ON THE ACCURATE SIMULATION OF NEARSHORE AND DAM BREAK PROBLEMS INVOLVING DISPERSIVE BREAKING WAVES

JOSÉ S. ANTUNES DO CARMO, JOSÉ A. FERREIRA AND LUÍS PINTO

ABSTRACT: The ability of numerical models to deal with wave breaking processes and dry areas is of paramount importance for applications in coastal zones and dam breaks. The mathematical models commonly used in such real problems are usually based on Boussinesq-type equations and, to a small extend, on Serre equations. However, these standard models are weakly dispersive and must be appropriately modified to deal with breaking waves and dry areas. Indeed, nearshore and dam break problems involve complex wave dynamics and highly dispersive wave processes can easily arise. In those cases, it is well known that weakly dispersive models like the ones based on classical Boussinesq or Serre equations are unreliable for an accurate simulation of the phenomena involved.

In this work we extend the applicability of an improved Serre model, herein denoted by Serre$_{\alpha,\beta}$, to include the wave breaking process, the broken waves propagation, and dry areas. We provide a comprehensive set of numerical examples involving wave propagation over exposed and submerged structures, as well as dam break problems. The numerical experiments show the accuracy and robustness of the proposed model. Particular attention is given to bottom friction modeling, where the standard Manning’s assumption is compared with a more realistic formulation. Also noteworthy is the simulation of wave breaking problems with highly dispersive effects. The advantages of the Serre$_{\alpha,\beta}$ model over the standard Serre model for these challenging cases are clear.

KEYWORDS: Extended Serre equations; Saint-Venant equations; Dispersive waves; Breaking waves; Improved bottom friction; Wave-structure interaction.

1. Introduction

Accurate simulation of water waves propagation is very important in the coastal regions management. One relevant example is the wave’s interaction with seawalls or underwater bars. These are structures build near the coast that have the function of protecting it against coastal erosion and flooding. Another context in which water wave models play a crucial role is in the
simulation of flooding due to dam breaking events. In both cases, an accurate simulation allows not only the design of more efficient structures, but also the prevention and mitigation of hazards.

In order to deal with the complex nearshore water dynamics, a coastal model must be able to reproduce wave breaking and dry areas. Most of the approaches for wave breaking consist in the introduction of additional dissipative terms in existing wave’s models. The role of these terms is to account for the decay of the water wave height inside the breaking region. Strategies to localize the breaking point are also required. In addition, the presence of dry areas means that we need to consider bottom friction effects. This brings additional numerical challenges since the friction term is stiff and dominant in dry or almost dry regions. In these conditions it is also important to use numerical schemes that have the positivity preserving property.

Another important aspect is the presence of dispersive and nonlinear effects. Wave dispersion strength is measured by the parameter $\rho = h_0/l$, where $h_0$ is the reference depth and $l$ the wavelength. The nonlinearity strength is measured by $\epsilon = a/h_0$, with $a$ the typical wave amplitude. It is well known that nearshore wave dynamics is strongly nonlinear [2]. This is particularly true at the end of the shoaling zone and also in the surf and swash zones. As the wave shoals up a slope and in very shallow water conditions, it steepens accumulating higher harmonics (dominant nonlinearity) that are released on wave breaking or on a downslope (dominant dispersive effects). After this process, the wave can eventually recover producing an irregular pattern, more or less complex. It is important to highlight that in the literature dispersive effects are usually analyzed separately from breaking, see, e.g., [16] and references therein. In this work, we show that a complete model must include simultaneously nonlinearity, wave breaking, and dispersive effects. Moreover, dispersion is so relevant, that models with weakly dispersive properties are not accurate enough.

Numerical models that solve the Saint-Venant equations [12], the Boussinesq equations [4], and the Serre or Green & Naghdi equations [41] are commonly used in coastal wave modeling. These models can be classified based on their dispersion and nonlinear properties. The Saint-Venant equations are weakly nonlinear and non-dispersive, i.e., some nonlinear terms and all terms of dispersive origin are ignored. Therefore, the Saint-Venant equations are suitable for wave propagation in the breaking zone. The Boussinesq equations are weakly dispersive and weakly nonlinear, i.e., it is assumed that
\( \epsilon \ll 1 \) and \( \varrho \ll 1 \). Due to these properties the standard Boussinesq equations have a limited applicability for nearshore applications. Several modifications to the Boussinesq equations have been proposed. For example, in [32, 52, 36], the dispersive properties were improved by adding terms of dispersive origin and in [53, 21], fully nonlinear Boussinesq-type models are presented. The standard Serre equations, like Boussinesq, are weakly dispersive, however, they are also fully nonlinear.

Despite being fully nonlinear Serre equations are inadequate for breaking zones. The problem is that near breaking points a singularity appears and the assumptions made to derive the equations are invalid [2]. To overcome this limitation some recent studies have proposed a hybrid approach [49, 39, 20, 31]. Since shock discontinuities are well resolved by Saint-Venant, this approach portrays breaking waves as shock waves and in the breaking zone a switch from Serre to Saint-Venant is made. Other limitation of the standard Serre equations is the weak dispersivity. This is a significant drawback for complex wave dynamics and several modifications have been proposed to overcome it [14, 52, 21, 6, 31, 10]. For example, following the approach used in [32] for Boussinesq, dispersion characteristics were improved in [14] considering additional terms of dispersive origin. An efficient numerical scheme to solve this improved Serre system, denoted \( \text{Serre}_{\alpha,\beta} \), was recently presented in [16].

