Extendible and Efficient Python Framework for Solving Evolution Equations with Stabilized Discontinuous Galerkin Methods

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Abstract This paper discusses a Python interface for the recently published DUNE-FEM-DG module which provides highly efficient implementations of the Discontinuous Galerkin (DG) method for solving a wide range of non-linear partial differential equations (PDE). Although the C++ interfaces of DUNE-FEM-DG are highly flexible and customizable, a solid knowledge of C++ is necessary to make use of this powerful tool. With this work easier user interfaces based on Python and the Unified Form Language are provided to open DUNE-FEM-DG for a broader audience. The Python interfaces are demonstrated for both parabolic and first order hyperbolic PDEs.

Keywords DUNE, DUNE-FEM, Discontinuous Galerkin, Finite Volume, Python, Advection-Diffusion, Euler, Navier-Stokes

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In this paper we introduce a Python layer for the DUNE-FEM-DG1 module [13] which is available open-source. The DUNE-FEM-DG module is based on DUNE [4] and DUNE-FEM [19] in particular and makes use of the infrastructure implemented by DUNE-FEM for seamless integration of parallel-adaptive Finite Element based discretization methods. DUNE-FEM-DG focuses exclusively on Discontinuous Galerkin (DG) methods for various types of problems. The discretizations used in this module are described by two main papers, [16] where we introduced a generic stabilization for convection dominated problems that works on generally unstructured and non-conforming grids and [8] where we introduced a parameter independent DG flux discretization for diffusive operators.

DG methods have been studied intensively by many other groups and many software packages exist. However, most of these packages do not combine the following features: unstructured grids for 2, and 3 space dimensions, grid adaptivity, parallel computing capabilities, and open-source licenses. Besides DUNE-FEM-DG a few alternatives exists, for example, deal.II ([31]), feel++ ([26]), or Nektar++ ([32]) or FLEXI ([30]).

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1https://gitlab.dune-project.org/dune-fem/dune-fem-dg.git
DUNE-FEM-DG has been used in several applications (see [13] for a detailed list), most notably a comparison with the production code of the German Weather Service COSMO has been carried out for test cases for atmospheric flow ([7,45]). The focus of the implementation is on Runge-Kutta DG methods using mainly a matrix-free approach to handle implicit time discretizations which is a method especially used for convection dominated problems.

A strength of the DUNE-FEM-DG module is the general application area, i.e. convection dominated as well as diffusion dominated problems, for 1d, 2d, and 3d models, including parallelization and local grid adaptivity. The model includes the implementation of several discretizations for the second order terms such as Interior Penalty (and variants), Compact Discontinuous Galerkin 1 and 2, and Bassi-Rebay 1 and 2 as well as Local Discontinuous Galerkin. For the first order terms different numerical flux functions can be used and various limiter based stabilizations are available. For the time discretization a method of lines approach is adopted and a number of implicit, explicit, and IMEX schemes are available. Overall the module thus offers not only a strong base for building state of the art simulation tools but it also allows for the development of new methods and comparative studies of different method. Parallelization and adaptivity, including $hp$ adaptivity, can be used seamlessly with all methods.

A shortcoming so far has been the template heavy and relatively complicated C++ user interfaces, leading to a steep learning curve for implementing new models and applications or coupling of such. Recent development (e.g. [18]) has therefore been focused on adding a Python layer on top of DUNE-FEM-DG allowing user friendly model description based on the Unified Form Language (UFL) [1] and code generation. While quite a few applications using UFL and also consider DG approximation, e.g., the FEniCS package [41], only a few examples exists that consider convection dominated evolution equations. For example, in [31] UFL is used to describe weak forms for the compressible Euler and Navier-Stokes equations but only in the stationary setting. Another example is [47] where shallow water applications are considered. A package that also provides Python bindings (not using UFL) but focuses on hyperbolic problems is [42], where higher order Finite-Volume schemes are the method of choice. To our best knowledge this is the first package which combines high level scripting support with efficient stabilized Discontinuous Galerkin methods for solving the full range of pure hyperbolic systems, through advection dominated problems, to diffusion equations.

In DUNE low level Python bindings were introduced for the DUNE grid interface in [22] and a detailed tutorial providing high level access to DUNE-FEM is also available [15]. In this work, we describe how these bindings can now be used. The new interface, together with the efficient and flexible C++ implementation of the DG methods available in DUNE-FEM-DG, make the rapid prototyping of new methods straightforward and provides a flexible framework to solve very complex coupled nonlinear evolution equations. Other works in this direction

Motivation and aim of this paper

This paper describes a collaborative effort to establish a test and research environment for DG methods based on DUNE and DUNE-FEM for advection-diffusion problems ranging from advection dominated to diffusion only problems. The aim here is to provide easy access to a comprehensive collection of existing methods and techniques to serve as a starting point for new developments and comparisons without having to reinvent the wheel. This is combined with the easy access to advanced features like the availability of different grid element types, not limited to dimensions less than or equal to three, $hp$ adaptation, and parallel computation with both distributed (e.g. MPI) and shared memory (e.g. OpenMP)
parallelization, with excellent scalability [36] even for large core counts and very good on-
core efficiency in terms of floating point operations per second [13]. Python has become a
widespread programming language in the scientific computing community including the use
in industry we see great potential in improving the scientific development of DG methods
by providing access to state of the art research tools which can be used through the Python
scripting language, backed up by an efficient C++ backend.

The focus of this paper is to describe the different parts of the framework and how they
can be modified by the user in order to tailor the code to their needs and research interests.
In this paper we will mainly look at advection dominated evolution equations because most
diffusion dominated problems can be often completely described within a variational frame-
work for which the domain specific language UFL is very well suited and the standard code
generation features available in DUNE-FEM are thus sufficient. For advection dominated
problems additional ingredients are often required, e.g., for stabilization, that do not fit the
variational framework. This aspect of the algorithm will be a central part of this paper. The
development and improvement of the DG method in general or the high performance com-
puting aspect of the underlying C++ framework, which has been investigated previously, are
outside the scope of this work.

The paper is organized as follows. In Section 1 we briefly recall the main building blocks
of the DG discretization of advection diffusion problems. In Section 2 we introduce the
newly developed Python based model interface. In Section 3 we investigate the performance
impact of using Python scripting and conclude with discussing the extensibility of the ap-
proach in Section 4. Additional code examples are available in Appendix A.

1 Governing Equations and Discretization

We consider a general class of time dependent nonlinear advection-diffusion-reaction prob-
lems for a vector valued function $U : (0,T) \times \Omega \rightarrow \mathbb{R}^r$ with $r \in \mathbb{N}^+$ components of the form

$$\partial_t U = \mathcal{L}(U) := -\nabla \cdot (F_c(U) - F_v(U, \nabla U)) + S_i(U) + S_e(U) \quad \text{in} \quad (0,T] \times \Omega \quad (1)$$

in $\Omega \subset \mathbb{R}^d$, $d = 1,2,3$. Suitable initial and boundary conditions have to be added. $F_c$
describes the convective flux, $F_v$ the viscous flux, $S_i$ a stiff source term and $S_e$ a non-stiff
source term. Note that all the coefficients in the partial differential equation are allowed to
depend explicitly on the spatial variable $x$ and on time $t$ but to simplify the presentation
we suppress this dependency in our notation. Also note that any one of these terms is also
allowed to be zero.

For the discretization we use a method of lines approach based on first discretizing the
differential operator in space using a DG approximation and then solving the resulting
system of ODEs using a time stepping scheme.

1.1 Spatial Discretization

Given a tessellation $\mathcal{T}_h$ of the domain $\Omega$ with $\bigcup_{K \in \mathcal{T}_h} K = \Omega$ we introduce a piecewise
polynomial space $V_h = \{ v \in L^2(\Omega, \mathbb{R}^r) : v|_K \in [P_k(K)]^r, K \in \mathcal{T}_h \}$ for some $k \in \mathbb{N}$, where
$P_k(K)$ is a space containing all polynomials up to degree $k$.

Furthermore, we denote with $\Gamma$ the set of all intersections between two elements of the
grid $\mathcal{T}_h$ and accordingly with $\Gamma$ the set of all intersections, also with the boundary of the
domain $\Omega$. 

