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Well-balanced high order 1D schemes on non-uniform grids and entropy residuals

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Abstract This paper is concerned with the construction of high order schemes on irregular grids for balance laws, including a discussion of an a-posteriori error indicator based on the numerical entropy production. We also impose well-balancing on non uniform grids for the shallow water equations, which can be extended similarly to other balance laws, obtaining schemes up to fourth order of accuracy with very weak assumptions on the regularity of the grid. Our results show the expected convergence rates, the correct propagation of shocks across grid discontinuities and demonstrate the improved resolution achieved with a locally refined non-uniform grid. The schemes proposed in this work naturally can also be applied to systems of conservation laws. They may also be extended to higher space dimensions by means of dimensional splitting.

The error indicator based on the numerical entropy production, previously introduced for the case of systems of conservation laws, is extended to balance laws. Its decay rate and its ability to identify discontinuities is illustrated on several tests.

 $\mathbf{Keywords} \hspace{0.1 cm} \text{high order finite volumes} \cdot \text{non-uniform grids} \cdot \text{entropy} \cdot \text{well balancing}$

Mathematics Subject Classification (2000) 65M08 · 76M12

1 Introduction

Many problems arising from engineering applications involve the ability to compute flow fields on complex domains, governed by hyperbolic systems of balance laws. Often, many scales are involved and this prompts the need for algorithms that are able to modify the scheme and/or the underlying grid following the evolution of the flow. Several wide purpose codes are available and many of them are

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based on finite volume schemes, see e.g. Fluent [14] or ClawPack [8]. Usually these codes are second order accurate with high order versions, if available, in progress. On the other hand they provide the user with the flexibility of an adaptive grid, which is extremely useful to tackle highly non-homogeneous solutions.

At the same time, high order finite volume schemes are well established in the literature: from the early review in [36] to the more recent paper [12], extensive studies have been conducted on the construction of high order finite volume schemes. In this paper we carry out a detailed study of the issues arising in finite volume algorithms on irregular grids. Non-uniform grids of the type analyzed in this work arise naturally in several applications, already in one space dimension. We mention the grids generated by moving mesh algorithms [37] or by an h-adaptive strategy [32] or the grids that arise when discretizing problems with moving boundaries [7]. In this last case, as for example in the piston problem [13], cells close to the boundary are created or destroyed and resized according to the location of the boundary in each timestep.

In particular we construct finite volume high order WENO schemes, including the treatment of source terms and addressing the issue of well balancing for steady state solutions. We concentrate on the one-dimensional case, since most problems already arise in this setting. These results can be naturally extended to multidimensional problems discretized with Cartesian grids by means of dimensional splitting. Schemes based on Cartesian grids can be easily parallelized and boundary conditions for complex domains can be implemented with the immersed boundary method as in [17].

Adaptive grids can be constructed either by defining a single non uniform grid on which all degrees of freedom are located, as in most unstructured grid managers, or superposing several patches of uniform Cartesian grids of different levels of refinement as in the ClawPack solver [8]. In this latter approach the different patches must communicate and the enforcement of conservativity and well balancing for steady states are not straighforward [11]. High order schemes for the AMR approach can be found in [2,34]. For applications to the shallow water equations, see the software GeoClaw [8] and [16].

In our case we consider a single highly non-uniform grid. Such grids commonly arise in h-adaptive methods [18], or also when using moving mesh methods [38,37]. In one space dimension, when the grid size varies smoothly, one can remap the problem to a uniform grid as in [13], but this cannot be expected to work in more space dimensions or when the grid size can jump abruptly as in dyadic/quadtree/octree grid refinement. These latter discretization techniques start from a conforming, often uniform, partitioning of the simulation domain and allow the local refinement of each control volume by splitting it in 2^d parts in dspace dimensions, as in [20] for simplices and [41] for quads. Lower order schemes on such grids were employed by the authors in [32] in one space dimension and in [33] in two space dimensions for general conservation laws. Two-dimensional applications to the shallow water system may be found in [28], or in [27].

The construction of a fifth order WENO scheme for conservation laws on onedimensional non-uniform grids, based on the superposition of three parabolas, has been conducted in [42]. Here we extend this construction to the case of balance laws, showing how to obtain positive coefficients in the quadrature of the source term. Moreover we also construct a third order scheme based on [26], characterized by a stencil of three cells. This reconstruction is particularly suited for two-dimensional problems due to its very compact stencil, see [9].

A first key ingredient of this work is the use of semidiscrete schemes which permit to decouple the space from the time discretization: in this fashion the nonuniformity of the grid boils down to an interpolation problem to reconstruct the boundary extrapolated data which interact through the numerical fluxes. Secondly, the use of the Richardson extrapolation as in [29] is crucial for the preservation of steady states on a non uniform grid, since it allows to enforce equilibrium at the level of each single cell, thus avoiding the need to account for the non-uniformity of the grid. This yields automatic well-balancing over the whole grid, unlike in the block-structured AMR case, where well-balancing has to be enforced not only on each grid patch but also in the projection and interpolation operators that relate the solution on different grid levels [11].

Moreover, we extend the entropy indicator of [32] to the case of balance laws. We show that the numerical entropy production provides a measure of the local error on the cell also in the case of balance laws on non-uniform grids. This complements the introduction of high order schemes on non-uniform grids: the entropy indicator can be used to drive an adaptive mesh refinement algorithm, but also as a monitor function in moving mesh algorithms. Moreover, the entropy can be used to locally modify a scheme in the presence of shocks, with p adaptive strategies. Other possible applications are discussed in [32].

Before giving the outline of the paper, we briefly introduce the setting and the notation used in the bulk of this work. We consider balance laws with a geometric source term of the form

$$u_t + \nabla \cdot f(u) = g(u, x) \tag{1}$$

and we seek the solution on a domain Ω , with given initial conditions. The computational domain Ω is an interval, discretized with cells $I_j = [x_{j-1/2}, x_{j+1/2}]$, such that $\cup I_j = \Omega$. The amplitude of each cell is $\delta_j = x_{j+1/2} - x_{j-1/2}$, with cell center $x_j = (x_{j-1/2} + x_{j+1/2})/2$.

We consider semidiscrete finite volume schemes and denote with $\overline{U}_j(t)$ the cell average of the numerical solution in the cell I_j at time t. The semidiscrete numerical scheme can be written as

$$\frac{\mathrm{d}}{\mathrm{d}t}\overline{U}_j = -\frac{1}{\delta_j}\left(F_{j+1/2} - F_{j-1/2}\right) + G_j(\overline{U}, x).$$
(2)

The numerical fluxes are computed starting from the boundary extrapolated data, namely

$$F_{j+1/2} = \mathcal{F}(U_{j+1/2}^{-}, U_{j+1/2}^{+}), \tag{3}$$

where \mathcal{F} is a consistent and monotone numerical flux, evaluated on two estimates of the solution at the cell interface $U_{j+1/2}^{\pm}$. These values are obtained with a high order non oscillatory reconstruction, as described in detail in §2. Finally, G_j is a consistently accurate discretization of the cell average of the source term on the cell I_j , see §3.

