## **DGANN** solver

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**DGANN** is a RKDG-solver written in MATLAB, which is also capable of using an artificial neural network trained to serve as a troubled-cell detector. The main source code for the RKDG solver is based on the solvers available with the following texts:

- 1. Nodal Disontinuous Galerkin methods, by Jan S. Hesthaven and Tim Warburton.
- 2. Numerical Methods for Conservation Laws: From Analysis to Algorithms, by Jan S. Hesthaven.

Details about the design of the Multilayer Perceptron (MLP) troubled-cell indicator have been published in the paper <u>"An artificial neural network as a troubled-cell indicator"</u>. Details about the 2D network are available <u>here</u>.

NOTE: If the math symbols do not display properly in README.md, have a look at README.html or README.pdf instead.

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# Running the code

After cloning the git repository, execute **mypath.m** from the parent directory in MATLAB. This will set all the neccesary paths to use the solver. The various test cases need to be run from the **Examples** directory or its sub-directories.

### Scripts for 1D problems

Currently, the 1D solver supports linear advection, Burgers' equation, the shallow water equations and the compressible Euler equations.

#### Scalar 1D

The basic structure of the example script is as follows.

```
CleanUp1D;
clc
clear all
close all
model = 'Advection';
test_name = 'Sine';
u_{IC} = 0(x) \sin(10*pi*x);
bnd_1 = 0.0;
bnd_r = 1.4;
mesh_pert = 0.0;
bc\_cond = \{'P', 0.0, 'P', 0.0\};
FinalTime = 1.4;
CFL = 0.2;
      = 100;
K
Ν
         = 4;
Indicator = 'MINMOD';
nn_model = 'MLP_v1';
Limiter = 'MINMOD';
plot_iter = 50;
save_soln = true;
save_ind = true;
save_plot = true;
ref_avail = true;
ref_fname = 'ref_soln.dat';
rk_comb = false;
var_ran = [-1.2, 1.5];
% Call code driver
ScalarDriver1D;
```

- CleanUp1D removes temporary file paths added, which is especially important if the previous run of the script terminated prematurely. This must be the first line of every script. DO NOT REMOVE IT!
- The model flag sets the type of scalar model which is being solved. The following scalar models are currently available:

```
• 'Advection': Linear advection equation with the advection speed set to 1.
```

- 'Burgers' : Burgers equation with the flux  $u^2/2$ .
- $\circ$  'BuckLev': Buckley-Leverett equation with flux contant set to 0.5.
- test\_name is used to declare the name of the test. This is used for creating various save files.
- u IC is used to set the initial condition for the problem.
- bnd\_1 and bnd\_r define the left and right domain boundaries, which is descritized using Nelem number of elements. The degree of the polynomial in each element/cell is set using N.
- mesh\_pert is used to randomly perturb the interior cell-interfaces using the following algorithm

```
x_{i+\frac{1}{2}} \to x_{i+\frac{1}{2}} + \text{mesh\_pert } h \omega_{i+\frac{1}{2}}, \qquad \omega_{i+\frac{1}{2}} \in \mathcal{U}[-0.5, 0.5], \qquad i = 1, \dots, K-1
```

where h is the mesh size of the initial uniform grid.

- bc\_cond is used to set the left and right boundary conditions. It is a cell of the form

  {LEFT BC TYPE, LEFT BC VAL, RIGHT BC TYPE, RIGHT BC VAL} . The BC TYPES can be set as:
  - 'P': Periodic boundary conditions. In this case, both boundary type must be set to 'P'.
  - 'N': Neumann boundary condition.