The goal of this paper is to extend the \( \text{Serre}_{\alpha,\beta} \) model to include wave breaking and dry regions. By doing so, we obtain an accurate model covering all the nearshore wave processes. The relevance of improved dispersion properties is highlighted. This paper is organized as follows. In Section 2 we briefly deduce the \( \text{Serre}_{\alpha,\beta} \) model. Following [16], we write the model in a decoupled dispersive-hyperbolic form suitable for numerical implementation. Next, in Section 3, we present the strategies used to embed wave breaking and bottom friction. Here, we deviate from the standard Manning’s assumption and use a more realistic formulation for friction. The numerical scheme is presented in Section 4. We propose a hybrid approach combining finite volume and finite difference. Numerical experiments are given in Section 5 and we finish in Section 6 with some conclusions.

2. The \( \text{Serre}_{\alpha,\beta} \) Model

As illustrated in Figure 1, we consider the one-dimensional horizontal case and represent by \( b(x) \) the time constant bathymetry, by \( u(x,t) \) the flow
velocity, and by \( h(x, t) = h_0 + \eta(x, t) - b(x) \) the water height, with \( \eta(x, t) \) the free surface elevation and \( h_0 \) the reference depth. Here, \( x \) is the spatial variable and \( t \) is the time variable.

![Figure 1. Sketch of the domain and main notation.](image)

Following [16, 17] the one-dimensional version of the classical Serre equations with bottom friction can be written in the following compact form

\[
\begin{align*}
& h_t + (hu)_x = 0 \\
& (I + hT_1)( (hu)_t + (hu^2 + \frac{1}{2}gh^2)_x + ghb_x) - hT(g\eta_x) + hQ(u) = -\frac{\sigma}{\rho},
\end{align*}
\]

(1)

where \( g \) is the gravitational acceleration, \( I \) is the identity operator, \( Q \) is defined by

\[
Q(v) = h(2h_x + b_x)(v_x)^2 + \frac{4}{3}h^2v_xv_{xx} + hb_{xx}v_x + (\eta_xb_{xx} + \frac{h}{2}b_{xxx})v^2
\]

(2)

and \( T_1 = T(v) \) with \( T \) given by

\[
T(v) = -\frac{h^2}{3}v_{xx} - hh_xv_x + (b_x\eta_x + \frac{h}{2}b_{xx})v.
\]

(3)

Still in (1), at the right-hand side of the momentum equation we have the friction term \( -\sigma/\rho \), with \( \rho \) the fluid density and \( \sigma \) the friction at the bottom.

To obtain an efficient numerical scheme for Serre system (1) we use a splitting approach to decouple the dispersive and nonlinear terms. Let \( S(t) \) be the solution operator of (1) and let us assume for now that

\[
S(t) = S_d(t) \circ S_{nl}(t)
\]

(4)

where \( S_{nl}(t) \) is the solution operator associated with the nonlinear part

\[
\begin{align*}
& h_t + (hu)_x = 0 \\
& (hu)_t + (hu^2 + \frac{1}{2}gh^2)_x + ghb_x = -\frac{\sigma}{\rho},
\end{align*}
\]

(5)
and \( S_d(t) \) is the solution operator of the dispersive part

\[
\begin{align*}
  h_t &= 0 \\
  (I + hT_1)((hu)_t) - hT(g\eta_x) + hQ(u) &= 0.
\end{align*}
\]

(6)

We observe that system (5) corresponds to the classical Saint-Venant system with friction at bottom level.

To obtain the proposed Serre_{\alpha,\beta} system we follow [14] and add and subtract to the momentum equation of (6) the term

\[
  hT_{\frac{1}{h}}^{\alpha,\beta}((hu)_t),
\]

with

\[
  T^{\alpha,\beta}(v) = -\beta \frac{h^2}{3} v_{xx} - \alpha hh_x v_x + \alpha(b_x\eta_x + \frac{h}{2} b_{xx}) v.
\]

(7)

Using also the classical approximation \( u_t = -g\eta_x \) and the fact that \( h_t = 0 \) we get

\[
\begin{align*}
  h_t &= 0 \\
  (I + hT_1 + hT_{\frac{1}{h}}^{\alpha,\beta})((hu)_t) - h(T - T^{\alpha,\beta})(g\eta_x) + hQ(u) &= 0.
\end{align*}
\]

(8)

Denoting the solution operator of (8) by \( S_d^{\alpha,\beta} \) we define our Serre_{\alpha,\beta} model as

\[
S^{\alpha,\beta}(t) = S_d^{\alpha,\beta}(t) \circ S_{nl}(t).
\]

(9)

**Remark 2.1** Serre and the proposed Serre_{\alpha,\beta} systems are equivalent up to order \( O(\sigma^2) \) and the classical one is recovered when \( \alpha = \beta = 0 \).

**Remark 2.2** Another well-known model [3] with improved dispersive properties can be obtained by setting

\[
S^\theta(t) = S^\theta_d(t) \circ S_{nl}(t),
\]

(10)

with \( S^\theta_d(t) \) given by

\[
\begin{align*}
  h_t &= 0 \\
  (I + h\theta T_1)((hu)_t) - hT(g\eta_x) + hQ(u) &= 0,
\end{align*}
\]

with \( \theta \geq 0 \). For non-breaking and non-friction problems with strong dispersive effects it was shown in [16] that the proposed model (9) offers noticeable
improvements over (10). Naturally, the advantages over traditional Serre (1) are even more significant.