In order to obtain a fully discrete scheme, we apply a Runge-Kutta method with Butcher's tableau (A, b), obtaining the evolution equation for the cell averages

$$\overline{U}_{j}^{n+1} = \overline{U}_{j}^{n} - \frac{\Delta t}{\delta_{j}} \sum_{i=1}^{s} b_{i} \left(F_{j+1/2}^{(i)} - F_{j-1/2}^{(i)} \right) + \Delta t \sum_{i=1}^{s} b_{i} G_{j}^{(i)}.$$
 (4)

Here $F_{j+1/2}^{(i)} = \mathcal{F}\left(U_{j+1/2}^{(i),-}, U_{j+1/2}^{(i),+}\right)$ and the boundary extrapolated data $U_{j+1/2}^{(i),\pm}$ are computed from the stage values of the cell averages

$$\overline{U}_{j}^{(i)} = \overline{U}_{j}^{n} - \frac{\Delta t}{\delta_{j}} \sum_{k=1}^{i-1} a_{ik} \left(F_{j+1/2}^{(k)} - F_{j-1/2}^{(k)} \right) + \Delta t \sum_{k=1}^{i-1} a_{ik} G_{j}^{(k)}$$

We point out that the spatial reconstruction procedures of §2 and the well-balanced quadratures for the source term of §3 must be applied for each stage value of the Runge-Kutta scheme. In this paper we consider a uniform timestep over the whole grid. A local timestep keeping a fixed CFL number over the grid can be enforced using techniques from [32, 22].

We will also consider the preservation of steady state solutions and we will illustrate these techniques on the shallow water system, namely

$$u = \begin{pmatrix} h \\ q \end{pmatrix} \qquad f(u) = \begin{pmatrix} q \\ q^2/h + \frac{1}{2}gh^2 \end{pmatrix} \qquad g(u, x) = \begin{pmatrix} 0 \\ -ghz_x \end{pmatrix}.$$
 (5)

Here *h* denotes the water height, *q* is the discharge and z(x) the bottom topography, while *g* is the gravitational constant (see also Figure 4). The preservation of steady states depends heavily on the structure of the equilibrium solution one wishes to preserve. Here we will concentrate on the lake at rest solution of the shallow water equation, given by H(t,x) = h(t,x) + z(x) = constant and q(t,x) = 0. Many works have been dedicated to this problem since the paper [3] shed light on the importance of well-balancing (or C-property). For example, see [44] in the finite difference setting, [45,29,30] in the finite volume setting, [45,43,5] in the Discontinuous Galerkin framework and [40,6] for the ADER methods.

The structure of the paper is as follows: in §2 we introduce the third order accurate C-WENO (Compact WENO) reconstruction on non uniform grids, generalizing the results of [26], and we extend the fifth order accurate WENO reconstruction on non uniform grids of [42], adding the evaluation of the reconstruction at the centre of cells which is needed in the computation of the source term. In §3 we extend the construction of well-balanced schemes of [1,29] to the non-uniform grid setting. Next, in §4 we extend the notion of numerical entropy production to non uniform grids for balance laws. Finally, §5 contains numerical tests, which illustrate the consistency between accuracy of the schemes and rate of convergence of the numerical entropy production, for several types of grids.

2 High order reconstructions on non uniform grids

The mission of reconstruction algorithms is to give estimates of a function at some chosen points, starting from discrete data. In particular, for finite volume schemes for balance laws, the starting data are the cell averages of a function v, and we wish to estimate v at the cell interfaces, and, if needed, at some other internal points, using a finite dimensional approximation, such as a piecewise polynomial interpolator. Typically, estimates of v at internal points within a cell are needed to compute the cell averages of the source term through a quadrature formula. Thus, the reconstruction will be described as an interpolation algorithm.

Suppose then that we are given the cell averages

$$\overline{V}_j = \frac{1}{\delta_j} \int_{I_j} v(x) \, \mathrm{d}x$$

of a smooth function v(x). In order to fix ideas, we consider a piecewise polynomial reconstruction \mathcal{R} such that

$$\mathcal{R}(\overline{V}, x) = \sum_{j} \chi_{I_j}(x) P_j(x),$$

which gives the boundary extrapolated data as

$$V_{j+1/2}^- = P_j(x_{j+1/2}), \qquad V_{j+1/2}^+ = P_{j+1}(x_{j+1/2}).$$
 (6)

The reconstruction must be conservative, i.e.

$$\frac{1}{\delta_j} \int_{I_j} \mathcal{R}(\overline{V}, x) \, \mathrm{d}x = \overline{V}_j,$$

and high order accurate at the cell interfaces for smooth data, in the sense that

$$V_{j+1/2}^{\pm} = v(x_{j+1/2}) + O(\delta_j^p), \qquad V_{j-1/2}^{\pm} = v(x_{j-1/2}) + O(\delta_j^p).$$

Moreover, the reconstruction should be non-oscillatory, preventing the onset of spurious oscillations. Finally, for accuracy of order higher than 2, the evaluation of the cell average of the source term requires the reconstruction of the point values of v at the nodes of the well-balanced quadrature formula. For schemes of order 3 and 4, it is enough to reconstruct v at the cell centers, thus we will require that, for smooth v(x),

$$V_j = v(x_j) + O(\delta_j^p).$$

$First \ order \ reconstruction$

In this case, the reconstruction is piecewise constant, and we have

$$V_{j+1/2}^- = \overline{V}_j, \qquad V_{j-1/2}^+ = \overline{V}_j.$$

 $Second \ order \ reconstruction$

Here, the reconstruction is piecewise linear, and we have

$$V_{j+1/2}^- = \overline{V}_j + \frac{1}{2}\sigma_j\delta_j, \qquad V_{j-1/2}^+ = \overline{V}_j - \frac{1}{2}\sigma_j\delta_j,$$

where σ_j is a limited slope, i.e., chosen a limiter Φ , define the interface slopes as

$$\sigma_{j+1/2} = \frac{\overline{V}_{j+1} - \overline{V}_j}{x_{j+1} - x_j} = \frac{\overline{V}_{j+1} - \overline{V}_j}{\frac{1}{2}(\delta_j + \delta_{j+1})},\tag{7}$$

then the limited slope within the I_j cell is given by

$$\sigma_j = \Phi\left(\sigma_{j-1/2}, \sigma_{j+1/2}\right).$$

For a collection of limiting functions, see [24]. In our tests, we have chosen the MinMod limiter.

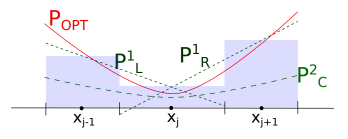


Fig. 1 Compact WENO reconstruction

Third order reconstruction

The third order reconstruction is based on the compact WENO (C-WENO) technique introduced in [26]. This reconstruction is characterized by a particularly compact stencil, which is very important when dealing with adaptive grids. Moreover, unlike the classical WENO third order reconstruction based on the combination of two linear functions, the C-WENO reconstruction contains also a parabola and it remains uniformly third order accurate throughout the interval I_j on smooth flows. To our knowledge, the reconstruction presented here is the first extension of the C-WENO reconstruction to the case of non-uniform grids. Fig. 1 illustrates the polynomials composing this reconstruction.

The interpolant is piecewise quadratic, and the parabola reconstructed in each cell is the convex combination of two linear functions P_L^1 , P_R^1 , and a parabola, P_C^2 . In order to simplify the notation we describe the reconstruction on a reference cell, labelled with the index j = 0. The two linear functions interpolate v in the sense of cell averages on the stencils $\{I_{-1}, I_0\}$ and $\{I_0, I_{+1}\}$. Each of these functions approximates v with order $O(\delta_0^2)$ accuracy uniformly on I_0 . Further, the parabola P_{OPT}^2 is introduced by the requirement that

$$\frac{1}{\delta_0} \int_{I_0} P_{\text{OPT}}^2(x) \, \mathrm{d}x = \overline{V}_0, \qquad \frac{1}{\delta_{\pm 1}} \int_{I_{\pm 1}} P_{\text{OPT}}^2(x) \, \mathrm{d}x = \overline{V}_{\pm 1}$$

This parabola approximates v with order $O(\delta_0^3)$ accuracy uniformly on I_0 . Next, the parabola P_C^2 is introduced, defined as

$$P_{\rm OPT}^2 = \alpha_0 P_C^2 + \alpha_{+1} P_R^1 + \alpha_{-1} P_L^1$$

with $\alpha_0 = \frac{1}{2}$, $\alpha_{\pm 1} = \frac{1}{4}$. The reconstruction is given by

$$P^{2}(x) = \omega_{0}P_{C}^{2} + \omega_{+1}P_{R}^{1} + \omega_{-1}P_{L}^{1}$$

When the function v is smooth, one would like that $\omega_k = \alpha_k + O(\delta_0^2)$, to ensure that P^2 has the same accuracy of P_{OPT}^2 , otherwise, the non linear weights ω_k are designed to switch on only the contribution coming from the one-sided stencil on which the function is smooth.