We seek $U_h \in V_h$ be discretizing the spatial operator $\mathcal{L}(U)$ in (1) with either Dirichlet, Neumann, or Robin type boundary conditions by defining for all test functions $\phi \in V_h$,

$$\langle \phi, \mathcal{L}_h(U_h) \rangle := \langle \phi, K_h(U_h) \rangle + \langle \phi, I_h(U_h) \rangle$$

with the element integrals

$$\langle \phi, K_h(U_h) \rangle := \sum_{e \in \mathcal{E}_h} \int_e \left( (F_c(U_h) - F_c(U_h, \nabla U_h)) : \nabla \phi + \mathcal{S}(U_h) \cdot \phi \right),$$

with $\mathcal{S}(U_h) = S_i(U_h) + S_c(U_h)$ and the surface integrals (by introducing appropriate numerical fluxes $\tilde{F}_c, \tilde{F}_v$ for the convection and diffusion terms, respectively)

$$\langle \phi, I_h(U_h) \rangle := \sum_{e \in \mathcal{E}_h} \int_e \left( \langle \{F_c(U_h, [U_h]_e)\}^{T} : \nabla \phi \rangle + \langle \{F_c(U_h, \nabla U_h)\} \rangle : \{\phi\}_e \right)$$

$$- \sum_{e \in \mathcal{E}_h} \int_e (\tilde{F}_c(U_h) - \tilde{F}_v(U_h, \nabla U_h)) : \{\phi\}_e,$$

with $\{\{U\}_e, \{\nabla U\}_e \}$ denoting the classic average and jump of $U$ over $e$, respectively. The convective numerical flux $\tilde{F}_c$ can be any appropriate numerical flux known for standard finite volume methods e.g. $\tilde{F}_c$ could be simply the local Lax-Friedrichs flux function (also known as Rusanov flux)

$$\tilde{F}_c^{\text{L-LLF}}(U_h)_e := \langle \{F_c(U_h)\} \rangle_e + \lambda_e \frac{\|U_h\|}{2}$$

where $\lambda_e$ is an estimate of the maximum wave speed on intersection $e$. One could also choose a more problem tailored flux (i.e. approximate Riemann solvers). Different options are implemented in DUNE-FEM-DG (cf. [13]). A wide range of diffusion fluxes $\tilde{F}_v$ can be found in the literature and many of these fluxes are available in DUNE-FEM-DG, for example, Interior Penalty and variants, Local DG, Compact DG 1 and 2 as well as Bassi-Rebay 1 and 2 (cf. [8,13]).

1.2 Temporal discretization

To solve the time dependent problem (1) we use a method of lines approach in which the DG method described above is first used to discretize the spatial operator and then a solver for ordinary differential equations is used for the time discretization, see [13]. After spatial discretization, the discrete solution $U_h(t) \in V_h$ has the form $U_h(t, x) = \sum U_i(t) \phi_i(x)$. We get a system of ODEs for the coefficients of $U(t)$ which reads

$$U'(t) = f(U(t)) \text{ in } (0, T]$$

with $f(U(t)) = M^{-1} \mathcal{L}_h(U_h(t))$, $M$ being the mass matrix which is in our case is block diagonal or even the identity, depending on the choice of basis functions. $U(0)$ is given by the projection of $U_0$ onto $V_h$. A range of different ODE solvers are available most based around Strong Stability Preserving Runge-Kutta methods (SSP-RK) (for details see [13]). In Section 2 we show how to easily implement other Runge-Kutta methods into the code. The results and implementation techniques presented in this paper can be applied to explicit, implicit, or semi-implicit methods and mostly the non-linear system arising from any implicit treatment is solved using a matrix-free Newton-Krylov method.
When using semi-implicit time stepping, the operator $\mathcal{L}$ is split such that $\mathcal{L}(U) = \mathcal{L}_e(U) + \mathcal{L}_i(U)$ with

$$
\mathcal{L}_e(U) := -\nabla \cdot F_e(U) + S_e(U) \quad \text{and} \quad \mathcal{L}_i(U) := \nabla \cdot F_i(U, \nabla U) + S_i(U)
$$

where $\mathcal{L}_e$ is treated explicitly and $\mathcal{L}_i$ is treated implicitly. Of course other splittings are possible but currently not available through the Python bindings.

1.3 Stabilization

The RK-DG method is stable when applied to linear problems such as linear hyperbolic systems; however for nonlinear problems spurious oscillations occur near strong shocks or steep gradients. In this case the RK-DG method requires some extra stabilization. In fact, it is well known that only the first order scheme ($k = 0$) produces a monotonic structure in the shock region. Many approaches have been suggested to make this property available in higher order schemes, without introducing the amount of numerical viscosity, which is such a characteristic feature of first order schemes. Several approaches exist, among those are slope limiters [12, 34, 16, 40, 25] and for a comprehensive literature list we refer to [46]. Another popular stabilization technique is the artificial diffusion (viscosity) approach [27, 35, 44, 29, 23] and others. Further techniques exists, such as a posteriori techniques to stabilize the DG method [20] or order reduction [24].

The construction of the general stabilization approach implemented in DUNE-FEM-DG is described in detail in [16] and follows the slope limiting approach combined with a troubled cell indicator suggested in [40]. However, since multiple discretization techniques for the diffusion operator are available within DUNE-FEM-DG, the implementation of an artificial diffusion approach would be straightforward.

In the following we briefly recall the main steps since these are necessary to understand the code design decision later on.

A stabilized discrete operator is constructed by concatenation of the DG operator $\mathcal{L}_h$ from (2) and a stabilization operator $\Pi_h$, leading to a modified discrete spatial operator $\tilde{\mathcal{L}}_h(U_h):= (\mathcal{L}_h \circ \Pi_h)(U_h)$. The stabilizations considered in this work can be computed element wise based only on data from neighboring elements thus not increasing the stencil of the operator. Given a DG function $U_h$ we call $U_h^\ast = \Pi_h(U_h)$ the stabilized DG function and we call $U_E = U_h^\ast|E$ the restriction of a function $U_h$ on element $E$ and denote with $\bar{U}_E$ its average. Furthermore, we call $e$ the intersection between two elements $E, K$ for $K \in \mathcal{N}_E$, with $\mathcal{N}_E$ being the set of neighbors of $E$ and $\mathcal{I}_E$ the set of intersections of $E$ with it’s neighbors.

The stabilized solution should fulfill the following requirements:

1. Conservation property, i.e. $\bar{U}_E = \bar{U}_E^\ast$
2. Physicality of $U_E^\ast$, i.e. values of $U_E^\ast$ belong to the set of states, i.e. positive density etc. at least for all quadrature points used to compute element and surface integrals.
3. Identity in "smooth" regions, i.e. in regions where the solution is "smooth" we have $U_h^\ast = U_h$. This requires an indicator for the smoothness of the solution.
4. Consistency for linear functions, i.e. if the average values of $U_h$ on $E$ and its neighbors are given by the same linear function $L_E$ then $U_E^\ast = L_E$ on $E$.
5. Minimal stencil, i.e. the stabilized DG operator shall have the same stencil as the original DG operator (only direct neighboring information in this context).
6. Maximum-minimum principle and monotonicity, i.e. in regions where the solution is not smooth the function $U^*_E$ should only take values between $\min_{K \in \mathcal{N}_E} \bar{U}_K$ and $\max_{K \in \mathcal{N}_E} \bar{U}_K$.

These requirements are built into the stabilization operator in different ways. For example, requirement 2 and 3 form the so called troubled cell indicator that triggers whether a stabilized solution has to be computed. Requirement 5 simply limits the choice of available reconstruction methods, e.g. higher order finite volume reconstructions could not be used because of the potentially larger stencil that would be needed which is not desired here.

The stabilization is then defined in two steps

1. We define the set of troubled cells by
   
   $$TC(U_h) := \{ E \in \mathcal{T}_h : J_E(U_h) > TOL \text{ or } U_E \text{ has unphysical values} \}$$

   with $J_E$ being a smoothness indicator and $TOL$ a threshold that can be influenced by the user (see Figure 9 in Section 2.3). As default we use $TOL = 1$ and a smoothness indicator based on the superconvergence result from [40]: this indicator accumulates the integral of the jump of some scalar quantity derived from $U_h$ denoted with $\phi(U_E, U_K)$ over all inflow boundaries of $E$ (defined as the intersection where some given velocity $v$ and the normal $n$ have opposite signs), i.e.

   $$J_E(U_h) := \sum_{e \in \mathcal{F}_E, v \cdot n < 0} \left( \int_e \phi(U_E, U_K) ds \right) \left( \frac{\alpha_d(k) h_E^{(k+1)/4} |e|}{\alpha_d(k) h_E^{(k+1)/4} |e|} \right),$$

   $\alpha_d(k) = \frac{2}{125} d_5^5$ denotes a scaling factor, $h_E$ is the elements diameter and $|e|$ the area of the intersection between the two elements $E$ and $K$. Please note the slight derivation from the notation in [16] by denoting the smoothness indicator in (9) with $J_E$ instead of $S_E$ like in [16].