3. Wave Breaking and Bottom Friction Modeling

In this section we discuss the strategies adopted to model wave breaking and bottom friction. These two features are crucial in any coastal wave model. We start by explaining how breaking is embedded in Serre$_{\alpha,\beta}$.

3.1. Wave Breaking. When a wave approaches the shoreline the water depth decreases and the wave becomes very step. Simultaneously, the crest velocity increases, and when it exceeds the overall wave velocity the crest breaks and separates from the wave. This turbulent phenomenon, characterized at large scales by strong energy dissipation and wave height reduction, is called wave breaking. Equations specifically designed for turbulent behavior, e.g., RANS (Reynolds-Average Navier-Stokes), have been employed to simulate this process [24]. However, due to the heavy computational cost, depth-averaged equations, such as Serre, are still preferred. The problem is that these equations are invalid in the breaking zone and they need to be complemented with wave breaking methodologies.

Most of the available techniques are based on local ad-hoc dissipative terms. These terms are added to the wave model (e.g., Serre or Boussinesq-type equations), with the goal of capturing energy dissipation in the breaking zone. The popular eddy viscosity approach [27] considers the term

$$D_{hu} = \partial_x (\nu(hu)_x),$$

(11)

where $\nu$ is the so-called eddy viscosity. The term (11) is to be added at the wave models momentum equation. In our case, the second equation of (1). The major criticism to this approach is the lack of physical meaning of the parameters used to express the eddy viscosity. In order to obtain a better approximation some authors have used $k$-$l$ turbulent models [36, 26, 55]. In general the eddy viscosity is written using the Prandtl and Kolmogorov relation, which assumes that $\nu$ can be written as a product of a turbulent kinetic energy ($k$) and a turbulent length scale ($l$). The several approaches differ in the way that $k$ and $l$ are calculated. For example, in [26], $l$ is assumed proportional to the water height $h$ and $k$ is the solution of a partial differential equation.
In [9], the authors used energy principles to claim that a more physically sounded model is obtained when an additional term is also added to the continuity equation. The proposed model is based on the two breaking terms

\[ D_h = \partial_x (d_h h_x), \quad (12) \]
\[ D_{hu} = \partial_x (d_{hu} (hu)_x), \quad (13) \]

where (13) is included in the momentum equation and (12) is the extra term that appears in the continuity equation. Exploiting arguments developed in [11, 13, 44] the authors also suggest expressions for the functions \(d_h\) and \(d_{hu}\). Basically, they are of exponential type, centered around the moving wave crest and dependent on the local still water depth. Estimates for the parameters involved are also given.

Let us also mention the friction-type wave breaking models, which are based on terms of the form \(f_d u\), with \(f_d\) a dissipation function dependent on \(h\) [51, 50, 17]. The rationale behind this approach is that energy dissipation due to wave breaking can be related to surface friction. Finally, we refer the roller-type models [45, 5, 13], based on terms of type \(R_x\), with \(R\) a function that depends on \(u\) and \(h\). This physically solid approach relies on the concept of “surface-roller”, the air-water mixing region formed in front of the breaking wave. In this context, energy dissipation occurs due to friction between the wave and the roller. In both models the extra terms are included in the momentum equation.

A numerical comparison against experimental data was performed in [42] using eddy viscosity [36], roller [5], and friction [50] models. It was concluded that the three models were equally accurate at predicting wave height. In [9], the numerical experiments favored the proposed model (based on (12)-(13)) over the eddy viscosity model given in [27]. In both cases the authors used Boussinesq-type equations.

In this work we adopt a different wave breaking strategy. Our approach, presented for the first time in [49], consists is switching locally from \(\text{Serre}_{\alpha,\beta}\) to the Saint-Venant system when the wave is about to break. The motivation is to explore the splitting strategy (9) presented in Section 2. Recall that \(\text{Serre}_{\alpha,\beta}\) is decoupled in a dispersive part \(S_{d}^{\alpha,\beta}(t)\) and a nonlinear part \(S_{nl}(t)\) which is exactly the Saint-Venant system. Therefore, breaking can be easily incorporated by simply ignoring the contribution from the dispersive term \(S_{d}^{\alpha,\beta}(t)\). This methodology has proven to be efficient and has gained
considerable popularity [25, 20, 31, 48, 26, 49]. This technique is also physically meaningful. In fact we can see breaking waves fronts as shocks and it is well known that the Saint-Venant system admit shock waves solutions that dissipate energy. Moreover, it doesn’t requires additional ad-hoc dissipative terms. Naturally, like the models previously presented, this is still a limited view of the complex wave breaking phenomenon.

In [26], this hybrid strategy was compared with a eddy viscosity model using Serre and Boussinesq-type equations. The numerical results suggest that the performance of the two strategies is identical. One drawback of the hybrid approach is the appearance of numerical instabilities when the switch is made, particularly for fine meshes. We refer that similar issues occur, e.g., in some eddy viscosity models when the diffusive terms are activated [27]. Here, we achieve a smooth transition by using the sigmoidal function

\[ f(t) = \frac{1}{1 + \exp(-4(10t - 1))}, \quad t \geq 0. \]

A different approach, where the transition is embedded as a kind of boundary condition is proposed in [20].

