for the performance of the model reduction than the model. The performance of structured balanced truncation and the related structured balanced gains may be improved in terms of stability(-preservation) either by a variant of the technique by [78], or a postprocessing as in [16].

Lastly, we invite researchers, engineers and users to provide their reductors, solvers, networks and scenarios for expansion and testing with `morgen` for a broader view of this comparison.

¹⁴<https://www.gas.scigrd.de>

Code Availability

The Matlab language source code of the **morgen** platform is licensed under BSD 2-CLAUSE LICENSE, can be obtained from:

`doi:10.5281/zenodo.4288510`

and is authored by: C. HIMPE and S. GRUNDEL.

Appendix

Model Fact Sheet

Basis: Euler equations for cylindrical pipes

Assumptions

Long pipes: One spatial dimension

Kinetic term: Removed due to slow subsonic velocities

Boundary values: Low-frequency (sum of) step functions

Simplifications

Temperature: Isothermal (temperature is parameter)

Gas composition: Constant global (specific gas constant is parameter)

Compressibility: Constant global (derived from steady-state)

Compressors: Affine / Additive

Modularization

Friction: Hofer | Nikuradse | Altshul | Schifrinson | PMT1025 | IGT

Compressibility: Ideal | DVGW-G-2000 | AGA88 | Papay

Discretization

Spatial: 1st order upwind finite differences

Temporal: RK-4 | IMEX-1 | IMEX-RK-2 | Rosenbrock-2

Vector Field Splitting

Implementationwise, the linear transformation of the steady-state and input are included into the (lumped) nonlinear vector field component:

$$E\dot{x}(t) = A(x(t) - \bar{x}) + (A\bar{x} + Bu(t) + f(x(t), u(t))).$$

Funding

This work is supported by the German Federal Ministry for Economic Affairs and Energy, in the joint project: “**MathEnergy** – Mathematical Key Technologies for Evolving Energy Grids”, sub-project: Model Order Reduction (Grant number: 0324019B),

and by the National Science Foundation under Grant number: DMS-1439786 while the author was in residence at the Institute for Computational and Experimental Research in Mathematics in Providence, RI, during the “**Model and dimension reduction in uncertain and dynamic systems**” program.

Competing Interest

The authors declare that they have no competing interests.

Abbreviations

CFL	Courant-Friedrichs-Levy
CSV	Comma Separated Value
DAE	Differential Algebraic Equation
DEIM	Discrete Empirical Interpolation Method
DIRK	Diagonally Implicit RK
DMD	Dynamic Mode Decomposition
DMDc	DMD with Control
EVD	Eigenvalue Decomposition
IMEX	Implicit-Explicit
MOR	Model Order Reduction
ODE	Ordinary Differential Equation
PDAE	Partial DAE
PDE	Partial Differential Equation
PDIRK	Passive DIRK
POD	Proper Orthogonal Decomposition
RK	Runge-Kutta
ROM	Reduced Order Model
SDIRK	Singly DIRK
SRSN	System with Repeated Scalar Nonlinearities
SSP	Strong Stability Preserving
SVD	Singular Value Decomposition
tEVD	truncated EVD
tSVD	truncated SVD

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