2. (a) If $E \notin TC(U_h)$ then $U'_E = U_E$, i.e. the operator $\Pi_h$ is just the identity.
   (b) Construction of an admissible DG function if $E \in TC(U_h)$. In this case the DG solution on $E$ needs to be altered until $E$ is no longer a troubled cell. In this work we guarantee this by restricting ourselves to the reconstruction of limited linear functions based on the average values of $U_h$ on $E$ and it’s neighbors. Two reconstruction methods are implemented, one described in [16] and extended to general polyhedral cells in [37] and an optimization based strategy described in [9] based in ideas from [43]. Both strategies correspond to 2nd order MUSCL type finite volume schemes when used with piecewise constant basis functions.

1.4 Summary of building blocks and limitations

- Grid structure: with discontinuous Galerkin methods we can use any form of grid elements, from (axis aligned or general) cubes, simplices, to general polyhedrons). The best choice will depend on the application. In addition different local adaptation strategies can be considered as part of the underlying grid structure, e.g., conforming refinement using red-green or bisection strategies, for example, or simple nonconforming refinement. From a parallelization point of view DUNE-FEM-DG is restricted to DG methods that can be implemented using direct neighboring information only. On the one hand this is beneficial for many core architectures and on the other hand unstructured grids in DUNE only implement this ghost cell approach and arbitrary overlap is only available for Cartesian grids.
– **Space:** an advantage of discontinuous Galerkin methods is that there is little restriction on the combination of grids and discrete spaces one can use. For approximation purposes these spaces are generally chosen so that they contain the full polynomials of a given order on each element. This order is perhaps the most important property of the spaces. But even after having fixed the space, the actual set of basis functions to use to represent the space, can have a huge impact on the performance of the scheme. Central here is the structure of the local mass matrix on each element of the grid. This has to be inverted in each time step so efficiency here can be crucial. So a possible choice is to use a polynomial space with basis functions orthonormalized over the reference elements in the grid. If the geometric mapping between physical grid elements and these reference element is affine then the resulting local mass matrix is a very simple diagonal matrix. If the mapping is not affine then the mass matrix depends non trivially on the element under consideration and will often be dense. In this case another approach is to use Lagrange type basis functions where the interpolation points coincide with a suitable quadrature for the mass matrix. The case of non affine mapping is especially relevant for cube grids and here tensor product quadrature points can be used to construct the Lagrange type space. A possible approach is to use a tensor product Gaussian rule which results in a diagonal mass matrix for each element in the grid with a trivial dependency on the element geometry. Another typical approach found in the literature is often referred to a spectral discontinuous Galerkin method based on Lobatto-Gauss-Legendre quadrature \cite{39,38}. Although this quadrature is not precise enough to exactly integrate products of the basis functions, it has been shown that diagonalizing (lumping) the mass matrix in this way leads to a well posed scheme.

– **Numerical fluxes:** the discretization of both the diffusion and advective terms, involve boundary integrals where the discrete solution is not uniquely defined. Consequently, numerical fluxes have to be used to evaluate these terms. We consider fluxes here which depend on the traces of the discrete solution on both sides of the interface. A wide range of choices are available. In the package presented here a large number of fluxes for the diffusion are available, but since we are focusing on advection dominated problems we will not discuss this further. The simplest choice for the advective flux is the Local-Lax-Friedrichs or Rusanov flux from equation (5) which requires little additional input from the user and in combination with higher order schemes generally gives good results.

– **Time evolution:** after the spatial discretization, the resulting ordinary differential equation is solved using a Runge-Kutta as already mentioned. We will only briefly discuss this aspect of the method in the following.

– **Stabilization:** as was discussed in some detail in the previous section, our stabilization approach is based on a troubled cell indicator (combining a smoothness indicator with a check on physicality at quadrature points) and a reconstruction process of the solution in troubled cells.

If not mentioned otherwise, the results presented in the following are obtained using a Cartesian grid with a forth order polynomial space spanned by orthonormal basis function. We use the local Lax Friedrichs flux (5) for the advection term and the CDG2 method \cite{8} for the problems with diffusion. The troubled cell indicator is based on a smoothness indicator described above. The reconstruction is computed based on a linear programming problem as described in \cite{9}. Finally, we use a standard third order SSP Runge-Kutta method for the time evolution \cite{28}.

In the following we will describe how the different building blocks making up the DG method can be provided by the user, starting with some very simple problems, requiring little
input from the user and slowly building up the complexity. We are not able to provide the full code listings in each case but a tutorial with all the test cases presented here is available [17].

2 Customizing the DG method using the Python Interface

First we need to choose the grid structure and the computational domain. For simplicity we concentrate here on problems defined on 2D intervals \([x_l, x_r] \times [y_b, y_t]\) and use an axis aligned cube grid capable of non-conforming refinement, with \(n_x, n_y\) elements in \(x\) and \(y\) direction, respectively. The space consists of piecewise polynomials of degree \(p = 4\) spanned by an orthonormal basis over the reference element \([0, 1]^2\). This can be setup with a few lines of Python code:

```python
from dune.alugrid import aluCubeGrid as grid
from dune.fem.view import adaptiveLeafGridView as view  # needed for adaptive simulation
from dune.fem.space import dgonb

gridView = view(grid(Model.domain))  # needed for adaptive simulation
gridView.hierarchicalGrid.globalRefine(3)  # refine a possibly coarse initial grid
space = dgonb(gridView, dimRange=Model.dimRange, order=4)
U_h = space.interpolate(Model.U0, name="solution")
```

The `Model` class will be specified in the following. It contains information on the PDE but as seen above also provides a description of the domain, the number of components of the PDE to solve and the initial conditions \(U_0\) defined as a UFL expression. Examples will be provided in the next sections. After setting up the grid, the space, and the discrete solution, we define the DG operator and the default Runge-Kutta time stepper:

```python
from dune.femdg import femDGOperator
from dune.femdg.rk import femdgStepper

operator = femDGOperator(Model, space, limiter=None)
stepper = femdgStepper(order=3, operator=operator)
```

Again we use the `Model` class to provide all required information to the DG operator. The constructor of the DG operator takes a number of additional parameters which will be described throughout this section as required (see also Appendix A). In our first example we will require no stabilization so the `limiter` argument is set to `None`. In the final line of code, we pass the constructed operator to the time stepper together with the desired order (as pointed out we use order 3 for all presented simulations). Finally, a simple loop is used to evolve the solution from the starting time (assumed to be 0) to the final time \(T\) (which is again a property of the `Model` class):

```python
t = 0
while t < Model.endTime:
    dt = stepper(U_h)
    t += dt
```

The call method on the stepper evolves the solution \(U_h\) from the current to the next time step and returns a suggested size \(\Delta t\) for the next time step. In the following we will describe the `Model` class in detail and show how the above code snippets have to be modified to include additional features like stabilization for nonlinear advection problems or adaptivity. We will not describe each problem in detail, the provided code should give all the required information.
2.1 Linear advection problem

In the following we solve a linear advection problem and explain various options for stabilization and local grid adaptivity.

2.1.1 Model description

```python
from dune.ufl import cell
from dune.grid import cartesianDomain
from ufl import *

x = SpatialCoordinate(cell(2))  # setup a 2d problem
y = x - as_vector([0.5,0.5])
velocity = as_vector([-y[1],y[0]])  # velocity

# setup for 'three body problem'
# body 1: cube
cube = conditional(x[0]>0.6, 1., 0.) * conditional(x[0]<0.8, 1., 0.)
# body 2: slotted cylinder with center (0.5,0.75) and radius 0.1
slotted_cylinder = conditional((x[0] - 0.5)**2 + (x[1] - 0.75)**2 < 0.01, 1.0, 0.0)
# body 3: smooth hump with center (0.25,0.5)
hump = conditional((x[0] - 0.25)**2 + (x[1] - 0.5)**2 < 0.01, 1.0, 0.0)

class Model:
    dimRange = 1
    endTime = pi
    U0 = [cube + slotted_cylinder + hump]
    domain = cartesianDomain((0,0),(1,1),(10,10))

def F_c(t,x,U):
    return as_matrix([*(velocity*U[0])])

def maxLambda(t,x,U,n):
    return abs(dot(velocity,n))

# simple 'dirchlet' boundary conditions on all boundaries
boundary = {range(1,5): lambda t,x,U: as_vector([0])}
```

Note that to use the local Lax-Friedrichs flux we need to define the maximum wave speed of the hyperbolic problem. This function is also used to provide an estimate for the time step based on the CFL condition, which is returned by the stepper. The stepper also provides a property `deltaT` which allows the user to fix a time step to use for the evolution. A result for the described *three body problem* is presented in Fig. 1 (left) and Fig. 2 (left).