- 'D': Dirichlet boundary condition, with the imposed Dirichlet value given by BC\_VAL. Note that BC\_VAL is ignored if BC\_TYPE is not set to 'D'.
- The final simulation time is set using FinalTime, while the time step is chosen using CFL.
- K is the number of elements/cells in the mesh.
- N sets the order of the basis.
- The troubled-cell indicator is set using Indicator. The following options are currently available:
  - 'NONE': No cells are flagged.
  - 'ALL': All cells are flagged.
  - 'MINMOD': Uses the basic minmod limiter.
  - LIVE : Uses the modified minmod-type TVB limiter. If this is chosen, then one also needs to set the variable TVBM to a positive number. Note that if this constant is set to 0, then the limiter reduces to the usual minmod limiter.
  - 'NN': Uses the trained neural network. The network name is set using the variable nn\_model. The available networks are described below in the section **Using the MLP indicator**.
- The limiter usd to reconstruct the solution in each troubled-cell, is set using Limiter. The following options are currently available:
  - 'NONE' : No limiting is applied.
  - 'MINMOD': MUSCL reconstruction using minmod limiter.
- The flag plot\_iter is used for visualization purposes. The solution plots are shown after every plot\_iter number of iterations during the simulation. This number also controls the frequency with which the time-history of the flagged troubled-cells is saved to file (see save ind).
- - Column 1: x-coordinates
  - Column 2: solution value at corresponding x-coordinate
- If the time-history of the troubled-cells needs to be viewe/saved, the flag save\_ind must be set to true. If this is not the case, set this flag to false. The time-history files are also save in the directory OUTPUT. The filename has the format:

  <model>1D\_<test\_name>\_P<N>\_N<K>\_IND\_<Indicator>\_LIM\_<Limiter>\_tcells.dat. If mesh perturbation is used, then the filename will end with the tag pert. The data in the file has the following format:
  - Each row of the file contains the time, followed by the mid-points of the cells flagged as troubled-cells at that time.
  - The first row corresponds to the cell flagged at time t = 0. This is essentially done for the initial condition.
  - Following the first row, the rows can be grouped in sets of size r, to list the cells flagged after each sub-stage of the r-stage time-integration scheme. Since the current implementation of the code uses SSP-RK3, the rows will be grouped as triplets.
- If save\_plot is set to true, then the solution plots are generated and saved in **OUTPUT**. NOTE: This needs save\_soln and save\_ind to be set as true.
- If a reference/exact solution data file is available, then set ref\_avail to true and the (relative) file name of the referene solution in ref fname.
- If the troubled-cell plots should be saved for each individual RK-stage, then set <code>rk\_comb</code> to <code>false</code>. To save a unified plot, set this to <code>true</code>.
- var\_ran is used to set the ylim for the solution plot. This should be a array of size (1,2).
- The main driver script ScalarDriver1D is called once all the flags have been set.

#### **Shallow Water 1D**

The basic structure of the example script is as follows.

```
CleanUp1D;
clc
clear all
close all
Globals1D DG:
Globals1D_MLP;
model = 'SWE';
gravity = 1.0;
test_name = 'Dambreak';
depth_IC =@(x) 3*(x<0.0) + 1*(x>=0.0);
velocity_IC =@(x) @*x;
bnd_l
        = -3.0;
bnd_r = 3.0;
mesh_pert = 0.0;
bc_cond = \{'N', 0.0, 'N', 0.0;
           'N',0.0,'N',0.0}; % For conserved variables
FinalTime = 1;
CFL = 0.4:
         = 100;
Κ
         = 4;
Ν
Indicator = 'TVB'; TVBM = 100;
ind_var
           = 'depth';
           = 'MLP_v1';
nn_model
Limiter
            = 'MINMOD';
lim_var
            = 'con';
plot_iter = 10;
save_soln = true;
save_ind = true;
save_plot = true;
ref_avail = true;
ref_fname = 'ref_soln.dat';
rk_comb = true;
var_ran = [1,3.2; 0,1];
% Call code driver
SWEDriver1D;
```

Most of the structure is similar to the Scalar 1D script. The differences are described below.

- The model needs to be set as 'SWE'. This should not be changed.
- The acceleration due to gravity is set using gravity.
- The initial depth and velocity are set using depth\_IC and velocity\_IC.
- The bc\_cond has the same format as earlier, although now it has two rows of parameters. The first row corresponds to depth, while the second corresponds to the discharge (depth\*velocity). Note that the boundary condition are set for the conserved variables.
- For systems of conservation laws, there are various choices for the variables to be used for troubled-cell detection. For the shallow water equations, this choice is made via the flag <u>ind\_var</u>, with the following options (the troubled-cells flagged for each variable is pooled together):

