

A Central Rankine Hugoniot Solver for Hyperbolic Conservation Laws

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Abstract

A numerical method in which the Rankine-Hugoniot condition is enforced at the discrete level is developed. The simple format of central discretization in a finite volume method is used together with the jump condition to develop a simple and yet accurate numerical method free of Riemann solvers and complicated flux splittings. The steady discontinuities are captured accurately by this numerical method. The basic idea is to fix the coefficient of numerical dissipation based on the Rankine-Hugoniot (jump) condition. Several numerical examples for scalar and vector hyperbolic conservation laws representing the inviscid Burgers equation, the Euler equations of gas dynamics, shallow water equations and ideal MHD equations in one and two dimensions are presented which demonstrate the efficiency and accuracy of this numerical method in capturing the flow features.

Key words: Rankine-Hugoniot jump condition, central scheme, low numerical diffusion, exact capturing of shocks and contact surfaces

PACS:

1 Introduction

Numerical simulation of hyperbolic conservation equations, which model the physics of gas dynamics, magnetohydrodynamics (MHD) or shallow water flows, is non-trivial and is still a topic of intense research, despite being attempted for the last five decades, especially by the researchers in Computational Fluid Dynamics (CFD). The challenge is to devise a numerical method

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that is stable, conservative, respects the hyperbolicity, satisfies the entropy condition, is robust and yet accurate. Since the non-linear hyperbolic partial differential equations give rise to discontinuities like shock waves, contact layers, slip surfaces and also develop expansion waves with possible sonic points, any numerical method developed for such systems must be able to capture these features with sufficient accuracy without losing robustness.

The numerical methods developed for solving hyperbolic conservation laws can be broadly classified into *central discretization methods* and *upwind methods*. While the central discretization methods are simpler than the upwind methods, they often contain *tuning parameters* which are problem dependent. Because of this drawback, upwind methods became more popular from 1970's. Reviews of these methods are available in several books, for example, by Hirsch [1,2], Toro [3], Laney [4] and Leveque [5,6]. Of all the upwind methods, the approximate Riemann solver of Roe [7] is the most popular, as it can capture steady discontinuities exactly, without any numerical dissipation. However, this low dissipation comes at the cost of the loss of robustness, leading to a list of problems like carbuncle shocks, kinked Mach stems and odd-even decoupling [8], apart from the violation of the entropy condition. Many of these failures seem to be common to most of the Riemann solvers and, therefore, some of the researchers started looking for numerical methods that are free of Riemann solvers. The reader is referred to the following literature for some of the numerical methods developed as interesting alternatives : Degond *et al.* [12], Nessyahu and Tadmor [9], Kurganov and Tadmor [13], Swanson and Turkel [10], AUSM family of schemes of Meng-Sing Liou [21,22] and the references therein etc.,.

Of all the central discretization methods, Lax-Friedrichs method is the simplest (it is also one of the first successful numerical methods for hyperbolic conservation laws) [15,14]. This simple central discretization method has been the building block for several improved algorithms developed later. In a finite volume method, with a staggered grid interpretation of the Lax-Friedrichs method, the cell-interface flux consists of an average of the fluxes on the left and right states, augmented by a diffusive flux which represents the numerical dissipation. The coefficient of numerical diffusion in the Lax-Friedrichs method has a fixed magnitude. In Rusanov's method [16,6], which is also called as Local Lax-Friedrichs method (LLF), the dissipation coefficient in the Lax-Friedrichs method is modified according to the local information (maximum of the left and right states of a cell-interface) and consists of the spectral radius of the flux Jacobian matrix. Since the numerical dissipation in either the basic Lax-Friedrichs method or the LLF is large and unacceptable for practical problems, higher order accurate formulations in this format were introduced by Nessyahu and Tadmor [9] which are free of Riemann solvers and complicated flux splittings yet simple and accurate. For further developments of this method, the reader is referred to the work of Kurganov and Tadmor

[13] and to the references therein.

For the history of the other classical alternative, the Lax-Wendroff method [19], and various related second order central schemes, the reader is refer to the book of Hirsch [2]. One popular central scheme of interest is due to Jameson, Schmidt and Turkel(JST) [20] in which spatial discretization is a blend of second and fourth central differences. A shock sensor is used to switch over from one type of difference to another. In smooth flows, a linear fourth difference will operate, damping the high frequencies, while near the large gradients, the second difference is switched on. Another popular scheme is of Swanson and Turkel [10] (see also Swanson, Radespiel and Turkel [11]), the scalar coefficient of numerical dissipation is replaced by a matrix, mimicking the upwind schemes, still retaining the simpler format of central discretization. The method presented in this work is closer in spirit to the first group of central schemes mentioned above, as the basic building block is the Local Lax-Friedrichs method and the higher order accuracy is obtained by reconstruction at the cell-interfaces, as in the Kurganov and Tadmor [13] scheme. *The motive for the present chapter is to develop a simple central discretization procedure based on a direct enforcement of Rankine-Hugoniot conditions on the LLF method and hence to enhance the resolution of discontinuities.*

The exact shock capturing feature of Roe scheme [7] (in steady state) is due to the enforcement of the conservation condition, which is related to the Rankine-Hugoniot condition. Another scheme which can capture the steady shock waves with a single interior point is due to Jameson [24] (a sequel to [23]). In this method too, the Rankine-Hugoniot condition is enforced in the discretization process, due to the special discrete shock structure being forced in such a way that entire shock is captured in one cell. The low dissipative nature of these two methods is owing to the Rankine-Hugoniot (jump) condition being used in the discretization process, though not enforced directly. Motivated by the idea of using the Rankine-Hugoniot condition directly in the discretization process, Raghurama Rao and Balakrishna [25] developed an accurate Relaxation Scheme which can capture steady shocks exactly. In this work, the jump condition is enforced directly in the discretization of the hyperbolic conservation laws.

In any conservative numerical method (usually a finite volume method), the interface flux can be written as a sum of an average of the fluxes on the left and right sides of the interface and a diffusive flux. The average flux, if present alone, leads to a central discretization. The diffusive flux represents the numerical diffusion or the artificial viscosity. In the central discretization methods developed earlier, the diffusive fluxes were added explicitly and since the amount of artificial viscosity to be added was not known *a priori*, tuning parameters became necessary. It was in this background that upwind methods became popular, as they had inherent artificial viscosity and thus did not require tun-

ing parameters. However, the right amount of artificial viscosity, required to capture discontinuities accurately and yet make the numerical method satisfy the entropy condition for avoiding unphysical expansion shocks, is complex to obtain. These two features actually demand opposite requirements on the artificial viscosity : the accurate capturing of discontinuities requires as less artificial viscosity as possible whereas capturing expansion waves with sonic points without violating the entropy condition requires a finite amount of artificial viscosity. Many of the upwind methods developed for vector conservation laws have either too much of artificial viscosity (which makes them robust but inaccurate) or too little artificial viscosity (which makes them accurate but less robust). Most of the upwind methods developed so far are either robust or accurate and it seems difficult to obtain both these features simultaneously. Another feature of most of the upwind methods is rather sophisticated procedures of separating the information coming from different sides, especially for vector conservation laws, leading to quite complicated flux splittings. In the method proposed here, the simplicity of the central discretization is retained, thereby avoiding the framework of Riemann solvers and complicated flux splittings, and yet the resolution of the shocks and contact discontinuities is enhanced, by using the Rankine-Hugoniot (jump) condition as a basis for deriving the diffusive flux. Thus, the numerical method presented in this work can be considered as a natural extension of both the central and upwind discretization methods, retaining the advantages of both. Since the Rankine-Hugoniot condition is used in discretization, and thus the optimum value of artificial viscosity is enforced, we term this method as the *Method of Optimal Viscosity for Enhanced Resolution of Shocks (MOVERS)*. The basic idea of the new method for the scalar Burgers equation is demonstrated in the next section. In the following sections, this procedure is extended for the vector conservation laws (Euler equations of Gas Dynamics, shallow water equations and ideal MHD equations). The idea of enforcing the jump condition at the discrete level to the vector case is presented with three different variations. Two of these variants of *MOVERS* require an entropy fix while the third variant does not need it as it is based on a novel use of a limiter in switching over from *MOVERS* to *LLF* method. The results for several bench-mark test problems for scalar and vector conservation laws are presented in the later sections, followed by some remarks about the *MOVERS* and its relation to upwinding strategy, with the conclusions presented in the last section.

2 Method of Optimal Viscosity for Enhanced Resolution of Shocks (MOVERS) for scalar hyperbolic conservation law

Consider the one dimensional scalar hyperbolic conservation law

$$\frac{\partial u}{\partial t} + \frac{\partial f(u)}{\partial x} = 0 \quad (1)$$

An explicit scheme in conservation form on a three point stencil and with a piecewise polynomial approximation for the conserved variable is given by

$$u_j^{n+1} = u_j^n - \frac{\Delta t}{\Delta x} (f_{j+\frac{1}{2}}^n - f_{j-\frac{1}{2}}^n) \quad (2)$$

where Δt and Δx are respectively time and space steps, j represents the cell centroid and $j \pm \frac{1}{2}$ represent the cell interfaces. In the finite-volume formulation (2), the differences among all the numerical schemes lie essentially in the definition of numerical flux $f_I = f_{j+\frac{1}{2}}$ evaluated at the cell interface. The upwind flux splitting schemes define the interface flux in terms of split fluxes of left and right states as

$$f_I = f_L^+ + f_R^- \quad (3)$$

The interface flux in a central scheme is written in the form

$$f_I = \frac{1}{2}(f_L + f_R) - \frac{1}{2}\alpha\Delta u \quad (4)$$

where L and R represent the left and right states of the interface, the first term on the right hand side (average flux) represents the central discretization of the flux terms and the second term is the diffusive flux, with α being the coefficient of numerical diffusion. The interface flux for the upwind method (3) can also be recast in the form (4) by using the definitions of the split fluxes f^\pm . Then the coefficient of numerical dissipation α will be fixed as per the chosen scheme. We can see that the central discretization method is simpler than the upwind method, but the amount of α required to produce a stable and accurate scheme is unknown *a priori*.

The basic idea in this work is to fix α based on a relevant criterion to enhance the resolution of shocks. Note that α must have the dimension of the wave-speed, $a(u) = \frac{\partial g(u)}{\partial u}$ (e.g., for the case of Burgers equation, $f(u) = \frac{u^2}{2}$ and $a(u) = u$) in (4) as $\alpha\Delta u$ must represent a diffusive flux. In Local Lax-Friedrichs method [16], the value of α is taken as the maximum value of the wave-speed of the left and right states of a cell-interface. This makes the LLF method

very simple but not so accurate. A better choice will be to use some physically relevant speed for choosing α . Since the motivation in this work is to enhance the resolution of the shocks, the most relevant physical speed is the shock speed (the speed with which a discontinuity travels) which is given by the Rankine-Hugoniot (jump) condition. Since the contacts also obey the jump conditions the sense of shock speed mentioned in this paper includes both shocks and contacts. Here, the Rankine-Hugoniot condition is enforced in the discretization process, by choosing a relevant expression for α .

2.1 Rankine Hugoniot Condition and Cell Interface Flux

For a shock wave with a left state and right state given by u_L and u_R , the Rankine-Hugoniot (jump) condition is given by

$$\Delta f = s\Delta u \text{ where } \Delta f = f_R - f_L \text{ and } \Delta u = u_R - u_L \quad (5)$$

To enforce this jump condition in the numerical method, the best way is to obtain the interface flux from the same criteria. The jump condition does not directly give the interface flux, as it describes only the relation between left and right states of the jump with the shock speed. In the finite volume method, because of the piece-wise polynomial approximation, a discontinuity in the conserved variable is present at every cell interface. Let us apply the above R-H (jump) condition at the cell interface, $j + \frac{1}{2}$.

$$f_{j+1} - f_j = s_{j+\frac{1}{2}} (u_{j+1} - u_j) \quad (6)$$

To obtain the interface flux, we split the above R-H condition into two parts (see [26]). Suppose there is a discontinuity between j and $j + 1$ moving with a speed $s_{j+\frac{1}{2}}$ as shown in figure 1.

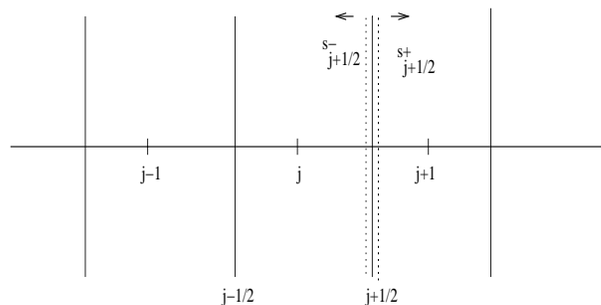


Fig. 1. Shock moving to left or right

This discontinuity can move to the left or right. The jump condition does not indicate the direction of the shock movement. To distinguish the positive and

negative speeds, let us split the shock speed into two parts : a positive one and a negative one.

$$s_{j+\frac{1}{2}} = s_{j+\frac{1}{2}}^+ + s_{j+\frac{1}{2}}^- \quad (7)$$

where

$$s_{j+\frac{1}{2}}^+ = \frac{s_{j+\frac{1}{2}} + |s_{j+\frac{1}{2}}|}{2} \text{ and } s_{j+\frac{1}{2}}^- = \frac{s_{j+\frac{1}{2}} - |s_{j+\frac{1}{2}}|}{2} \quad (8)$$

We have

$$s_{j+\frac{1}{2}}^+ - s_{j+\frac{1}{2}}^- = \frac{1}{2}s_{j+\frac{1}{2}} + \frac{1}{2}|s_{j+\frac{1}{2}}| - \frac{1}{2}s_{j+\frac{1}{2}} + \frac{1}{2}|s_{j+\frac{1}{2}}| = |s_{j+\frac{1}{2}}| \quad (9)$$

Since the left and right moving shocks are separated, we can now split the Rankine-Hugoniot condition into two parts as

$$\begin{aligned} f_{j+1} - f_{j+\frac{1}{2}} &= s_{j+\frac{1}{2}}^+ (u_{j+1} - u_j) \\ f_{j+\frac{1}{2}} - f_j &= s_{j+\frac{1}{2}}^- (u_{j+1} - u_j) \end{aligned} \quad (10)$$

and which then leads to

$$f_{j+\frac{1}{2}} = \frac{1}{2} (f_j + f_{j+1}) - \frac{1}{2} |s_{j+\frac{1}{2}}| (u_{j+1} - u_j) \quad (11)$$

This is the interface flux obtained from the Rankine-Hugoniot condition. Let us now consider the cell interface flux (4) obtained from the finite volume method, given by

$$f_{j+\frac{1}{2}} = \frac{1}{2} (f_j + f_{j+1}) - \frac{1}{2} \alpha_{j+\frac{1}{2}} (u_{j+1} - u_j) \quad (12)$$

Comparing this flux (12) with the interface flux obtained from the Rankine-Hugoniot condition (11), we obtain

$$\alpha_{j+\frac{1}{2}} = |s_{j+\frac{1}{2}}| \quad (13)$$

2.2 Wave-speed Evaluation and Update Formula

We can evaluate the value of $s_{j+\frac{1}{2}}$ numerically as

$$s_{j+\frac{1}{2}} = \begin{cases} \frac{f_{j+1}-f_j}{u_{j+1}-u_j} & \text{if } u_j \neq u_{j+1} \\ \frac{\partial f}{\partial u}|_j & \text{if } u_j = u_{j+1} \end{cases} \quad (14)$$

Similarly, the flux at the cell interface $j - \frac{1}{2}$ can be computed.

