Time Integration of Reacting Flows and Energy-Stable Hybrid Spatial Discretizations

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1 Time Integration of Reacting Flows

1.1 Motivation & Background

In the simulation of reacting flows, the compressible Navier-Stokes equations are augmented to include equations governing the rate of change of the mass fraction of each species in the chemical mechanism at hand based on momentum, mass diffusion, and reaction kinetics. What results is a set of equations that can feature both a wide range of timescales and a number of comparatively stiff terms that can have an adverse effect on maximum allowable timestep size and, as a result, simulation performance and fidelity. In particular, the timescales of the chemical reactions are often multiple orders of magnitude lower than those of the fluid motion in practice. One approach to effectively time-marching these problems at the fluid-relevant timescale, then, is to introduce an operator-splitting technique ([44], [46], [26], [30], [42], [24]) in which the advection and diffusion terms in the right-hand-sides are advanced explicitly, with the chemical source term(s) being advanced implicitly and in a separate substep, typically by software packages such as CVODE [8]. However, this approach results in a scheme that is typically at best first- or second- order accurate in time globally.

To more accurately handle such situations, implicit-explicit (IMEX) time integration schemes have been developed in order to partition the governing equations and treat the stiff terms implicitly while treating the nonstiff terms explicitly. In 2003, Kennedy and Carpenter [20] introduced additive Runge-Kutta schemes for convection-diffusion-reaction equations, demonstrating a higher order of temporal accuracy than the operator splitting approach is capable of attaining. In 2011, Svärd and Mishra [48] also introduced more simplistic one-step IMEX schemes mimicking single stages in these additive-RK schemes, applying them to the reactive Euler equations and demonstrating first order with a need for numerical diffusion as the problem becomes less stiff.

Another approach to handling stiffness, and in particular the resulting disparity in timescales between different simulation components, is the use of multi-rate time integration. Gear and Wells [15] were the initial authors of work on multi-step multi-rate methods, with similar contributions provided more recently by Constantinescu and Sandu [9] for hyperbolic conservation laws. Constantinescu and Sandu later extended this work to develop implicit multi-rate schemes [10], [11], but failed to introduce a scheme featuring both implicitly and explicitly advanced components within a multi-rate framework. Klöckner [23] also conducted an empirical investigation into the effects of the various design choices present in a two-component scheme based on the work of Gear and Wells, with Mikida et. al [35] applying this framework to fluid simulation on overset grids using provably stable interpolation to form the coupling between grid-determined timescales.

The primary goal of this section of the thesis is demonstration of a novel time integration scheme driving a demonstrative reacting flow problem with improved efficacy identified by either superior accuracy and stability and/or faster performance relative to existing methods. The integration project being undertaken aims to augment the capabilities of the multi-rate integrators developed in [35], adding the ability to treat certain right-hand

side components implicitly as well as adapt either the macro-timestep or the step ratio of the scheme based on local truncation error estimates. These additions are explicitly designed to handle the stiffness introduced to the flow problems to which these integrators were originally applied via the introduction of chemical source terms, as well as to efficiently integrate reacting flow simulations in which the stiffness varies. This latter phenomenon is especially common in ignition simulations, where the high-stiffness qualities of the kinetics during ignition contrast with surrounding simulation time in which the mass fractions are at or near chemical equilibrium.

A critical component in our effort will be comparison of the resulting scheme with CVODE as a proxy for the state of the art in time integration of chemically reacting flows, with the goal of achieving superiority in terms of observed temporal accuracy of the entire system, or in terms of performance (most accurately measurable by a reduction in the number of chemistry right-hand-side evaluations required to reach a given solution time).

Meanwhile, an aspect of the algorithms being implemented that has been identified early as beyond our bounds is the presence of multi-variable solves in a number of potential adaptive implicit integration scenarios, namely:

- Coupled implicit right-hand-sides
- Use of adaptive timestep controllers where the local error estimate of all solution components makes use of implicit and explicit state estimates.

With these goals and bounds in mind, the proposed thesis has the potential to have a significant impact on the simulation of reacting flows, of which there are countless real-world applications including (but not limited to) subsonic and supersonic combustion for propulsion (gas turbines, ramjets/scramjets, rockets), furnaces and residential heating, and solution of problems relating to atmospheric sciences.

1.2 Governing Equations

The problem at hand involves the simulation of chemically reacting flows, the governing equations of which amount to the Navier-Stokes equations with (in our form):

- The addition of a governing equation for the rate of change of the mass fractions of each species in the chemical mechanism simulated
- The addition of governing equations specifying the rate of change of temperature and pressure, derived from an assumption of constant internal energy and volume in a reaction substep (for the former) and the ideal gas law (for the latter).

These equations are given below, along with a brief explanation of the notation/nomen-clature used.

$$\frac{\partial \rho}{\partial t} + \frac{\partial}{\partial x_j}(\rho u_j) = 0 \tag{1}$$

$$\frac{\partial}{\partial t}(\rho u_i) + \frac{\partial}{\partial x_i}(\rho u_i u_j + P\delta_{ij} - \tau_{ij}) = 0$$
(2)

$$\frac{\partial}{\partial t}(\rho E) + \frac{\partial}{\partial x_i}((\rho E + P)u_j + q_j - u_i \tau_{ij}) = 0$$
(3)

$$\frac{\partial}{\partial t}(\rho Y_k) + \frac{\partial}{\partial x_j}(\rho Y_k u_j - \varphi_{ki}) = W_k \dot{\omega}_k \tag{4}$$

$$\frac{\partial T}{\partial t} + \frac{\sum_{k=1}^{N_{sp}} U_k(T) \frac{\partial \rho Y_k}{\partial t} \frac{1}{W_k}}{\sum_{k=1}^{N_{sp}} [C]_k C_{v,k}(T)} = 0$$
 (5)

$$\frac{\partial P}{\partial t} - \frac{R}{V} \left(T \frac{\partial n}{\partial t} + \frac{\partial T}{\partial t} n \right) = 0 \tag{6}$$

In these equations, ρ is the fluid density, u_i is the velocity in the *i*-th direction, and E is the total energy density, calculated as

$$E = U + \frac{1}{2} \sum u_i^2 \tag{7}$$

where U is the total internal energy of the gas mixture. δ_{ij} is the Kronecker delta, P is the pressure, T is the temperature, R is the universal gas constant, and Y_k is the mass fraction of species k. $[C]_k$ is the concentration of species k in the gas mixture, N_{sp} is the number of species in the chemical mechanism, W_k is the molecular weight of species k, n is the number of moles of the gas mixture, V is the volume of the gas, and $\dot{\omega}_k$ is the net production rate of species k. τ_{ij} is the viscous stress tensor, φ_{ki} are the diffusion fluxes, and q_i are the heat fluxes, defined by

$$q_j = -\frac{\partial(\lambda T)}{\partial x_j} + h_k \varphi_{kj}. \tag{8}$$

In this expression, λ is the thermal conductivity, and h_k is the enthalpy of species k. As for the diffusion fluxes, these are defined using a mixture-averaged approach with a correction term to ensure mass conservation ([43], [41], [19]). $C_{v,k}(T)$ is the specific heat at constant volume of species k at temperature T, and $U_k(T)$ is the internal energy of species k at temperature T. These quantities are defined using the NASA 9-coefficient polynomial parameterization [33].