For a non-uniform grid, the coefficients of the two linear interpolants on the cell ${\cal I}_0$ are

$$P_R^1(x) = \overline{V}_0 + \sigma_{+1/2}(x - x_0),$$

$$P_L^1(x) = \overline{V}_0 + \sigma_{-1/2}(x - x_0),$$

where $\sigma_{\pm 1/2}$ have been defined in (7). The optimal parabola is

$$\begin{split} P_{\rm OPT}^2 &= a + b(x - x_0) + c(x - x_0)^2, \\ c &= \frac{3}{2} \frac{\sigma_{+1/2} - \sigma_{-1/2}}{\delta_{-1} + \delta_0 + \delta_{+1}}, \\ b &= \frac{(\delta_0 + 2\delta_{-1})\sigma_{+1/2} + (\delta_0 + 2\delta_{+1})\sigma_{-1/2}}{2(\delta_{-1} + \delta_0 + \delta_{+1})} \\ a &= \overline{V}_0 - \frac{1}{12} c \, \delta_0^2. \end{split}$$

As in WENO-like reconstructions, the non linear weights ω_k are computed as

$$\tilde{\omega}_k = \frac{\alpha_k}{(\epsilon + \mathrm{IS}_k)^2}, \qquad \omega_k = \frac{\tilde{\omega}_k}{\sum_{l=-1}^1 \tilde{\omega}_l}$$

starting from the smoothness indicators IS_k defined in [36]. In this case, they are given by

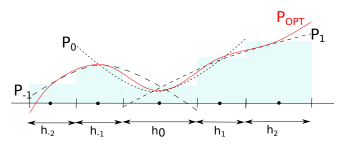
$$\begin{split} \mathrm{IS}_{-1} &= \delta_0^2 \sigma_{-1/2}^2, \\ \mathrm{IS}_1 &= \delta_0^2 \sigma_{+1/2}^2, \\ \mathrm{IS}_0 &= \frac{1}{\alpha_0^2} \left[\left(b - \alpha_{-1} \sigma_{-1/2} - \alpha_{+1} \sigma_{+1/2} \right) \delta_0^2 + \frac{13}{3} c^2 \delta_0^4 \right]. \end{split}$$

Since P_{OPT}^2 is uniformly third order accurate on the whole interval, the boundary extrapolated data and the value V_0 at the cell center are all computed evaluating the same quadratic polynomial at the corresponding points inside the cell.

Fourth order reconstruction

The fourth order reconstruction is based on the fifth order WENO reconstruction computed from the convex combination of three parabolas, as in [36]. The coefficients of the combination of the three parabolas are computed in order to yield fifth order accuracy at the boundary of the cell, see Fig 2. It is tedious but straightforward to see that positive coefficients can be found to result in fifth order accuracy at the cell interfaces even on non uniform grids (see below and [42]). However, there is no set of positive coefficients resulting in fifth order accuracy at the cell center, see [29]. This issue has already prompted the development of the CWENO reconstruction for central schemes [25] and the generalization of the classical WENO procedure described in [35]. Here we show that it is possible to find three positive coefficients giving *fourth* order accuracy at the center of the cell, which is sufficient to construct a fourth order accurate scheme that employs the reconstruction at three quadrature nodes located at the boundary and at the center of the cells.

For the sake of completeness, we review the coefficients of the reconstruction on non uniform grids, as in [42], using the notation established in Fig. 2. Again we consider a reference cell with index 0. The goal of the reconstruction is to mimic the quartic polynomial $P_{\rm OPT}$ interpolating the data $\overline{V}_l, l = -2, \ldots, 2$ in the sense of cell averages. Clearly, $P_{\rm OPT}$ would provide fifth order accuracy uniformly in the interval I_0 , in the case of smooth data.



 $\mathbf{Fig. \ 2} \ \mathsf{Parabolic} \ \mathsf{WENO} \ \mathsf{reconstruction}$

For each point \hat{x} in which the reconstruction is needed, we look for three positive coefficients d_{-1}, d_0, d_1 that add up to 1 and such that

$$P_{\rm OPT}(\hat{x}) = \sum_{l=-1}^{1} d_l P_l(\hat{x}), \tag{8}$$

where the P_l 's are the three parabolas, interpolating in the sense of cell averages the data $\overline{V}_{l-1}, \overline{V}_l, \overline{V}_{l+1}$. The coefficients of the three parabolas can be found in [42]. Here we give the linear weights that permit to reconstruct the left and right boundary extrapolated data. To simplify the notation, we write

$$\delta_l^k = \sum_{i=l}^k \delta_i,\tag{9}$$

then the coefficients for the boundary extrapolated data $V^-_{\pm 1/2}$ are

$$d_{1} = \frac{\delta_{-1}(\delta_{-2} + \delta_{-1})}{\delta_{-2}^{2}\delta_{-1}^{2}},$$

$$d_{0} = \frac{\delta_{0}^{2}(\delta_{-2} + \delta_{-1})(\delta_{-2}^{1} + \delta_{-1}^{2})}{\delta_{-2}^{2}\delta_{-1}^{2}\delta_{-2}^{1}},$$

$$d_{-1} = \frac{\delta_{0}^{2}(\delta_{0} + \delta_{1})}{\delta_{-2}^{2}\delta_{-2}^{1}}.$$

Note that, if $\delta_{-2} = \delta_{-1} = \delta_0 = \delta_1 = \delta_2$, then $d_{-1} = \frac{3}{10}, d_0 = \frac{3}{5}, d_1 = \frac{1}{10}$, as in the usual uniform grid case. Similarly, the coefficients for the reconstruction of $V^+_{-1/2}$ are

$$d_{-1} = \frac{\delta_1(\delta_1 + \delta_2)}{\delta_{-2}^2 \delta_{-2}^1},$$

$$d_0 = \frac{\delta_{-2}^0(\delta_1 + \delta_2)(\delta_{-2}^1 + \delta_{-1}^2)}{\delta_{-2}^2 \delta_{-1}^2 \delta_{-2}^1},$$

$$d_1 = \frac{\delta_{-2}^0(\delta_{-1} + \delta_0)}{\delta_{-2}^2 \delta_{-1}^2}.$$

We remark that the coefficients d_k are positive and add up to 1, so that (8) is a convex combination, for all possible values of the local grid size $\delta_{-2}, \ldots, \delta_2$.

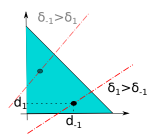


Fig. 3 Reconstruction of the point value in the cell center for WENO. Locus of positive linear weights (dash-dot lines) and the coefficients chosen by (10) (black dots).