2.1.2 Stabilization

It is well known that the DG method with a suitable numerical flux is stable when applied to linear advection problems like the one studied here. But it does produce localized over and undershoots around steep gradients. While this is not necessarily a problem for linear equations, it can be problematic in some instances, e.g., when negative values are not acceptable. More crucially, this behavior leads to instabilities in the case of nonlinear problems. Therefore, stabilization mechanism have to be provided. As mentioned above we use a reconstruction approach combined with a troubled cell indicator. The standard troubled cell indicator is described in Section 1.3 and requires the user to provide some additional information in the `Model` class, i.e., a description of the jump of the solution \( \phi(U,V) \) between of two elements \( E \) and \( K \) and a suitable velocity \( v \) required to compute the smoothness indicator given in equations (9):
Fig. 1: Three body advection problem at time $t = \pi$. Left without stabilization and right with stabilization. Color range based on minimum and maximum values of discrete solution.

Fig. 2: Three body advection problem at time $t = \pi$. Left without stabilization and right with stabilization. Values above $1 + 10^{-3}$ or below $-10^{-3}$ are colored in grey and black, respectively.

We now need to construct the operator to include stabilization:

```python
operator = femDGOperator(Model, space, limiter="minmod")
U_h = space.interpolate(Model.U0, name="solution")
operator.applylimiter(U_h)  # apply limiter to initial solution
```

A result for three body problem including a stabilization is presented in Fig. 1 (right) and Fig. 2 (right).

Although over- and undershoots are clearly reduced by the stabilization approach there are still some oscillations clearly visible. There can be removed by basing the troubled cell indicator on a physicality check, e.g., requiring that the solution remains in the interval $[0, 1]$.

So (possibly instead of the `jump, velocity`) attributes we add

```python
Model.physical = lambda t,x,U, V: conditional(U[0]>-1e-8, 1.0, 0.0) *
```

Fig. 3: Three body advection problem at time \( t = \pi \). On the left using physicality check to reduce oscillations and on the right using scaling limiter. Values above \( 1 + 10^{-3} \) or below \( -10^{-3} \) are colored in grey and black, respectively.

```
conditional( U[0]<1.0+1e-8, 1.0, 0.0 )
```

to the code. Another alternative is to use the approach detailed in [48, 11] which required to add a tuple with upper and lower bounds for each component in the solution vector to the Model and to change the limiter parameter in the operator constructor:

```
Model.lowerBound = [0]  # the scalar quantity should be positive
Model.upperBound = [1]  # the scalar quantity should be less than one
operator = femDGOperator(Model, space, limiter="scaling")
```

The resulting plots using the scaling limiter and simply limiting in cells where the solution is below zero or above one, are shown in Fig. 3.

### 2.1.3 Adaptivity

Due to the regions of steep gradients and low regularity in the solution, advection dominated problems benefit a lot from the use of local grid refinement and coarsening. Although this is a more general feature of the DUNE-FEM framework on which the package is based, adaptivity is such an important tool to improve efficiency of schemes for the type of problems discussed here, that we will describe how to use it in some detail. The grid modification requires an indicator computed between time steps which provides the information for each cell whether to keep it as it is, refine it, or coarsen it. DUNE-FEM provides a very simple function

```
dune.fem.markNeighbors(indicator, refineTol, coarsenTol, minLevel, maxLevel)
```

Here indicator provides a number \( \eta_E \geq 0 \) on each element \( E \) which is used to determine if an element is to be refined or coarsened. An element \( E \) is refined, if it’s level in the grid hierarchy is less than \( \text{minLevel} \) and \( \eta_E > \text{refineTol} \); it is coarsened if it’s level is greater than \( \text{maxLevel} \) and \( \eta_E < \text{coarsenTol} \); otherwise it remains unchanged. In addition if a cell is to be refined all its neighboring cells are refined as well, so that important structures in the solution do not move out of refined regions during a time step. If the initial grid is suitably refined, then one can take \( \text{maxLevel}=0 \). The choice for \( \text{minLevel} \) will depend on the problem and has to be chosen carefully since increasing \( \text{minLevel} \) by one can potentially double the runtime simply due to the reduction in the time step due to the CFL condition. At the moment we do not provide any spatially varying time step control. We usually choose \( \text{coarsenTol} \) simply as a fraction of \( \text{refineTol} \). In our simulation we use
coarsenTol=0.2*refineTol. This leaves us needing to describe our choice for indicator and refineTol.

The simulations shown here use a residual type indicator based on a scalar quantity and flux provided in a subclass Indicator of the Model class. We consider the residual of an auxiliary PDE

\[ \partial_t \eta(t,x,U(t,x,U)) + \nabla \cdot (F(t,x,U,U)) = S(t,x,U,U) \]  

where for example \( \eta,F,S \) could simply be taken from one of the components of the PDE or could be based on an entropy, entropy flux pair. We now define the element residual to be

\[ \begin{align*}
\eta^2_E &= h^2_E \| R_{\text{vol}} \|^{2}_{L^2(E)} + \frac{1}{2} \sum_{e \in \partial E} h_e \| R_{e2} \|^{2}_{L^2(e)} + \frac{1}{h_e} \| R_{e1} \|^{2}_{L^2(e)} \\
\end{align*} \]

where \( R_{\text{vol}} \) is a discretized version of the interior residual indicating how accurate the discretized solution satisfies the auxiliary PDE at every interior point of the domain for two timesteps \( t^n \) and \( t^{n+1} \).

\[ R_{\text{vol}} := \frac{1}{t^{n+1} - t^n} (\eta(t^{n+1},U^{n+1}) - \eta(t^n,U^n)) \]

\[ + \frac{1}{2} \nabla \cdot (F(t^{n},U^n) - F(t^{n+1},U^{n+1})) - \frac{1}{2} (S(t^n,U^n) - S(t^{n+1},U^{n+1})) \]

and we have jump indicators

\[ R_{e2} := \frac{1}{2} \| F(t^n,\ldots,F(t^{n+1},\ldots) \|_e \quad R_{e1} := \frac{1}{2} \| \eta(t^n,\ldots,\eta(t^{n+1},\ldots) \|_e. \]

This indicator is just one of many possible choices, e.g., simply taking the jump of \( U_h \) over intersections will often lead to very good results as well. We have had very good experiences with this indicator as shown here see also \[14\] where we used a similar indicator and \[20\] where a similar indicator was derived.

For the advection problem we will simply use the original PDE:

```python
class Indicator:
    def eta(t,x,U):
        return U[0]
    def F(t,x,U,DU):
        return Model.F_c(t,x,U)[0]
    def S(t,x,U,DU):
        return 0
Model.Indicator = Indicator
```

The indicator is now computed by applying an operator mapping the DG space onto a finite volume space resulting an elementwise constant value for the indicator which we can use to refine/coarsen the grid:

```python
# previous version used three global refinement steps
maxLevel = 3  # maximal allowed level for grid refinement
un = U_h.copy()  # to store solution at previous time

from dune.fem.space import finiteVolume
from dune.ufl import Constant
from dune.fem import markNeighbors, adapt
from dune.fem.operator import galerkin

indicatorSpace = finiteVolume(gridView)
indicator = indicatorSpace.interpolate(0, name="indicator")
dt, t = Constant(1,"dt"), Constant(0,"t")
x, n = SpatialCoordinate(space), FacetNormal(space)
hT, he = MaxCellEdgeLength(space), avg(CellVolume(space)) / FacetArea(space)
eta, F, S = Model.Indicator.eta, Model.Indicator.F, Model.Indicator.S
eta_new, eta_old = eta(t+dt,x,u), eta(t,x,un)
```
etaMid = ( eta(t,x,un) + eta(t+dt ,x,u) ) / 2
FMid = ( F(t,x,un,grad(un)) + F(t+dt ,x,u,grad(u)) ) / 2
SMid = ( S(t,x,un,grad(un)) + S(t+dt ,x,u,grad(u)) ) / 2
Rvol = ( eta_new - eta_old )/dt + div(FMid) - SMid
estimator = hT**2 * Rvol **2 * phi * dx +
             he * inner ( jump ( FMid ), n('+'))**2 * avg ( phi ) * dS +
             1/he * jump ( etaMid )**2 * avg ( phi ) * dS
residualOperator = galerkin ( estimator )

In the following we refine the initial grid to accurately resolve the initial condition. We also use the initial indicator to fix a suitable value for the refineTol:

maxSize = gridView . size (0)*2**(gridView . dimension*maxLevel)

# initial refinement
for i in range (maxLevel +1):
    un. assign ( U_h )
    dt = stepper ( U_h )
    residualOperator . model .dt = dt
    timeTol = gridView . comm . sum ( sum ( indicator . dofVector ) ) / Model . endTime / maxSize
    hTol = timeTol * dt
    markNeighbors ( indicator , refineTolerance=hTol , coarsenTolerance=0.2*hTol ,
                   minLevel =0, maxLevel =maxLevel )
    adapt ( U_h )
    U_h . interpolate ( Model .U0)
    operator . applyLimiter ( U_h )

Within the time loop we now also need to add the marking and refinement methods, we repeat the full time loop here:

Results on a dynamically adapted grid with and without stabilization are shown in Fig. 4 and Fig. 5.