1.3 Numerical Methods

1.3.1 Adams Methods

For clarity, here we provide a brief derivation of a standard Adams-Bashforth (AB) integrator, as described in [3], starting from a model IVP given by

$$\frac{\mathrm{d}y}{\mathrm{d}t} = F(t, y), \quad y(0) = y_0. \tag{9}$$

Applying a method of lines (MOL) approach to spatially discretize and solve the reactive Navier-Stokes equations (1) - (6) ultimately provides this form. We approximate the time dependency of the right-hand side function with a polynomial with coefficients α (formed by interpolating past values of F(t, y)), extrapolate with that polynomial approximation, and integrate the extrapolant. To obtain the coefficients α used to form this history-interpolating polynomial, we construct a linear system using a Vandermonde matrix:

$$V^{T} \cdot \boldsymbol{\alpha} = \int_{0}^{\Delta t} \tau^{i} d\tau = \frac{(\Delta t)^{i}}{i+1}, \quad i = 1, 2..., n, \quad V = \begin{bmatrix} 1 & t_{1} & \dots & t_{1}^{n-1} \\ 1 & t_{2} & \dots & t_{2}^{n-1} \\ \vdots & \vdots & \ddots & \vdots \\ 1 & t_{n} & \dots & t_{n}^{n-1} \end{bmatrix}, \quad (10)$$

where $\int_0^{\Delta t} \tau^i d\tau$ is a vector evaluating the integral of the interpolation polynomial, and V is the Vandermonde matrix with monomial basis and nodes $t_1, t_2, \ldots t_n$, corresponding to past time values. In (10), n is equal to the order of the integrator, and t_i are the time history values, with $0 \le t_1 < t_2 \ldots < t_n$. The coefficients α are used to extrapolate to the next state via

$$y(t_{i+1}) = y(t_i) + \alpha_1 F(t_{i-n}, y_{i-n}) + \alpha_2 F(t_{i-n-1}, y_{i-n-1}) + \dots + \alpha_n F(t_i, y_i).$$
(11)

1.3.2 Multi-rate Adams Methods

We now describe a multi-rate generalization of the scheme, making use of the algorithm introduced in [15]. We consider the following model system with "fast" and "slow" solution components:

$$\frac{\mathrm{d}}{\mathrm{d}t} \begin{pmatrix} f(t) \\ s(t) \end{pmatrix} = \begin{pmatrix} a_f(f,s) \\ a_s(f,s) \end{pmatrix}. \tag{12}$$

With this in mind, we can set a slow (larger) time step H for a_s such that we maintain stability in the integration of the slow component. We also set a fast time step h for a_f such that H is an integer multiple of h, and define the ratio between the two, SR = H/h, as the step ratio of the multi-rate scheme. While the reactive flow scheme presented here makes use of only two separate state components, each with its own right-hand side function and

independent rate, the theory is readily extensible to any number of rates.

Within this two-component scheme, a few design choices are available:

- The order in which we evaluate and advance the solution components. Namely, two primary options are advancing the fast-evolving solution component through all of its micro-timesteps h and waiting to perform the single macro-timestep H required for the slow component until the end (a "fastest-first" scheme, per the nomenclature of [15]), or pursuing an algorithm in which the slow component is instead advanced first.
- For slowest-first evaluation schemes, the choice of whether or not to re-extrapolate the slow state after additional state and right-hand side information is gathered at the micro-timestep level.

Empirical observations on the effects of these choices are made in [23], with a step-by-step walkthrough of the evaluations present in a single macrostep of a fastest-first scheme given by [35]. The code implementing this method, as well as other methods within the multi-rate Adams framework, is generated by the Python packages Leap [22] and Dagrt [21], and all time integration scheme development is occurring via implementation in Leap.

To fully realize the potential multi-rate formulation for reacting Navier-Stokes problems (for which the matrix of right-hand sides is not as simple as in [35], where both right-hand sides and state components are realized as per-overset-grid quantities), we need to introduce additive terms to (12), which we express as terms coupling the fast and slow states (a_{fs} and a_{sf}) and self-influencing terms (a_{ff} and a_{ss}):

$$\frac{\mathrm{d}}{\mathrm{d}t} \begin{pmatrix} f(t) \\ s(t) \end{pmatrix} = \begin{pmatrix} a_{ff}(f,s) + a_{fs}(f,s) \\ a_{sf}(f,s) + a_{ss}(f,s) \end{pmatrix}. \tag{13}$$

In the reacting flow formulation with which we are concerned (Equations (1) - (6)), we define the "fast" component of our Navier-Stokes solution as a vector of the mass fractions of the species as well as the temperature and pressure, $[\rho Y_k, T, P]^T$. The "slow" component is then defined as the set of typical conserved variables in a non-reacting simulation, $[\rho, (\rho \vec{u}), (\rho E)]^T$. As a result, one possible definition of the additive terms (13) can be given

as follows:

$$a_{ff}(f,s) = \begin{pmatrix} W_k \dot{\omega}_k \\ -\frac{\sum_{k=1}^{N_{sp}} U_k(T) \frac{\partial \rho Y_k}{\partial t} \frac{1}{W_k}}{\sum_{k=1}^{N_{sp}} [C]_k C_{v,k}(T)} \\ \frac{R}{V} \left(T \frac{\partial n}{\partial t} + \frac{\partial T}{\partial t} n \right) \end{pmatrix}$$

$$a_{fs}(f,s) = \begin{pmatrix} -\frac{\partial}{\partial x_j} (\rho Y_k u_j - \varphi_{ki}) \\ 0 \\ 0 \end{pmatrix}$$

$$a_{sf}(f,s) = \begin{pmatrix} 0 \\ -\frac{\partial}{\partial x_j} (\rho Y_k u_j) \\ -\frac{\partial}{\partial x_j} (\rho U_j) \end{pmatrix}$$

$$a_{ss}(f,s) = \begin{pmatrix} -\frac{\partial}{\partial x_j} (\rho u_j) \\ -\frac{\partial}{\partial x_j} (\rho u_i u_j - \tau_{ij}) \\ -\frac{\partial}{\partial x_j} ((\rho E) u_j + q_j - u_i \tau_{ij}) \end{pmatrix}$$

As our treatment of this set of equations with multi-rate integration develops, so too will our approach to defining the additive terms of (13), as well as which state variables are deserving of fast or slow treatment.

1.3.3 Implicit Adams Methods and Extension to Multi-rate

The concepts in the above sections, both single- and multi-rate, are easily extended to include implicit integration methods. We define an implicit Adams (Adams-Moulton [36]) method in the same manner as above, starting with the same simplified IVP (9) and expressing the scheme as

$$y(t_{i+1}) = y(t_i) + \beta_1 F(t_{i-n+1}, y_{i-n+1}) + \beta_2 F(t_{i-n}, y_{i-n}) + \dots + \beta_n F(t_{i+1}, y_{i+1}).$$
(14)

The coefficients $\boldsymbol{\beta}$ are constructed in a manner identical to the Adams-Bashforth coefficients $\boldsymbol{\alpha}$ (see Equation 10), using temporal nodes shifted Δt forward in time.