- 'prim': The primitive variables i.e., depth and velocity, are used.
- 'con': The conserved variables i.e., depth and discharge, are used.
- As was the case with detection, there are several options for the variables which can be reconstructed. This is set using the flag lim var, with the following options:
  - 'prim': The primitive variables i.e., depth and velocity, are recontructed.
  - 'con': The conserved variables i.e., depth and discharge, are recontructed.
  - 'char\_cell': The local characterictic variables are reconstructed. These are obtained cell-by-cell using the linearized transformation operators. More precisely, the transformation matrix in each cell is evaluated using the cell-average value, following which the conserved variables are transformed to the characteristic variables in that cell. The same transformation is used to retrieve the conserved variables after limiting the characteristic variables.
  - 'char\_stencil': The local characterictic variables obtained cell-by-cell can introduce spurious oscillations in the solution.
    One can also obtain the local characteristic variables, stencil-by stencil. More precisely, for a given reconstruction stencil of 3-cells, the transformation matrix is evaluated using the cell-average value of the central cell, following which the conserved variables are transformed to the characteristic variables in every cell of that stencil. The transformed variables are used to obtain the reconstructed characteristic variables in the central cell. Note that this approach can be 3 times more expensive than the 'char cell' approach.
- The solution filename has the format:

- Column 1: x-coordinates
- Column 2: the value of depth and velocity (in that order) at corresponding x-coordinate.
- The troubled-cell time-history filename has the format:

- var\_ran is used to set the ylim for the solution plots, with the format [depth\_min,depth\_max; velocity\_min, velocity\_max].
- The main driver script SWEDriver1D is called once all the flags have been set.

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#### **Euler 1D**

The basic structure of the example script is as follows.

```
CleanUp1D;
clc
clear all
close all
Globals1D_DG;
Globals1D_MLP;
model = 'Euler';
gas\_const = 1.0;
gas\_gamma = 1.4;
test_name = 'ShockEntropy';
rho_IC =0(x) (x<-4)*3.857143 + (x>=-4).*(1 + 0.2*sin(5*x));
vel_IC = @(x) (x<-4)*2.629369;
pre_IC =@(x) (x<-4)*10.33333 + (x>=-4)*1.0;
        = -5.0;
bnd_l
      = 5.0;
bnd_r
mesh_pert = 0.0;
bc_cond = \{'D', 3.857143, 'N', 0.0;
            'D', 10.141852, 'D', 0.0;
           'D',39.166661,'N',0.0}; % For conserved variables
FinalTime = 1.8;
CFL = 0.1;
K
         = 200;
        = 4;
Indicator = 'NN';
ind_var = 'prim';
nn_model = 'MLP_v1';
Limiter = 'MINMOD';
lim_var = 'con';
plot_iter = 100;
save_soln = true;
save_ind = true;
save_plot = true;
ref_avail = true;
ref_fname = 'ref_soln.dat';
rk_comb = true;
var_ran = [0,1.2; -0.2,1.5; 0,1.2];
% Call code driver
EulerDriver1D;
```

Most of the structure is similar to the shallow water 1D script. The differences are described below.

- The model needs to be set as 'SWE'. This should not be changed.
- The gas constant and ratio of specific heats is set using <code>gas\_const</code> and <code>gas\_gamma</code>.
- The initial density, velocity and pressure are set using <code>rho\_IC</code>, <code>vel\_IC</code> and <code>pre\_IC</code>.
- The bc\_cond has the same format as earlier, although now it has three rows of parameters. The first row corresponds to density, the second corresponds to the momentum, and the third to energy. Note that the boundary condition are set for the conserved variables.
- For the Euler equations, ind\_var can be set as (the troubled-cells flagged for each variable is pooled together):
  - $\circ$   $\ \ \ ' \ \ \ \ ' \ \ \ \ \ \ '$  : Only the density is used
  - 'velocity': Only the velocity is used
  - 'pressure' : Only the pressure is used

- 'prim': The primitive variables i.e., density, velocity and pressure, are used.
- 'con': The conserved variables i.e., density, momentum and energy, are used.
- var\_ran is used to set the ylim for the solution plots, with the format [rho\_min,rho\_max; velocity\_min, velocity\_max; pressure\_min, pressure\_max].
- The main driver script EulerDriver1D is called once all the flags have been set. The troubled-cells flagged for each variable is pooled together.