For the fifth order reconstruction at cell center x_0 , one finds negative coefficients even for uniform meshes. In fact, see [29], $d_{-1} = -\frac{9}{80}$, $d_0 = \frac{49}{40}$, $d_1 = -\frac{9}{80}$. Since the well balanced quadrature based on the three points $x_{\pm 1/2}$, x_0 (see the next section) is only fourth order accurate, even a fifth order accurate reconstruction at the middle quadrature point would not give overall fifth order accuracy: the fourth order limitation is given by the quadrature formula and a more accurate quadrature formula would require the reconstruction at extra quadrature points, which would make the scheme much more complicated. Thus, we look for *positive* coefficients d_0 , $d_{\pm 1}$ such that $1 = \sum d_l$, and V_0 is fourth order accurate,

$$V_0 = \sum_{l=-1}^{1} d_l P_l(x_0) = v(x_0) + O(\delta_0^4).$$

After tedious computations, we find that d_1 and d_{-1} must satisfy

$$\delta_{-2}^1 d_{-1} - \delta_{-1}^2 d_1 = \delta_1 - \delta_{-1}.$$

Since we wish all coefficients to be positive, the solution must be sought in the simplex shown in Fig. 3. Clearly, the solution is over-determined, we pick the values that maximize the size of the minimum coefficient, that is

$$\operatorname{If} \delta_{1} > \delta_{-1} \begin{cases} d_{1} = \frac{1}{2} \frac{\delta_{-2} + 2\delta_{-1} + \delta_{0}}{\delta_{-2}^{1} + \delta_{-1}^{2}} \\ d_{-1} = \frac{\delta_{1} - \delta_{-1} + d_{1}\delta_{-1}^{2}}{\delta_{-2}^{1}} \\ d_{0} = 1 - d_{-1} - d_{1} \end{cases} \quad \text{else} \begin{cases} d_{-1} = \frac{1}{2} \frac{\delta_{2} + 2\delta_{1} + \delta_{0}}{\delta_{-2}^{1} + \delta_{-1}^{2}} \\ d_{1} = \frac{\delta_{-1} - \delta_{1} + d_{-1}\delta_{-2}^{1}}{\delta_{-1}^{2}} \\ d_{0} = 1 - d_{-1} - d_{1} \end{cases} \tag{10}$$

where again we have used the convention (9).

3 Well-balanced schemes

It is important to perform numerical integration of a system of balance laws with schemes that preserve the steady states exactly at a discrete level (well-balanceed schemes), since only these allow to distinguish small perturbations of these states from numerical noise [3].

In this section we describe a technique to obtain well-balanced schemes on non-uniform grids for the shallow water equations, with particular attention to the lake at rest solution. Recall that the equilibrium variables are the momentum q and the energy $E = \frac{q^2}{2h^2} + g(h + z)$. The lake at rest solution corresponds to the particular choice q = 0, for which the equilibrium energy E is stationary if and only if the total water height H = h + z remains constant. We use and generalize to non-uniform meshes the techniques of [1] for obtaining well-balanced schemes irrespectively of the chosen numerical fluxes and of [29] to obtain high order accuracy through Richardson extrapolation.

There are two sources of error in non well-balanced schemes. We illustrate them with a very simple example. We consider a first order reconstruction with the Lax-Friedrichs numerical flux on the lake at rest solution (see Fig. 4 for notation), thus we suppose that for every index j, $q_j^n = 0$ and $h_j^n + z_j = H$. The discretized equation on a uniform grid would be

$$h_{j}^{n+1} = h_{j}^{n} + \frac{\lambda}{2} \alpha \left(h_{j+1}^{n} - 2h_{j}^{n} + h_{j-1}^{n} \right),$$

$$q_{j}^{n+1} = -\frac{\lambda}{4} g \left((h_{j+1}^{n})^{2} - (h_{j-1}^{n})^{2} \right) + \frac{\lambda}{2} g h_{j}^{n} \left(z_{j+1} - z_{j-1} \right),$$

where we have already substituted $q_j^n = 0$. It is easy to see that in the first equation, h does not remain stationary because the artificial diffusion term introduces a perturbation whenever z(x) is not constant, thus impliying that the equilibrium variable would change, i.e. $H_j^{n+1} = z_j + h_j^{n+1} \neq z_j + h_j^n = H_j^n$. In order to prevent this kind of perturbation it is enough to reconstruct along equilibrium variables or to ensure that the boundary extrapolated values at the interface are continuous when equilibrium occours. In the second equation, the perturbation due to the artificial diffusion does not appear exactly because q is an equilibrium variable for the lake at rest equilibrium. However there is a lack of balance between the source and the fluxes at the discrete level: in fact one finds that $q_j^{n+1} = -\frac{\lambda}{4}(z_{j+1}^2 - 2z_j z_{j+1} + 2z_j z_{j-1} - z_{j-1}^2)$, which is in general nonzero, unless the bottom is flat.

For these reasons we use the hydrostatic reconstruction of [1] which ensures that the reconstruction is continuous across interfaces when the system is in equilibrium and moreover preserves positivity of the water height. Given a reconstruction algorithm \mathcal{R} with accuracy of order p, reconstruct the equilibrium variables Hand q, obtaining the boundary extrapolated data as in equation (6). In order to ensure that the water height appearing in the fluxes remains non-negative, one locally modifies the bottom by computing boundary extrapolated data also for hand defining

$$z_{j+1/2}^{\pm} = H_{j+1/2}^{\pm} - h_{j+1/2}^{\pm}$$

and these are used to compute the bottom topography at the interface

$$z_{j+1/2} = \max(z_{j+1/2}^+, z_{j+1/2}^-).$$

Once these are known, the interface values of h are corrected giving new values

$$\widehat{h}_{j+1/2}^{\pm} = \max(H_{j+1/2}^{\pm} - z_{j+1/2}, 0).$$

Note that $\hat{h}_{j+1/2}^{\pm} \geq 0$ and that at equilibrium $\hat{h}_{j+1/2}^{+} = \hat{h}_{j+1/2}^{-}$. The numerical fluxes (3) are then applied to the states

$$U_{j+1/2}^{\pm} = \left[\hat{h}_{j+1/2}^{\pm}, \ \hat{h}_{j+1/2}^{\pm} v_{j+1/2}^{\pm} \right].$$

Here $v_{j+1/2}^{\pm}$ denotes the velocity, obtained as $v_{j+1/2}^{\pm} = q_{j+1/2}^{\pm}/\hat{h}_{j+1/2}^{\pm}$ or through a desingularization procedure as proposed in [21]. In this fashion, the reconstruction is continous at equilibrium, for lake at rest data. Thus, for each consistent numerical flux, one has $\mathcal{F}(U_{j+1/2}^{-}, U_{j+1/2}^{+}) = f(U_{j+1/2}^{\pm})$ at equilibrium. In this fashion Audusse et al. are able to ensure well-balancing independently of the particular numerical flux used [1].

In order to complete the semidiscrete scheme (2) we still need to specify the discretization of the source term. For a first order scheme it is enough to choose

$$G_j = \frac{g}{2} \begin{pmatrix} 0\\ (\hat{h}_{j+1/2}^-)^2 - (\hat{h}_{j-1/2}^+)^2 \end{pmatrix}.$$
 (11)

Note that at equilibrium, the above expression exactly cancels out the numerical fluxes and thus the lake at rest solution is preserved at the discrete level. Consistency is obtained through the dependence of \hat{h} on z. For instance, if z(x) is monotonically increasing, then $z_{j+1/2} = z_{j+1}$ and $z_{j-1/2} = z_j$. Thus

$$\widehat{h}_{j-1/2}^+ = \max(h_j, 0) \qquad \widehat{h}_{j+1/2}^- = \max(h_j + z_j - z_{j+1}, 0)$$

and substituting into G_j one recovers the term ghz_x . For more details, see [1].

At second order, the second component of the source term is

$$G_{j,2} = \frac{g}{2} \left(\left(\hat{h}_{j+1/2}^{-} \right)^2 - \left(h_{j+1/2}^{-} \right)^2 + \left(h_{j-1/2}^{+} + h_{j+1/2}^{-} \right) \left(z_{j-1/2}^{+} - z_{j+1/2}^{-} \right) \right) + \left(h_{j-1/2}^{+} \right)^2 - \left(\hat{h}_{j-1/2}^{+} \right)^2 \right).$$
(12)

On the lake at rest solution, the two \hat{h} terms cancel the numerical fluxes, while the other terms add up to zero, again giving a well-balanced scheme [1]. On the other hand, off equilibrium, the first and the last two terms cancel by consistency and the middle term is consistent with the cell average of the source. Clearly, equation (12) must be applied to both stages of the second order Runge-Kutta method needed to achieve second order accuracy also in time.