Extension of this implicit method into the multi-rate framework is straightforward, given that the driving principle of the Adams-Moulton method (namely, maintenance of right-hand side history values through which to interpolate) is the same as that of Adams-Bashforth, and amounts to allowing the histories of each right-hand side term to maintain their own timestep intervals.

In the case that implicit time integration is used, it will be used to drive $a_{ff}(f, s)$. In order to perform the implicit solve required to advance this right-hand side via Adams-Moulton, a Jacobian matrix is required. While other implementations such as CVODE's adaptive BDF algorithm for stiff problems often make use of finite-difference generated

Jacobians, we will instead use PyJac-V2 [12] to obtain the analytical Jacobian for a given chemical mechanism via code generation, which will then be used by a simple Newton algorithm to perform the implicit solve. More details on the construction of this Jacobian are given by Curtis et. al, but we document the basic form, in addition to an alternate differential-algebraic formulation, below.

The chemical kinetics Jacobian can be written as the matrix

$$J = \frac{\partial f_i}{\partial \Phi_i} \tag{15}$$

where f is the set of right-hand sides directly pertaining to the kinetics and reactive source term:

$$f = \left(\frac{dT}{dt}, \frac{dP}{dt}, \frac{dn_1}{dt}, \frac{dn_2}{dt}, \dots \frac{dn_{N_{sp}-1}}{dt}\right). \tag{16}$$

Here, n_i is the number of moles of species i in the mixture. While the temperature and pressure right-hand sides are given by (5) and (6), note that the molar source terms pertain only to the net production rates of the species:

$$\frac{dn_i}{dt} = V\dot{\omega}_i. \tag{17}$$

Note also (based on the structure of PyJac-V2) that the state vector for the Jacobian is defined using the number of moles of each species in the chemical mechanism rather than the mass fractions:

$$\Phi = (T, P, n_1, n_2, \dots n_{N_{sp}-1}). \tag{18}$$

The number of moles in of the last species in the mixture is calculated via the mass fractions summing to 1.

With these definitions in mind, we can expand (15) and express it in true matrix form:

$$J = \begin{bmatrix} \frac{d\dot{T}}{dT} & \frac{d\dot{T}}{dP} & \frac{d\dot{T}}{dn_{1}} & \cdots & \frac{d\dot{T}}{dn_{N_{sp}-1}} \\ \frac{d\dot{P}}{dT} & \frac{d\dot{P}}{dP} & \frac{d\dot{P}}{dn_{1}} & \cdots & \frac{dP}{dn_{N_{sp}-1}} \\ \frac{d\dot{n}_{1}}{dT} & \frac{d\dot{n}_{1}}{dP} & \frac{d\dot{n}_{1}}{dn_{1}} & \cdots & \frac{d\dot{n}_{1}}{dn_{N_{sp}-1}} \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ \frac{d\dot{n}_{N_{sp}-1}}{dT} & \frac{d\dot{n}_{N_{sp}-1}}{dP} & \frac{d\dot{n}_{N_{sp}-1}}{dn_{1}} & \cdots & \frac{d\dot{n}_{N_{sp}-1}}{dn_{N_{sn}-1}} \end{bmatrix}$$

$$(19)$$

Providing this matrix to the implicit solver should serve to reduce the number of chemical right-hand side calls required to time-march the system relative to methods that employ a finite-difference generated Jacobian, which will directly improve performance.

A critical consideration in the formulation of both the governing equations and the chemical Jacobian will be maintaining a solution that lies on the ideal gas constraint manifold. This has been identified in early work as a potential concern when fully differential approaches

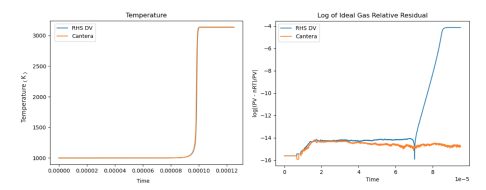


Figure 1: Temperature profiles (left) and log of relative ideal gas law residual (right) for advancement of homogeneous H2-O2 autoignition example (employing the nine-species San Diego mechanism) using a fully differential approach (blue) versus a differential-algebraic approach (orange) where the constant internal energy and ideal gas law constraints are fulfilled by a Newton solve within Cantera.

to solving the governing equations that time-march the rates-of-change of temperature and pressure (as opposed to differential-algebraic approaches that directly satisfy the ideal gas and constant internal energy constraints) are used. To avoid drift off of the ideal gas law manifold, a chemical kinetic Jacobian modified to include the algebraic constraints explicitly can be used. This approach, detailed in Hung [18], modifies the molar source terms to incorporate this constraint while removing the temperature and pressure source terms (which we then replace with a temperature Newton solve based on the constant internal energy constraint via Cantera [16]), as well as a direct application of the ideal gas law to obtain the pressure. This modified Jacobian can be expressed as

$$J^* = \begin{bmatrix} \frac{d\dot{n}_1^*}{dn_1} & \cdots & \frac{d\dot{n}_1^*}{dn_{N_{sp}-1}} \\ \vdots & \ddots & \vdots \\ \frac{d\dot{n}_{N_{sp}-1}^*}{dn_1} & \cdots & \frac{d\dot{n}_{N_{sp}-1}^*}{dn_{N_{sp}-1}} \end{bmatrix}$$
(20)

where

$$\frac{d\dot{n}_i^*}{dn_i} = \frac{d\dot{n}_i}{dn_i} - \frac{U_j}{C_v} \frac{d\dot{n}_i}{dT}.$$
 (21)

In this expression, U_j is the contribution of the jth species to the internal energy, and C_v is the specific heat of the mixture at constant volume.

Which of these Jacobians is ideal for our purposes is a consideration worth monitoring as our implicit integration techniques develop. In particular, a third relatively unexplored approach would use the fully differential method with pressure and temperature source terms, while also using penalization to force the solution towards the constraint manifold. This attenuation approach is described in some detail (albeit for smaller model problems) by Ascher and Petzold [2].

1.3.4 Timestep Control Algorithm

Early implicit-explicit runs of small-scale reacting flow problems quickly identified a need to adapt the timestep of the integrator based on relative and/or absolute local error demands. We therefore employ a timestep control algorithm which, in its current form, is similar to that of CVODE, using a weighted root-mean-square norm to more adequately handle states with components that feature a large variation in magnitude. The weights are calculated as

$$w_i = \frac{1}{\text{ATOL} + \text{RTOL}|s_{q,i}|}$$
 (22)

with the weighted root-mean-square norm then being given by

$$||E||_{wrms} = \left(\sum_{i=1}^{N} \frac{1}{N} (w_i (s_{q+1,i} - s_{q,i}))^2\right)^{1/2},$$
(23)

where $s_{q,i}$ is the *i*-th component of the state estimate obtained using an order-q scheme. The timestep is *decreased* if $||E||_{wrms}$ is greater than 1 via

$$\Delta t = 0.9 \Delta t (\|E\|_{wrms})^{-1/q} \tag{24}$$

and increased if $||E||_{wrms} < 1$ via

$$\Delta t = 0.9\Delta t (\|E\|_{wrms})^{-1/(q+1)} \tag{25}$$

As mentioned in Section 1.1, this added capability presents a number of questions in terms of how best to accomplish this error control. In particular, step ratio adjustment in situ, rather than timestep adjustment, to meet error needs provides an implementation challenge in terms of code generation of these integrators, whereas the choice of which solution component(s) are involved in error estimation/timestep control also presents a design decision that could have notable effect on numerical performance.