In the numerical evaluation of the shock speed as shown above, the denominator involves the differences of the conserved variables across a cell interface. When the left and right states of the cell interface are very close, this denominator can be very small and the shock speed can become nonphysically large. To avoid this problem, we restrict its numerical value within the minimum and maximum of the wave-speed in the computational domain.

$$\begin{aligned} & \text{if } s_{j+\frac{1}{2}} > a_{max} \text{ then } s_{j+\frac{1}{2}} = a_{max} \\ & \text{if } s_{j+\frac{1}{2}} < a_{min} \text{ then } s_{j+\frac{1}{2}} = a_{min} \\ & \text{where } a = \frac{\partial f(u)}{\partial u} \text{ is the wave-speed} \end{aligned} \quad (15)$$

The cell interface fluxes derived above can now be used in the finite volume update formula

$$u_j^{n+1} = u_j^n - \frac{\Delta t}{\Delta x} \left[f_{j+\frac{1}{2}}^n - f_{j-\frac{1}{2}}^n \right] \quad (16)$$

Since the coefficient of numerical dissipation is now fixed in such a way that the Rankine-Hugoniot condition is satisfied in the expressions for the cell interface fluxes, steady shocks is expected to be resolved accurately by this numerical method.

The final update formula for the above central scheme, which we name as the *Method of Optimal Viscosity for Enhanced Resolution of Shocks (MOVERS)*, for a scalar conservation law is equivalent to that of the approximate Riemann solver of Roe [7], despite the derivation and implementation being different. Though the approximate Riemann solver of Roe was derived first for vector conservation laws, it can be simplified for scalar conservation laws (see [2,3]) and the above comparison illuminates how the present method is both similar to and yet different from the popular upwind method of Roe. Note that this method is not based on Roe linearization and will be advantageous for use in the case of vector conservation laws, as will be seen in the next section. The numerical method proposed here will be simple and yet accurate and neither the characteristic decomposition will be used in the approximate Riemann solvers nor the Roe averages be needed. The coefficient of numerical dissipation, $\alpha_{j\pm\frac{1}{2}}$ is linearly proportional to the numerical wave-speed and thus the

dissipation goes to zero when the wave-speed goes to zero. This happens near the sonic points where the wave-speed changes sign. Zero dissipation leads to the violation of the entropy condition, resulting in unphysical solutions such as expansion shocks which violate the entropy condition. To overcome this problem, Harten’s entropy fix [27], like in the case of Roe’s scheme, was used in [25]. Alternatively, for a scalar equation we can sense the diverging of waves at the expansion and locally avoid the R-H condition, taking for α the numerical value of the local maximum of the wave-speeds. This strategy ensures avoiding of unphysical solutions, while capturing the steady shocks exactly. In the later sections, we also present a variation of this method in which the entropy fix is avoided by a novel use of limiter, which can also be used for hyperbolic scalar conservation laws. In the next section, we extend the *MOVERS* to the hyperbolic vector conservation laws.

3 Method of Optimal Viscosity for Enhanced Resolution of Shocks (MOVERS) for Hyperbolic System of Equations

Consider the hyperbolic system of conservation laws in one dimension, as

$$\frac{\partial U}{\partial t} + \frac{\partial F}{\partial x} = 0 \tag{17}$$

where U is vector of conserved variables, $U_i, i = 1, \dots, n$ and F is the flux vector which may have a non-linear functional relationship with U . The hyperbolicity can be seen in the quasi-linear form of the above equations, given by

$$\frac{\partial U}{\partial t} + A \frac{\partial U}{\partial x} = 0 \tag{18}$$

where A is the flux Jacobian matrix, $\frac{\partial F}{\partial U}$ (a $n \times n$ matrix). The system is hyperbolic if the eigenvalues of the flux Jacobian matrix A , denoted by $\lambda_i, i = 1, \dots, n$ are real and the corresponding eigenvectors are linearly independent. If the eigenvalues are also distinct, the system is strictly hyperbolic. For a hyperbolic system, we can diagonalize the matrix as $A = RDR^{-1}$, where the diagonal elements of D are the characteristic values or eigenvalues of A . However, the numerical method presented in this work is not dependent on the eigenvectors and neither the evaluation of the eigenvectors nor the projection of the solution on to the space of eigenvectors is required for this scheme, unlike in the case of Riemann solvers. An explicit scheme in conservation form on a three point stencil and with piecewise polynomial approximation for conserved

variable is given by

$$U_j^{n+1} = U_j^n - \frac{\Delta t}{\Delta x} (F_{j+\frac{1}{2}}^n - F_{j-\frac{1}{2}}^n) \quad (19)$$

where Δt and Δx are respectively time step and the grid-spacing, j represents the cell centroid and $j \pm \frac{1}{2}$ represent the cell interfaces.

The interface flux is written as in a central scheme in the form

$$F_I = \frac{1}{2}(F_L + F_R) - \frac{1}{2}\alpha(U_R - U_L) \quad (20)$$

where L and R represent the left and right states of the cell interface. Note that α is the coefficient of numerical dissipation which will be chosen later based on the Rankine-Hugoniot condition. The Rankine-Hugoniot condition is

$$\Delta F = s\Delta U \quad (21)$$

Note that while conserved variable vector U and the flux vector F are vectors, the shock speed s is a scalar. Proceeding in the same way as was done for the scalar case, we can obtain the coefficient of numerical dissipation, based on enforcing the Rankine-Hugoniot jump condition in the discretization, as

$$\alpha_{j+\frac{1}{2}} = |s_{j+\frac{1}{2}}| \quad (22)$$

So far, the situation is similar to that of the scalar case. Now, evaluating s from the Rankine-Hugoniot condition, $\Delta F = s\Delta U$, is not straightforward, as U and F are vectors but s is a scalar. We present three different possibilities of evaluating the shock speed, leading to three different variations of the *Method of Optimal Viscosity for Enhanced Resolution of Shocks* (**MOVERS**), which we name as (1) *MOVERS- n* - with a n -wave approximation for the wave-speed, (2) *MOVERS-1* - with a one wave approximation for the wave-speed and (3) *MOVERS-L* - a limiter based central solver which switches over from R-H condition based dissipation of 1(or n)-wave approximation method to that of LLF. The first two variants will require an entropy fix, while the last one does not need it. All the three variants will capture steady discontinuities (shocks and shear layers) with enhanced resolution, resolving them exactly wherever appropriate.

3.1 *MOVERS-n*

One possibility is to choose s and, therefore, α as diagonal matrices. Thus, in the relation $\Delta F = s\Delta U$, since U and F are $n \times 1$ vectors, we can choose s as a $n \times n$ matrix. The simplest of such $n \times n$ matrices is a diagonal matrix with n diagonal elements.

$$\begin{bmatrix} \Delta F_1 \\ \Delta F_2 \\ \vdots \\ \Delta F_n \end{bmatrix} = \begin{bmatrix} s_{1,j+\frac{1}{2}} & 0 & \cdots & 0 \\ 0 & s_{2,j+\frac{1}{2}} & \cdots & 0 \\ \vdots & \vdots & \cdots & \vdots \\ 0 & 0 & \cdots & s_{n,j+\frac{1}{2}} \end{bmatrix} \begin{bmatrix} \Delta U_1 \\ \Delta U_2 \\ \vdots \\ \Delta U_n \end{bmatrix} \quad (23)$$

We can now evaluate the shock speeds as

$$s_{m,j+\frac{1}{2}} = \frac{\Delta F_m}{\Delta U_m}, \quad m = 1, 2, \dots, n \quad (24)$$

As an example, let us consider the case $n = 3$. Then, we obtain

$$\begin{bmatrix} \Delta F_1 \\ \Delta F_2 \\ \Delta F_3 \end{bmatrix} = \begin{bmatrix} s_{1,j+\frac{1}{2}} & 0 & 0 \\ 0 & s_{2,j+\frac{1}{2}} & 0 \\ 0 & 0 & s_{3,j+\frac{1}{2}} \end{bmatrix} \begin{bmatrix} \Delta U_1 \\ \Delta U_2 \\ \Delta U_3 \end{bmatrix} \quad (25)$$

We can now calculate the shock speeds (three of them) as

$$\begin{bmatrix} s_{1,j+\frac{1}{2}} \\ s_{2,j+\frac{1}{2}} \\ s_{3,j+\frac{1}{2}} \end{bmatrix} = \begin{bmatrix} \frac{(F_1)_R - (F_1)_L}{(U_1)_R - (U_1)_L} \\ \frac{(F_2)_R - (F_2)_L}{(U_2)_R - (U_2)_L} \\ \frac{(F_3)_R - (F_3)_L}{(U_3)_R - (U_3)_L} \end{bmatrix} \quad (26)$$

This is an approximation based on ‘ n ’ waves for a hyperbolic system with ‘ n ’ equations ($n = 3$ in the illustrated example considered above, as in the case of 1-D Euler equations). The Rankine-Hugoniot condition then becomes a simple diagonal matrix and can be easily extended from the scalar case to conservation laws of complex physics involving multiple characteristics such as the magneto-hydrodynamics equations. We term this method as *MOVERS-n*, representing *n-waves* approximation for the shock speed evaluation.

3.2 *MOVERS-1*

A simpler alternative to n -wave approximation to shock speed in *MOVERS* is a one-wave approximation. This changes the dissipation from a diagonal matrix form to a scalar form. One way of choosing a scalar dissipation, satisfying the Rankine-Hugoniot condition as derived before, is to choose one among all the n -waves representing the shock speed. A simple guide line for this choice is to find out which of the n -waves has the maximum possible information. It is still better if this information also reflects the expected behavior of characteristics converging at the shocks and diverging in expansion regions. Such a central Rankine-Hugoniot solver with one-wave approximation for the shock speed is named as *MOVERS-1*. To illustrate this idea, we consider the system of Euler equations in one dimension as follows.

3.2.1 *MOVERS-1 for 1-D Euler Equations*

The Euler equations has three values of the shock speed as in the equation (26). For the n -wave approximation (*MOVERS- n*) each of the diagonal elements becomes the co-efficient of dissipation for each of the equations in the system which makes α a matrix. However, here we approximate with a single wave as a representation of multiple waves and hence seek a scalar dissipation coefficient.

To choose a single wave, we use a criterion based on suitability of the numerical wave-speed to represent the actual physical requirements for a shock wave and an expansion wave. The characteristics must converge on to the shock wave and must diverge at an expansion wave. Since the coefficient of numerical diffusion represents a wave-speed, it is preferable for it to mimic these properties. We also consider the amount of information carried by each of the terms while choosing the single wave, as the best term will carry the maximum information. Numerical experiments for the 1-D shock tube problem and the quasi 1-D nozzle problem have shown that out of all the three components in (26), the third coefficient (for the energy equation) has the desirable properties. The mass wave does not represent merging of characteristics at shocks while the momentum wave, the second component of (26) can lead to problem of denominator becoming zero for equal left and right flux (for a steady discontinuity) even when a Riemann problem persists. The energy term also has information of all the primitive variables and responds to all the flow features.

The plot of Wave speed for quasi 1D nozzle flow shown in figure 2a changes sign at two points (at $X \approx 1.5$ and $X \approx 2.1$) where an expansion and a shock occur, respectively. It can be seen that wave speeds diverge at $X = 1.5$ and converge at $X = 2.1$, indicating that the choice of the third component (corresponding to energy equation) for the wave speed in the 1-wave approximation

is the desired choice. The wave speed behavior for the unsteady flow in shock tube (figure 2b) also shows the same behavior, by a sign change from negative to positive for an expansion at ($X \approx -4.5$) and changing sign from positive to negative at the shock at ($X \approx +4.5$).

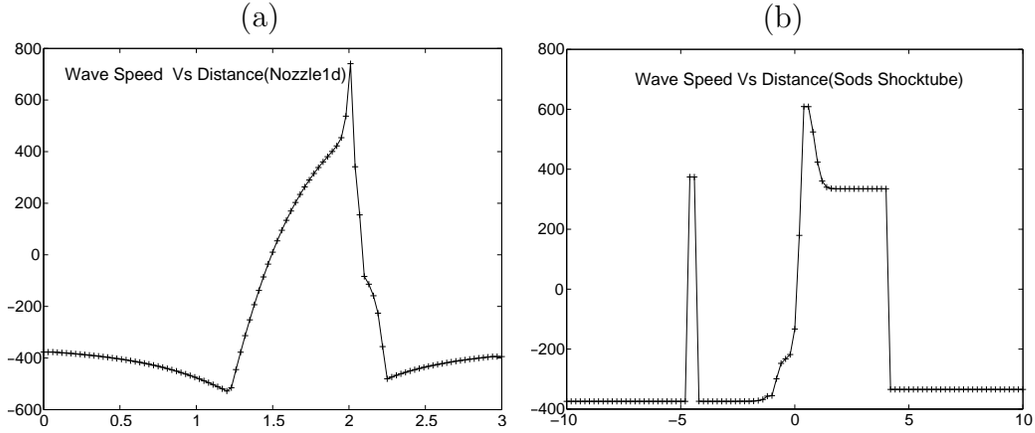


Fig. 2. Plot of Wavespeed (α) Vs Distance(X) for (a) quasi 1D converging diverging nozzle steady flow (b) for Sod's shocktube problem

Therefore, the third component in (26) is chosen as the scalar coefficient of numerical dissipation for all the three equations. Thus, we can write

$$\begin{bmatrix} s_{1,j+\frac{1}{2}} \\ s_{2,j+\frac{1}{2}} \\ s_{3,j+\frac{1}{2}} \end{bmatrix} = \begin{bmatrix} \frac{(F_3)_R - (F_3)_L}{(U_3)_R - (U_3)_L} \\ \frac{(F_3)_R - (F_3)_L}{(U_3)_R - (U_3)_L} \\ \frac{(F_3)_R - (F_3)_L}{(U_3)_R - (U_3)_L} \end{bmatrix} \quad (27)$$

This effectively means choosing a scalar numerical dissipation coefficient based on Rankine-Hugoniot condition as a representative numerical wave speed

$$s_{j+\frac{1}{2}} = \frac{(F_3)_R - (F_3)_L}{(U_3)_R - (U_3)_L} \quad (28)$$

even for a system of equations.

For system of equations representing ideal MHD flow, we again choose the energy term for the same reasons as described for Euler equations and for shallow water equations, we choose the momentum equation which has the maximum information.

The variants of the new scheme proposed here, namely *MOVERS-1,n* require entropy fix to avoid unphysical expansion shocks near the sonic points. Another variant *MOVERS-L*, in which the numerical dissipation is switched from that in the Local Lax-Friedrichs method to the numerical dissipation based on the Rankine-Hugoniot condition, based on a limiter, is presented here. The co-efficient of numerical diffusion of this hybrid scheme is written as

$$\alpha = \alpha_{MOVERS} + \phi(r) [\alpha_{LLF} - \alpha_{MOVERS}] \quad (29)$$

where α_{LLF} is the local maximum of the spectral radius of the flux Jacobian matrix and α_{MOVERS} is the corrected wave speed obtained using *MOVERS-n(or 1)* (see the expressions (33) to (36) in the following subsection), using the R-H condition. Here, ϕ is a standard limiter. This is to enforce the *MOVERS* to work only at the discontinuities while reverting to the dissipative (entropy condition satisfying) Local Lax-Friedrichs method elsewhere. We use the *minmod* limiter suggested by Yee [45] as defined below

$$\phi_{j+\frac{1}{2}}(r_{j+\frac{1}{2}}) = \text{minmod}(1, r_j^+, r_{j+1}^-) \quad (30)$$

The ratio of solution differences are defined as

$$r_j^+ = \frac{U_j - U_{j-1}}{U_{j+1} - U_j} \quad \text{and} \quad r_j^- = \frac{U_{j+1} - U_j}{U_j - U_{j-1}} \quad (31)$$

If the denominator in the above expression is small (when $U_j \approx U_{j+1}$), then we modify r to prevent numerical overflow as

$$r_j^+ = \text{sign}(U_{j+1} - U_j) \frac{U_j - U_{j-1}}{\delta} \quad \text{if } |U_{j+1} - U_j| < \delta \quad (32)$$

where δ is a small parameter. The same approach is used for r_j^- also. The limiter based *Central Rankine-Hugoniot Solver*, *MOVERS-L*, therefore, can capture steady discontinuities exactly and still avoid expansion shocks.