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#### Scripts for 2D problems

Currently, the 1D solver supports linear advection, Burgers' equation and the compressible Euler equations. For each problem, we need the following files:

- A main script (with the default name Main.m ) to set the various problem parameters.
- The initial condition is defined using the script/function (with the default name IC.m).
- When physical boundary conditions are used, an additional script/function (must be named BC.m) needs to be created. This file is not needed when all boundary conditions are periodic.
- Finally, a mesh file is needed, which is currently generated using <u>Gmsh</u>. Each example is already provided with the Gmsh geometry file (with the extension <u>.msh</u>), you could either use the Gmsh GUI or create the file non-interactively from the terminal as \$ gmsh -2 mymeshfile.geo

#### Scalar 2D

The basic structure of the various scripts are as follows:

#### Main.m

```
CleanUp2D;
close all
clear all
clc
              = 'Advection';
model
AdvectionVelocity = [1,1]; % Used for linear advection only
test_name = 'Shapes';
FinalTime = 1;
CFL
             = 0.6;
            = 1.0e-3;
% fixed_dt
             = 2;
tstamps
              = 1;
% Set type of indicator
Indicator = 'TVB'; TVBM = 10; TVBnu = 1.5;
Filter_const = true;
nn_model = 'MLP_v1';
Limiter = 'BJES';
% Mesh file
msh_file = 'square_trans.msh';
% Output flags
plot_iter = 50;
show_plot = true;
xran = [-1,1];

yran = [-1,1];
yran = [-1,1];
clines = linspace(-0.98,0.98,30);
save_soln = true;
% Call main driver
ScalarDriver2D;
```

- CleanUp2D removes temporary file paths added, which is especially important if the previous run of the script terminated prematurely. This must be the first line of every script. DO NOT REMOVE IT!
- The model flag sets the type of scalar model which is being solved. The following scalar models are currently available:
  - 'Advection' : Linear advection equation with the advection speed set set using AdvectionVelocity .
  - "Burgers": Simple extension of the Burgers' equation to 2D, with the flux  $u^2/2$  in each coordinate direction. NOTE: AdvectionVelocity not required for this model.
- test name is used to declare the name of the test. This is used for creating various save files.
- InitialCond is used to set the initial condition for the problem (see the description of IC.m below).
- BC\_cond is used to specify the boundary conditions for each face of the domain. This must be a MATLAB cell array of size  $NBfaces \times 2$ , where NBfaces is the number of boundary curves specified in the mesh geometry file (with file-extension \_\_.geo\_). The first element of each row must be the physical face tag of a boundary curve in the geometry file, while the second element must be one of the following flags
  - 'P': Periodic boundary conditions. Note there must be an even number of periodic boundary faces.
  - 'D': Dirichlet boundary condition. The actual Dirichlet conditions are specified in BC.m
  - 'Sym': Symmteric boundary conditions.

Boundary conditions require the creation of ghost cells, with the value of the solution in these cells depending on the type of boundary condition.