Concerning breaking modeling we still need to present the mechanisms used to activate and deactivate the breaking process. There are several strategies and the optimal one is problem dependent [20, 25, 26, 48, 9, 29]. In this work, we consider that a wave starts to break when the front slope angle exceeds a predefined threshold, i.e.,

\[ |\eta_x| \geq \phi_i. \] \hfill (14)

For breaking termination we use one of following two conditions, the front slope angle satisfies

\[ |\eta_x| \leq \phi_f, \] \hfill (15)
or the wave’s Froude number \((F)\) verifies \(F \leq F_f\), with
\[
F = \left( \frac{(2H_{wc}/H_{wt} + 1)^2 - 1}{8} \right)^{1/2}
\]  
and where \(H_{wc}\) and \(H_{wt}\) (see Figure 2) are the water depth at the wave’s crest and trough, respectively. We consider two breaking termination conditions because in some experimental tests one gives better results, or is more robust, than the other. For breaking initiation we found condition (14) accurate enough for all the experiments. The optimal values of \(\phi_i\), \(\phi_f\), and \(F_f\) depend on several aspects like, type of breaking model, forms of wave breaking (spilling, plunging, or surging), and employed numerical scheme.

Some efforts have been made in order to predict the optimal wave breaking parameters \(\phi_i\), \(\phi_f\), and \(F_f\). One such case is [54] where, using extensive experimental data, the authors propose two empirical formulas for \(\phi_i\). One such formula expresses \(\phi_i\) as a function of local wave parameters, while the other relates \(\phi_i\) with the relative water depth and bed slope. The extensive data analyzed in that work also shows that \(\phi_i\) can be in the range of 0.15 to 0.8. Note that condition (14) is based on the fact that when a wave approaches shallow waters the wave front angle increases and beyond a critical threshold the wave crest becomes unstable and breaks. The breaking termination condition (15) is closely related with the breaking initiation condition (14). Naturally, for the same experiment, the value \(\phi_f\) must be lower than the one adopted for \(\phi_i\). Let us also note that (16) can be obtained from the ratio of conjugate depths in a hydraulic jump. In this sense the value \(F = 1\) means that there is no jump and as \(F\) increases we observe a transition from a smooth undular jump to a undular jump characterized by shock waves and breaking phenomena. The application of Froude number to wave breaking is based on the relation between the increasing Froude number and the development of wave breaking mechanism in a hydraulic jump. Using this analogies some authors have estimated that the transition from non-breaking wave to breaking wave occurs for \(F\) equal 1.36 [38].

If a breaking wave is identified we also need to define the extension of the region, denoted \(L_{nl}\), around the breaking point that is governed by \(S_{nl}(t)\), i.e., the Saint-Venant system. The common practice is to use a value proportional to the wave front height, \(L_{nl} = CH_w = C(H_{wc} - H_{wt})\), for \(C > 1\). The value of \(C\) must be sufficient larger to avoid that non-physical dispersion effects influence the break region, we have set \(C = 10\). In Figure 2 we give...
an illustration of the notation used in the breaking model. The proposed methodology can be summarized as follows:

- If (14) is false; solve \( S^{\alpha,\beta}(t) = S_{d}^{\alpha,\beta}(t) \circ S_{nl}(t) \).
- If (14) is true; a wave is breaking.
  Calculate \( L_{nl} \) and solve \( S^{\alpha,\beta}(t) = S_{nl}(t) \) in \( L_{nl} \) and \( S^{\alpha,\beta}(t) = S_{d}^{\alpha,\beta}(t) \circ S_{nl}(t) \) elsewhere.
- If (15) (or (16)) is false; the wave is still breaking.
  Calculate \( L_{nl} \) and solve \( S^{\alpha,\beta}(t) = S_{nl}(t) \) in \( L_{nl} \) and \( S^{\alpha,\beta}(t) = S_{d}^{\alpha,\beta}(t) \circ S_{nl}(t) \) elsewhere.
- If (15) (or (16)) is true; the wave stopped breaking.
  Solve \( S^{\alpha,\beta}(t) = S_{d}^{\alpha,\beta}(t) \circ S_{nl}(t) \).

Naturally, more than one breaking point can be identified in the computational domain. In that case all the breaking wave fronts are followed and the breaking algorithm is applied to each one of them.

3.2. Bottom Friction. Bottom friction is an energy dissipation process consequence of the interaction between a viscous fluid (water) and a rough surface (sea-bottom). This process has a strong impact on erosion and sediment transport, two crucial aspects in coastal management. Friction is strongly influenced by several factors like sea-bottom roughness and morphology, water depth, and velocity. Another friction-inducing factor that is frequently overlooked by wave models is the current. This may not be relevant for laboratory experiments, but in real life applications any complete coastal model must take it into account. For instance, it is known that breaking waves, particularly surging breaking, induce strong currents in the surf zone.

In this work, we express bottom friction using the usual quadratic dependence on velocity, \( \sigma/\rho = \tau_{b}u|u| \), with \( \tau_{b} \) a friction coefficient. This parameterization can be seen as a particular case of the more general, \( \tau_{l}u + \tau_{t}hu|u| \), with \( \tau_{l} \) and \( \tau_{t} \) laminar and turbulent coefficients, respectively.