2.2 Compressible Euler equations

We continue our presentation with a system of evolution equations. We focus on simulations of the compressible Euler (and later Navier-Stokes) equations. We show results for two standard test cases: the interaction of a shock with a low density region and the simulation of a Kelvin-Helmholtz instability with a density jump.

2.2.1 Model description

We start with the methods needed to describe the PDE, the local Lax-Friedrichs flux and time step control, the troubled cell indicator, and the residual indicator:
Fig. 4: Three body advection problem at time $t = \pi$ using an adaptive grid. Left without stabilization and right with stabilization. Color range based on minimum and maximum values of discrete solution.

Fig. 5: Three body advection problem at time $t = \pi$ using an adaptive grid. Left without stabilization and right with stabilization. Top shows solution with values above $1 + 10^{-3}$ or below $-10^{-3}$ are colored in grey and black, respectively. Bottom row shows corresponding grid with solution colored according to their respective minimum and maximum values.
The next step is to fix the initial conditions and the end time for the two problems we want to study. First for the shock bubble problem

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To complete the description of the problem we need to define boundary conditions. For the advection problem we used Dirichlet boundary conditions which are used as second state
for the numerical flux over the boundary segments. For this problem we will use Dirichlet boundary conditions on the left and right boundary but want to use no flow boundary conditions on the top and bottom boundary (which will have an identifier of 3 on this domain):

```python
def noFlowFlux(u,n):
    p = Model.toPrim(u)
    return as_vector([0] + [c*n[2] for c in n] + [0])
```

Next we describe initial conditions and boundary conditions for the Kelvin-Helmholtz instability between two layers with a density jump where we use periodic boundary conditions in the horizontal direction and reflective boundary conditions on the vertical boundaries:

```python
sigma = 0.05/sqrt(2)
rho, pres = conditional(abs(x[1]-0.5)<0.25, 2, 1), 2.5
u = conditional(abs(x[1]-0.5)<0.25, 0.5, -0.5)
v = 0.1*sin(4*pi*x[0])*(exp(-(x[1]-0.25)**2/(2*sigma**2))
Model.U0 = Model.toCons([rho, u, v, pres])
```

Now that we have set up the model class, the code presented for the advection problem for evolving the system and adapting the grid can remain unchanged.

Results for both test cases on locally adapted grids are shown in Fig. 6 and the left column of Fig. 8. It turns out that in the Kelvin-Helmholtz case the stabilization is almost completely determined by the physicality check since the smoothness indicator shown here is based on the pressure which in this case is continuous over the discontinuity. To increase the stabilization of the method it is either possible to reduce the tolerance in the troubled cell indicator (results shown on right of Fig. 9) or to use a different smoothness indicator all together, which is discussed in Section 2.3.

### 2.2.2 Adding source terms

So far we have in fact simulated the interaction of a shock wave with a column of low density gas and not in fact a bubble. To do the later would require to either extend the problem to 3D (discussed in the next section) or to simulate the problem using cylindrical coordinates. This requires adding a geometric source term to the right hand side of the Euler equations. As pointed out in the introduction our model can take two types of source term depending upon the desired treatment in the time stepping scheme. Here we want to treat the source explicitly so need to add an $S_e$ method to the Model class:

```python
# geometric source term for cylindrical coordinates
def source(t,x,U,DU):
    v, p = Model.toPrim(U)
    return as_vector([-U[0]*v[1]/x[1],
                      -U[1]*v[1]/x[1],
                      -U[2]*v[1]/x[1],
                      -(U[3]+p)*v[1]/x[1]])
Model.S_e = source
```

### # additional source term for residual indicator
def indicatorSource(t,x,U,DU):
    a = Model.Indicator.eta(t,x,U)
    return a - U[1]/U[0]/x[1]
Model.Indicator.S = indicatorSource
2.2.3 Adding diffusion

Finally, we discuss the steps needed to add a diffusion term. This requires adding an additional method to the Model class. So to solve the compressible Navier-Stokes equations instead of the Euler equations the following method needs to be added (given a viscosity parameter $\mu$):

```python
def F_v(t, x, U, DU):
    assert dim == 2
    Pr = 0.72
    rho, rhoU, rhoE = U[0], as_vector([U[j] for j in range(1, dim+1)]), U[dim+1]
    grad_rho = DU[0, :]
    grad_rhou = as_matrix([[DU[j, :]
                            for j in range(1, dim+1)]])[:, 0]
    grad_rhoE = DU[dim+1, :]
    grad_u = as_matrix([[grad_rhou[j, :]* rho - rhou[j]* grad_rho]
                        for j in range(dim)])[:, 0]
```

Results are shown in Fig. 7.
Fig. 8: Diffusive kelvin-Helmholtz instability with \( \mu = 0 \) (Euler), \( \mu = 0.0001 \), and \( \mu = 0.001 \) (from left to right). Top row shows density and bottom row the grid levels of the adaptive simulation. The smoothing effect of the viscosity is clearly visible compared to the results shown for the Euler equations. Consequently, the small scale instabilities are completely suppressed and a coarser grid is used.

\[
\begin{align*}
\text{grad}_E &= \frac{\text{grad}_E \rho + \rho \text{grad}_E \rho - \rho \text{grad}_E \rho}{\rho^2} \\
\tau &= \mu \left( \text{grad}_u + \text{grad}_u^T - \frac{2}{3} \text{tr} \left( \text{grad}_u \right) \text{Identity}(\text{dim}) \right) \\
K\_{\text{grad}_T} &= \mu \frac{\text{Model} \cdot \gamma}{\text{Pr}} \left( \text{grad}_E - \frac{\text{dot}(\rho u, \text{grad}_u)}{\rho} \right) \\
\text{return } \text{as} \text{matrix}(\left[
\begin{array}{c}
0.0, \\
\tau[0,0], \\
\tau[1,0], \\
\text{dot}(\tau[0,:], \rho u)/\rho + K\_{\text{grad}_T}[0],
\end{array}\right])
\end{align*}
\]

As an example we repeat the simulation of the Kelvin-Helmholtz instability (see Fig. 8). Note that with the default setting for the stepper, an IMEX scheme is used where the diffusion is treated implicitly and the advection explicitly with a time step given by the CFL condition.

2.3 User defined smoothness indicator

To exchange the smoothness indicator the construction of the operator has to be slightly changed. Assuming that the indicator is defined in a source file `modalindicator.hh` which defines a C++ class `ModalIndicator` which takes the C++ type of the discrete function \( U_h \) as template argument. This class needs to be derived from a pure virtual base class and override a single method. Then the construction of the operator needs to be changes to

```
from dune.typeregistry import generateTypeName
from dune.femdg import smoothnessIndicator

# compile and load the module for the smoothness indicator - need the correct C++ type
clsName, includes = generateTypeName("ModalIndicator", U_h)
# the ModalIndicator class has a default constructor (ctor) without arguments
indicator = smoothnessIndicator(clsName, ["modalindicator.hh"]+includes, U_h, ctorArgs=[[]])
# construct the operator
operator = femDGOperator(Model, space, limiter=["minmod", indicator])
```