- The final simulation time is set using FinalTime, while the time step is chosen using a constant CFL or a setting a step fixed dt. NOTE: Either CFL can be mentioned or the fixed time-step fixed dt, but not both at the same time.
- tstamps is used to specify the number of uniform time-instances (exluding the initial time) at which the solution files are saved. This must be a positive integer. For instance if tstamps=2 and FinalTime = 2, then the solution evaluated at times closest (less than half the local time-step) to t = 0, 1, 2 are saved. NOTE: It might happen that pen-ultimate time instance in the simulation is very close to FinalTime, with the difference being much smaller than the pen-ultimate time-step. In this case, the solver saves the solution at this pen-ultimate as the solution representing the final solution. Thus, we also save the solution at the actual FinalTime, leading to a total of tstamps +2 save points in time. This becomes crucial when comparing the troubled-cells flagged at the final time-step, since the number of cell flagged is sensitive to the size of the time-step taken.
- N sets the order of the basis.
- The troubled-cell indicator is set using Indicator. The following options are currently available:
  - 'NONE': No cells are flagged.
  - 'ALL' : All cells are flagged.
  - 'TVB': Uses the 2D minmod-type TVB limiter. If this is chosen, then one also needs to set the variables TVBM and TVBnu as positive numbers. See the paper for details.
  - 'NN': Uses the trained neural network. The network name is set using the variable nn\_model. The available networks are described below in the section **Using the MLP indicator**.
- Set Filter\_const to true, if you want to avoid flagging almost constant cells. This is especially important for speeding up the performance with the network. See the paper for details.
- The limiter usd to reconstruct the solution in each troubled-cell, is set using Limiter. The following options are currently available:
  - 'NONE': No limiting is applied.
  - 'BJES': Barth-Jespersen limiter.
- The flag plot\_iter is used for visualization/verbose purposes. The solution plots (and simulation time etc) are shown after every plot iter number of iterations during the simulation.
- Set show\_plot to true if you want solution plots to be generated during the simulation.
- xran and yran are used to crop the x and y axes of the solution plots (if show\_plot is set to true). This is particularly helpful when an extended domain is used to impose artificial boundary conditions, for instance when solving 2D Riemann problem.
- clines is used to set the contour lines for the solution contour plots (if show plot is set to true).
- The solution variables are saved in the **OUTPUT** directory if save\_soln is set to true. The filename has the format:
  <model>2D\_<test\_name>\_P<N>\_IND\_<Indicator>\_LIM\_<Limiter>\_DATA.mat. If Filter\_const is set to true,
  then ConstFilt also appears in the filename. The following MATLAB variables are saved:
  - The x and y coordinates of all degrees of freedom in the mesh.
  - Inverse of the Vandermonde matrix invV
  - Save times is the array of exact time-instances at which the solution is saved. This is of length tstamps +2.
  - The solutions evaluated at the time-instances listed in Save\_times is stored in the MATLAB-cell Q\_save .
  - The troubled-cells flagged in the mesh at each time-instance listed in Save times is stored in the MATLAB-cell ind save .
  - ptc\_cell stores the time-history of the percentage of the total number mesh cells flaged as troubled-cells. t\_hist stores all the time-instances attained during the simulation. Both these arrays have the same size.
  - sim time stores the full simulation time (excludes time taken to generate mesh data structures).
- The main driver script | ScalarDriver2D | is called once all the flags have been set.

#### IC.m

The initial condition function must take as input the arrays x and y, where each of these is of the shape  $m \times n$ . The output should be a array of dimension  $m \times n \times 1$ . Almost always, m will denote the number of DOFs per cell in the mesh, while n will be the number of cells. We give an example of this function below:

```
function Q = IC(x, y)
Q(:,:,1) = sin(4*pi*x).*cos(4*pi*y);
return;
```

#### BC.m

When using non-periodic boundary conditions, a function script called BC.m is also needed. An example function is as follows:

where the function takes in as input a boundary physical tag ckey and the curent simulation-time. The output is a function of x and y (and perhaps implicitly of time).

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#### **Euler 2D**

The basic structure of the various scripts are as follows:

#### Main.m

```
% Remove NN diretory paths if they still exist
CleanUp2D;
close all
clear all
clc
%Model
model = 'Euler';
gas_const = 1.0;
gas_gamma = 1.4;
test_name = 'DoubleMach';
InitialCond = @IC;
            = {101, 'I'; 102, '0'; 103, '0'; 104, 'S'; 105, 'D'};
BC_cond
FinalTime = 0.2;
fixed_dt
              = 2e-5;
tstamps
               = 1;
               = 1;
N
% Set type of indicator
Indicator = 'NN';
ind_var = 'con';
nn_model = 'MLP_v1';
Limiter = 'BJES'
lim_var = 'con';
             = 'BJES';
Filter_const = true;
% Mesh file
msh_file = 'domain.msh';
plot_iter = 50;
show_plot = true;
xran = [0,4];

yran = [0,1];
plot_var = {'density','pressure'};
clines = \{linspace(0.255,1.9,30), linspace(0.36,2.15,30)\};
save_soln = true;
% Call main driver
EulerDriver2D;
```

Most of the structure is similar to the scalar 2D script. The differences are described below.

```
• The model flag must be set to Euler .
```

- The gas constant and ratio of specific heats is set using gas const and gas gamma.
- The following flags

```
• 'P': Periodic boundary conditions.
```

- 'D': Dirichlet boundary condition.
- 's': Slip boundary conditions.
- 'I': Inflow boundary.
- o 'o': Outflow boundary
- The troubled-cells are detected using the variables mentioned in ind var. The following options are available :

```
• 'density'
```

o 'pressure'