For higher orders, we use Richardson extrapolation as in [29]. This technique is particularly useful on non-uniform grids because it concentrates all the computational effort for the source term within one cell. In fact, the subcell resolution required to compute the quadrature of the source term with high order accuracy can be naturally applied introducing uniformly distributed nodes within each cell. Thus the high order evaluation of the source term is performed entirely within one cell and the coefficients of the quadrature formula will not be affected by the non-uniformity of the mesh. The source can be rewritten as

$$G_{j} = \frac{g}{2} \begin{pmatrix} 0 \\ (\hat{h}_{j+1/2}^{-})^{2} - (\hat{h}_{j+1/2}^{-})^{2} + \tilde{G}_{j} + (\hat{h}_{j-1/2}^{+})^{2} - (\hat{h}_{j-1/2}^{+})^{2} \end{pmatrix}.$$
 (13)

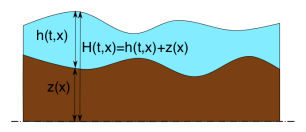


Fig. 4 Shallow water set up.

At second order,

$$\widetilde{G}_{j} = \frac{1}{2} (h_{j-1/2}^{+} + h_{j+1/2}^{-}) (z_{j-1/2}^{+} - z_{j+1/2}^{-}) = -\int_{x_{j-1/2}}^{x_{j+1/2}} h z_{x} \mathrm{d}x + O(\delta_{j}^{2}).$$

Following [29] that showed that the well-balanced property of the above quadrature is maintained under numerical extrapolation, for order up to four, it is enough to perform one Richardson extrapolation step, namely

$$\widetilde{G}_{j} = \frac{4}{3} \left((h_{j-1/2}^{+} + h_{j})(z_{j-1/2}^{+} - z_{j}) + (h_{j} + h_{j+1/2}^{-})(z_{j} - z_{j+1/2}^{-}) \right) \\ - \frac{1}{3} (h_{j-1/2}^{+} + h_{j+1/2}^{-})(z_{j-1/2}^{+} - z_{j+1/2}^{-}),$$

where h_j and z_j denote the reconstruction at the center of the cell, which is why we have developed high order reconstructions for the point values of the solution in x_j . Again, equation (13) will be applied to all stages of the Runge-Kutta method used in the fully discrete scheme. For higher orders, more Richardson steps must be applied [29] and this in turn introduces additional quadrature points, which must be computed with high order reconstructions.

4 Numerical entropy production for balance laws

We wish to devise an error indicator for driving adaptive schemes for balance laws. In particular we extend the notion of numerical entropy production proposed in [31,32] to the case of balance laws with a geometric source term.

In the homogeneous case, that is for systems of hyperbolic conservation laws, the entropy is defined as a convex function $\eta(u)$ for which there exists a function $\psi(u)$ (called entropy flux) such that $\nabla^T \eta f' = \nabla^T \psi$ where f' denotes the Jacobian of the flux function f. Then, on smooth solutions,

$$\partial_t \eta + \partial_x \psi = 0,$$

while on entropic shocks

$$\partial_t \eta + \partial_x \psi \le 0$$

in a weak sense, thus singling out the correct unique solutions [10]. One can exploit this structure at the discrete level to devise a regularity indicator for finite volume schemes for conservation laws. A fully discrete finite volume conservative scheme for a hyperbolic system can be written in the form

$$\overline{U}_{j}^{n+1} = \overline{U}_{j}^{n} - \lambda \left(F_{j+1/2} - F_{j-1/2} \right).$$

Here

$$F_{j+1/2} = \sum_{i=1}^{s} b_i \mathcal{F}\left(U_{j+1/2}^{(i),-}, U_{j+1/2}^{(i),+}\right)$$

 \mathcal{F} is a consistent and monotone numerical flux and $U_{j+1/2}^{(i),\pm}$ denote the boundary extrapolated data computed on the *i*-th stage value.

Choosing a numerical entropy flux \mathcal{P} , consistent with the exact entropy flux ψ , we *define* the numerical entropy production as

$$S_j^n = \frac{1}{\Delta t_n} \left[\overline{\eta(U^{n+1})}_j - \overline{\eta(U^n)}_j + \lambda \left(P_{j+1/2} - P_{j-1/2} \right) \right], \tag{14}$$

where

$$P_{j+1/2} = \sum_{i=1}^{s} b_i \mathcal{P}\left(U_{j+1/2}^{(i),-}, U_{j+1/2}^{(i),+}\right)$$

and $\overline{\eta(U)}_j$ denotes the average of $\eta(U(x))$ on the *j*-th cell. $\overline{\eta(U)}_j$ can be computed as $\eta(\overline{U}_j)$ up to second order accuracy or, for higher orders, with the help of the reconstruction and of a sufficiently accurate quadrature formula.

In [32] we proved that the scaling of the numerical entropy production S_j^n with respect to the cell size h is

$$S_j^n = \begin{cases} O(h^p) & \text{on smooth flows} \\ O(1) & \text{on contacts} \\ \sim C/h & \text{on shocks,} \end{cases}$$

where C does not depend on h and p is the order of accuracy of the scheme. Thus, on regular solutions, S_i^n is of the same order of the local truncation error.

Moreover, if the numerical flux can be written in viscous form as

$$\mathcal{F}(U^{-}, U^{+}) = \frac{1}{2}(f(U^{-}) + f(U^{+})) - \frac{1}{2}Q(U^{-}, U^{+})(U^{+} - U^{-}),$$
(15)

we choose the numerical entropy flux as

$$\mathcal{P}(U^{-}, U^{+}) = \frac{1}{2}(\psi(U^{-}) + \psi(U^{+})) - \frac{1}{2}Q(U^{-}, U^{+})(\eta(U^{+}) - \eta(U^{-})).$$
(16)

In [32] we have proved that, with this choice, the entropy S_j^n is negative definite on monotone profiles for the upwind and the Lax Friedrichs numerical fluxes applied to first order schemes in the scalar case, while it can exhibit very small positive overshoots (which are $O(h^4)$) only near local extrema. Without taking into account the viscous term in (16), the resulting S_j^n has the same convergence rate, but it is much more oscillatory and thus less suited to drive an h-adaptive scheme. Further, numerical evidence shows that the numerical entropy production S_j^n designed with the viscous correction is less oscillatory also in the case of systems of conservation laws and for higher order schemes. We wish to extend this construction to systems of m balance laws. We consider balance laws of the form

$$g(u,x) = \sum_{j=1}^{M} s_j(u,x) z'_j(x),$$
(17)

(with $s_j : \mathbb{R}^m \times \mathbb{R} \to \mathbb{R}^m$). This idea follows from the definition of separable balance laws in the sense of [45], where the authors write the source in equilibrium variables, obtaining the form (17) with M = 2 even for the lake at rest solution, because then the equilibrium variables are H = h+z and q, and the source becomes $g(H,q) = -gHz_x + g/2(z^2)_x$. Another interesting case in which the source is splitted in the sum of two different greometric terms can be found in [39], where the first term is the standard force term given by the slope in the bottom topography, while the second term is due to variations in the orthogonal sections of a channel.