For a second order accurate version of *MOVERS-L*, as the left and right states of a cell interface are obtained by reconstruction, the scheme switches from a second order accurate *LLF* method in smooth regions to second order accurate *MOVERS* in high gradient regions. Two kinds of limiters will be operating in this case - one the standard limiter used for second order reconstruction of conserved variables as seen in equations (38,39) of section 3.6. This makes the scheme first order accurate at high gradients and retains second order

accuracy in smooth regions. The other limiter(29,30) is specific to this scheme for switching over from *MOVERS-n(or 1)* to LLF method (here we use only minmod limiter for a convex combination). For practical applications, since second order accurate LLF method is used in smooth regions, the solution is not degraded by the limiter in these regions.

3.4 Wave-speed correction

In both *MOVERS-n(and 1)*, while calculating the wave-speeds (expressions (26) and (28) respectively), if the denominator ΔU becomes small, the condition refers to an isentropic case and we assign the minimum in the characteristic speed spectrum of the flux Jacobian as the wave speed. When there is a steady shock ($s = 0$), since the discrete update formula $U_j^{n+1} = U_j^n - \frac{\Delta t}{\Delta x} [F_{j+\frac{1}{2}}^n - F_{j-\frac{1}{2}}^n]$ approximates the Euler equations in a consistent and stable manner, all the above representative numerical wave speeds of (26) and (27) reduce to the speed of the shock, which is zero in the steady case. Thus any version of the Central Rankine-Hugoniot Solver, *MOVERS*, will capture steady shocks exactly, without any numerical dissipation. Because of the form of the dissipation we have assumed, even when there is a steady contact discontinuity, this algorithm will capture it exactly, without numerical dissipation (which is demonstrated in the section on results). This low dissipative algorithm is employed only close to the discontinuities. Elsewhere, when the denominator ΔU is too small, the evaluated value of s can be non-physically high. We use a wave speed correction algorithm to bring the wave speeds to physically admissible values. The wave speed correction algorithm is a min-max wave speed limiter with the absolute values of discrete wave speed being restricted within the spectrum of characteristic values (eigenvalues) of the hyperbolic system while still retaining the same direction of the computed uncorrected discrete wave speed.

$$s_{j+1/2} = \begin{cases} \left| \frac{F_{j+1} - F_j}{U_{j+1} - U_j} \right| & \text{if } U_j \neq U_{j+1} \\ \lambda_{min,j+1/2} & \text{if } U_j = U_{j+1} \end{cases} \quad (33)$$

$$s_{j+\frac{1}{2}} = \lambda_{max} \quad \text{if } |s_{j+\frac{1}{2}}| \geq \lambda_{max} \quad (34)$$

$$s_{j+\frac{1}{2}} = \lambda_{min} \quad \text{if } |s_{j+\frac{1}{2}}| \leq \lambda_{min} \quad (35)$$

With these corrections, the steps to update the vector conservation laws involve computation of representative numerical wave speeds at all the points

in the domain and correction of such wave speeds to make them lie within the bounds of spectral radius. Here, λ_{max} and λ_{min} refer to the local maxima and minima of the magnitudes of characteristic speeds defined by the eigenvalues of the flux Jacobian matrix. As an example, for the Euler equations in one dimension, we define

$$\begin{aligned}\lambda_{max} &= \text{Max}(\text{max}(|u|, |u - a|, |u + a|)_R, \text{max}(|u|, |u - a|, |u + a|)_L) \\ \lambda_{min} &= \text{Max}(\text{min}(|u|, |u - a|, |u + a|)_R, \text{min}(|u|, |u - a|, |u + a|)_L)\end{aligned}\quad (36)$$

since the eigenvalues in this case are $u - a$, u and $u + a$ with u being the fluid velocity and a representing the sound speed.

3.5 Entropy Correction

The numerical diffusion of the two variants *MOVERS-n* (and 1) take to low values at the expansive sonic points where wave speeds change the sign. To avoid any expansion shocks, just like Harten's entropy correction[27] with using the maxima of the characteristic speeds as given in (37).

$$s_j = \frac{s_j^2 + \delta^2}{2\delta} \quad \text{if } s_j < \delta \quad \& \quad s_j = s_j \quad \text{otherwise} \quad (37)$$

where $\delta = \kappa \lambda_{max}$ and κ is a fractional numerical value from 0 to 1. As Local Lax-Friedrichs method is a good alternative to low dissipative schemes like Roe scheme or *MOVERS* in the regions of expansion fans with sonic points, it is also used as an entropy fix [6]. In the numerical experiments done in this work, it was found that reducing the numerical dissipation of the LLF method by half(i.e., $\kappa = 0.5$) works reasonably well avoiding expansion shocks and other non-smoothness. Since, this entropy correction adds numerical diffusion everywhere, loss of accuracy in shocks can be expected. A suitable correction to this scheme can then be to retain sufficient entropy in all regions, especially at the sonic points except the weak solution. This becomes the motivation for the limiter based variant of the scheme *MOVERS-L*.

3.6 Second Order Accurate Schemes

The second order accurate schemes are constructed by assuming piecewise linear approximation for the conserved variables, as

$$U(x, t^n) = \left[U_j^n + (U_x)_j^n (x - x_j) \right]_{[x_{j-1/2}, x_{j+1/2}]}, \quad x_{j\pm 1/2} = x_j \pm \frac{\Delta x}{2} \quad (38)$$

where U_j^n is the computed cell average, and $(U_x)_j^n$ are approximations to the exact derivatives, $U_x(x, t^n)$, evaluated from the computed cell averages. The non-oscillatory behavior of the scheme depends on the appropriate choice of approximate derivatives, and we use the one parameter family of minmod limiter for this purpose.

$$(U_x)_j = \text{minmod} \left(\beta \frac{U_j - U_{j-1}}{\Delta x}, \frac{U_{j+1} - U_{j-1}}{2\Delta x}, \beta \frac{U_{j+1} - U_j}{\Delta x} \right), \quad 1 \leq \beta \leq 2 \quad (39)$$

The numerical values of fluxes and the primitive variables required for the update are computed at the cell interfaces from the above reconstruction procedure. Several benchmark problems of gas dynamics, shallow water flows and ideal magneto-hydrodynamics are solved and results are presented in the next section.

4 Results and Discussion

4.1 Results with *MOVERS* for Burgers equation

MOVERS is first tested for the inviscid Burgers equation in one dimension, given by

$$\frac{\partial u}{\partial t} + \frac{\partial f(u)}{\partial x} = 0 \quad \text{where } f(u) = \frac{1}{2}u^2 \quad (40)$$

The test case used models a shock wave and an expansion fan [4]. The results with the Local Lax-Friedrichs method are also presented for comparison, in figure (3). The shock wave is captured exactly with *MOVERS* and the expansion fan is captured well along with a Harten's entropy fix.

The results obtained with *MOVERS* for the 2-D Burgers equation, given by

$$\frac{\partial u}{\partial t} + \frac{\partial f_1(u)}{\partial x} + \frac{\partial f_2(u)}{\partial y} = 0 \quad \text{with } f_1(u) = \frac{1}{2}u^2 \quad \text{and } f_2(u) = u \quad (41)$$

are shown in the figure (4). This test case is taken from [28]. The shock is again captured exactly.

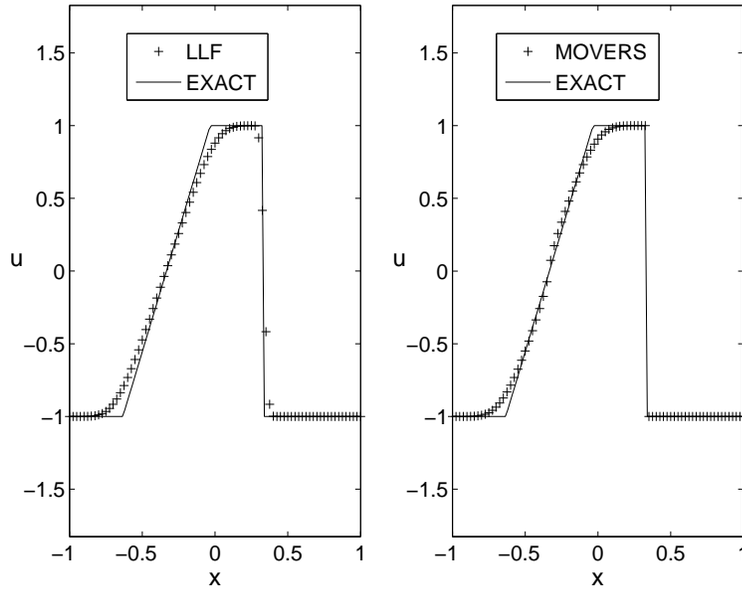


Fig. 3. 1D Burgers equation solution : First order accurate results with *MOVERS*

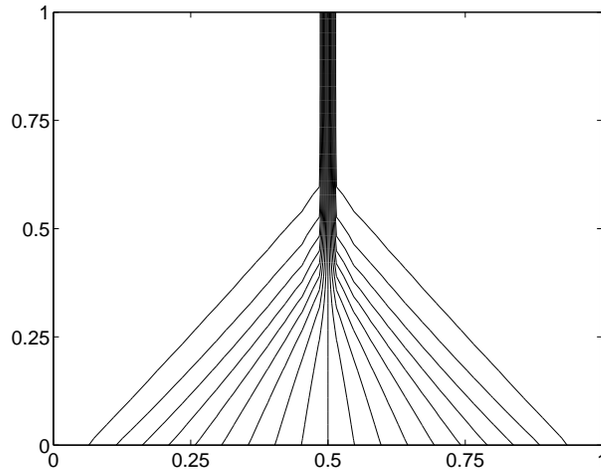


Fig. 4. 2D Burgers equation solution with first order accurate *MOVERS* using 32x32 points: Velocity Contour plot

4.2 Results with *MOVERS* for Euler equations

The Euler equations in one dimension are given by

$$\frac{\partial U}{\partial t} + \frac{\partial F}{\partial x} = 0 \tag{42}$$

where

$$U = \begin{bmatrix} \rho \\ \rho u \\ \rho E \end{bmatrix} \text{ and } F = \begin{bmatrix} \rho u \\ p + \rho u^2 \\ pu + \rho u E \end{bmatrix} \quad (43)$$

where U is the vector of conserved variables and F is the flux vector, with ρ representing the density, u the fluid velocity, p the pressure and E the total energy, defined by $E = \frac{p}{\rho(\gamma-1)} + \frac{1}{2}u^2$. This system of equations is hyperbolic, with the eigenvalues given by $\lambda_1 = u - a$, $\lambda_2 = u$ and $\lambda_3 = u + a$. The quasi-1D and multi-dimensional forms of the Euler equations are given in [3]. Numerical solution of inviscid compressible flows for the 1-D case of steady contact discontinuity in a shock tube, the quasi-one dimensional nozzle flow [29], oblique shock reflection, supersonic flow over a ramp in a channel [30], supersonic flow over a forward-facing step in a channel [31], double Mach reflection [31], shear flow of Mach 3 over Mach 2 [32] and hypersonic flow across a half-cylinder [33] are obtained using *MOVERS-n*. The results consistently show high accuracy of the new solver in capturing flow features. It is worth noting that this is achieved without any Riemann solvers and complicated flux splittings, in a simple format of a central scheme.

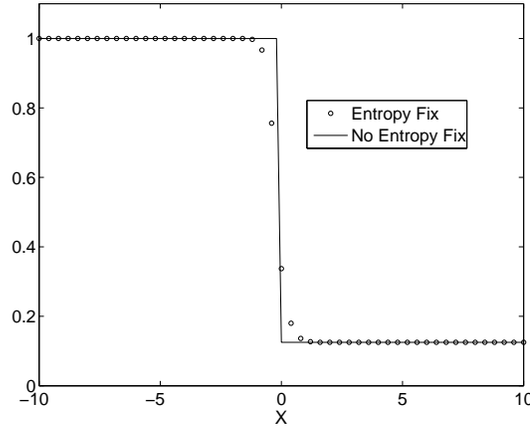


Fig. 5. Steady contact discontinuity in a shock tube with first order accurate *MOVERS-n*, with and without entropy fix; Density plot

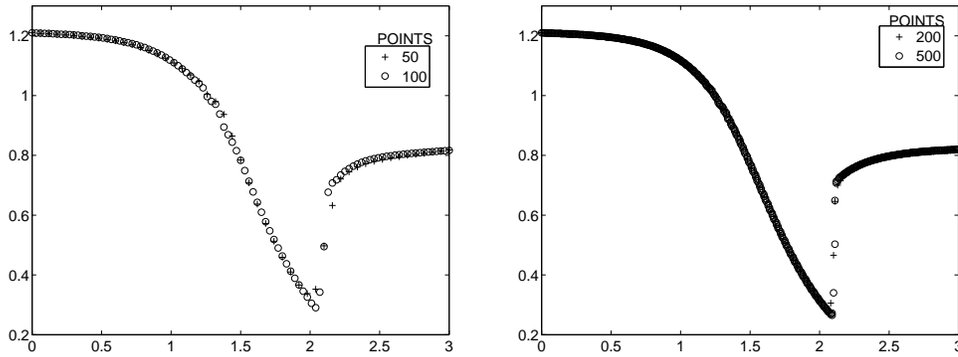


Fig. 6. Nozzle flow : Density plot with first order accurate *MOVERS-n*

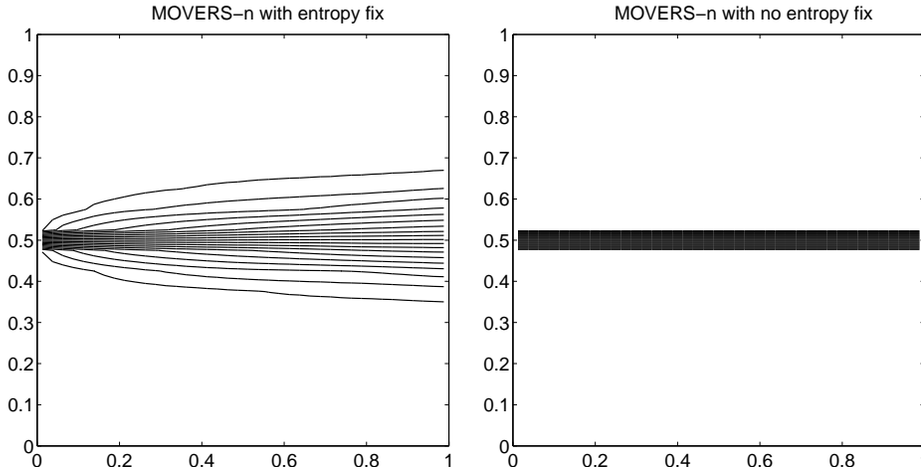


Fig. 7. Horizontal Shear flow (Mach 3 flow over Mach 2 flow), Mach contours with *MOVERS-n* (first order) on a 20x20 grid