- 'velocity': Both components of velocity are use.
- 'prim': The primitive variables are used.
- 'con': The conserved variables are used.
- The limiting variables are set using the flag lim\_var, with the following options being available:

```
o 'prim'
o 'con'
```

• plot var is a MATLAB-cell which listing the variables that should be plotted. The following options are available:

```
'density''pressure'velx': x-velocity component.'vely': y-velocity component.'energy'
```

- clines is also a MATLAB-cell array, with each element setting the contour lines for each variable mentioned in plot var.
- The solution variables are saved in the **OUTPUT** directory, with the filename now also mentioning the lim\_var and ind\_var used. The gas\_const and gas\_gamma variables are also saved in the file. Furthermore, the variable Q\_save saves all the conserved variables.
- The main driver script EulerDriver2D is called once all the flags have been set.

#### IC.m

The initial condition function must take as input the arrays [x], [y],  $[gas\_gamma]$ , and  $[gas\_const]$ , where [x] and [y] are of the shape  $m \times n$ . The output should be the conserved variables array [g] of dimension  $m \times n \times 4$ . Once again, m generally denotes the number of DOFs per cell in the mesh, while n will be the number of cells. We give an example of this function below:

```
function Q = IC(x, y, gas_gamma, gas_const)

xc = 1/6; yc = 0; aos = pi/3;

p1 = 116.5; rho1 = 8.0; u1 = 7.14471; v1 = -4.125;
p2 = 1.0; rho2 = 1.4; u2 = 0.0; v2 = 0;

% Initial profile
    pre = p1*(y>(x-xc)*tan(aos) + yc) + p2*(y<=(x-xc)*tan(aos) + yc);
        u = u1*(y>(x-xc)*tan(aos) + yc) + u2*(y<=(x-xc)*tan(aos) + yc);
        v = v1*(y>(x-xc)*tan(aos) + yc) + v2*(y<=(x-xc)*tan(aos) + yc);
        rho = rho1*(y>(x-xc)*tan(aos) + yc) + rho2*(y<=(x-xc)*tan(aos) + yc);

Q(:,:,1) = rho;
Q(:,:,2) = rho.*u;
Q(:,:,3) = rho.*v;
Q(:,:,4) = Euler_Energy2D(rho,u,v,pre,gas_gamma);

return;</pre>
```

where the function <code>Euler\_Energy2D</code> is an available custom function used to determine the total energy from the primitive variables. Similarly, the function <code>Euler\_Pressure2D(Q,gas\_gamma)</code> is available to evaluate the pressure from the conserved variables.

#### BC.m

When using non-periodic boundary conditions, a function script called BC.m is also needed. An example function is as follows:

```
function [rhoG,uG,vG,preG] = BC(ckey,time,gas_gamma,gas_const)
    xc = 1/6; yc = 0; aos = pi/3; M = 10;
    pre1 = 116.5; rho1 = 8.0; u1 = 7.14471; v1 = -4.125;
                  rho2 = 1.4;
                               u2 = 0.0;
    pre2 = 1.0;
                                               v2 = 0;
    xs = time*M/sin(aos);
    if(ckey == 101 || ckey == 103)
        rhoG =@(x,y) rho1*ones(size(x));
        uG = 0(x,y) u1*ones(size(x));
           = @(x,y) v1*ones(size(x));
       preG =@(x,y) pre1*ones(size(x));
    elseif(ckey == 102)
        rhoG =@(x,y) rho2*ones(size(x));
        uG =@(x,y) u2*ones(size(x));
        vG = @(x,y) v2*ones(size(x));
        preG =@(x,y) pre2*ones(size(x));
    elseif(ckey == 104)
        rhoG = @(x,y) @*x;
        uG =@(x,y) @*x;
        vG = @(x,y) 0*x;
        preG = @(x,y) 0*x;
    elseif(ckey == 105)
         \text{rhoG = @(x,y) rho1*(y>(x-xc-xs)*tan(aos) + yc) + rho2*(y<=(x-xc-xs)*tan(aos) + yc); } \\
        uG = (x,y) u1*(y>(x-xc-xs)*tan(aos) + yc) + u2*(y<=(x-xc-xs)*tan(aos) + yc);
           = (x,y) v1*(y>(x-xc-xs)*tan(aos) + yc) + v2*(y<=(x-xc-xs)*tan(aos) + yc);
        preG = (x,y) pre1*(y < (x-xc-xs)*tan(aos) + yc) + pre2*(y < (x-xc-xs)*tan(aos) + yc);
    end
return;
```

where the function takes in as input a boundary physical tag ckey, the curent simulation-time, <code>gas\_gamma</code> and <code>gas\_const</code>. The output is four functions to determine each of the primitive variables on the boundary.