When the source is written in the form (17), the balance law can be rewritten as an homogeneous system of m+M equations. For the case M = 1, denoting with A(u) the $m \times m$ Jacobian matrix of the flux f, one has

$$\partial_t \begin{pmatrix} u \\ z_1 \end{pmatrix} + \begin{pmatrix} A(u) \ s_1(u, x) \\ 0 \ 0 \end{pmatrix} \partial_x \begin{pmatrix} u \\ z_1 \end{pmatrix} = 0.$$
(18)

Exploiting this structure one can extend the notion of entropy. In fact the entropyentropy flux pair for the balance law must satisfy

$$\left[\nabla_u^T \eta A(u), \ \nabla_u^T \eta \cdot s_1(u, x)\right] = \left[\nabla_u^T \psi, \ \partial_{z_1} \psi\right].$$
(19)

Note that the z-derivative of η does not appear in the compatibility condition above, and thus convexity with respect to z is not required. This construction can be easily extended for M > 1.

Thus we still have entropy conservation for the balance law in the smooth case, provided the entropy-entropy flux pair satisfies (19), and the entropy residual defined in (14) gives a measure of the local error of the numerical scheme.

In the shallow water case, the entropy pair can be chosen as

$$\eta(h,v) = \frac{1}{2} \left(hv^2 + gh^2 \right) + ghz \qquad \psi(h,v) = \eta(h,v)v + \frac{1}{2}gh^2v, \tag{20}$$

where v is the water velocity (see [4]). Note that the function η represents the total energy of the system including the potential energy due to the bottom topography. In the following section we will show that the entropy residual converges with the expected rate on smooth flows and detects the presence of shocks in the solution.

5 Numerical tests

The following tests assess the accuracy of the high order reconstructions on nonuniform grids proposed in this work, the well-balancing properties of the fully discrete schemes for the shallow water equations, the resolution of discontinuities on non-uniform grids and the performance of the entropy residual as an error indicator.

In all tests we used the local Lax-Friedrichs numerical flux, i.e. we choose $Q(U^-, U^+) = \max(|v^-| + g\sqrt{h^-}, |v^+| + g\sqrt{h^+})$ in the viscous forms (15) and (16).

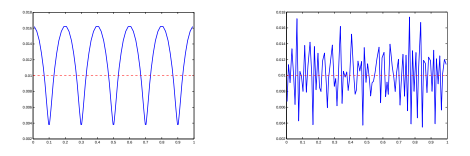


Fig. 5 Grid spacing δ as a function of x for the non-uniform grids used in the numerical tests, shown for the case of 100 points in [0, 1]. *Quasi-regular* grids (left) and *random* grids (right). The dashed line is the spacing of the uniform grid with the same number of points.

Grids In the numerical tests we use several grids that will be referred to as uniform, quasi-regular, random and locally refined. For simplicity we define them on the reference interval [0, 1]. The quasi regular grid is obtained as the image of a uniform grid with spacing $\delta = 1/N$ under the map

$$\varphi(x) = x + 0.1 * \sin(10\pi x)/5.$$

The resulting grid spacing is depicted in the left panel of Figure 5: we point out that $\delta_j \sim \frac{1}{N} \varphi'(j/N)$ and thus

$$(1 - \frac{\pi}{5})\frac{1}{N} \le \delta_j \le (1 + \frac{\pi}{5})\frac{1}{N}.$$

Next, we consider non-uniform rough grids that are obtained moving randomly the interfaces of a uniform grid, namely starting from a uniform grid with spacing δ we consider grids with interfaces at

$$\tilde{x}_{j+1/2} = j\delta + \frac{3}{5}\xi_j\delta_j$$

where ξ_j are random numbers uniformly distributed in [-0.5, 0.5]. A realization of such a grid is shown in the right panel of Figure 5. Here it is easily seen that

$$\frac{2}{5}\frac{1}{N} \le \delta_j \le \frac{8}{5}\frac{1}{N}.$$

We use this grid for the purpose of illustration even if of course one would not use such an irregular grid in an application. This grid will be referred to as *random* grid. We point out that the ratio of the maximum to the minimum cell size is approximately 4 in both quasi-uniform and random grids. On the other hand, the ratio of the size of nearby cells is approximately 4 on random grids, but slightly lower than 2 in quasi-uniform grids.

In some tests we need a grid which is locally refined around a given point w_C . For this purpose we consider a grid which, on the standard domain [0, 1], is a map of a uniform grid under a monotone function $\varphi : [0,1] \to [0,1]$ such that $0, w_C, 1$ are fixed points and $\varphi''(w_C) = 0$, so that the grid spacing is minimum at w_C . In particular we consider the quartic polynomial defined by $\varphi(0) = 0, \varphi(1) = 1, \varphi(w_C) = w_C, \varphi''(w_C) = 0$ and $\varphi'(w_C) = \alpha$, where α is a parameter which gives the amplitude of the grid at w_C , and it must be chosen in order to ensure that the

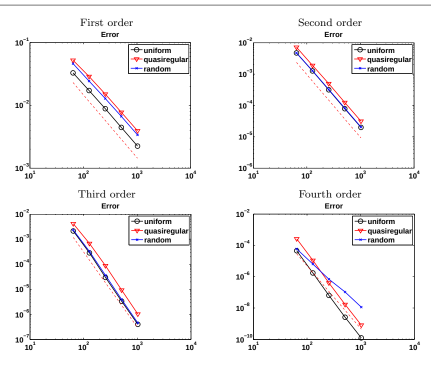


Fig. 6 Error decay under grid refinement for first (top-left), second (top-right), third (bottom-left) and fourth (bottom-right) order schemes. The dashed line indicates the expected decay in each case.

map remains monotone. In our case, we pick $\alpha = 0.25$, which gives a grid with a ratio between maximum and minimum spacing of order 4. We find

$$\varphi(x) = x \left[1 + \frac{1-\alpha}{w_C} (x - w_C) \left[-1 + \frac{1}{w_C} (x - w_C) \left[1 + (x - w_C) \frac{2w_C - 1}{(1 - w_C)^2} \right] \right] \right].$$
(21)

5.1 High order schemes on non-uniform grids

Convergence tests Following [44], we compute the flow with initial data given by

$$z(x) = \sin^2(\pi x) \qquad h(0, x) = 5 + e^{\cos(2\pi x)} \quad q(0, x) = \sin(\cos(2\pi x)), \tag{22}$$

with periodic boundary conditions on the domain [0, 1]. At time t = 0.1 the solution is still smooth and we compare the numerical results with a reference solution computed with the fourth order scheme and 16384 cells. The 1-norm of the errors appears in Figure 6 and the maximum entropy production is shown in Figure 13 for all schemes and the three grid types considered.

All schemes have the expected accuracy, except for the fourth order scheme on the random grids, where the accuracy is decreased due to the extreme irregularity of the grid. We point out however that, despite the reduced decay rate, the actual values of the error of the fourth order scheme even on the random grid are orders of magnitude smaller than those obtained with the third order scheme with the

Well-balanced high order 1D schemes on non-uniform grids and entropy residuals

	$\ \Delta(h+z)\ _{\infty}$				$\ q\ _{\infty}$			
Smooth	100	200	400	800	100	200	400	800
p = 1	0	0	0	0	4.51e-16	5.55e-16	5.00e-16	7.68e-16
p = 2	0	2.22e-16	2.22e-16	2.22e-16	3.82e-16	8.47e-16	7.36e-16	$1.54e{-}15$
p = 3	0	4.44e-16	4.44e-16	6.66e-16	6.87e-16	1.47e-15	1.67e-15	2.47e-15
p = 4	8.88e-16	6.66e-16	1.55e-15	1.55e-15	9.89e-16	1.82e-15	1.67e-15	1.90e-15
Random								
p = 1	2.22e-16	2.22e-16	2.22e-16	2.22e-16	2.08e-16	6.24e-16	6.77e-16	9.65e-16
p = 2	2.22e-16	2.22e-16	2.22e-16	2.22e-16	2.91e-16	7.25e-16	8.95e-16	9.99e-16
p = 3	2.22e-16	6.66e-16	6.66e-16	6.66e-16	5.63e-16	8.47e-16	9.94e-16	1.28e-15
p = 4	6.66e-16	8.88e-16	1.33e-15	1.11e-15	8.68e-16	7.94e-16	1.11e-15	1.43e-15

Table 1 Lake at rest test: well-balancing errors with rough bottom.