Most of the formulations used to express \( \tau_{b} \) are based on empirical laws. For example, from laboratory experiments with uniform turbulent flow in open-channel, Manning [33] presented the following law

\[
S = \frac{n_{k}^{2}u|u|}{R^{4/3}},
\]

where \( S \) is the open-channel slope, \( n_{k} \) is the so-called Manning’s coefficient, related with the channel roughness, and \( R \) is the hydraulic radius, the ratio
between the water area $A$ and the wetted perimeter $P$. To realize how $S$ is related with bottom friction let us consider a open-channel with bottom angle $\zeta$ and length $L$, and let us assume that the only forces are the water impelling force $F_g$, given by,

$$F_g = g\rho AL \sin(\zeta),$$

and the opposite resisting force $F_b$ due to bottom friction,

$$F_b = \sigma PL.$$

Since the flow is uniform the two forces are of equal magnitude and we easily deduce

$$\sigma = g\rho R \sin(\zeta).$$

Using Manning’s equation (17) to evaluate $S$, together with the usual approximation $R \approx h$ and the small-angle condition $\sin(\zeta) \approx S$, we get,

$$\frac{\sigma}{\rho} = g \frac{n_k^2}{h^{1/3}} u |u| = \tau_b u |u|. \quad (18)$$

The friction coefficient $\tau_b$ defined by (18) is the most widely used in the literature.

More well founded parameterizations can however be obtained. One such option is presented next. Following [17], we set

$$\frac{\sigma}{\rho} = \frac{1}{2} \tau_{cw} u |u|,$$

with $\tau_{cw}$ a coefficient that accounts for current ($\tau_c$) and wave ($\tau_w$) friction. Denoting by $u_c$ the current velocity we define $\tau_{cw}$ by

$$\tau_{cw} = \frac{|u_c|}{|u_c| + |u|} \tau_c + \frac{|u|}{|u_c| + |u|} \tau_w.$$

This expression can be extended to accommodate the friction arising from the pressure gradient acting on the bottom roughness ($\tau_p$). In this case, it reads

$$\tau_{cw} = \frac{|u_c|}{|u_c| + |u|} \tau_c + \frac{|u|}{|u_c| + |u|} (\tau_w + \tau_p).$$

Several expressions have been presented for the coefficients $\tau_c$, $\tau_w$, and $\tau_p$. We adopt the ones of [19]. They are built on the work developed in [47, 23, 43], which is devoted to the analysis of a two-equation turbulent boundary layer.
model of the \( k-l \) type [23]. For a single current and rough turbulent flow the suggest parameterizations are

\[
\tau_c = \frac{k^2}{\left( \ln \left( \frac{h}{z_0} \right) - 1 \right)^2}, \quad \tau_w = c_1 \exp \left( c_2 \left( \frac{K_N}{A} \right)^{c_3} \right), \quad \text{and} \quad \tau_p = c_4 \left( \frac{K_N}{A} \right),
\]

with \( k \) the von Kármán’s constant \( (k = 0.4) \), \( z_0 \) the roughness length \( (z_0 = K_N/30) \), \( A = T |u|/2\pi \) with \( T \) the wave period, and \( c_i, i = 1, \ldots, 4 \), empirical coefficients. After experimental fitting they were set to \( c_1 = 0.00140 \), \( c_2 = 4.5840 \), and \( c_3 = 0.1340 \). Regarding \( c_4 \), it was set to 0.48 as given in [43]. For our laboratory like experiments we ignore current and pressure friction and we get term

\[
\frac{\sigma}{\rho} = \frac{1}{2} \tau_w |u| = \frac{1}{2} c_1 \exp \left( c_2 \left( \frac{K_N}{A} \right)^{c_3} \right) |u| |u|.
\]

The optimal value of the parameter \( K_N \) is problem dependent. It is usually assumed [15] that \( K_N \) and \( n_k \) (the Manning’s coefficient) are related by the equation

\[
n_k = \frac{1}{21.1} \left( \frac{K_N}{2.5} \right)^{1/6}.
\]

4. Numerical Scheme

The splitting strategy (9) decouples the proposed Serre\( \alpha,\beta \) model in a hyperbolic part \( S_{nl}(t) \) and a dispersive part \( S_{d\alpha,\beta}(t) \). This separation allows us to use different methodologies suitable for each term. A brief description of our strategy is presented next.

4.1. Spatial Discretization. To solve the hyperbolic part we resort to the efficient high-order finite volume method presented in [8], see also [30]. This is a positivity-preserving scheme in the sense that preserves the positivity of water depth \( h \). This crucial physical restriction brings additional challenges for the numerical methods particularly near dry or almost dry conditions, i.e., \( h \sim 0 \).