As an example we use here a smoothness indicator based on studying the decay properties of the modal expansion of the solution on each cell following the ideas presented in [35]. For the following implementation we assume that we are using a modal basis function...
Fig. 9: Density (top) and grid levels (bottom) for Kelvin-Helmholtz instability at time \( t = 1.5 \). In the initial setup the density in the upper layer is 2 and 1 in the lower layer. The middle figure shows the default setting for the troubled cell indicator. Since the indicator uses the pressure to determine troubled cells, very little stabilization is applied in this case. Consequently very fine structures develop with considerable under- and overshoots developing. The minimum and maximum density are around 0.6 and 2.6, respectively. To increase the sensitivity of the smoothness indicator, we can change the tolerance by passing a suitable parameter to the constructor of the operator. On the right this is shown with \( TOL = 0.2 \) (default is 1). The simulation clearly shows less structure while the under- and overshoots are reduced to 0.8 and 2.4. On the left we show a simulation using the indicator from [35] based on the modal expansion of the density. Here minimum and maximum densities are 0.95 and 2.1, respectively. As reference for the Navier-Stokes simulation shown on the right of Fig. 8 density throughout the simulation was in the range 0.97 and 2.1.

set orthonormalized over the reference element. Then the C++ code required to compute the smoothness indicator based on the modal expansion of the density is given in the next snippets.

```cpp
template <class DiscreteFunction>
struct ModalIndicator
    : public Dune::Fem::TroubledCellIndicatorBase<DiscreteFunction> {
    using LocalFunctionType = DiscreteFunction::LocalFunctionType;
    ModalIndicator() {}
    double operator()( const DiscreteFunction& U, const LocalFunctionType& uEn ) const override {
        double modalInd = smoothnessIndicator( uEn );
        return std::abs( modalInd ) > 1e-14? 1. / modalInd : 0.;
    }
};
```

The actual computation of the indicator is carried out in the method `smoothnessIndicator` but is slightly too long to include here directly but is shown in Appendix B. It is important to note that it requires very little knowledge of the DUNE programming environment or even C++ since it relies mainly on the local degrees of freedom vector provided by the argument `uEn`.

2.4 User defined time stepping schemes

The DUNE-FEM package provides a number of standard strong stability preserving Runge-Kutta (SSP RK) solvers including explicit and implicit methods and IMEX schemes method.
of degree one to four. In the literature there is a wide range of additional suitable RK methods (having low storage or better CFL constants using additional step for example). Furthermore, multistep methods can be used. Also there are a number of other packages providing implementations of timestepping methods. Since the computationally critical part of a DG method of the type described here lies in the computation of the spatial operator, the additional work needed for the timestepper can be carried out on the Python side with little impact. Furthermore, as pointed out in the introduction, it is often desirable to use Python for rapid prototyping and to then reimplement the finished algorithm in C++ after a first testing phase to avoid even the slightest impact on performance. The following code snippet shows how a multistage third order RK method taken from [33] can be easily implemented in Python and used to replace the stepper used so far:

```python
class ExplSSP3:
    def __init__(self, stages, op, cfl=0.45):
        self.op = op
        self.n = int(sqrt(stages))
        self.stages = self.n * self.n
        self.r = stages - self.n
        self.q2 = op.space.interpolate(op.space.dimRange*[0], name="q2")
        self.tmp = self.q2.copy()
        self.cfl = cfl * stages * (1 - 1 / self.n)
        self.dt = None

    def c(self, i):
        if i <= (self.n + 2) * (self.n - 1) / 2 + 1:
            return (i - 1) / (self.n * self.n - self.n)
        else:
            return (i - self.n - 1) / (self.n * self.n - self.n)

    def __call__(self, u, dt=None):
        if dt is None and self.dt is None:
            self.op.stepTime(0, 0)
            self.op(u, self.tmp)
            dt = self.op.timeStepEstimate[0] * self.cfl
        elif dt is None:
            dt = self.dt
            self.dt = 1e10
            fac = dt / self.r
            i = 1
            while i <= (self.n - 1) * (self.n - 2) / 2:
                self.op.stepTime(self.c(i), dt)
                self.op(u, self.tmp)
                self.dt = min(self.dt, self.op.timeStepEstimate[0] * self.cfl)
                u.aspy(fac, self.tmp)
                i += 1
            self.q2.assign(u)

            while i <= self.n * (self.n - 1) / 2:
                self.op.stepTime(self.c(i), dt)
                self.op(u, self.tmp)
                self.dt = min(self.dt, self.op.timeStepEstimate[0] * self.cfl)
                u.aspy(fac, self.tmp)
                i += 1
            u.as_numpy[:] *= (self.n - 1) / (2 * self.n - 1)
            u.asopy(self.n / (2 * self.n - 1), self.q2)

            while i <= self.stages:
                self.op.stepTime(self.c(i), dt)
                self.op(u, self.tmp)
                self.dt = min(self.dt, self.op.timeStepEstimate[0] * self.cfl)
                u.aspy(fac, self.tmp)
                i += 1
            self.op.applyLimiter(u)
            self.op.stepTime(0, 0)
            return dt
```

# use a four stage version of this stepper
stepper = ExplSSP3(4, operator)
Fig. 10: Adaptive 3D simulation of shock bubble interaction problem shown from two different sides. The grid levels and the density distribution at time $t = 0.5$ is shown together with the isosurface $\rho = 1.5$. The simulation was performed on a workstation using 8 processors. The corresponding globally refined grid would have contained 1.5M elements resulting in 52M degrees of freedom. The grid shown here consists of 150,000 elements with about 5.2M degrees of freedom.

Again the remaining code can remain unchanged. Results with this time stepping scheme are included in some of the comparisons shown in the next section (see Fig. 12 and Fig. 13).

2.5 Different grids and spaces

One of the strengths of the DUNE framework on which we are basing the software presented here, is that it can handle many different types of grid structures. Performing a 3D simulation can be as simple as changing the domain attribute in the Model class (Fig. 10).

It is also straightforward to perform the simulations on for example a simplicial grid instead of the cube grid used so far:

```python
from dune.alugrid import aluSimplexGrid as grid
gridView = view(grid(Model.domain))
gridView.hierarchicalGrid.globalRefine(3) # refine a possibly coarse initial grid
```

It is even possible to use a grid consisting of general polygonal elements, by simply importing the correct grid implementation:

```python
from dune.polygrid import polygonGrid as grid
```

A wide range of other grid types are available and a recent overview is given in [5] some examples are shown in Fig. 11.

As pointed out in the previous section, the choice of the basis function set used to represent the discrete solution can strongly influence the efficiency of the simulation. So far we have used an orthonormal basis function set for the polynomial space over the reference cube $[0,1]^d$. If the grid elements are all affine mapping of $[0,1]^d$ (i.e. parallelograms) then this is good choice, since it has the minimal number of degrees of freedom for a desired approximation accuracy. Also the mass matrix will be a very simple diagonal matrix on all elements in this case. These properties always hold for simplicial elements when an orthonormal polynomial basis over the reference simplex is used. As soon as the mapping between the reference element and a given element in the grid becomes non affine, both properties can be lost. In this case it might be necessary, to achieve the right approximation properties, to use a tensor product polynomial space, increasing the number of degrees of
freedom per element considerably. Also an orthonormal set of basis function over the reference cube, will still lead to a dense mass matrix. This lead to a significant reduction in the efficiency of the method, if the local mass matrix on each element can not be stored due to memory restrictions. A possible solution to this problem is to use a Lagrange type set of basis functions with interpolation points coinciding with a quadrature rule over the reference cube. The obvious choice for this quadrature is a tensor product Gauss-Legendre rule. Since this quadrature is accurate up to order $2k + 1$, it can be used to exactly compute the mass matrix and consequently, the mass matrix \[ \int \phi_i \phi_j = \sum_q \omega_q \phi_i(x_q) \phi_j(x_q) = \sum_q \omega_q \delta_{ij} = \omega_i \delta_{ij} \]
is a diagonal matrix where the diagonal element only depend on the integration element of the reference mapping at the quadrature points. In addition, if all the other element integrals needed to evaluate the spatial operator use the same quadrature, the interpolation property of the basis functions can be used to further speedup evaluation. All of this is well known and for example investigated in [39]. In the software framework presented here, it is straightforward to switch to this representation of the discrete space by replacing the construction of the space object by

```python
from dune.fem.space import dglagrange
space = dglagrange(gridView, dimRange=Model.dimRange, order=4, pointType="gauss")
```

If the operator is constructed using this space, the suitable Gaussian quadrature is chosen automatically. Note that this space is only well defined over a grid consisting of cubes.