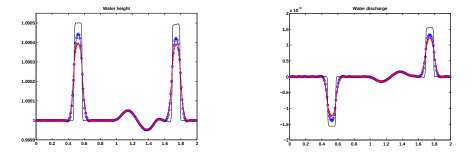


Fig. 7 LeVeque's test (23). Third order scheme on a uniform (blue circles) and *quasi-regular* grid (red crosses) on top of a reference solution (black solid line).

same number of degrees of freedom. We also point out that a real life application of adaptive grids would produce a non-uniform grid which is refined especially around shocks, which are isolated discontinuities along which accuracy degrades anyhow because of the non regularity of the solution. What matters in this case is to keep a high resolution, i.e. a small local error.

Well-balancing We show a well-balancing test on the lake at rest solution using a bottom topography described by a uniformly distributed random variable sampled between 0 and 1, with water height at h(x) + z(x) = 1.5. Table 1 shows the well-balancing errors in the total water height and momentum, in the case of smooth non-uniform grids and random grids. Here $\Delta(h+z)_{j+1/2} = (h+z)_{j+1} - (h+z)_j$. All data are close to machine precision, as expected.

Small perturbation of a lake at rest The domain is $x \in [0, 2]$, the bottom and initial total height are given by

$$z(x) = \begin{cases} 0.25(1 + \cos(10\pi(x - 0.5))) & 1.2 \le x \le 1.4 \\ 0 & \text{otherwise,} \end{cases}$$
(23)
$$H(x, 0) = 1 + 0.001\chi_{[1,1,1,2]}(x).$$

This test was first used by LeVeque in [23] with a second order scheme, but here we use it with a smaller perturbation for the third and fourth order schemes,

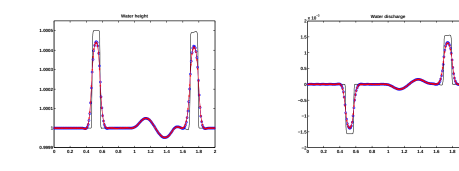


Fig. 8 LeVeque's test (23). Third order scheme on uniform (blue circles) and *random* grid (red crosses) on top of a reference solution (black solid line)

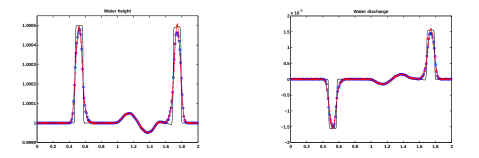


Fig. 9 LeVeque's test (23). Fourth order scheme on uniform (blue circles) and *quasi-regular* grid (red crosses), on top of a reference solution (black solid line)

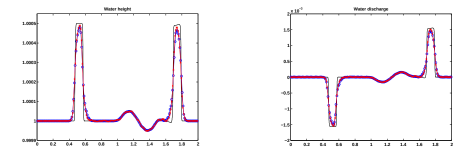


Fig. 10 LeVeque's test (23). Fourth order scheme on uniform (blue circles) and *random* grids (red crosses), on top of a reference solution (black solid line).

as in [29]. This test requires a well-balanced scheme to resolve correctly the small perturbations which otherwise would be hidden by numerical noise. The solutions with N = 200 cells are shown in Fig. 7 and 8 for the third order scheme and Fig. 9 and 10 for the fourth order one. In each of the figures the numerical solution obtained with the uniform grid is compared with the one obtained on a non-uniform mesh. It can be seen that the pulse is well-resolved in all cases and the results obtained with a non-uniform grid can be perfectly superposed on those computed with the uniform mesh. As already pointed out in [29], for this test the

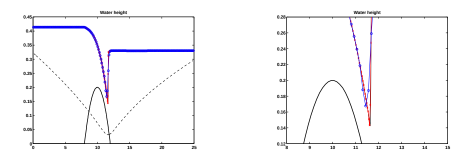


Fig. 11 Steady solution with transcritical shock, approximated with a third order scheme (uniform and adapted grids). The dashed line in the left panel is the local grid size in the non-uniform grid (it should be read against the axis on the left).

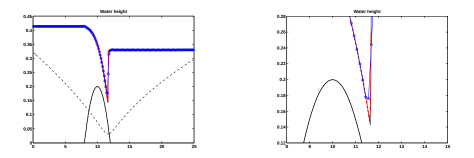


Fig. 12 Steady solution with transcritical shock, approximated with a fourth order scheme (uniform and adapted grids). The dashed line in the left panel is the local grid size in the non-uniform grid (it should be read against the axis on the left).

parameter ϵ in the nonlinear weights of the WENO schemes has to be reduced; we have chosen 10^{-12} for the fourth order scheme (see [29]) and 10^{-9} for the third order case.

Moving water equilibria Since our schemes are well-balanced around the lake-atrest equilibrium, one does not expect them to compute moving water equilibria at machine precision. Here we show two tests. In the first case we consider a transcritical steady state with a shock, over the parabolic hump

$$z(x) = \begin{cases} (0.2 - 0.05 * (x - 10)^2) & 8 \le x \le 12\\ 0 & \text{otherwise} \end{cases}$$

in the domain [0,25]. We consider the steady state solution with q(x) = 0.18, with Dirichlet boundary conditions q = 0.18 at x = 0 and h = 0.33 at x = 25. The flow starts off subcritical, turns supercritical on the hump and goes back to the subcritical regime through a steady shock located at $x_s = 11.6655042815533$. The computation was initialized with the exact steady state solution (computed as explained in the Appendix A of [19]) and the numerical integration was performed until t = 50.

	p = 1		p = 2		p = 3		p = 4	
Uniform	error	rate	error	rate	error	rate	error	rate
100	1.96e-1	_	5.54e-2	_	2.02e-2	_	2.92e-3	_
200	1.17e-1	0.74	1.42e-2	1.96	4.26e-3	2.24	1.40e-4	4.38
400	6.35e-2	0.89	3.29e-3	2.11	4.87e-4	3.13	5.12e-6	4.77
800	3.26e-2	0.96	8.08e-4	2.03	3.89e-5	3.65	1.60e-7	5.00
Adapted								
100	9.20e-2	_	6.96e-3	_	9.78e-4	_	4.54e-5	_
200	4.67e-2	0.97	1.71e-3	2.02	7.97e-5	3.62	1.36e-6	5.07
400	2.34e-2	0.99	4.25e-4	2.01	6.57e-6	3.60	3.87e-8	5.13
800	1.17e-2	1.00	1.06e-4	2.01	5.63e-7	3.55	1.25e-9	4.95

Table 2 Well-balancing errors for the subcritical steady state with gaussian bottom.

We show the solutions computed with uniform grids and with a grid refined ad-hoc around the shock position (as in Eq (21), with $w_C = x_s/25$) with the scheme of order three (Figure 11) and four (Figure 12). The figures report with a dashed line the local cell size of the non-uniform grid, which is refined close to the shock. The right panels of each figure show a zoom on the shock. It is clear that the adapted solution (in red with crosses) approximates better the exact solution (thin black line) than the solution obtained with a uniform grid with the same number of points (blue line with dots), with no spurious oscillations.