Let us consider the one-dimensional domain \( \Omega = [x_L, x_R] \) and the uniform cell-centered finite volume mesh, \( \bar{x}_{i+1} = \bar{x}_i + \Delta x, i = 1, \ldots, N - 1, \) with \( \bar{x}_1 = x_L + \Delta x/2 \) and \( \bar{x}_N = x_R - \Delta x/2 \). Let us also define \( \bar{x}_{i+1/2} = \bar{x}_i + \Delta x/2, i = 1, \ldots, N, \) with \( \bar{x}_{1/2} = x_L \) and \( \bar{x}_{N+1/2} = x_R \). Taking the friction term
(19) and the change of variables $w = h + b$ and $q = hu$, we rewrite the hyperbolic system (5) in the conservative form

$$U_t + F(U, b)_x = N(U, b) + M(U, b),$$

with $U = (w, q)^T$,

$$F(U, b) = \left(q, \frac{q^2}{w-b} + \frac{1}{2}g(w-b)^2\right)^T, \quad N(U, b) = (0, -g(w-b)b_x)^T,$$

$$M(U, b) = \left(0, \frac{1}{2}c_1 \exp \left(c_2 \left(\frac{2\pi K_N(w-b)}{T|q|}\right)^{c_3}\right) \frac{1}{(w-b)^2} q|q|\right)^T,$$

with $c_1 = 0.00140$, $c_2 = 4.5840$, and $c_3 = 0.1340$. If we use Manning’s friction term (18), we get

$$M(U, b) = \left(0, -g n_k^2 \frac{1}{(w-b)^{7/3}} q|q|\right)^T.$$

With this notation, the proposed finite volume method takes the form

$$\frac{d}{dt} U_i(t) = -\frac{H_{i+1/2}(t) - H_{i-1/2}(t)}{\Delta x} + N_i(t) + M_i(t), \quad (21)$$

where $U_i(t)$ represents the numerical approximation at $\bar{x}_i$ and $H_{i+1/2}(t)$ are the numerical fluxes

$$H_{i+1/2}(t) = \frac{a^+_{i+1/2} F(U^-_{i+1/2}, b_{i+1/2}) - a^-_{i+1/2} F(U^+_{i+1/2}, b_{i+1/2})}{a^+_{i+1/2} - a^-_{i+1/2}}$$

$$+ \frac{a^+_{i+1/2} a^-_{i+1/2}}{a^+_{i+1/2} - a^-_{i+1/2}} \left(U^+_{i+1/2} - U^-_{i+1/2}\right). \quad (22)$$

Here, $U^+_{i+1/2} = \tilde{U}_i(x_{i+1/2})$ and $U^-_{i+1/2} = \tilde{U}_i(x_{i+1-1/2})$ with $\tilde{U}_i(x)$ defined as

$$\tilde{U}_i(x) = U_i + (U_x)_i(x - \bar{x}_i), \quad x_{i-1/2} < x < x_{i+1/2},$$

where $(U_x)_i$ denotes the numerical derivative calculated with the generalized minmod limiter. In (22), the local propagation speeds $a^\pm_{i+1/2}$ are determined by

$$a^+_{i+1/2} = \max\{u^+_{i+1/2} + \sqrt{gh^+_{i+1/2}}, u^-_{i+1/2} + \sqrt{gh^-_{i+1/2}}, 0\},$$

$$a^-_{i+1/2} = \min\{u^+_{i+1/2} - \sqrt{gh^+_{i+1/2}}, u^-_{i+1/2} - \sqrt{gh^-_{i+1/2}}, 0\}.$$
Note that \( u = q/h \), therefore to obtain an accurate approximation for \( u \) when \( h \sim 0 \) we use the formula

\[
    u = \frac{2(w - b)q}{(w - b)^2 + \max((w - b)^2, \epsilon^2)},
\]

with \( \epsilon = \min((\Delta x)^4, 1e-5) \). This formula changes the computation of \( u \) only when \( h < \epsilon \). Other formulas could be used without affecting the method robustness [30, 8]. At last, the source terms \( N_i(t) \) and \( M_i(t) \) are approximated by

\[
    N_i(t) = \left( 0, -g \frac{(b_{i+1/2} - b_{i-1/2})((w^-_{i+1/2} - b_{i+1/2}) + (w^+_{i-1/2} - b_{i-1/2}))}{2\Delta x} \right)^T
\]

and

\[
    M_i(t) = \left( 0, -\frac{1}{2}c_1 \exp \left( c_2 \left( \frac{2\pi K_N(w_i - b_i)|q_i|}{T(q_i^2 + \max(q_i^2, \epsilon^2))} \right)^{c_3} \right) \left( \frac{w_i - b_i}{(w_i - b_i)^2 + \max((w_i - b_i)^2, \epsilon^2)} \right)^2 q_i|q_i| \right)^T,
\]

or

\[
    M_i(t) = \left( 0, -g n_k^2 \left( \frac{w_i - b_i}{(w_i - b_i)^2 + \max((w_i - b_i)^2, \epsilon^2)} \right)^{7/3} q_i|q_i| \right)^T,
\]

for Manning’s friction. Note the use of a formula analogous to (23) to compute \( 1/(w - b) \) and \( 1/|q| \). In the following we denote by Serre\textsuperscript{M}_{\alpha,\beta} the combination of the proposed model with Manning’s friction.

\textbf{Remark 4.1} This is a second-order accurate scheme that reduces to first-order near non-smooth regions. In some cases it may be necessary to recalculate \( w_{i+1/2}^\pm \) to guarantee the positivity of the scheme. For ease of presentation, we refer again to [30, 8] for further details. Information about the linear dispersion properties of similar finite difference and finite volume methods can be found in [20, 56].