It is also common practise to use the points of a tensor product Lobatto-Gauss-Legendre (LGL) quadrature (see e.g. [38]). In this case the evaluation of the intersection integrals in the spatial operator can also be implemented more efficiently. To still retain the simple structure of the mass matrix requires to use the same Lobatto-Gauss-Legendre rule to compute \( \int \phi_i \phi_j \). Since this rule is only accurate up to order \( 2k - 1 \) this results in an underintegration of the mass matrix similar to mass lumping; this does not seem to influence the accuracy of the scheme. It is straightforward to use this space:

```python
space = dglagrange(gridView, dimRange=Model.dimRange, order=4, pointType="lobatto")
```

and again using this space in the construction of the spatial operator will result in the correct LGL quadrature being used. We compare $L^2$ errors on a sequence of non affine cube grids

Fig. 11: Shock bubble interaction problem at $t = 0.1$ using different grid structures. A structured simplex grid (left), a non affine cube grid (right), a polygonal grid (middle) consisting of the dual of the simplex grid on the left.
using different sets of basis functions in Fig. 12, 13, and 14. We show both the error vs. the number of degrees of freedom (left) and vs. the runtime (right).

2.6 Reactive advection diffusion problem

We conclude with an example demonstrating the flexibility of the framework to combine different components of the DUNE-FEM package to construct a scheme for a more complex problem. As a simple example we use a chemical reaction type problem with linear advection and diffusion where the velocity field is given by discretizing the solution to an elliptic problem in a continuous Lagrange space. As mentioned this is still a simple problem but can be seen as a template for coupled problems, e.g., transport in porous media setting or where the flow is given by solving incompressible Navier Stokes equations.

Let us first compute the velocity given as the curl of the solution to a scalar elliptic problem:

```python
streamSpace = lagrange(gridView, order=order)
Psi = streamSpace.interpolate(0, name="streamFunction")
u,v = TrialFunction(streamSpace), TestFunction(streamSpace)
x = SpatialCoordinate(streamSpace)
form = (inner(grad(u),grad(v)) - 5* sin(x[0])* sin(x[1]) * v) * dx
streamScheme = galerkin([form == 0, DirichletBC(streamSpace,0)])
streamScheme.solve(target=Psi)
transportVelocity = as_vector([-Psi.dx(1),Psi.dx(0)])
```
Fig. 13: Sod’s Riemann problem simulated from \( t = 0 \) to 0.2 using different representation for the discrete space with polynomial order 4. The simulations were performed on a sequence of grids consisting of non affine cubes (split into 2 simplices for the simplicial simulation). Shown are the errors measured in the \( L^2 \) norm versus the number of degrees of freedom (left) and the runtime (right). The right plot also includes results from a simulation with the LGL method and the SSP3 time stepper implemented in Python as discussed previously. The results of this simulation are not included on the left since they are broadly in line with the LGL simulation using the 3 stage RK method. Due to the larger CFL constant this method is more efficient than the 3 stage method of the same order. The simulation on the simplicial grid produces a significantly better error on the same grid but requires twice as many degrees of freedom. Overall it is more efficient than the LGL or GL simulations using the same three stage RK method. Note that the runtime with the ONB basis is significantly larger due to the additional cost of computing the inverse mass matrix on each element of the grid as discussed above. On an affine cube grid the runtime is comparable to the GL scheme on the same grid but this is not shown here.

Fig. 14: Sod’s Riemann problem simulated from \( t = 0 \) to 0.2 using different troubled cell indicators. The simulations were performed on a sequence of grids consisting of non affine cubes (split into two triangles for simplicial simulations) using piecewise polynomials of order 4 with the orthonormal basis. Shown are the errors measured in the \( L^2 \) norm versus the number of degrees of freedom (left) and the runtime (right).
We use this velocity field to evolve three chemical components reacting through some non-linear reaction term and include some small linear diffusion:

```python
from ufl import *
from dune.ufl import DirichletBC
from dune.fem.space import lagrange
from dune.fem.scheme import galerkin

class Model:
    dimRange = 3

    def S_e(t,x,U,DU):
        # reaction term
        r = 10*as_vector([U[0]*U[1], U[0]*U[1], -2*U[0]*U[1]])
        # source for component one and two
        P1 = as_vector([0.2*pi, 0.2*pi])  # midpoint of first source
        P2 = as_vector([1.8*pi, 1.8*pi])  # midpoint of second source
        f1 = conditional(dot(x-P1,x-P1) < 0.2, 1, 0)
        f2 = conditional(dot(x-P2,x-P2) < 0.2, 1, 0)
        f = conditional(t<5, as_vector([f1,f2,0]), as_vector([0,0,0]))
        return f - r

    def F_v(t,x,U,DU):
        return 0.02*DU

    def F_c(t,x,U):
        return as_matrix([[*(Model.velocity(t,x,U)*u)] for u in U])

    def maxLambda(t,x,U,n):
        return abs(dot(Model.velocity(t,x,U),n))

    boundary = {range(1,5): as_vector([0,0,0])}

    U0 = [0,0,0]
    endTime = 10
```

Note that the source term includes both the chemical reaction and a source for the first two components. The third component is generated by the first two interacting.

Due to the diffusion we do not need any stabilization of the form used so far. However, in this case a reasonable assumption is that all components remain positive throughout the simulation, so the physicality check described above is still a useful feature. We can combine this with the scaling limiter already described for the advection problem. To this end we need to add bounds to the model and change to the construction call for the operator:

```python
Model.lowerBound = [0,0,0]
operator = femDGOperator(Model, space, limiter="scaling")
```

Note that by default the stepper switches to a IMEX Runge-Kutta scheme if the Model class contains both an advective and a diffusive flux. This behavior can be changed by using the rkType parameter in the constructor call for the stepper. A final remark concerning boundary conditions: here we use simple Dirichlet boundary conditions which are then used for both the diffusive and advective fluxes on the boundary as outside cell value. We saw in a previous example that we can also prescribe the advective flux on the boundary directly. In that example we used

```python
Model.boundary[3] = lambda t,x,u,n: noFlowFlux(u,n)
```

If this type of boundary conditions is used we also need to prescribe the flux for the diffusion term, so for an advection-diffusion problem we pass in a pair of fluxes at the boundary, e.g.,

```python
Model.boundary[3] = [lambda t,x,u,n: inner(Model.Fc(t,x,Ubnd), n),
                    lambda t,x,U,DU,n: inner(Model.Fv(t,x,Ubnd,DUbnd), n)]
```

Fig. 15 shows the results for the chemical reaction problem.
Fig. 15: The three components of the chemical reaction system (left to right) at \( t = 4.5 \) (top row) and \( t = 10 \) (bottom row). Velocity field included in middle figure.

Table 1: Runtime comparison of the C++ and the Python code for a simple test example solving the Euler equations in 2d with explicit time stepping and second order polynomials.

| Code | SPGrid \#el | 1024 | 4096 | 16384 |
|------|------------|------|------|-------|
| C++  | 7.19       | 57.45| 464.85|       |
| Python | 7.04    | 56.29| 457.23|       |
| ALUGrid \#el | 1024 | 4096 | 16384 |
| C++   | 12.72      | 106.08| 884.28|       |
| Python | 13.32     | 110.48| 924.81|       |

3 Efficiency of Python based auto-generated models

While Python is easy to use, its flexibility can lead to some deficiencies when it comes to performance. In DUNE-Python [22] and DUNE-FemPy [15] a just-in-time compilation concept is used to create Python modules based on the static C++ type of every object used, i.e. the models described in the previous section are translated into C++ code based on the UFL descriptions in the various model methods which is then compiled and loaded as Python modules. This way we avoid virtualization of the DUNE interfaces and consequently one would expect very little performance impact as long as calls between Python and C++ are only done for long running methods. To verify this, we compare the performance of the approach shown here with the previously hand-coded pure C++ version described in [13].

As a test example we choose a standard Riemann problem for the Euler equations solved on a series of different grid resolutions using quadratic basis functions, a minmod limiter, and explicit RK3 time stepping. We use two different grid implementation, a dedicated Cartesian grid (SPGrid) and a fully unstructured grid (ALUGrid). The results are shown in Table 1.

We observe that for the Cartesian grid (SPGrid) using the Python frontend leads consistently to a small improvement of 2%. For the unstructured grid (ALUGrid) we observe a performance decrease of about 4-5%. This can be explained with the fact that for SPGrid all code can be inlined in the just-in-time compiled Python module. For ALUGrid, where
a library exists, this is not so straight forward. In the future we will experiment with link time optimization and try to reduce implementation of small code snippets in the ALUGrid library.

4 Conclusion and outlook

In this paper we presented a comprehensive framework for the Discontinuous Galerkin (DG) Method with an easy to use Python interface for model description and solver configuration. The framework covers different variants of existing DG methods and can be easily extended to include improvements and new development of the methodology.