1.4 Validation Plan

The efficacy of the time integrators constructed via the above methodology will be assessed using a reacting mixing layer problem with a hyperbolic tangent velocity profile for the baseflow (as used by [34] and [4]), with the San Diego 9-species chemical mechanism [51] employed for the chemical kinetics. By perturbing this baseflow with the mode having the highest growth rate to instability (reproduced based on the inviscid analyses of [34] and [4]), and by varying the temperature in the domain to induce ignition local to the mixing layer, we can establish a fast-evolving solution component (the chemical kinetics of the ignition) and a slow-evolving solution component (the fluid motion), both possessing accessible means of rate modification (for the chemistry, we can accomplish this by scaling the pre-exponential factors in the Arrhenius reaction rates, and for the fluid, we can scale

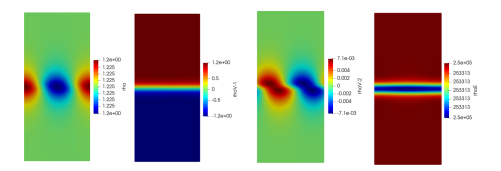


Figure 2: Perturbed initial condition of the hyperbolic tangent example - sea-level air.

the magnitude of the initial crossflow velocity). In doing so, what should result is a problem with ample testbed capabilities for our new multi-rate integrators, and also a problem firmly rooted in physical utility in that it provides the simplest model for scramjet/ramjet combustion processes.

1.5 Outlook

1.5.1 Current Status

At present, implicit-explicit time integration of reacting flows using Runge-Kutta based methods in conjunction with code generation of source terms and Jacobians for the chemical mechanisms is implemented and undergoing testing with the fluid solver application. Implicit Adams methods (with single-rate Adams-Moulton being the baseline) for the purposes of chemistry integration are also being implemented, along with an application of the error-informed timestep control algorithm to both single-rate and multi-rate Adams methods. Construction of implicit-explicit multi-rate Adams methods with error-informed adaptivity are well underway, with testing of fully-explicit adaptive Adams multi-rate integrators on a stiff small-scale Cantera [16] two-reactor system in progress. A schematic of this problem, which is based in large part on an example problem bundled with the software, is given in Figure 3. While the governing equations for this problem are not the same as those generated by PyJac, our hope is that it will serve as a suitably small stiff test problem to be used as a stepping stone in verification of the error-informed timestep controller for explicit methods, as well as further exploration of its efficacy.

As for the reacting crossflow validation problem, an initial setup employing cold flow (no autoignition) with the perturbed baseflow applied to sea level air is complete (see Figure 2), with validation via the growth rate of the unstable mode ongoing. Also implemented in the reacting flow solver is a one-dimensional laminar free flame problem, which may also be used as a first-pass validation (via comparison to the Cantera-estimated steady-state flame speed) for new integration methods for reacting flows as they become available. A basic description of this case, as well as flame speed data generated via a Leap IMEX integrator using PyJac analytical Jacobians, is given in the next section.

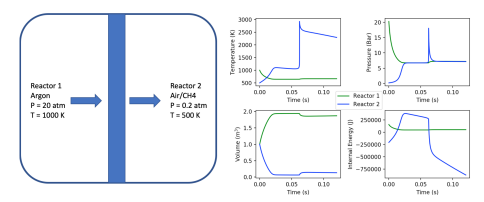


Figure 3: Cantera two-reactor example problem. The solution shown is obtained using Cantera's internal reactor system integrator, which is based on the same BDF algorithm employed by CVODE. The piston velocity is initially zero, with ignition eventually occurring in the air-methane reactor.

1.5.2 Risk Mitigation

Assuming demonstration of improved performance or accuracy of the *multi-rate* integrators over the state-of-the-art is for reasons unforseen impossible, a "fallback" takeaway would be demonstration of improvement over a low-order splitting approach via Leap implementation of an existing IMEX Runge-Kutta based scheme ([20]) that treats the chemistry implicitly via code-generated analytical Jacobians. Given the use of a coupled implicit-explicit scheme (for which high order has already been proven in other circumstances), as well as a chemical Jacobian that is more accurately obtained than via finite differences (the method of CVODE), this outcome should be attainable at minimum.

As a step towards proving this, we have implemented a small-scale one-dimensional free-flame test, wherein we use a code-generated implementation of the fourth-order additive Runge-Kutta scheme of Kennedy and Carpenter to drive a simulation of a premixed, freely propagating hydrogen flame with an unburned gas temperature of 300 K at a pressure of one atmosphere. Comparing the estimated laminar flame speed to that of a steady-state analysis performed by Cantera at a number of equivalence ratios ϕ , we see suitable agreement of less than 3% error, indicating that the reacting flow equations - and in particular the chemical source terms and Jacobians - are correctly implemented and ready for both further experimentation with multi-rate timescale splitting and a more thorough analysis of performance.

$\overline{\phi}$	Leap S_L (m/s)	Cantera S_L (m/s)	% Error
0.75	1.585566	1.599170	0.85
1.0	2.351950	2.421837	2.89
1.5	3.1734067	3.153888	0.62

Table 1: Comparing the laminar flame speeds estimated by Leap-PyJac chemistry integration and by Cantera's steady-state analysis.

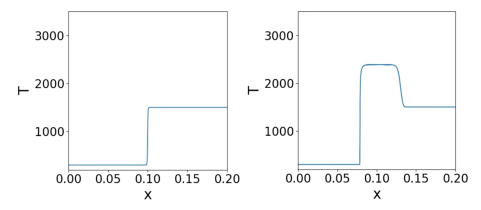


Figure 4: Figures showing the temperature profile of the one-dimensional free-flame example at initial condition (left) and after 0.0025 seconds (right).

2 Energy-Stable Hybrid Spatial Discretizations

2.1 Motivation & Background

When simulating problems in computational fluid dynamics involving complex geometries, numerical difficulties are often encountered that require employing one of a number of different approaches to mitigate loss of accuracy and/or stability due to poor spatial resolution. While finite-difference approaches to spatial discretization are popular due to their simplicity and efficiency when working with smooth solutions, their geometric flexibility is typically limited in spite of the well-documented ability to use coordinate transforms to create intricate curvilinear meshes, as well as the ability to use overset meshes ([6], [39]) to produce higher resolution in particularly troublesome sections of the spatial domain.

In particular, the SBP methodology pioneered by Strand [45], which mimics integration-by-parts in the discrete sense and is provably stable via the energy method when coupled with weak enforcement of boundary conditions (simultaneous approximation term; see [47]), is an attractive one when it comes to finite differences, and its geometric flexibility has recently been further improved by a large body of work in creating accurate and stable interfaces between nonconforming blocks. In 2010, Mattsson and Carpenter [31] introduced interpolation operators that maintained the stability and high-order accuracy of the underlying spatial scheme on each block, but the resulting scheme is based on a fixed refinement ratio and the requirement that the blocks conform at their corners (the latter of these characteristics was later removed by Nissen et. al in 2015 [37]). The use of SBP-SAT finite difference techniques in conjunction with nonconforming grid interfaces has been further extended to use with the second-order wave equation [50], 3D elastic wave simulations [14], the Schrödinger equations [38], and the advection-diffusion equation [28].