Now we address the dispersive part (8). We define the approximation points, \( x_{i+1} = x_i + \Delta x, i = 1, \ldots, N \), with \( x_1 = x_L \) and \( x_{N+1} = x_R \). Note
that these nodes coincide with the finite volume nodes $\bar{x}_{i \pm 1/2}$. Since $h_t = 0$, system (8) reduces to the equation

$$(I + T + T^{\alpha,\beta})u_t = g(T - T^{\alpha,\beta})(\eta_x) - Q(u),$$

with $T$, $T^{\alpha,\beta}$, and $Q$ the differential operators given by (3), (7), and (2), respectively. Applying standard second-order centered finite differences,

$$v_x(x_i) \approx \frac{v_{i+1} - v_{i-1}}{2\Delta x}, \quad v_{xx}(x_i) \approx \frac{v_{i+1} - 2v_i + v_{i-1}}{\Delta x^2}, \quad \text{and}$$

$$v_{xxx}(x_i) \approx \frac{-v_{i-2} + 2v_{i-1} - 2v_{i+1} + v_{i+2}}{2\Delta x^3},$$

we get the system of ordinary differential equations (odes)

$$\frac{d}{dt}u(t) = M_{FD}(u(t)). \quad (24)$$

where $M_{FD}$ denotes a linear finite difference operator (see [16] for details).

The hybrid approach finite difference finite volume means that some relations are required to carry the numerical approximations from one mesh to the other. Whenever necessary we use the following fourth-order approximations,

$$\bar{v}_i = -\frac{1}{24} v_{i-1} + \frac{13}{24} v_i + \frac{13}{24} v_{i+1} - \frac{1}{24} v_{i+2}$$

for the switch from finite difference to finite volume, and

$$v_i = -\frac{1}{12} \bar{v}_{i-2} + \frac{7}{12} \bar{v}_{i-1} + \frac{7}{12} \bar{v}_i - \frac{1}{12} \bar{v}_{i+1}$$

for the switch from finite volume to finite difference.

In the following, reflecting boundary conditions are always assumed. They are implemented considering two additional fictitious nodes at the right and at the left end of the computational domain. At the right we set $u_{N+1} = -u_{N-1}$ and $u_{N+2} = -u_{N-2}$ for the finite difference approximation and $w_N = w_{N-1}$, $w_{N+1} = w_{N-2}$, $q_N = q_{N-1}$, and $q_{N+1} = q_{N-2}$ for the finite volume approximation. At the left end of the domain we proceed similarly. When required, we simulate absorbing boundary conditions by expanding the computational domain by a size large enough.
4.2. Time Discretization. In the time domain $t \in [0, T]$ we consider the time mesh $t_m = t_{m-1} + \Delta t_m, m = 1, \ldots, M - 1$, with $t_0 = 0$ and $t_M = T$. From the numerical point of view the splitting strategy (9) is approximated by a second-order Strang scheme. Thus, from time level $t_{m-1}$ to $t_m$ the numerical solution associated to $\text{Serre}_{\alpha, \beta}$ is given by

$$(h^m, u^m) = S_{nl}(\Delta t_m/2) \circ S^\alpha_{d}(\Delta t_m) \circ S_{nl}(\Delta t_m/2)(h^{m-1}, u^{m-1}).$$

After the spatial discretization $S_{nl}(\cdot)$ and $S^\alpha_{d}(\cdot)$ refer to the systems of odes (21) and (24), respectively.

To solve (24) we use a second-order strong stability preserving Runge-Kutta method [22]. For the generic odes system

$$\frac{d}{dt} w(t) = L(w(t)),$$

it is defined by

$$w^{(1)} = w^m + \Delta t_m L(w^m)$$

$$w^{m+1} = \frac{1}{2} w^m + \frac{1}{2} w^{(1)} + \frac{1}{2} \Delta t_m L(w^{(1)}).$$

For the system of odes related with the hyperbolic part (21) we resort to a new class of semi-implicit Runge-Kutta methods [7]. This class of methods is specifically design for odes systems of the form

$$\frac{d}{dt} w(t) = F(w(t)) + G(w(t))w(t),$$

where $F(w(t))$ and $G(w(t))$ represent a non-stiff and a possibly stiff term, respectively. These methods treat the non-stiff term $F(w(t))$ explicitly and the stiff term $G(w(t))w(t)$ in a mixed way, $G(w(t))$ explicitly and $w(t)$ implicitly. Note that (21) can be written in the form (25) and the stiffness arises in the friction term $M(t)$ when $h \to 0$. As detailed in [7] this new class of semi-implicit schemes has the advantage of preserving the positivity of the spatial discretization. It is also simpler to implement when compared with traditional implicit schemes that treat the term $G(w(t))w(t)$ implicitly. Moreover, like in the implicit ones, the associated time step restriction depends only on the explicit discretization of the non-stiff term $F(w(t))$. For
(25), the adopted semi-implicit second-order method reads

\[
\begin{align*}
    w^{(1)} &= \frac{w^m + \Delta t_m F(w^m)}{1 - \Delta t_m G(w^m)} \\
    w^{(2)} &= \frac{3}{4} w^m + \frac{w^{(1)} + \Delta t_m^{(1)} F(w^{(1)})}{4(1 - \Delta t_m^{(1)} G(w^{(1)}))} \\
    w^{(3)} &= \frac{1}{3} w^m + \frac{2(w^{(2)} + \Delta t_m^{(2)} F(w^{(2)}))}{3(1 - \Delta t_m^{(2)} G(w^{(2)}))} \\
    w^{m+1} &= \frac{w^{(3)} - (\Delta t_m)^2 F(w^{(3)}) G(w^{(3)})}{1 + (\Delta t_m G(w^{(3)}))^2}.
\end{align*}
\]