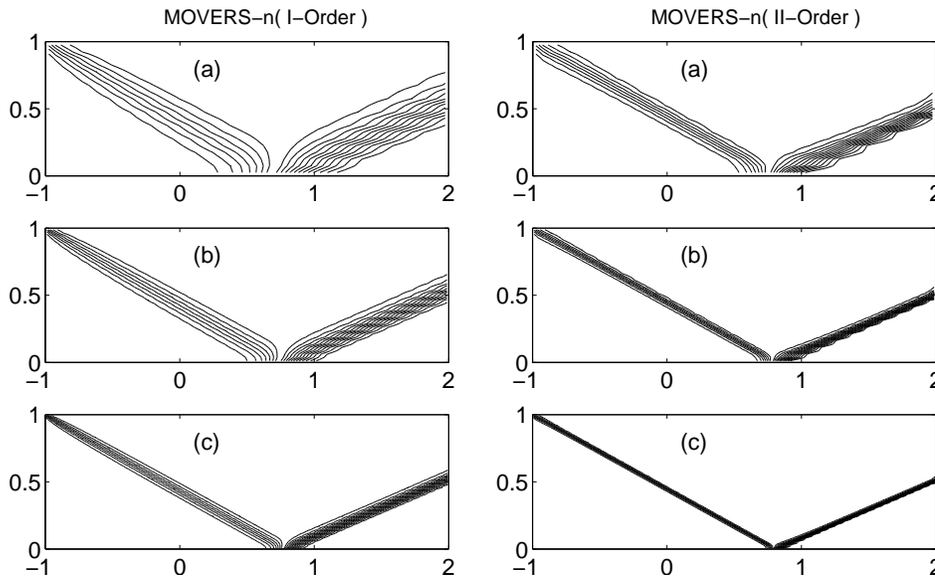


Fig. 8. Oblique Shock Reflection, Density contours (0.9:0.1:2.7) with *MOVERS-n* on grids (a) 60x20 (b) 120x40 and (c) 240x80

The results with *MOVERS-n* for the steady contact discontinuity in a shock tube (Fig. 5) and quasi 1-D nozzle flow (Fig. 6) show clear resolution of the flow and slip surface is captured exactly for a shear flow of Mach 3 over Mach 2 (Fig. 7). The two dimensional oblique shock reflection problem (Fig. 8) on crude, medium and fine grids, show a consistent improvement in the resolution of the solutions. The trend persists for unsteady problems as well, as depicted in figures 9, 10 and all the flow features are captured with high accuracy. Stagnation line pressure and Mach number plots for steady hypersonic flow over half-cylinder in figure 11 effectively highlight the improvement in solution over the *LLF* scheme. C_p distribution for two cases of transonic flow over NACA0012 aerofoil [42] are presented in figures 12-14. The solutions

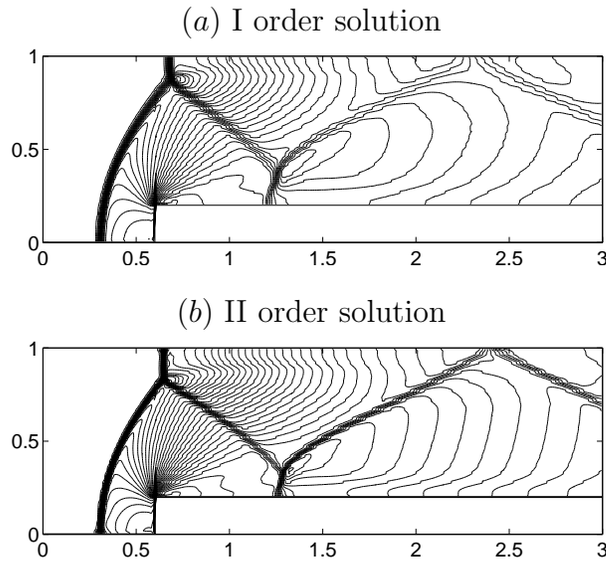


Fig. 9. Density contours at time=4 with *MOVERS-n* using structured mesh of (240x80) points for Mach 3 flow over a forward-facing step in channel

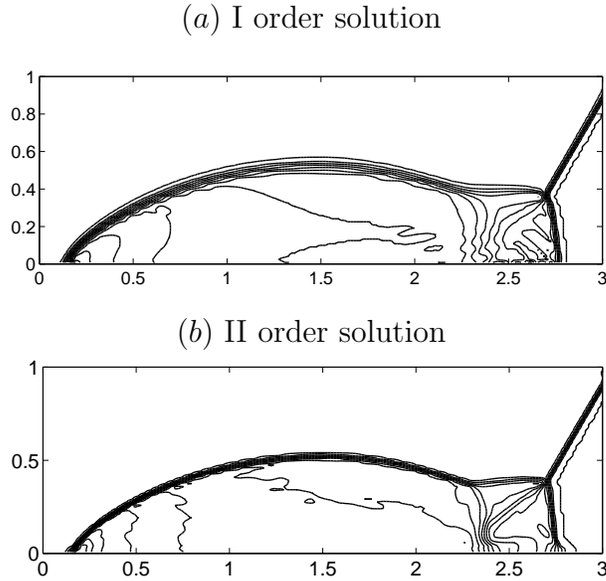


Fig. 10. Density contours at time=0.2 with *MOVERS-n* using structured mesh of (240x60) points for Double Mach Reflection due to Mach 10 shock moving through a 30° wedge

without any entropy correction resolve the shock accurately but for mild non-smoothness elsewhere. However, with an entropy fix using $\phi = 0.25$ in (37) the shocks are diffusive.

The results of the n-wave solver are compared with Roe's Solver in figures (15) and (16). The solution for ramp in channel flow problem is shown in figure 17 indicates the high accuracy resolution with *MOVERS-n* using unstructured grid computation. The results presented demonstrate the accuracy of this

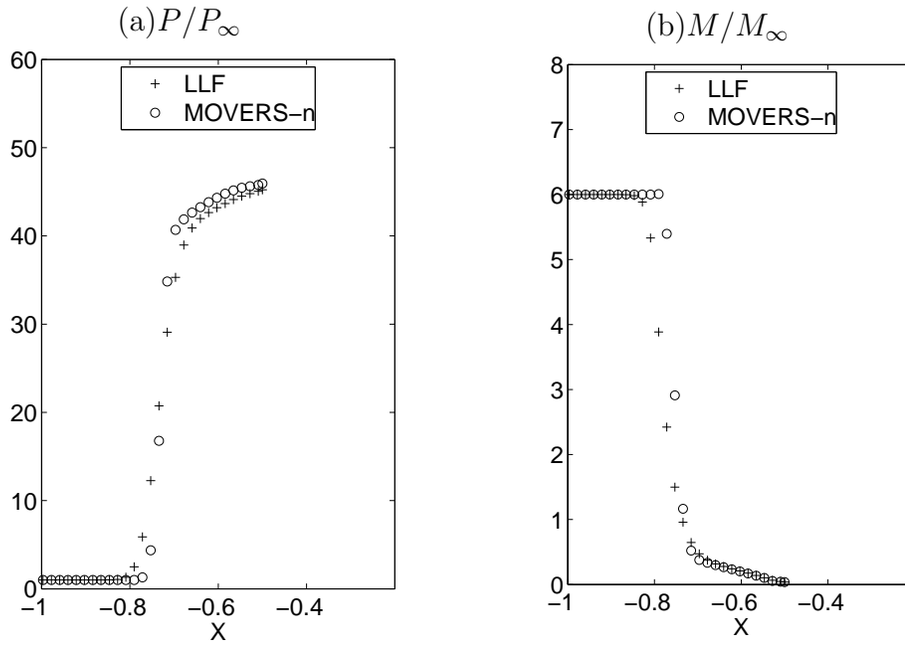


Fig. 11. Results along stagnation line for Mach 6 flow across half cylinder using structured mesh of 45x45 points with first order accurate *MOVERS-n*

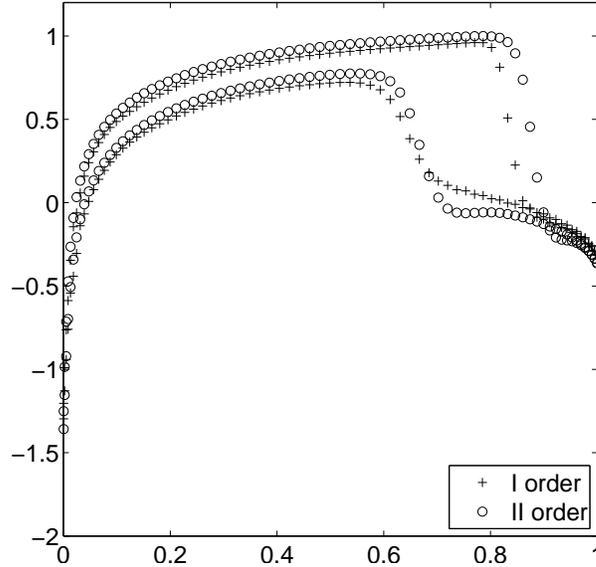


Fig. 12. Pressure co-efficient distribution on NACA0012 airfoil for Mach 0.85 flow at 1° angle of attack with *MOVERS-n*

version of the Central Rankine-Hugoniot Solver (*MOVERS-n*) in capturing the flow features accurately. The steady shocks aligned with grid lines are captured exactly, without any numerical dissipation. With the use of an entropy fix, the expansion fans are captured well.

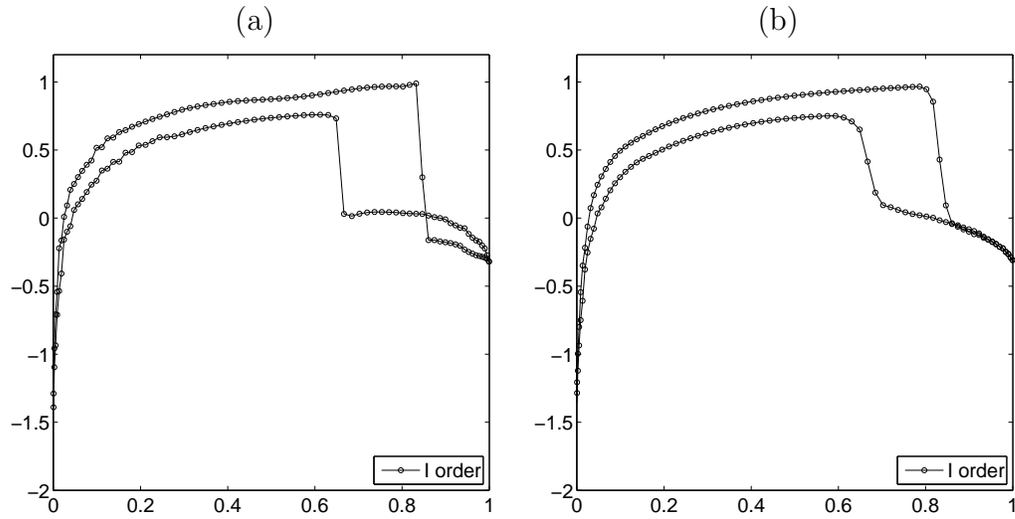


Fig. 13. Pressure co-efficient distribution on NACA0012 airfoil for Mach 0.85 flow at 1° angle of attack with *MOVERS-n* (a) without entropy fix (b) with entropy fix

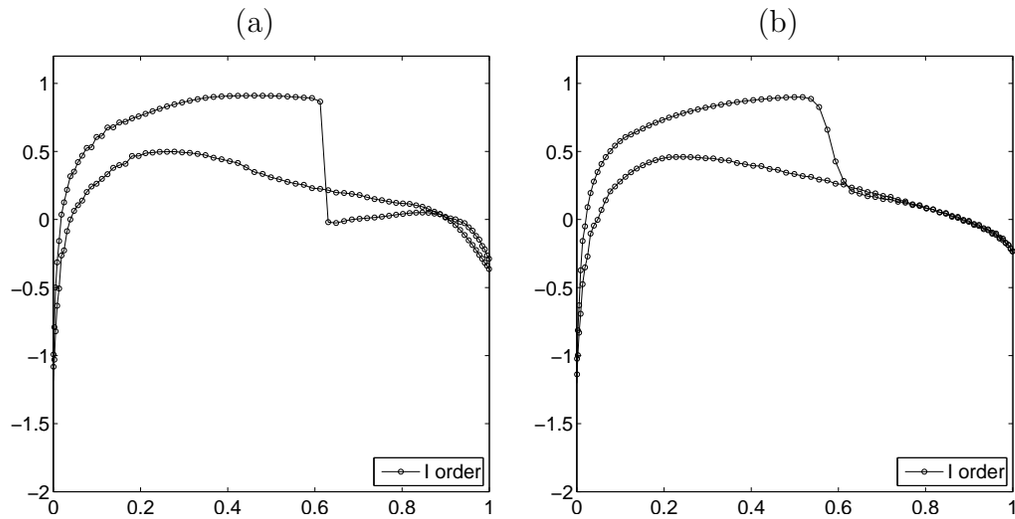


Fig. 14. Pressure co-efficient distribution on NACA0012 airfoil for Mach 0.8 flow at 1.25° angle of attack with *MOVERS-n*(a) without entropy fix (b) with entropy fix

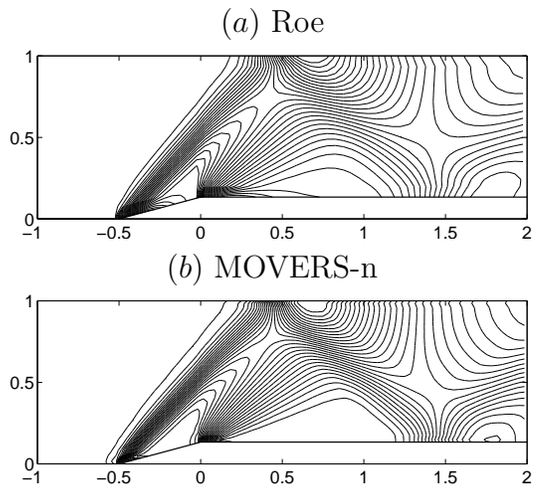


Fig. 15. Comparison of first order accurate solutions of Roe's scheme and *MOVERS-n* for supersonic flow over ramp in channel