Although the DUNE-FEM-DG framework serves as a good starting point for high-performance DG solvers, the aim of this work is to demonstrate how the Python interface simplifies the simulation of a wide range of evolution equations as well as enabling the rapid prototyping of new methods, their testing and their comparison with other methods. For example, in almost all the tests presented here, computations were done using the same setting for the indicator and tolerance for adaptivity and the troubled cell detection, demonstrating the robustness of the default setup. Of course the default setup will only be a suitable starting point and so most of the building blocks of the discretization can be straightforwardly replaced from within the Python script, often without requiring any or not much C++ knowledge. Also we would like to point out the implemented DG solver can be directly used as higher (2nd) order Finite Volume (FV) scheme by replacing the DG space with a FV space consisting of piecewise constant polynomials.

The Python interfaces are not yet fully reflecting the possibilities on the C++ side, for example, the reconstruction in primary variables for the Euler equations is possible, but not yet available on the Python side. Another missing feature is that it is currently not possible on the Python side to easily exchange the average operators in the discretization of the diffusive terms as, for example, needed in some applications [14].

Another missing feature in Python is the assembly of Jacobian matrices during the nonlinear solve. This could be desirable for some applications or to test different existing solvers. Although the feature is available in the C++ code it needs a few code alterations in the underlying infrastructure package. This would also make it straightforward to then use other solver packages available in Python, e.g., scipy. For the compressible applications, which were the focus of this work, it is much more feasible to work with the Jacobian free nonlinear solvers and there robust and efficient preconditioning techniques are still a very active research topic. We are currently focusing on including ideas presented in [6] based on subcell finite volume multigrid preconditioning.

The extension of the DGSEM methods to simplicial grids (see [10] for an overview) could be implemented with a few minor modifications. Based on earlier work [21] the necessary basis function implementations for arbitrary quadrature points are available but need to be integrated into the DUNE-FEM and DUNE-FEM-DG framework.

While DUNE-FEM provides a number of Runge-Kutta methods, we have shown here that it is straightforward to add other time stepping algorithms on the Python side, a feature which should also allow to use other packages which provide bindings for Python, such as the Assimulo package [2]. For IMEX schemes the splitting is at the moment still a bit restricted focusing on the important case of implicitly treating the diffusion (and part of the source term) while using an explicit method for the advection. In some applications other types of splitting are of interest and will be made available in future releases.
Conflict of interest

On behalf of all authors, the corresponding author states that there is no conflict of interest.

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A Interfaces

Signature of femdgOperator:

```python
def femdgOperator(Model, space, limiter=“default”,
advectionFlux=“default”, diffusionScheme=“cdg2”, parameters=None)
```

The Model class contains the description of the mathematical model to solve and is described below. The space parameter is one of the available Discontinuous Galerkin spaces available in the DUNE-FEM framework. We have limiter equal to None, “minmod”, “scaling”, taking advectionFlux equal to “default” to use Local-Lax-Friedrichs scheme, for Euler there are some other fluxes available, this parameter can also be used to pass in a user defined flux implementation. The parameter diffusionScheme can be used to change the diffusive flux. The parameters is a dictionary where additional information can be passed to the C++ code, e.g., the tolerance for the troubled cell indicator. Static methods on Model class passed to femdgOperator:

```python
class Model:
    dimRange = r  # source term (treated explicitly)
    def S_e(t,x,U,DU):
        # return R
    # source term (treated implicitly)
    def S_i(t,x,U,DU):
        # return R'
    # diffusion term
    def F_v(t,x,U,DU):
        # return R', d
    # advection term
    def F_c(t,x,U):
        # return R'
    # max advection speed used in LLF flux
    def maxLambda(t,x,U,n):
        # return >= 0
    # boundary conditions
    boundary = dict(...)  # for physicality check in troubled cell indicator
    def physical(t,x,U):
        # return bool
    # for jump smoothness indicator
    def jump(t,x,U,V):
        # return R'
    # for scaling limiter
    Model.lowerBound = R'
    Model.upperBound = R'
```

The final ingredient is the time stepper:

```python
def femdgStepper(*, order, operator, rkType=None, cfl=0.45, parameters=None)
```

If rkType is None or "default" then the type of Runge-Kutta method used will depend on the methods defined on the Model class used to construct the operator. If no diffusive flux $F_v$ was defined an explicit RK method is used, if a diffusive flux but no advective flux $F_c$ is defined an implicit method is used, while if both fluxes are present an IMEX scheme is employed. Again the parameter dictionary can be used to set further parameters read by the underlying C++ code.

B C++ code for modal troubled cell indicator

The full class for the modal indicator. The class needs to derive from a templated pure virtual base class Dune::Fem::TroubledCellIndicatorBase<DiscreteFunction> and override a single method. The main work is done in the private method smoothnessIndicator which sets up a least square problem to compute a smoothness indicator following [35]. We assume that the degrees of freedom $u_{En[i]}$ are modal. The modal decay is computed for the first component, i.e., $u_{En[0]}, u_{En[\text{dimRange}]}, u_{En[2+\text{dimRange}]}$, ...

```cpp
template <class DiscreteFunction>
struct ModalIndicator {
    using LocalFunctionType = DiscreteFunction::LocalFunctionType;
    ModalIndicator () {}

    double operator()( const DiscreteFunction& U, const LocalFunctionType& uEn)const override {
        double modalInd = smoothnessIndicator( uEn );
        return std::abs( modalInd ) > 1e-14 ? 1. / modalInd : 0.0;
    }
```

```cpp
class ModalIndicator : public Dune::Fem::TroubledCellIndicatorBase<DiscreteFunction> {
    using LocalFunctionType = DiscreteFunction::LocalFunctionType;
    ModalIndicator () {}

    double operator()( const DiscreteFunction& U, const LocalFunctionType& uEn)const override {
        double modalInd = smoothnessIndicator( uEn );
        return std::abs( modalInd ) > 1e-14 ? 1. / modalInd : 0.0;
    }
```
private:

    template <class LF>
    static double smoothnessIndicator(const LF& uLocal) {
        using ONB = Dune::Fem::OrthonormalShapeFunctions<LF::dimDomain>;
        const std::size_t R = LF::dimRange; // will be using first component
        const std::size_t P = uLocal.entity().order();
        const double area = uLocal.entity().geometry().volume();
        const double factor = 1/std::sqrt(area); // scaling (ONB over reference element)

        // compute a 1D moments vector by averaging
        double q[P+1], b2[P+1], f = 0;
        std::size_t k = ONB::size(0); // number of moments of zero degree (is 1)
        q[0] = uLocal[0]* uLocal[0]; // constant part not used
        double l2norm2 = 0;
        for (std::size_t i=1;i<=P;++i) {
            q[i] = 0;
            b2[i] = 0;
            double l2norm2 = 0;
            double nofMoments = ONB::size(i)-k; // number of moments of ith degree
            for (k<ONB::size(i);++k) { // averaging process
                q[i] += uLocal[k*R]* uLocal[k*R] / nofMoments;
                b2[i] += pow(1/(double(i)),2*P) / nofMoments;
                f += pow(1/(double(i)),2*P) / nofMoments;
            }
            for (std::size_t i=1;i<=P;++i)
                q[i] = std::sqrt(q[i] + l2norm2*b2[i]/f) / factor;
        }
        double maxQ = std::max(q[P], q[P-1]);
        // find first 'significant' mode
        std::size_t significant = 0;
        for (std::size_t i=P;i>=1;--i) {
            maxQ = std::max(maxQ, q[i]);
            if (maxQ>1e-14) {
                significant = i;
                break;
            }
        }
        if (significant==0) return 1000; // constant, i.e., very smooth indeed
        if (significant==1) return 100; // linear, not enough info to fit
        // least squares fit to obtain 'smoothness' exponent 's'
        Dune::DynamicMatrix<double> matrix(significant,2);
        Dune::DynamicVector<double> rhs(significant);
        for (std::size_t r=significant; r>=0; --r) {
            maxQ = std::max(maxQ, q[r+1]);
            rhs[r] = std::log( maxQ );
            matrix[r][0] = 1;
            matrix[r][1] = -std::log(double(r+1));
        }
        Dune::FieldMatrix<double,2,2> A; // LS matrix
        Dune::FieldVector<double,2> b; // rhs
        for (std::size_t r=0,r2<2;++r) {
            for (std::size_t c=0,c2<2;++c) {
                A[r][c] = 0;
                for (std::size_t k=0;k<significant;++k)
                    A[r][c] += matrix[k][r]*matrix[k][c];
            }
            b[r] = 0;
            for (std::size_t k=0;k<significant;++k)
                b[r] -= matrix[k][r]*rhs[k];
        }
        Dune::FieldVector<double,2> x;
        A.solve(x,b);
        return x[1];
    }