Meanwhile, another attractive approach to modeling complex geometries more efficiently with a localized spatial discretization change is to create a hybrid spatial discretization that requires an interface between a high-order finite difference method (used away from the complex boundary) and an unstructured mesh with improved geometric flexibility

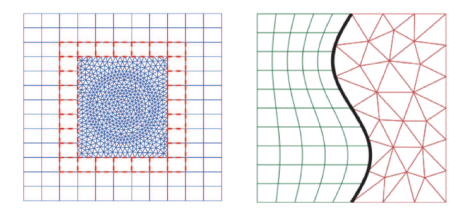


Figure 5: Two examples demonstrating use of nonconforming interfaces. Left: Zhu, Chen, Zhong, and Liu [52]. Right: Kozdon and Wilcox [25]

(used near the complex boundary). Nordström and Gong proposed such a scheme in 2006 [40], coupling a high-order finite difference method and a second-order finite volume scheme. More recently, in 2016, Kozdon and Wilcox [25] developed an interface method between SBP finite-difference blocks that was based on projection into an intermediate "glue" grid characterized by piecewise polynomials, with the method also extending to the coupling of SBP finite-difference methods and discontinuous Galerkin methods. In this work, the interface method was proven to be stable via the energy method, but the projection operators were not fully determined and were based in part on an optimization procedure focused on driving the product of the projection matrices closer to an identity operation. Furthermore, the method was not extended to three dimensions, and the method is reduced to first order accuracy if the continuous coordinate transforms on either side of the interface are dissimilar. Since, Friedrich et. al [13] extended the method(s) of Kozdon and Wilcox to make the projection process degree-preserving (a characteristic that had been previously shown to be problematic by Lundquist and Nordström [29]) as well as energy-stable and conservative, but their approach still results in projection operators that are not fully determined, relying upon an optimization procedure to produce the operators that give their results. In 2019, Almquist et. al [1] developed new orderpreserving interpolation operators that accomplished the same goal as Friedrich et. al without increasing the quadrature order, and which extended to problems with secondorder spatial derivatives, but their results again were confined to problems with a fixed 2:1 refinement ratio. Furthermore, unlike Kozdon and Wilcox, both the methods of Almquist and Friedrich fail to extend their method to hybrid structured-unstructured discretizations.

This section of the thesis targets a lack of fully-determined high-order accurate and provably stable interface conditions between structured and unstructured meshes for computational fluid dynamic simulations. The key contribution of this work will be construction of an interface between the structured (finite difference SBP) and the unstructured (DG) spatial discretizations that does not rely upon optimization or an unequal number of constraints and unknowns for construction of the solution projections. Currently, our plan is

to match the approach of [25], with the interface being defined by operators P_{a2b} and P_{b2a} that project the solution along the edge of discretization a into discretization b, and vice versa.

An additional goal to be pursued in conjunction with a unisolvent procedure while maintaining accuracy and stability is a resulting projection process that is able to provide "round-trip" fidelity of projection, meaning that projection from the structured grid to the unstructured grid and back results in a solution identical to the starting point. This characteristic is one that the operators of [25] lack, even in projection from one grid into the intermediate glue layer.

The strategy for pursuance of these goals begins with consideration of the *interior* projection operator only, given the simplicity of the existing interior stencil relative to the boundary (which features many more unknowns). Noting also that the compatibility condition, which we will later find to be the critical condition for proving a stable interface, disallows us from obtaining exact "round-trip" projection, other conditions may be considered based on further manipulation or restructuring of the edge energy dissipation relations, which will be derived for both structured and unstructured spatial discretizations in subsequent sections. Should a procedure for fully-determined interior interface conditions be applied successfully to a periodic problem, the boundary interface will subsequently be considered.

2.2 Governing Equations

The initial problem to which our new interface method will be targeted is the acoustic wave equation in two dimensions, in first-order form:

$$\rho \frac{\partial v_i}{\partial t} + \frac{\partial p}{\partial x_i} = 0 \quad (i = 1, 2), \tag{26}$$

$$\frac{\partial p}{\partial t} + \lambda \left(\frac{\partial v_1}{\partial x_1} + \frac{\partial v_2}{\partial x_2} \right) = 0 \tag{27}$$

As in [25], our aim will be to prove the semi-discrete hybrid discretization stable via the energy method by proving that

$$\frac{d\mathcal{E}}{dt} \le 0 \tag{28}$$

where the total energy of the semi-discrete system \mathcal{E} is obtained by summing the energy E of each discretization block:

$$\mathcal{E} = \sum_{blocks} E \tag{29}$$

The definitions of the energy dissipation rates for each block (each having its own spatial discretization), are given in subsequent sections.

2.3 Numerical Methods

2.3.1 Summation-by-Parts Operators

For the structured discretization that makes up one component of the proposed hybrid, we make use of several finite difference operators that possess the summation-by-parts (SBP) property. Taking two matrices H, Q, we here state that these two matrices are SBP matrices of order p provided

- $H^{-1}Qv$ is an order h^p approximation to $\partial/\partial x$, where h is the spatial step size in one dimension
- *H* is a symmetric positive-definite matrix.
- $Q + Q^T = diag(-1, 0, 0, ..., 0, 1).$

These conditions together ensure that the discrete version of the integration by parts property holds; that is,

$$\langle H^{-1}Qx, y \rangle_H = x_N y_N - x_1 y_1 - \langle x, H^{-1}Qy \rangle_H.$$

Note that these SBP operators do not alone guarantee strict stability for an initial boundary value problem. We must also apply the boundary conditions using a formulation that permits an energy estimate. More information on these boundary conditions, called simultaneous approximation term (SAT) boundary conditions, can be found at [47], [49], and [5]. Additionally, more information on SBP operators of various order, including the coefficients themselves, can be found in [45], [7], [32]. For this work, we will employ diagonal-norm SBP operators for the finite difference discretizations.

The SBP-SAT spatial discretization of the governing equations in the previous section (Equations (26) - (27)), in two dimensions and including the effect of coordinate transform from the physical space to a reference space $[-1,1] \times [-1,1]$, is given by

$$\rho J \frac{d\mathbf{v}_i}{dt} + \mathbf{D}_1 J \frac{\partial \xi_1}{\partial \mathbf{x}_i} \mathbf{p} + \mathbf{D}_2 J \frac{\partial \xi_2}{\partial \mathbf{x}_i} \mathbf{p} = -\mathbf{H}^{-1} \mathbf{F}_{v_i}, \quad i = 1, 2,$$
(30)

$$J\frac{\partial \boldsymbol{p}}{\partial t} + \lambda \left(J\frac{\partial \boldsymbol{\xi_1}}{\partial \boldsymbol{x_1}} \boldsymbol{D}_1 v_1 + J\frac{\partial \boldsymbol{\xi_2}}{\partial \boldsymbol{x_1}} \boldsymbol{D}_2 v_1 + J\frac{\partial \boldsymbol{\xi_1}}{\partial \boldsymbol{x_2}} \boldsymbol{D}_1 v_2 + J\frac{\partial \boldsymbol{\xi_2}}{\partial \boldsymbol{x_2}} \boldsymbol{D}_2 v_2 \right) = -\lambda \boldsymbol{H}^{-1} \boldsymbol{\mathcal{F}}_p.$$
(31)

Here, we have defined the matrices

$$H = H_{N_1} \otimes H_{N_2}, \qquad D_1 = D_{N_1} \otimes I_{N_2}, \qquad D_2 = I_{N_1} \otimes D_{N_2},$$
 (32)

where $\mathbf{D}_i = \mathbf{H}_i^{-1} \mathbf{Q}_i$ defines an SBP approximation of $\frac{\partial}{\partial x_i}$. We define corresponding solution vectors as, for example,

$$\mathbf{p} = \begin{bmatrix} p_{00} & p_{01} & \cdots & p_{0N_2} & p_{10} & \cdots & p_{N_1 N_2} \end{bmatrix}^T$$
 (33)

for a domain of size $(N_1 + 1) \times (N_2 + 1)$. Note also that in these equations, x_i refer to coordinates in the physical domain, whereas ξ_i refer to coordinates in the reference domain.