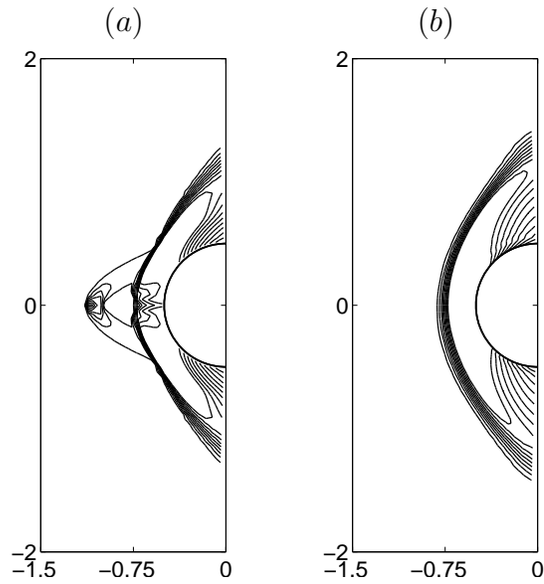
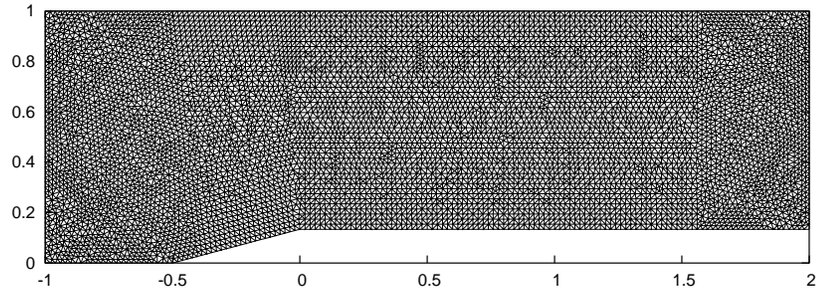
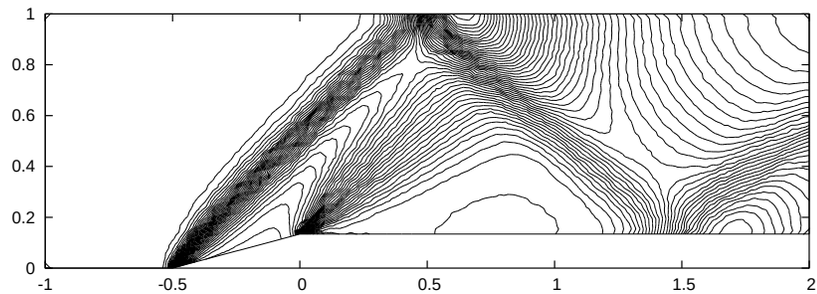


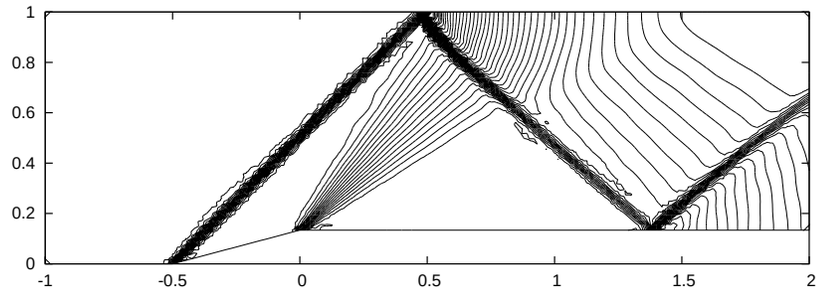
Fig. 16. Comparison of first order accurate solutions of (a) Roe's scheme with (b) *MOVERS-n* for Mach 6 flow across half cylinder



(a) Finite Volume Mesh



(b) I order solution



(c) II order solution

Fig. 17. Supersonic flow over a ramp in a channel: Pressure contours with *MOVERS-n* using unstructured mesh of 7503 points

4.3 Results with *MOVERS-n* for Shallow Water Equations

The one-dimensional form of shallow water equations in differential form can be written as

$$U_t + F(U)_x = S \quad (44)$$

where

$$U = \begin{bmatrix} h \\ hu \end{bmatrix}, \quad F = \begin{bmatrix} hu \\ hu^2 + \frac{1}{2}gh^2 \end{bmatrix}, \quad S = \begin{bmatrix} 0 \\ -ghB_x \end{bmatrix} \quad (45)$$

The system is hyperbolic with eigenvalues $\lambda_1 = u - \sqrt{gh}$ and $\lambda_2 = u + \sqrt{gh}$ for the one dimensional flux Jacobian matrix, given by

$$F'(U) = \begin{bmatrix} 0 & 1 \\ -u^2 + gh & 2u \end{bmatrix} \quad (46)$$

The non-homogeneous (source) term S is very common and quite often represents the water bed beneath. Numerical solution with *MOVERS-n* for some bench mark problems are presented below.

1D Dam-Break problem on a variable Depth River-bed

This is a test case [34] with variable river-bed and so the system of equations is not homogeneous, which means that a source term is present in the equation. For this test case, the river-bed is defined as

$$B(x) = \begin{cases} 0.125(\cos(10(\pi(x - \frac{1}{2}))) + 1) & \text{if } 0.4 \leq x \leq 0.6 \\ 0 & \text{otherwise} \end{cases}$$

and we have the following initial conditions

$$u(x) = 0$$

$$h(x, 0) = \begin{cases} 1 - B(x) & \text{if } 0 \leq x \leq \frac{1}{2} \\ 0.5 - B(x) & \text{if } \frac{1}{2} < x \leq 1 \end{cases}$$

The presence of discontinuity at $x = 0.5$ is like a barrier separating two river heights. Reflection boundary conditions are taken into consideration assuming walls at $x = 0$ and $x = 1$. Results presented in figure (18) shows all the flow features being captured well by *MOVERS-n*.

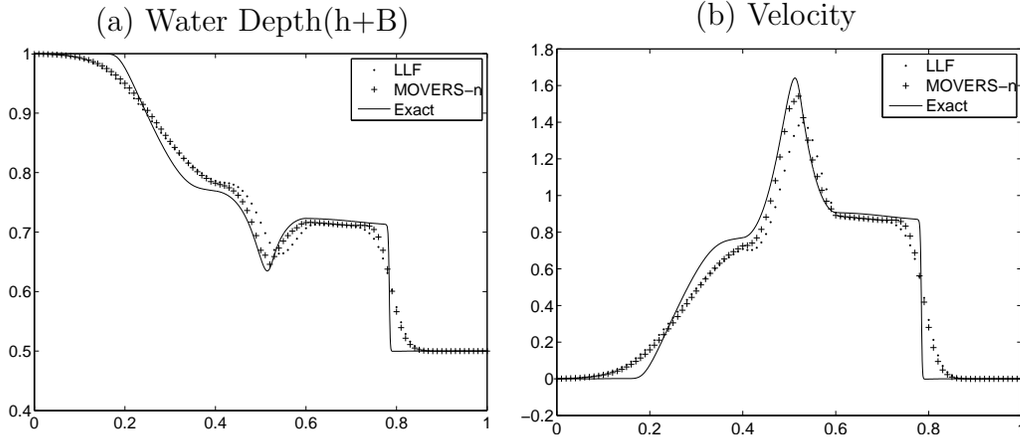


Fig. 18. Dam-break flow with bottom topography on 100 grid points using first order accurate *MOVERS-n* at time 0.1 sec

4.3.1 1D Steady State Trans-critical flow with Shock

In this benchmark test case [35] we study the convergence towards steady flow over a hump. We consider the system with initial conditions

$$\begin{aligned} u(x, 0) &= 0 \\ h(x, 0) + Z(x) &= H_0 \end{aligned}$$

where H_0 is the constant water level downstream provided by the boundary condition. We test our scheme to this steady state flow where the bathymetry is non-trivial and is given by

$$Z(x) = \begin{cases} 0.2 - 0.05(x - 10)^2 & \text{if } 8 \leq x \leq 12, \\ 0 & \text{otherwise} \end{cases}$$

in a channel of length $L = 25m$ and $H = 2m$, $CFL = 0.5$. In this test case we impose an upstream boundary condition for the discharge as $q = 0.18m^2/s$ and the downstream boundary condition for the water level is $H_0 = 0.33m$. The final water level at time 200 seconds computed with mesh size $\Delta x = 0.125m$

and residue for computation are compared with LLF method and displayed in the figure (19).

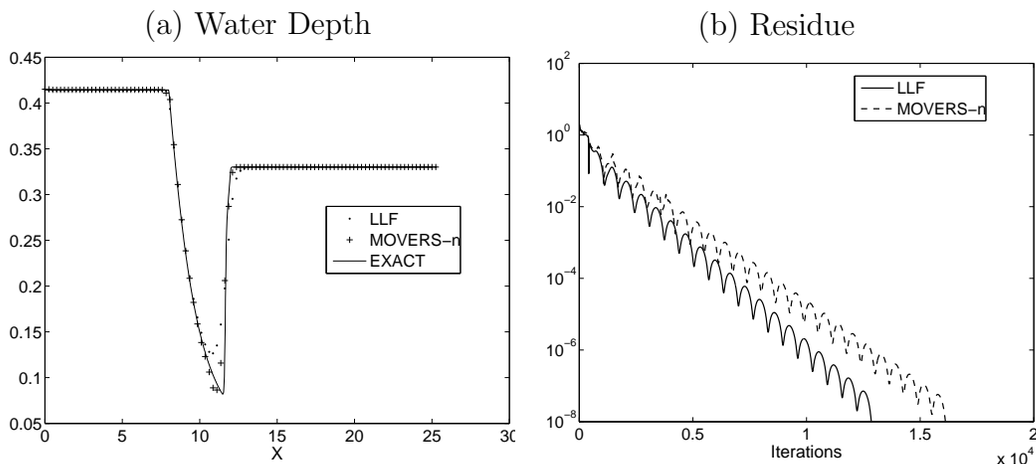


Fig. 19. Transcritical steady flow with hydraulic jump on 120 points at $t=200$ seconds: first order accurate *MOVERS-n* compared with that of LLF

The numerical simulation of several rapidly varying two-dimensional flows are presented in the following sub-section.

4.3.2 2D Partial Dam Break Problem

This problem corresponds to a partial breach [36]. Consider a 2D partial dam-break problem with an unsymmetrical breach. The computational domain is defined in a 200 m long and 200 m wide channel. The non-symmetrical breach or sluice gates are 75-m wide, and the thickness is 10 m. (see figure 20). A dam is assumed to fail instantaneously or the sluice gates are assumed to be opened instantly. The initial upstream water depth is $h_1 = 10m$ and downstream water depth is $h_0 = 5m$. A grid of 40×40 cells is used for this problem. This test case is similar to the shock tube problem for the Euler equations.

The Numerical results displaying 3D views of the water surface elevation after dam failure and velocity contours at time $t=7.2$ seconds are presented in figure (21). At the instant the dam breaks, water is released through the breach, forming a positive wave that propagates downstream and a negative wave that moves upstream.

This problem corresponds to a partial breach [36] and detailed in figure 20. The Numerical results displaying 3D views of the water surface elevation after dam failure and velocity contours at time $t=7.2$ seconds are presented in figure (21) and that for steady oblique hydraulic jump compared with LLF scheme is depicted in figure (22). The efficacy of the scheme in resolving all the flow features is demonstrated. Figure 23 presents the surface height plot variation for the oblique jump.

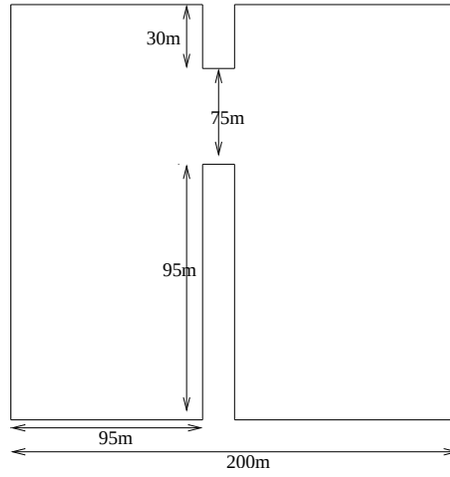


Fig. 20. Problem Sketch for partial dam breach

4.4 Results with *MOVERS-n* for Ideal Magneto-Hydrodynamic Equations

The one-dimensional ideal MHD equations are given by

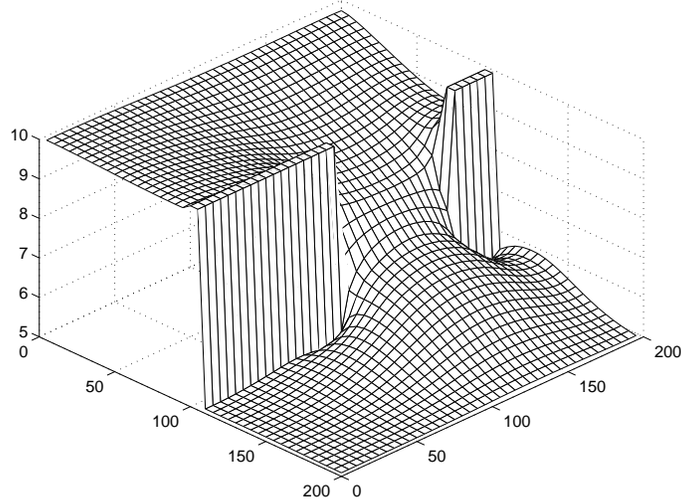
$$\frac{\partial}{\partial t} \begin{bmatrix} \rho \\ \rho u_1 \\ \rho u_2 \\ \rho u_3 \\ B_x \\ B_y \\ B_z \\ E \end{bmatrix} + \frac{\partial}{\partial x} \begin{bmatrix} \rho u_1 \\ \rho u_1^2 + p + \frac{1}{2} B^2 - B_x^2 \\ \rho u_1 u_2 - B_x B_y \\ \rho u_1 u_3 - B_x B_z \\ 0 \\ B_y u_1 - u_2 B_x \\ B_z u_1 - u_3 B_x \\ u_1 (E + p + \frac{1}{2} B^2) - B_x (u_1 B_x + u_2 B_y + u_3 B_z) \end{bmatrix} = 0 \quad (47)$$

Since the magnetic field in the flow direction is constant, both time and space derivatives of B_x are zero. The seven eigenvalues of the Jacobian matrix (see [37]) are

$$\begin{aligned} \lambda_1 &= u - c_f, \quad \lambda_2 = u - c_A, \quad \lambda_3 = u - c_s, \quad \lambda_4 = u, \\ \lambda_5 &= u + c_f, \quad \lambda_6 = u + c_A, \quad \lambda_7 = u + c_s \end{aligned} \quad (48)$$

Several standard problems in one and two dimensions are solved with the scheme *MOVERS-n* and results are presented here.

(a) Height Surface Plot



(b) Velocity Contours + Vector

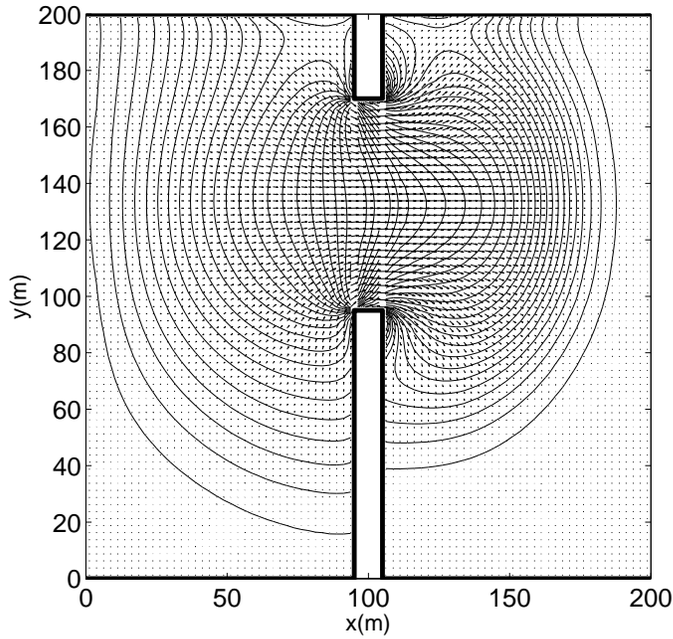


Fig. 21. Solution of partial dam break problem using first order accurate *MOVERS-n* at time 7.2 seconds with 40x40 points

4.4.1 (A) 1D Coplanar Problem

This 1D MHD problem with the initial Riemann conditions is given by [37]. The initial conditions $[\rho, u_1, u_2, p, B_y]$ on the left ($x \leq 0$) are $[1, 0, 0, 1, 1]$ and on right ($x \geq 0$) are $[0.125, 0, 0, 0.1, -1]$.