In order to quantify the improvement due to the adapted grid and the rate of convergence of the schemes on moving water equilibria, we consider a smooth test problem, namely a subcritical steady flow over the smooth bump $z(x) = 0.2e^{-(x-12.5)^2}$ on the domain [0,25]. The numerical scheme was initialized with the exact solution and the flow computed until t = 10. Since the behaviour of the errors on the water height and on momentum is very similar, only the former are reported in Table 2. The first and second order schemes show the expected rates of convergence, while the third and fourth order ones have convergence rates well above the expected values (respectively 3.60 and 5.00).

We also consider non-uniform grids that are finer on the hump and coarser on the flat portion of the bottom function, namely the grid is defined by Eq. (21) with $w_C = 12.5/25 = 0.5$. The errors on the adapted grids are much smaller than the corresponding results on uniform grids and the convergence rates are confirmed also on non-uniform grids. Note also that the improvement obtained with the adapted grid is stronger for the high order schemes.

5.2 Numerical entropy production

Rate of decay on smooth flows. Figure 13 shows the numerical entropy production in the smooth test (22) on several grid types. In particular, for each number of grid points, we plot the 1-norm of the numerical entropy production defined in equation (14) observed in the last timestep of each simulation. It is apparent that the decay rate, as expected, follows the order of accuracy of the corresponding schemes. Moreover, comparing this figure with Figure 6, we note that the entropy decay mimics exactly the behaviour of the error, even in the case of the slight deterioration of accuracy observed on the random grid for the fourth order scheme.

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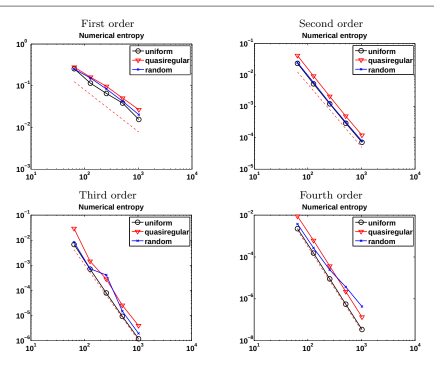


Fig. 13 Numerical entropy production decay under grid refinement for first (top-left), second (top-right), third (bottom-left) and fourth (bottom-right) order schemes. The dashed line indicates the expected decay in each case.

Two shocks. We set up initial data with a flat bottom, water at rest and $h(0,x) = e^{-50x^2}$ on the domain [-2, 2]. As the flow evolves, two shocks form and separate from each other: at t = 0.2 the computed water height is depicted in the top-left plot of Figure 14. Each of the other panels of Figure 14 shows the entropy residual obtained with four different grid sizes. The results for second, third and fourth order schemes appear in the top-right, lower left and lower right panels respectively. In all three cases it can be seen that the numerical entropy production on the two shocks increases under grid refinement like 1/h. On the other hand, the magnitude of the peak of the numerical entropy production does not depend on the order of the scheme. This is to be contrasted with the numerical entropy residuals of $O(h^p)$, where p is the order of the scheme.

Due to the different orders of magnitude of the numerical entropy production in the smooth regions of the flow and around shocks, it can be concluded that the entropy residual provides an effective discontinuity detector, expecially in the case of high order schemes.

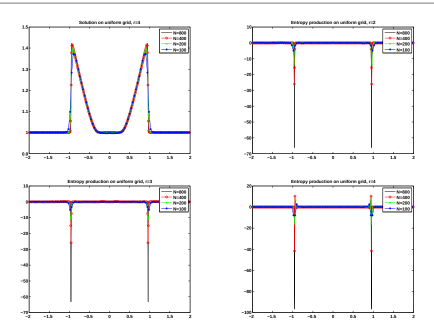


Fig. 14 Entropy production on shocks under grid refinement for several schemes. Top-left: water height. Top-right: second oder scheme. Bottom-left: third order scheme. Bottom right: fourth order scheme. N = 800 (black solid line), N = 400 (red line with circles), N = 200(green line with crosses), N = 100 (blue line with stars).

Stream on artificial river bed. In the domain [-0.5, 1.5] we consider the bottom topography and initial conditions:

$$z(x) = \begin{cases} x(1-x)\sin(10\pi x) & x \in [0,1] \\ 0 & \text{otherwise,} \end{cases}$$
(24)
$$H(0,x) = \begin{cases} 1.0 & x < -0.2 \\ 0.5 & x \ge -0.2, \end{cases} \quad q(0,x) = \begin{cases} \frac{1}{2}\sqrt{\frac{3}{2}g} & x < -0.2 \\ 0.0 & x \ge -0.2. \end{cases}$$

 $x \ge -0.2.$

We integrate with free flow boundary conditions until t = 0.4, when the shock originated from the Riemann problem has overcome the irregularity in the bottom topography (see the left panel of Figure 15). The right panel compares the numerical entropy production of the second order scheme with grids varying from 200 to 1600 points. The vertical axis is in logarithmic scale. The peaks in the numerical entropy production clearly show the location of the shocks and have the expected O(1/h) behaviour, while in regions of smoothness the numerical entropy production reduces with the grid size.

Finally, we wish to illustrate the importance of choosing the numerical entropy flux customized on the numerical flux used by the scheme, as in (16). Figure 16 shows the numerical entropy production on the test (24) computed with the numerical entropy flux of (16), which uses the same viscous form of the underlying scheme (black solid line), and with the numerical entropy flux

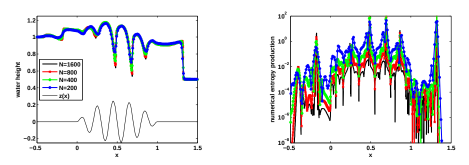


Fig. 15 Stream on artificial river bed, integrated with the second order scheme. Left: water height. Right: numerical entropy production. The black curve is the bottom topography.

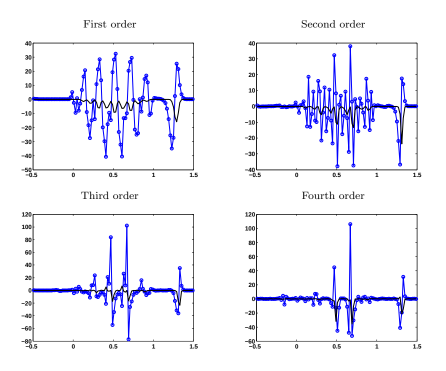


Fig. 16 Comparison of the numerical entropy production obtained with (black line) and without (blue line) the viscous correction.

 $\Psi(U^-, U^+) = \frac{1}{2}(\psi(U^-) + \psi(U^+))$ (blue line with circles). Note that both numerical entropy fluxes are consistent with the exact entropy flux ψ , and therefore they will both provide entropy residuals with the same rate of decay of the local error of the scheme.

However, for all the schemes considered here, it is clear from the example in Figure 16 that using the local Lax-Friedrichs flux for both the conservation law and the computation of the numerical entropy flux leads to much smaller positive overshoots in the numerical entropy production and thus a much more reliable error indicator.

6 Conclusions

In this work we have derived formulas for high order schemes for balance laws on non-uniform grids. More precisely, we give the extension of the third order compact WENO reconstruction of [26] to non-uniform grids and we also derive high order reconstructions to compute the cell averages of source terms, needed by high order finite volume schemes on balance laws. Further, we illustrate how well balancing on equilibrium solutions can be enforced for high order schemes on irregular grids.

We also include the extension of the entropy indicator we proposed in [32] and [31] to the case of balance laws. The proofs given in [32] carry over to the case of balance laws with geometric source terms, and prove that the entropy indicator provides a measure of the local truncation error on smooth flows, and it reliably selects the location of discontinuities.

Several numerical tests are included, to show the achievement of the expected accuracy of the schemes proposed, even on extremely irregular grids, and the improvement obtained with ad-hoc chosen grids.

Future work on this topic will be dedicated to the construction of locally refined Cartesian grids of octree type, driven by the entropy error indicator for balance laws, with particular attention on the enforcement of equilibrium solutions at the discrete level.

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