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# **Using the MLP indicator**

For those interested in using the trained indicator in their own solvers, we explain the various components of the network. The descriptor files for the various trained networks are available under the folder **Trained\_networks**. For each network, the following files exist:

- *model\_parameters.dat*: Lists the dimensions of the input (IDIM) and output (ODIM) layers for the network, the number of hidden layers (NHL), and network hyperparameters (the Leaky ReLU factor, etc.).
- w\_h{i}.dat: Weights for the i'th hidden layer.
- w\_out.dat: Weights for the output layer
- b\_h{i}.dat: Biases for the i'th hidden layer
- b\_out.dat. Biases for the output layer.
- Scaling.m: Script for scaling the input data before passing it through the network.

Consider the input X to be an array of size  $m \times n$ , where m is the dimension of each data sample, while n is the number of samples. The algorithm for the network with L hidden-layers having  $N_l$  neurons in the l-th hidden layer, is as follows:

```
X_{0} = \operatorname{Scaling}(X), \quad N_{0} = m
X_{l} = f_{\operatorname{activation}}(W_{l}X_{l-1} + b_{l}\mathbb{1}(n)), \quad W_{l} \in \mathbb{R}^{N_{l} \times N_{l-1}}, \quad b_{l} \in \mathbb{R}^{N_{l}}, \quad l = 1, \dots, L
Y = \operatorname{Softmax}(W_{out}X_{L} + b_{out}\mathbb{1}(n)), \quad W_{out} \in \mathbb{R}^{2 \times N_{L}}, \quad b_{out} \in \mathbb{R}^{2}
```

where the final output Y is of dimension  $2 \times n$ , and  $\mathbb{1}(n)$  is a row vector of n ones. The activation function is chosen as a suitable non-

linear function, such as the Logistic function, hyperbolic tangent, ReLU, etc. The Softmax function is used a the output function. The first row of Y gives the probability of the sample corresponding to a troubled-cell, while the second row gives the probability of the sample corresponding to a good-cell. Note that the sum along each column of Y equals 1. The indices of the troubled-cells can be obtained as

$$ind = find(Y(1, :) > 0.5).$$

The MATLAB scripts to read and run the networks can be found under the folder **MLP\_scripts**. For instance, the following scripts for 1D problems are available in the **1D** sub-directory:

- read\_mlp\_param1D.m reads the various weights and biases for a given network.
- ind MLP1D.m runs the network for a given input x.

The **Common** sub-directory contains additional scripts needed to run the networks, such as the implementation of the leaky ReLU activation function.

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## **Network for 1D problems**

The following is a list of the available networks for 1D problems. The latest recommended network is MLP\_v1.

• MLP\_v1: This network has an input layer of size 5. In particular, the input for the classification of cell i is  $X = [\overline{u}_{i-1}, \overline{u}_i, \overline{u}_{i+1}, u^+_{i-\frac{1}{2}}, u^-_{i+\frac{1}{2}}]$ , where the first 3 quantities are the cell averages of the solution in the cells i-1, i, i+1 and the last two entries are the left and right cell interface values of the approximating polynomial in cell i. There are 5 hidden layers, whose widths are 256, 128, 64, 32 and 16, going from the input to the output layer. The activation function is taken to be the Leaky ReLU activation function

$$f_{\text{activation}}(U) = max(0, U) - \nu(0, -U),$$

with the parameter  $\nu$ . The details and results with this indicator are published <u>here</u>.

#### **Network for 2D problems**

The following is a list of the available networks for 2D problems. The latest recommended network is MLP\_v1.

• MLP\_v1: This network has an input layer of size 12. In particular, the input is the linear modal coefficient of each triangle in a 4-cell patch. There are 5 hidden layers of width 20 each. The activation function is taken to be the Leaky ReLU activation function. The details and results with this indicator are available here.

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