200 grid points are used in a domain of $[-50, 50]$. The CFL number is chosen as 0.75 and the results given are at $t = 10$ with $B_x = 0.75$ and $\gamma = 2$. The results are shown for the domain $[-30:30]$ in figures (24, 25) in comparison with LLF

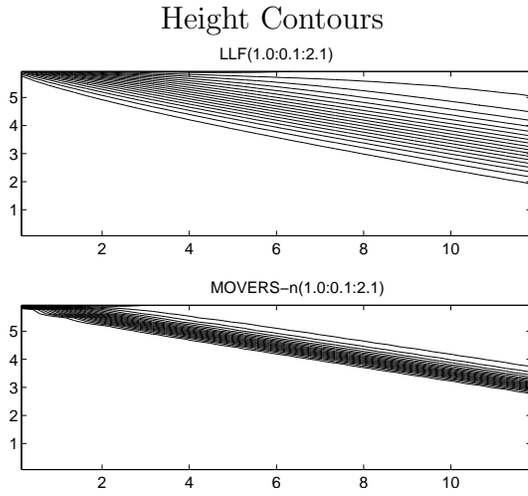


Fig. 22. Solution of oblique jump problem using first order accurate $MOVERS-n$ compared with LLF method

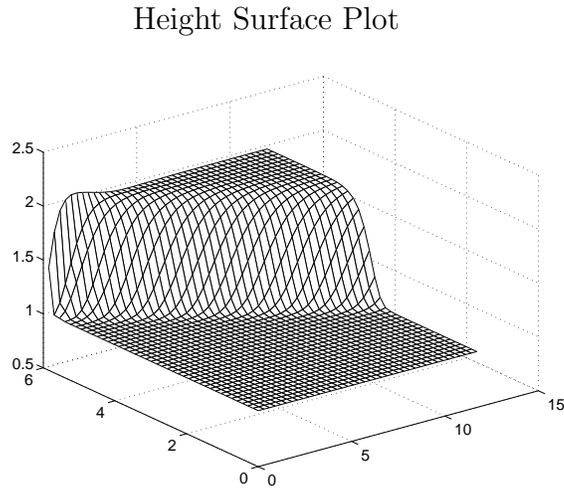


Fig. 23. Solution of oblique jump problem using first order accurate $MOVERS-n$ and HLLC [38], which is a Riemann solver and well known for its accuracy. The results here are deliberately given for a very coarse grid for this problem to compare the schemes. $MOVERS-n$ is seen to capture the discontinuities much better than LLF while it improves even over the HLLC scheme, although being simpler. When the grids are made finer with 800 points and the results are shown in the figure 26 where the results are in close agreement with HLLC Riemann solver.

4.4.2 (B) Riemann problem with Seven Discontinuities

This is a one dimensional problem which involves all the three components of velocity and magnetic field. It was suggested by Ryu *et al.* [39]. The spatial interval considered is $(-1, 1)$ on the x -axis. The initial conditions

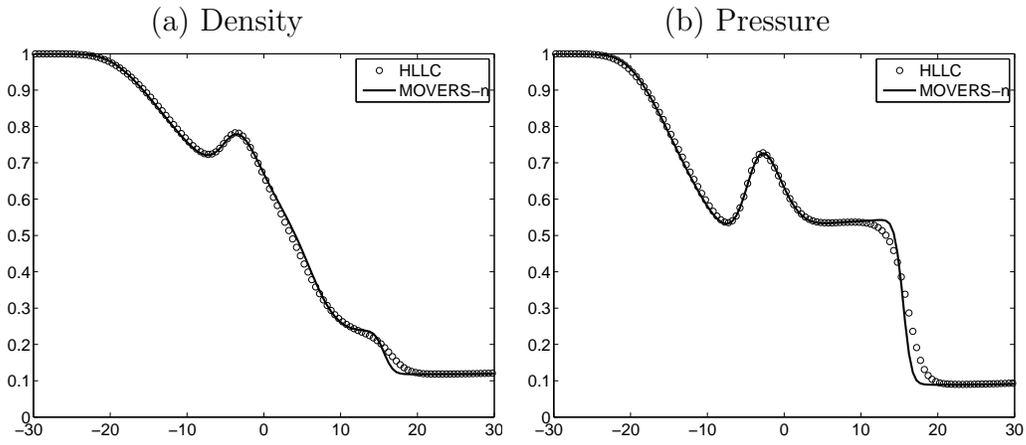


Fig. 24. Numerical Simulation of Ideal MHD problem (A) with I order accurate *MOVERS-n* and 200 points at time=10

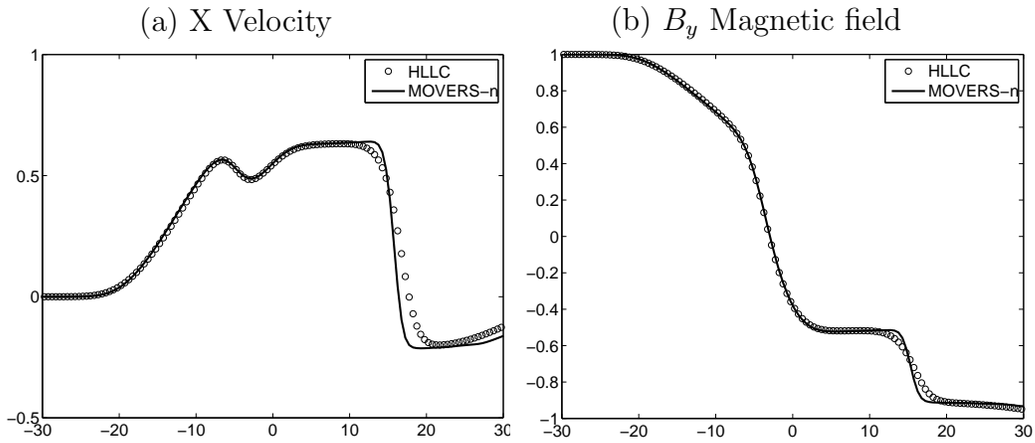


Fig. 25. Numerical Simulation of Ideal MHD problem (A) with I order accurate *MOVERS-n* and 200 points at time 10

$[\rho, u_1, u_2, u_3, p, B_y, B_z]$ on the left ($x \leq 0$) are $[1.08, 1.2, 0.01, 0.5, 0.95, 3.6, 2.0]$ and on the right ($x \geq 0$) are $[1, 0, 0, 0, 1, 4, 2]$.

Fast shocks, rotational discontinuities and slow shocks propagate from each side of the contact discontinuity. The rotation of the magnetic field across the initial discontinuity generates two rotational discontinuities. All the above features are seen to be captured by the scheme using 500 points (figures 27,28) with the solution falling close to a HLLC scheme but is much better than LLF method (not shown). In this problem, due to the complex interaction of multiple waves, more amount of numerical diffusion is used for entropy fix. The *MOVERS-n* predictions merge with that of HLLC with fine grid of 1500 points (29).

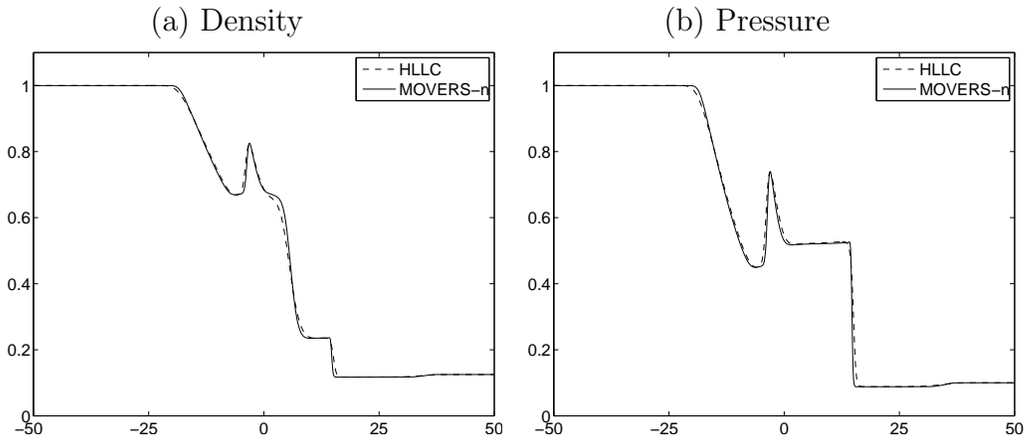


Fig. 26. Numerical Simulation of Ideal MHD problem (A) with I order accurate *MOVERS-n* on 800 points at time 10

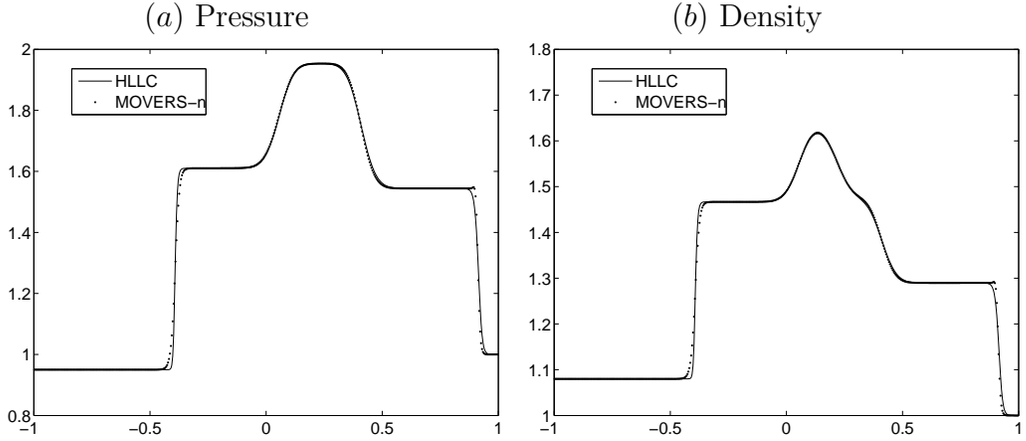


Fig. 27. Numerical Simulation of Ideal MHD problem (B) with I order accurate *MOVERS-n* at time 0.4 on 500 points

4.4.3 (C) Orszag Tang Vortex Problem

This is a problem of evolution of a compressible vortex system [40] involving interaction between several shock waves traveling at different speeds. The computational domain is $[0, 2\pi] \times [0, 2\pi]$ with periodic boundary conditions in both x and y directions using a uniform grid of 288×288 points. The initial conditions are

$$\begin{aligned} \rho(x, y, 0) &= \gamma^2, & p(x, y, 0) &= \gamma, & v_x(x, y, 0) &= -\sin y \\ v_y(x, y, 0) &= \sin x, & B_x(x, y, 0) &= -\sin y, & B_y(x, y, 0) &= \sin 2x \end{aligned}$$

Value of γ is taken as $5/3$.

The solutions for 1-D ideal MHD equations shown in figures (24-28) are com-

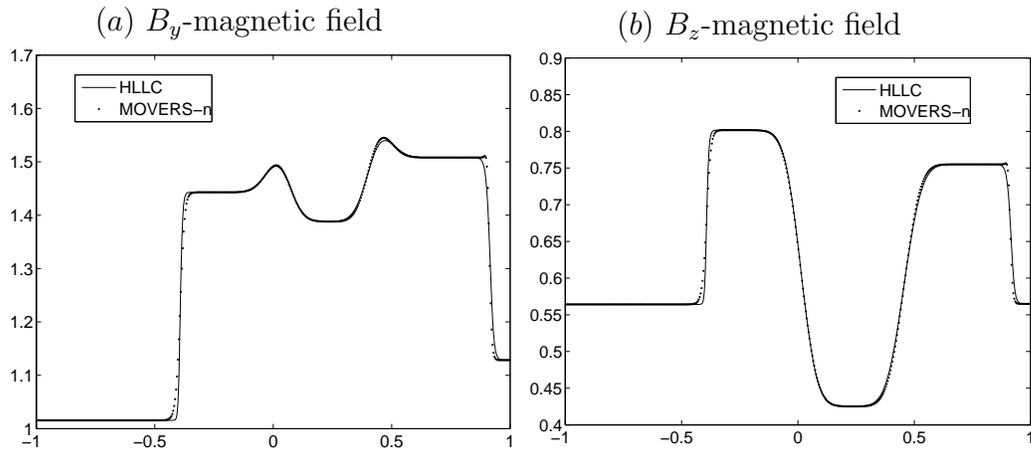


Fig. 28. Numerical Simulation of Ideal MHD problem (B) with I order accurate *MOVERS-n* at time 0.4 on 500 points

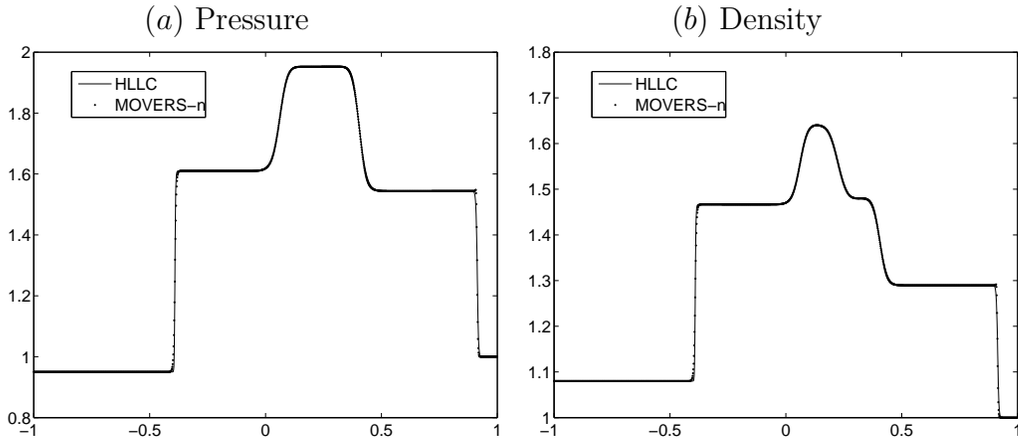


Fig. 29. Numerical Simulation of Ideal MHD problem (B) with I order accurate *MOVERS-n* at time 0.4 on 1500 points

pared with *Local Lax Friedrichs Scheme* (not shown) which is very dissipative and the *HLLC Solver* which is known to be very accurate. The solutions obtained with *MOVERS-n* are, as expected, comparable with the HLLC solutions although the former scheme is much simpler and free of Riemann solvers. The solution for 2D-ideal MHD equations for the Orszag Tang Vortex System in figures (30-33) are very accurate and are comparable with reported results [40].