The matrix J is a diagonal matrix

$$J = \operatorname{diag} \left[J_{00} \quad J_{01} \quad \dots \quad J_{0N_2} \quad J_{10} \quad \dots \quad J_{N_1 N_2} \right],$$
 (34)

where the elements J_{kl} correspond to Jacobian determinants at grid point (k, l):

$$J = \frac{\partial x_1}{\partial \xi_1} \frac{\partial x_2}{\partial \xi_2} - \frac{\partial x_2}{\partial \xi_1} \frac{\partial x_1}{\partial \xi_2}.$$
 (35)

The bolded coordinate transform matrices $\frac{\partial \xi_i}{\partial x_j}$ and $\frac{\partial x_i}{\partial \xi_j}$ are defined analogously.

The penalty terms in the above equations implement the boundary conditions via the SAT method, and are defined as follows:

$$\mathcal{F}_{v_i} = \left(\boldsymbol{e}_W \otimes \mathcal{F}_{v_i}^W\right) + \left(\boldsymbol{e}_E \otimes \mathcal{F}_{v_i}^E\right) + \left(\mathcal{F}_{v_i}^S \otimes \boldsymbol{e}_S\right) + \left(\mathcal{F}_{v_i}^N \otimes \boldsymbol{e}_N\right), \tag{36}$$

$$\boldsymbol{\mathcal{F}}_{p} = \left(\boldsymbol{e}_{W} \otimes \boldsymbol{\mathcal{F}}_{p}^{W}\right) + \left(\boldsymbol{e}_{E} \otimes \boldsymbol{\mathcal{F}}_{p}^{E}\right) + \left(\boldsymbol{\mathcal{F}}_{p}^{S} \otimes \boldsymbol{e}_{S}\right) + \left(\boldsymbol{\mathcal{F}}_{p}^{N} \otimes \boldsymbol{e}_{N}\right). \tag{37}$$

In these expressions, the subscripts E, W, N, S refer to the edges of the domain, with the \boldsymbol{e} vectors being equal to zero in every entry except for the first or last. The actual penalty vectors $\boldsymbol{\mathcal{F}}_{v_i}^W$ and $\boldsymbol{\mathcal{F}}_p^W$ are expressed as

$$\boldsymbol{\mathcal{F}}_{v_{i}}^{W} = \boldsymbol{H}_{2} \boldsymbol{S}_{JW} \boldsymbol{n}_{i}^{W} \left(\boldsymbol{p}_{W}^{*} - \boldsymbol{p}_{W} \right), \qquad \boldsymbol{\mathcal{F}}_{p}^{W} = \boldsymbol{H}_{2} \boldsymbol{S}_{JW} \lambda \left(\boldsymbol{v}_{W}^{*} - \boldsymbol{v}_{W} \right), \tag{38}$$

where the surface normals and Jacobians are given by

$$S_J = J \sqrt{\left(\frac{\partial \xi_i}{\partial x_2}\right)^2 + \left(\frac{\partial \xi_i}{\partial x_1}\right)^2}, \quad n_1 = \pm \frac{J}{S_J} \frac{\partial \xi_i}{\partial x_2}, \quad n_2 = \pm \frac{J}{S_J} \frac{\partial \xi_i}{\partial x_1}, \tag{39}$$

and the corresponding bolded matrices are constructed to be diagonal and contain values of these quantities at each grid point, as with J (see (34)). The starred vectors p^* and v^* are specified by the boundary or interface condition.

It can be shown (see [25]) that the energy dissipation rate of an SBP block in this case is given by

$$\frac{dE}{dt} = \sum_{K=\{W,E,S,N\}} \mathcal{D}_K,\tag{40}$$

$$\mathcal{D}_{K} = -\boldsymbol{v}_{K}^{T}\boldsymbol{H}_{K}\boldsymbol{S}_{JK}\boldsymbol{p}_{K}^{*} + \boldsymbol{v}_{K}^{T}\boldsymbol{H}_{K}\boldsymbol{S}_{JK}\boldsymbol{p}_{K} - (\boldsymbol{v}_{K}^{*})^{T}\boldsymbol{H}_{K}\boldsymbol{S}_{JK}\boldsymbol{p}_{K}. \tag{41}$$

Naturally, this dissipation rate, in conjunction with a definition of p^* and v^* based on the numerical characteristics of the interface, will be critical in proving stability via the energy method.

2.3.2 Discontinuous Galerkin Method

The discontinuous Galerkin method we will target our interface method(s) to will be briefly summarized here. As in Kozdon and Wilcox, we use a curvilinear triangle-based DG method, and we denote a DG element Ω_e with triangular reference element $\tilde{\Omega}$. To obtain a version of the governing equations (26) - (27) usable by DG, we must first obtain the variational form by introducing test functions w_i and φ for the velocities and pressure respectively, multiplying them into their respective equations, and integrating over the reference element. Applying integration by parts to the resulting equations to move the spatial derivatives to the test functions gives us the skew-symmetric form of the variational equations, which can then be spatially discretized with the DG method (see Hesthaven and Warburton [17] for more detail on this) to obtain:

$$\rho \boldsymbol{M}_{J} \frac{d\boldsymbol{v}_{i}}{dt} = \boldsymbol{D}_{1}^{T} \boldsymbol{M}_{1i} \boldsymbol{p} + \boldsymbol{D}_{2}^{T} \boldsymbol{M}_{2i} \boldsymbol{p} - \sum_{K=1}^{3} \boldsymbol{L}_{K}^{T} \boldsymbol{P}_{bc}^{T} \boldsymbol{n}_{iK} \boldsymbol{\Omega}_{bc} \boldsymbol{S}_{JK} \boldsymbol{p}_{K}^{*}, \tag{42}$$

$$\boldsymbol{M}_{J} \frac{d\boldsymbol{p}}{dt} = -\lambda \left(\boldsymbol{M}_{11} \boldsymbol{D}_{1} \boldsymbol{v}_{1} + \boldsymbol{M}_{21} \boldsymbol{D}_{2} \boldsymbol{v}_{1} + \boldsymbol{M}_{12} \boldsymbol{D}_{1} \boldsymbol{v}_{2} + \boldsymbol{M}_{22} \boldsymbol{D}_{2} \boldsymbol{v}_{2} \right)$$
$$-\sum_{K=1}^{3} \lambda \boldsymbol{L}_{K}^{T} \boldsymbol{P}_{bc}^{T} \boldsymbol{\Omega}_{bc} \boldsymbol{S}_{JK} \left(\boldsymbol{v}_{K}^{*} - \boldsymbol{v}_{K}^{-} \right). \tag{43}$$

These equations provide a semi-discretization on each element. The normals n_i and surface Jacobians S_J are the same as the ones defined by (39), and the starred quantities again refer to boundary and interface conditions implemented via penalization, in this instance better referred to as numerical fluxes. The vector \mathbf{v}_K^- is the normal component of velocity along edge K of the element evaluated at the cubature points:

$$\boldsymbol{v}_{K}^{-} = \boldsymbol{n}_{1K} \boldsymbol{P}_{bc} \boldsymbol{L}_{K} \boldsymbol{v}_{1} + \boldsymbol{n}_{2K} \boldsymbol{P}_{bc} \boldsymbol{L}_{K} \boldsymbol{v}_{2}. \tag{44}$$

Edge-projected pressures are similarly defined as

$$\boldsymbol{p}_{K}^{-} = \boldsymbol{P}_{bc} \boldsymbol{L}_{K} \boldsymbol{p}. \tag{45}$$

In these expressions, L_K and L_K^T are operators which take volume terms to edge K of the element and edge K terms to the volume respectively. D_1 and D_2 are the reference element differentiation matrices for the two reference coordinate directions. The projection matrices P_c and P_{bc} project from the volume and edge approximations to the volume and edge cubature points, respectively. The matrices Ω_c and Ω_{bc} are diagonal matrices of the integration weights for the volume and an edge, respectively, defined at the cubature locations. A critical condition assumed by [25] for stability of their method is that Ω_c and Ω_{bc} are positive definite. Finally, the element mass matrices in the discretization are defined as

$$\boldsymbol{M}_{J} = \boldsymbol{P}_{c}^{T} \boldsymbol{\Omega}_{c} \boldsymbol{J} P_{c}, \quad \boldsymbol{M}_{ij} = \boldsymbol{P}_{c}^{T} \boldsymbol{\Omega}_{c} \boldsymbol{J} \frac{\partial \boldsymbol{\xi}_{i}}{\partial \boldsymbol{x}_{i}} \boldsymbol{P}_{c}.$$
 (46)

Here the diagonal matrices J and $\frac{\partial \xi_i}{\partial x_j}$ are defined identically to (34) and defined at the cubature points. The diagonal matrices S_{JK} and n_{iK} are defined in the same way at the cubature points.

As with SBP, it can be shown that a single DG element has the energy dissipation rate

$$\frac{dE}{dt} = \sum_{K=1}^{3} \mathcal{D}_K,\tag{47}$$

$$\mathcal{D}_{K} = -\left(\boldsymbol{v}_{K}^{-}\right)^{T}\boldsymbol{\Omega}_{bc}\boldsymbol{S}_{JK}\boldsymbol{p}_{K}^{*} - \left(\boldsymbol{p}_{K}^{-}\right)^{T}\boldsymbol{\Omega}_{bc}\boldsymbol{S}_{JK}\left(\boldsymbol{v}_{K}^{*} - \boldsymbol{v}_{K}^{-}\right). \tag{48}$$

As in the case of SBP, this relation is naturally critical in establishing the energy stability of the interface, especially considering its direct incorporation of starred (interface-driven) quantities.

2.3.3 Interface Method

Clearly critical in the proof of energy stability for the marriage of the above methods is the definition of the terms p^* and v^* , most often referred to as the numerical fluxes (in the context of DG) or penalty terms (in the context of SBP). At the interface between the two discretizations, these quantities are naturally contingent on the nature of the interface method, which for the purposes of this discussion take a basic form based on projection operators that move the Grid a solutions to Grid b,

$$\boldsymbol{f}_a = \boldsymbol{P}_{b2a} \boldsymbol{f}_b \tag{49}$$

$$\boldsymbol{f}_b = \boldsymbol{P}_{a2b} \boldsymbol{f}_a, \tag{50}$$

where f_a is of course a function defined on Grid a and f_b is a function defined on Grid b. Note that it is nowhere stated (and in fact proves to be untrue for SBP-to-SBP projection using Kozdon and Wilcox's method) that $P_{a2b}P_{b2a} = I$.

While the somewhat unique definitions of p^* and v^* , which incorporate these projected quantities directly, are perhaps equally important to provable stability, for now we will focus on the projection operation itself, using the work of Kozdon and Wilcox as a starting point. As detailed in their paper, the operators of Kozdon and Wilcox rely primarily upon the principle of projecting solutions from either discretization first into an intermediate "glue" space of piecewise continuous polynomials before projecting the solution onto the other nonconforming discretization. Whereas the solution on the respective structured and unstructured grids is represented at the block boundaries on the grid points, the solution on the intermediate glue grid is represented on each interval by modal coefficient values multiplying Legendre polynomials defined on the interval [-1, 1].

The critical condition that allows proof of stability is referred to as the *compatibility condition*, and amounts to a simple relation between the grid-to-glue and glue-to-grid projection operators, the mass matrix of the piecewise polynomial description used in the glue layer,

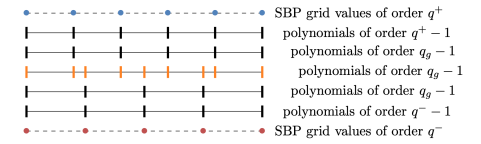


Figure 6: Image from Kozdon and Wilcox [25] demonstrating the glue layers present in grid-to-grid projection at a nonconforming interface.

and a norm operator H:

$$\boldsymbol{P}_{f2g}\boldsymbol{M} = \boldsymbol{P}_{q2f}^{T}\boldsymbol{H} \tag{51}$$

In the case of SBP finite difference discretizations, the norm \boldsymbol{H} is equivalent to the SBPnorm, which for the purposes of this work we will take to be a diagonal matrix. In the case of discontinuous Galerkin, this \boldsymbol{H} operator is the diagonal matrix of integration weights at the cubature points at a single DG element edge, Ω_{bc} . This compatibility condition, used in conjunction with the per-block semi-discrete expressions for energy dissipation (41) and (48), is what allows for energy stability to be proven.

Also critical in the projection from one spatial discretization to another via this method is the projection between glue spaces (see Figure 6). A compatibility condition similar to (51) is assumed for this projection,

$$\boldsymbol{M}_{a}\boldsymbol{P}_{gb2ga} = \boldsymbol{P}_{qa2qb}^{T}\boldsymbol{M}_{b} \tag{52}$$

but it is shown in [25] (see Lemma 2.3) that the operators P_{g2f} and P_{f2g} can also be generalized as projection operators to and from the finest glue layer. Given its clear importance, the compatibility condition (51) is used as an explicit constraint when constructing the projection operators P_{f2g} and P_{g2f} themselves. The construction, which is laid out in further detail in [25], defines interior and boundary accuracy conditions when projecting from grid to glue (and vice versa) that match those of corresponding SBP diagonal-norm operators of the same order. A pictorial representation of this process is given by Figure 7. These conditions, in conjunction with symmetry conditions, result in a non-square system featuring more unknowns than constraints, in particular near the boundary stencils. Given the underdetermined nature of the system, the authors then use an optimization procedure aimed at minimizing the distance between the nearest eigenvalues of $B = P_{f2g}P_{g2f}$ to produce the final projection operators for a given order. As mentioned in Section 2.1, our goal is to eliminate this optimization procedure, either by imposing more constraints (such as steps towards round-trip projection fidelity) or removing extraneous unknowns.