4.5 Results with *MOVERS-1* for Euler equations

The solutions for 1-D steady contact discontinuity problem, Sod's shock tube test case, quasi 1-d nozzle flow, supersonic flow over a step in a wind tunnel and hypersonic flow across half cylinder are given below. The solutions

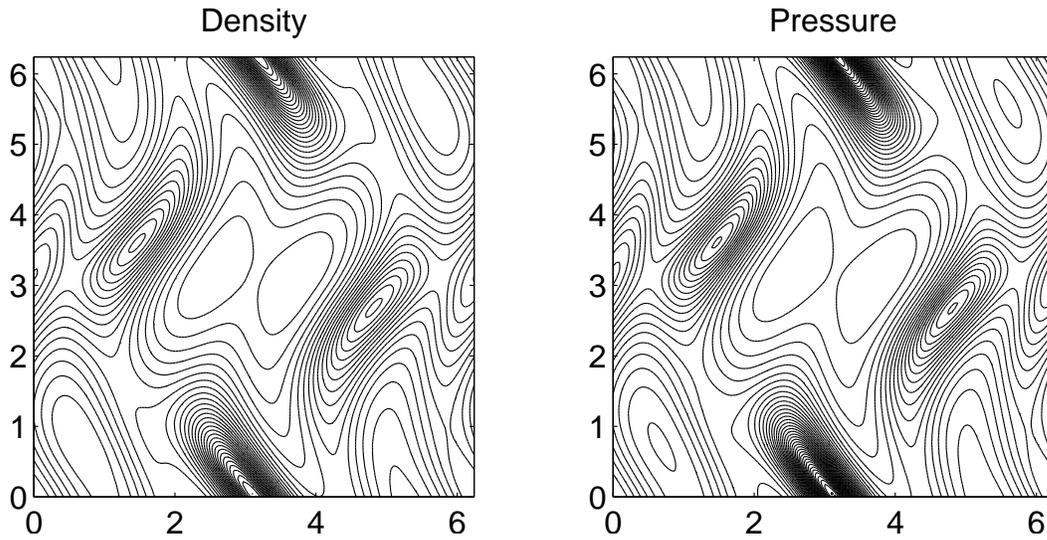


Fig. 30. Numerical Simulation of Orszag-Tang Vortex System with first order accurate *MOVERS-n* at time=1

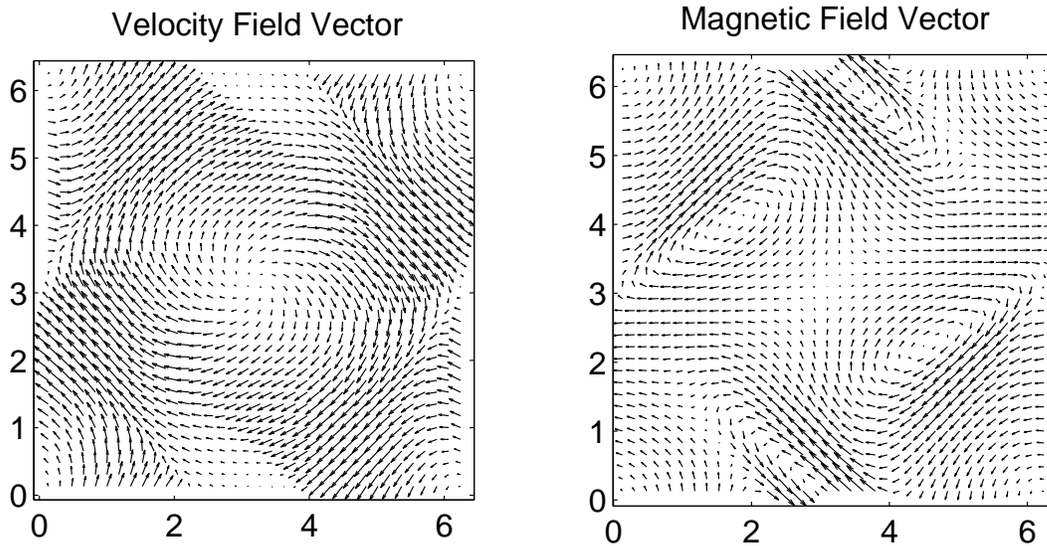


Fig. 31. Numerical Simulation of Orszag-Tang Vortex System with first order accurate *MOVERS-n* at time=1

are as good as *MOVERS-n* solutions or even better than some of them. As mentioned before, the choice of one of the three wave speeds (corresponding to the energy equation and which uses the maximum possible information) is made based on reasons mentioned in section (3.2). The shock solution of the quasi-1d nozzle flow consistently contains a maximum of only two points which is currently possible only by highly accurate Riemann solvers such as Roe scheme when only first order scheme is used. The solution for shear flow

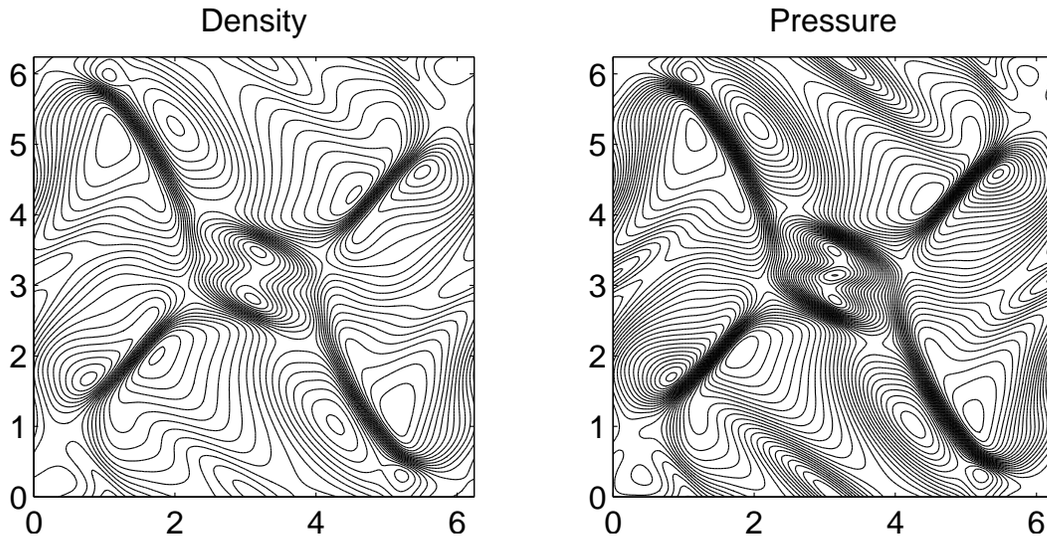


Fig. 32. Numerical Simulation of Orszag-Tang Vortex System with first order accurate *MOVERS-n* at time=3

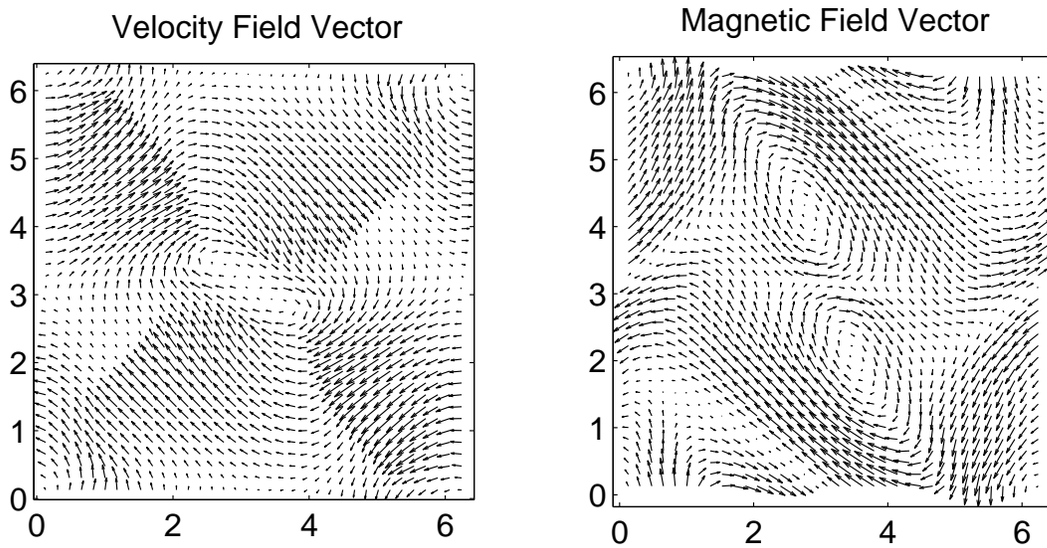


Fig. 33. Numerical Simulation of Orszag-Tang Vortex System with first order accurate *MOVERS-n* at time=3

problem with *MOVERS-1 variant* is almost identical to that of *n-variant* (figure 7) and hence is not reproduced here. The scheme is used to solve flow over a forward-facing step in wind tunnel and the slip lines starting at the Mach stem from right above the step are clearly seen in figure (36). The computation with unstructured mesh is also found accurate as evident in the solution for hypersonic flow across a half cylinder presented in figure (37).

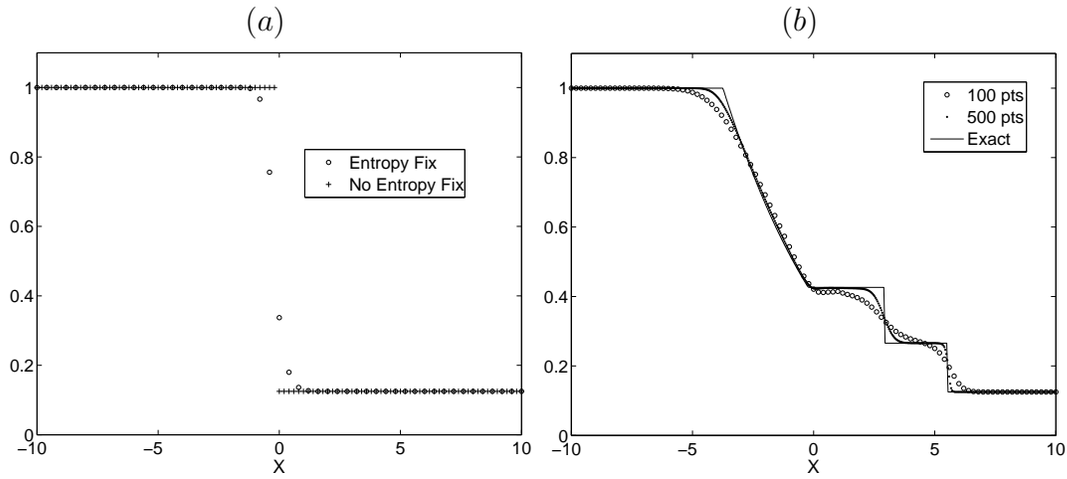


Fig. 34. Density plot with first order accurate *MOVERS-1* for (a) Steady 1D contact (b) Sod's 1D shock tube problem

It is found that the residue for the second order solution does not fall to very low values for this version of *MOVERS*. However, instead of calculating the wave speed at the interface directly with the immediate neighboring points as in equation (33), if the wave speeds are calculated at every cell center [18] as

$$s_j = \begin{cases} \frac{F_{j+1} - F_{j-1}}{U_{j+1} - U_{j-1}} & \text{if } U_{j-1} \neq U_{j+1} \\ \lambda_{min,j} & \text{if } U_{j-1} = U_{j+1} \end{cases} \quad (49)$$

$$s_j = \text{sign}(s_j) \lambda_{max,j} \quad \text{if } |s_j| \geq \lambda_{max,j} \quad (50)$$

$$s_j = \text{sign}(s_j) \lambda_{min,j} \quad \text{if } |s_j| \leq \lambda_{min,j} \quad (51)$$

and the interface wave speed is then taken to be the maxima of left and right states, the residual of the second order accurate scheme falls to much lower values. With such a change in computing the interface flux, the solutions agree closely with the parent methods *MOVERS-(n,1)* presented and hence only the convergence plots are given here in figure (38).

4.6 Results with *MOVERS-L*

The results for *MOVERS-L*, a limiter based Central Rankine-Hugoniot Solver in which the dissipation switches from the one in the *MOVERS-1(or n)* to that in the *LLF* method, obtained with a *minmod Limiter*, are presented in figures from 39 to 43. Steady discontinuities are resolved accurately and all features of the flow are captured well, as can be seen from the results. Note

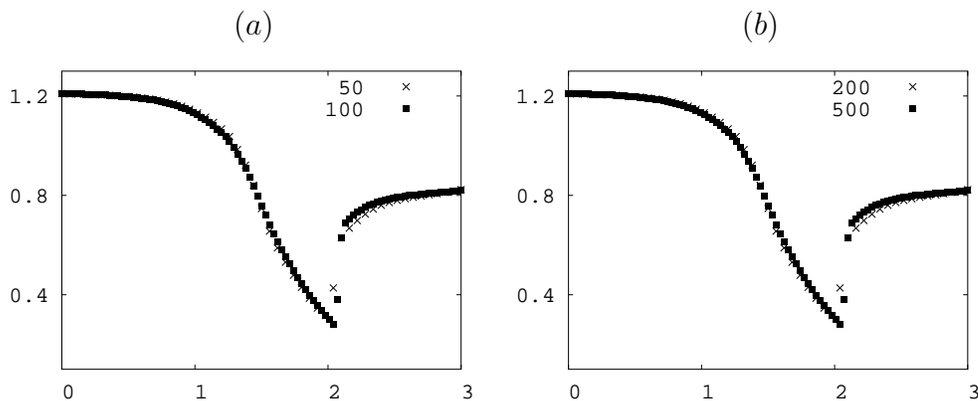


Fig. 35. Density plot with first order accurate *MOVERS-1* for Quasi 1-D nozzle flow (a) 50,100 points (b) 200,500 points

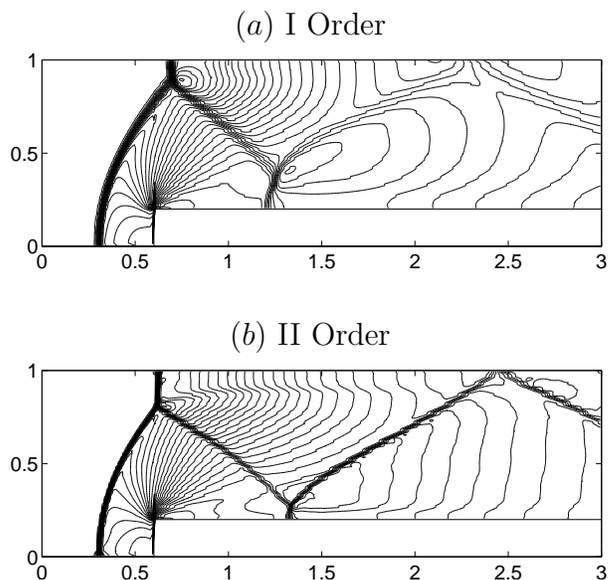


Fig. 36. Supersonic flow over forward-facing a step in channel : Density contours with *MOVERS-1* on 240x80 grid

that the high resolution is achieved here without the use of an entropy fix, based only on the flow variables.