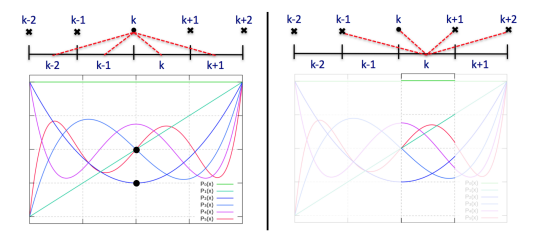


Figure 7: Images demonstrating the interior constraints for the projection mimicking SBP4-2 operators. Left: projection of modal coefficients of the piecewise polynomial solution on the four surrounding glue grid intervals to the central grid point in the stencil must result in a correct Legendre polynomial solution at point k up to the interior order. Right: projection of the solutions at the four points surrounding interval k to the modal coefficients on interval k must produce the correct piecewise approximation of Legendre polynomials up to interior order on that interval.

2.4 Validation Plan

The test problem for validation will match that of [25], modeling the two-dimensional acoustic wave equation in first-order form with the following initial condition:

$$p(x_1, x_2, 0) = \cos(k_1 x_1) \cos(k_1 x_2) + \sin(k_2 x_1) \sin(k_2 x_2), \tag{53}$$

$$v_i(x_1, x_2, 0) = 0, \quad i = 1, 2,$$
 (54)

where $k_1 = \pi/2$ and $k_2 = \pi$. All exterior boundary conditions are zero pressure (free-surface) conditions. The exact solution of this problem is

$$p(x_1, x_2, t) = \cos(\omega_1 t) \cos(k_1 x_1) \cos(k_1 x_2) + \cos(\omega_2 t) \sin(k_2 x_1) \sin(k_2 x_2), \tag{55}$$

$$v_1(x_1, x_2, t) = \frac{k_1}{\omega_1} \sin(\omega_1 t) \sin(k_1 x_1) \cos(k_1 x_2) - \frac{k_2}{\omega_2} \sin(\omega_2 t) \cos(k_2 x_1) \sin(k_2 x_2), \quad (56)$$

$$v_2(x_1, x_2, t) = \frac{k_1}{\omega_1} \sin(\omega_1 t) \cos(k_1 x_1) \sin(k_1 x_2) - \frac{k_2}{\omega_2} \sin(\omega_2 t) \sin(k_2 x_1) \cos(k_2 x_2), \quad (57)$$

where
$$\omega_j = k_j \sqrt{2}$$
 for $j = 1, 2$.

As for the spatial discretization, we will employ a square spatial domain with the left half using diagonal-norm SBP operators for finite differencing, and the right half using a discontinuous Galerkin method. The interior order of the SBP method, as well as the order of the projection across the interface, is specified to match the order of the discontinuous Galerkin method. Our success metrics for this test problem include:

• Demonstrating accurate recreation of the total energy of the system, as observed in

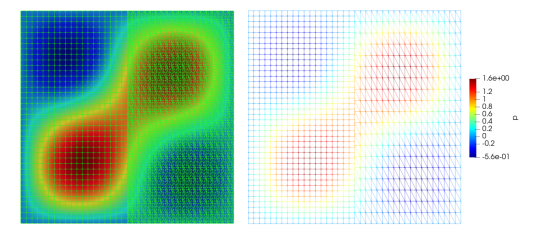


Figure 8: Pressure initial condition for test problem, also showing split-domain discretization. The right figure coarsens the wireframe to better show the individual triangular DG elements.

the numerical simulation of the problem, via a total energy relation derived from the spatial and interfacial schema.

- Demonstrating that this energy does not increase (condition for semi-discrete stability).
- Demonstrating the prescribed order of accuracy using L2 norms.

Upon fulfillment of these goals, extension to three dimensions will be considered.

2.5 Outlook

2.5.1 Current Status

Current work on this project has thus far amounted to reconstruction of the operators of Kozdon and Wilcox, application of these operators to simple, periodic SBP-to-SBP advection examples, and verification of their accuracy and energy-stability properties when focusing on the use of the interior projection stencil (see Figure 9). Having both verified and better understood the construction and capabilities of these operators, the question then becomes improving upon them, and the chief task to that end is better understanding the importance of the compatibility condition in both the construction of the operators and in the proof of energy stability. Namely, a question being asked via the use of symbolic evaluation tools such as Maxima [27] is whether there are alternative conditions on the grid-to-glue projection that will provide similar characteristics. In particular, one question that demands answer pertains directly to the *interior* operator construction, which in the grid-to-glue direction amounts to an equal number of degrees of freedom and accuracy constraints imposed upon the glue grid intervals (see Figure 7). In this case, the projection in the glue-to-grid direction can then be fully determined by the compatibility condition (Equation 51), but it is not clear why this also results in a glue-to-grid projection operator that fulfills its own accuracy constraints imposed on the grid points.

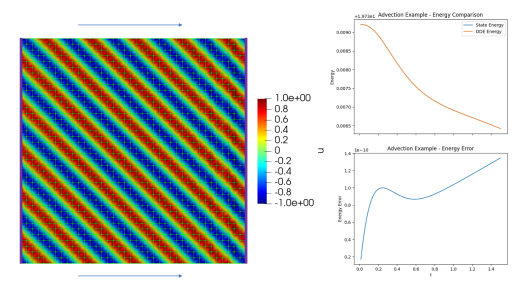


Figure 9: Application of the glue layer projection theory of [25] to a scalar advection example with an advection speed of 1. The purple line in the figure on the left is a glue layer to which we project the solution on the rightmost grid points at the periodic boundary in x. We then project from the glue layer to the leftmost set of grid points. The plots on the right show both an overall decrease in energy (energy stability) and a good agreement between the energy calculated from the time-integrated state and the energy calculated from time-integrating the energy dissipation rate.

Additionally, and related to these questions, another avenue of investigation is in the choices of the degree-of-freedom pattern in the projection from grid to glue layer and vice versa, as well as whether or not there are other polynomial bases for the glue layer that maintain positive-definite mass matrices while also introducing improved accuracy.

2.5.2 Risk Mitigation

Should our goal of a new and unisolvent stable interface method not come to fruition, a "fallback" goal of sorts is identified in that Kozdon and Wilcox's method has yet to be extended to three dimensions. This would present more of a technical challenge than an intellectual one, but a study of how the method responds to certain geometric scenarios untestable in two dimensions (namely three-dimensional corners) has potential to yield further information about how the optimized operators perform numerically, as well as shed light on optimization targets to yield better results in higher dimensions.

Furthermore, an additional unexplored topic that could prove to be a worthwhile "fallback" goal - and one that potentially ties the two topics of this proposal together - would be exploring the implications of the use of Kozdon and Wilcox's projection operators on the timestep limitations of a given physical problem, and (if a timestep disparity between the interior of a given subdomain and its interface with another subdomain is identified) exposing this challenge to multi-rate Adams integration.

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