5 MOVERS and Upwind Interpretation

The results presented in the previous sections show that the *Central Rankine Hugoniot Solver* developed in this work, called the *Method of Optimal Viscosity for Enhanced Resolution of Shocks* (*MOVERS*), is a very efficient scheme, capturing the steady discontinuities exactly when aligned with grid lines, and providing low diffusion solutions otherwise. The motivation with which this work started was to develop an accurate flow solver based on a simple central

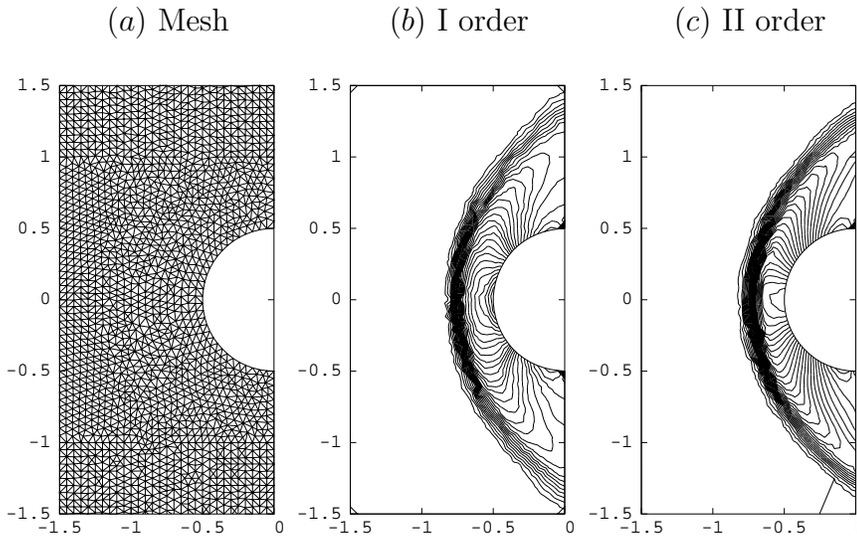


Fig. 37. Hypersonic ($M=6$) flow across a semi-cylinder : Pressure contours with *MOVERS-1* using unstructured mesh of 3897 points; (a) mesh (b) I order (c) II order

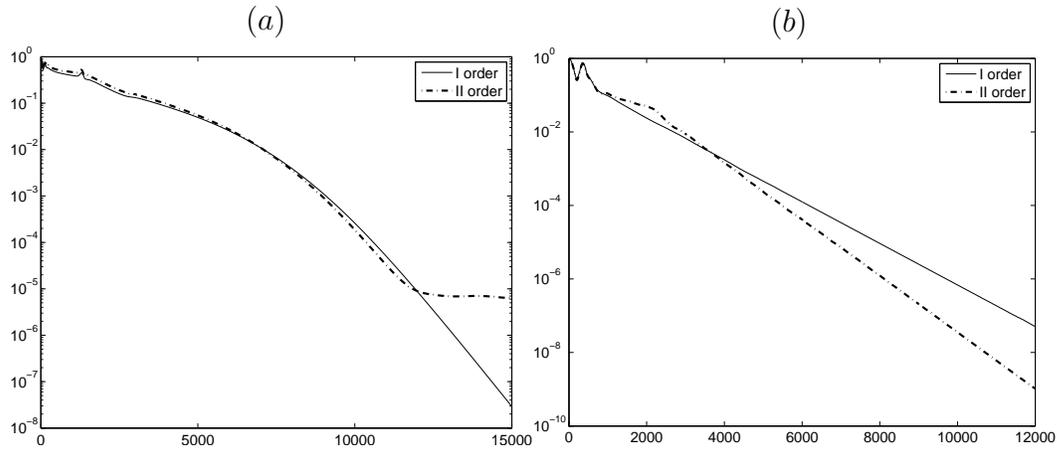


Fig. 38. Convergence History for *MOVERS-1* : (a) Supersonic flow ($M=2$) over a ramp in channel; (b) Hypersonic flow ($M=6$) across a semi-cylinder

discretization scheme, taking the Local Lax-Friedrichs method as the building block. It is worth noting that *MOVERS* can also be interpreted as an upwind scheme. An upwind interpretation may give a different perspective on this new algorithm and may be useful in modifying the existing upwind codes in a simple way in implementing *MOVERS*. In the following, an upwind interpretation for n -wave variant is presented. The cases of 1 -wave and L variants can be treated in a similar fashion.

Consider the case of a system of hyperbolic conservation laws in one dimension.

$$\frac{\partial U}{\partial t} + \frac{\partial F}{\partial x} = 0 \quad (52)$$

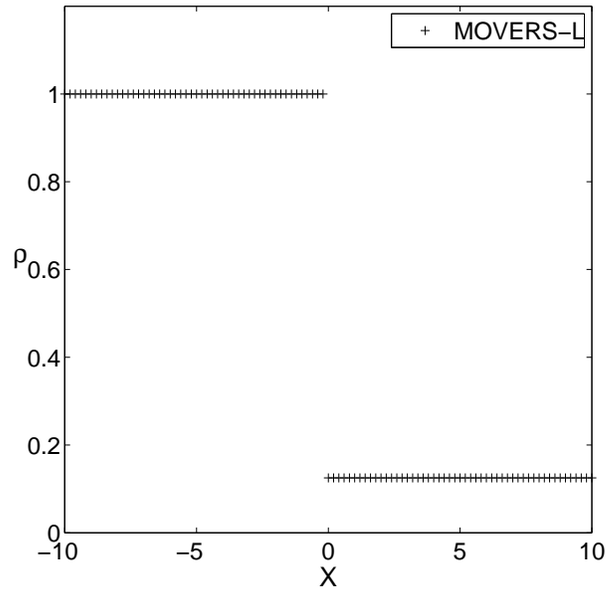


Fig. 39. Steady contact discontinuity in a shock tube with first order accurate *MOVERS-L*

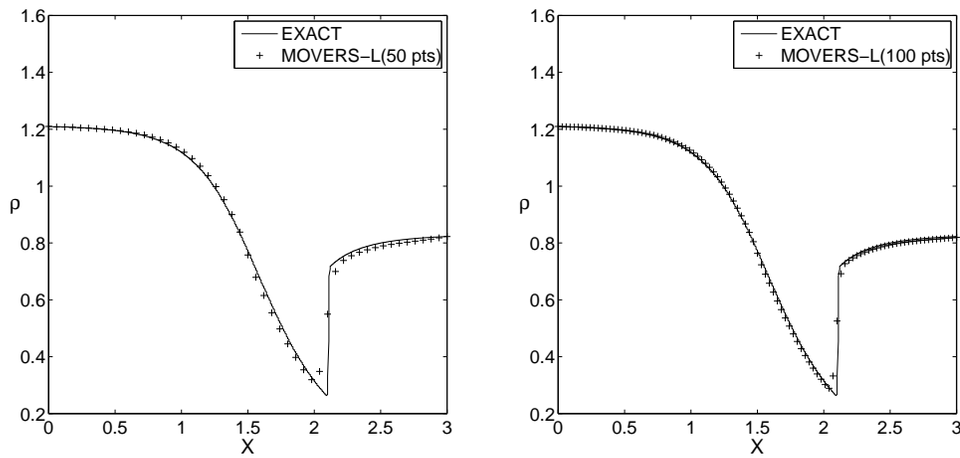


Fig. 40. Quasi 1-D nozzle flow with first order accurate *MOVERS-L*

Let us introduce a linearization of the flux vector as

$$F \approx \tilde{A}U + B \tag{53}$$

where \tilde{A} and B are constant matrices. We further simplify by assuming \tilde{A} to be a diagonal matrix.

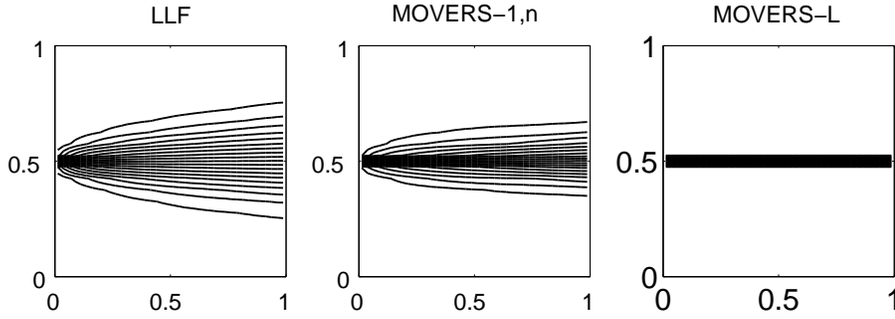


Fig. 41. Horizontal Shear flow (Mach 3 flow over Mach 2 flow), Mach contours with first order accurate *MOVERS-L* on 40x20 grid

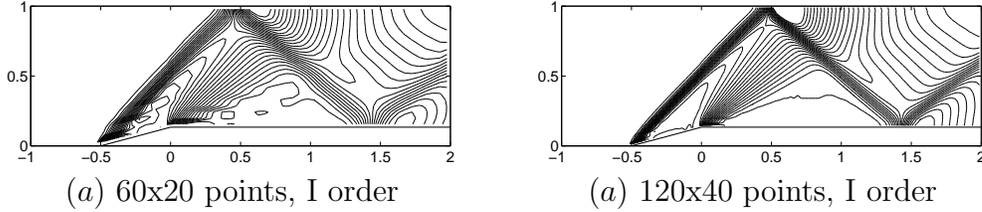


Fig. 42. Mach 2 flow over Ramp in channel: Density Contours with *MOVERS-L*

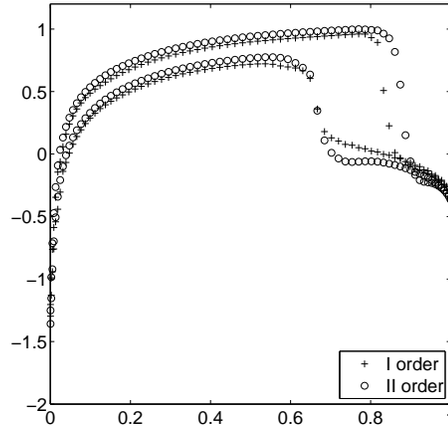


Fig. 43. Transonic flow across NACA0012 airfoil: Pressure Co-efficient Distribution obtained with *MOVERS-L* using O-type mesh with 160x60 points

$$\tilde{A} = \begin{bmatrix} \tilde{\lambda}_1 & 0 & \cdots & 0 \\ 0 & \tilde{\lambda}_2 & \cdots & 0 \\ \vdots & \vdots & \cdots & \vdots \\ 0 & 0 & \cdots & \tilde{\lambda}_n \end{bmatrix} \quad (54)$$

Then the system of conservation laws can be rewritten as

$$\frac{\partial U}{\partial t} + \frac{\partial (\tilde{A}U + B)}{\partial x} = \frac{\partial U}{\partial t} + \tilde{A} \frac{\partial U}{\partial x} = 0 \quad (55)$$

which is the approximate Riemann problem being solved at the cell interfaces, with the initial conditions as

$$U(x, t = 0) = \begin{cases} U_L \\ U_R \end{cases} \quad (56)$$

Expanding, we obtain

$$\frac{\partial U_m}{\partial t} + \tilde{\lambda}_m \frac{\partial U_m}{\partial x} = 0, \quad m = 1, \dots, n \quad (57)$$

The approximate Riemann problem consists of n linear convection equations. The n wave speeds need to be constants as per the linearization assumption made and this requirement can be satisfied by taking them as local averages (not necessarily arithmetic averages), across a cell interface. Now consider the flux Jacobian matrix for the original conservation laws, given by

$$A = \frac{\partial F}{\partial U} \quad (58)$$

which has n eigenvalues, $\lambda_1, \dots, \lambda_n$. If the physical situation presents a shock wave, then relevant wave speed will be the shock speed, defined by the Rankine-Hugoniot condition as

$$\Delta F = s \Delta U \quad (59)$$

Since our aim is to enhance resolution of the shocks, we approximate the wave speeds by relevant speeds of the shock coming from the jump condition as

$$\tilde{\lambda}_m = |s_{m,I}|, \quad m = 1, \dots, n \quad (60)$$

leading to

$$\tilde{A} = \begin{bmatrix} |s_{1,I}| & 0 & \cdots & 0 \\ 0 & |s_{2,I}| & \cdots & 0 \\ \vdots & \vdots & \cdots & \vdots \\ 0 & 0 & \cdots & |s_{n,I}| \end{bmatrix} \quad (61)$$

Here, the subscript I denotes the cell interface. Using the notations $\Delta U = U_R - U_L$ and $\Delta F = F_R - F_L$, we can now use the approximate Flux Jacobian

in the relation

$$\Delta F = \tilde{A} \Delta U \quad (62)$$

and the connection to *MOVERS-n* has now become apparent, if we use

$$s_{m,I} = \frac{\Delta F_m}{\Delta U_m}, m = 1, \dots, n \quad (63)$$

with the wave speed corrections introduced before.

If the evaluated wave speeds are positive

$$|s_{m,I}| = s_{m,I}, m = 1, \dots, n \quad (64)$$

Then $\tilde{A} = \tilde{A}$ and the cell interface flux becomes

$$\begin{aligned} F_I &= \frac{1}{2} (F_L + F_R) - \frac{1}{2} \tilde{A} (U_R - U_L) \\ &= \frac{1}{2} (F_L + F_R) - \frac{1}{2} (F_R - F_L) \\ &= F_L \end{aligned} \quad (65)$$

which is the upwind flux for positive wave speeds. If the evaluated wave speeds are negative

$$|s_{m,I}| = -s_{m,I}, m = 1, \dots, n \quad (66)$$

Then

$$\tilde{A} = -\tilde{A} \quad (67)$$

and the cell interface flux becomes

$$\begin{aligned} F_I &= \frac{1}{2} (F_L + F_R) + \frac{1}{2} \tilde{A} (U_R - U_L) \\ &= \frac{1}{2} (F_L + F_R) + \frac{1}{2} (F_R - F_L) \\ &= F_R \end{aligned} \quad (68)$$

which is again the upwind flux for negative wave speeds. Thus, *MOVERS-n* is also an upwind scheme, similar to the Flux Difference Splitting method of Roe, but the expressions are simpler and are not based on Roe averages. Then,

this scheme can be interpreted as an upwind method with the wave speeds being evaluated based on the Rankine-Hugoniot jump condition, with wave speed corrections. The interpretation of *MOVERS-1* (and *L*) can be done in a similar way, noting that *variant L* does not need an entropy fix and is a combination of *LLF* and *variant 1 or n*. It is also important to note that the upwind interpretation of *MOVERS-1* is possible only if we take a very broad outlook, like treating the Local Lax-Friedrichs method also as upwind scheme. To go beyond such interpretations, it is important to note that *simplicity* and *accuracy* in capturing flow features like discontinuities are worthy goals to strive for in developing new algorithms, and this is the motive behind the work presented in this work.

With the above interpretation, the central Rankine-Hugoniot solvers developed in this work (*MOVERS-n* and its variants 1 and L) can be thought of as an advancement over both the existing central discretization methods (in capturing the steady discontinuities exactly) and upwind methods (leading to simplification and being equally accurate and may also be thought of as uniting the two frameworks).

6 Conclusion

An accurate numerical solver for hyperbolic partial differential equations is presented here. The new solver is in the framework of central schemes and is based on the direct discretization of Rankine-Hugoniot condition. The solver is simple and easily adoptable for any hyperbolic system. For hyperbolic vector conservation laws, three different variants of the scheme are proposed, *viz.*, *MOVERS-(n,1 and L)*. *MOVERS-n* is based on the propagation of *n*-waves while *MOVERS-1* is based on an approximation of a single wave, for the shock speed in the jump condition, to represent the coefficient of numerical dissipation. *MOVERS-L* is a hybrid limiter based solver smoothly varying between Local Lax-Friedrichs method and the *MOVERS* based on the jump condition, thus avoiding the need for an entropy fix. These different variants of the scheme are used to solve various bench-mark test cases of gas dynamics, shallow water flows and ideal magneto-hydrodynamics. The results show high accuracy in capturing flow features, with grid-aligned shock waves and contact discontinuities being resolved accurately. The variants of *MOVERS* can also be interpreted as upwind methods but are simpler, with retaining the accuracy in shock capturing, and one of the variants avoids use of any entropy fix due to the flow based adaptation of the numerical dissipation.

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