For our finite volume method the Courant-Friedrich-L\'evy (CFL) time step stability condition is given by

\[
\Delta t_m \leq \frac{\Delta x}{2a}, \quad \text{with } a = \max_i \{a_i^+, -a_{i+1/2}^-\}.
\]

In (26), the notation \(\Delta t_m^{(j)}\) refers to the CFL condition associated with \(w^{(j)}\), \(j = 1, 2\). To ensure the positivity of the solution the time step \(\Delta t_m\) needs to satisfy \(\Delta t_m \leq \Delta t_m^{(j)}\), \(j = 1, 2\). Otherwise, we have to restart the time scheme with \(\Delta t_m = k\Delta t_m^{(1)}\) (or \(\Delta t_m = k\Delta t_m^{(2)}\), with \(k \in (0, 1)\); we used \(k = 0.9\).

5. Numerical Experiments

In this section we test our model as well as the proposed numerical implementation. We use a problem with known theoretical solution and several laboratory experiments. The laboratory experiments are divided in two groups: weakly dispersive waves and strongly dispersive waves.

In all the following experiments, and when friction is considered, the Manning coefficient is equal to 0.01 (m\(^{-1/3}\)s) or 0.02 (m\(^{-1/3}\)s). These are common values for the roughness of glass-type or concrete surfaces like the ones used in the experiments reported in this paper [20, 25, 40]. Regarding the breaking parameters, their estimation a priori is an unsolved problem and they have to be individually adjusted in order to reflect the particular characteristics of each experiment. Nevertheless, the values used in the following sections are similar to many values found in the literature [20, 25, 48, 29]. In what concerns the dispersion parameters \(\alpha\) and \(\beta\) their optimal value is also problem dependent. Based on our experimental work, the combination
$(\alpha, \beta) = (0.04, 0.07)$ seems to be a good initial guess. Some attempts have been made to estimate the optimal values of the dispersion parameters associated with improved Serre-type systems, see e.g., [14, 3]. However, the analysis is not problem-independent and several restrictions are made.

5.1. Solitary wave propagation. We start our numerical experiments with the propagation of a solitary wave over a flat bottom. For this simple case we can obtain an analytic solution for $\text{Serre}_{\alpha,\beta}$ with $\alpha$ and $\beta$ equal to zero, which corresponds to classical Serre. The solution is given by

$$h(x, t) = h_0 + a(\text{sech}(K(x - Ct - x_0)))^2, \quad u = C(1 - h_0/h),$$

where $a$ is the wave amplitude, $x_0$ is the initial position of the wave crest, $K = \sqrt{3a/(4h_0^2(h_0 + a))}$, and $C = c_0\sqrt{1 + a/h_0}$, with $c_0 = \sqrt{gh_0}$.

We use this example to illustrate the convergence rate of our numerical scheme. We define the domain $[0, 100]$ (m) and the constants $h_0 = 1$ (m), $a = 0.5h_0$, and $x_0 = 20$ (m). We run the simulation until $T = 15$ (s) using successively smaller mesh sizes, namely, $\Delta x = 0.2$ (m), $\Delta x = 0.1$ (m), $\Delta x = 0.05$ (m), and $\Delta x = 0.025$ (m). To measure the error we use the discrete $L^\infty$ norm $\|\cdot\|_\infty$. We denote by $E_\eta$ and $E_u$ the difference between the numerical solution and the theoretical solution of $\eta$ and $u$, respectively. Like in the following sections, the time step is fixed by the CFL condition (27).

In Figure 3 we plot the $\log(\|E_\eta\|_\infty)$ and $\log(\|E_u\|_\infty)$ versus $\log(\Delta x)$. The convergence rate is estimated by the slope of the best fitting least square line. The values obtained are 1.84 for $\eta$ and 1.82 for $u$. For this smooth example these values are in close agreement with the expected second order convergence rate.
5.2. **Weakly dispersive waves.** The next set of experiments illustrates the ability of our approach to accurately simulate wave breaking and propagation over dry areas. We focus on weakly dispersive waves, meaning that the parameters $\alpha$ and $\beta$ are set to zero. In all examples we use the breaking parameters $\phi_i = 0.5$ and $\phi_f = 0.4$. The parameter $F_f$ was not considered.

Also presented is a comparison between the results obtained with Manning’s friction formula (18) and those obtained with the proposed one (19). To perform this comparison we first tuned the Manning’s friction coefficient $n_k$. Afterwards, we used formula (20) to obtain the corresponding value of the parameter $K_N$. No attempt was made to optimize $K_N$ or the parameters $c_1$, $c_2$, and $c_3$ arising in (19).

5.2.1. Synolakis [46] - Solitary wave run-up and run-down over a sloping beach. Our first experiment refers to the run-up and run-down of a breaking solitary wave over a sloping beach [46]. The experimental setup is given in Figure 4, it shows the beach with a slope of 1 : 19.85 and the still water depth $h_0 = 1$ (m).

![Figure 4. Schematic representation of the setup used in the experiment of Synolakis (1987). The horizontal grey line corresponds to the initial still water depth, $h_0 = 1$ (m).](image)

The solitary wave with amplitude $a = 0.3$ (m) is initially centered around $x_0 = 25.5$ (m). Simulations were performed with $\Delta x = 0.1$ (m) and Manning’s coefficient $n_k = 0.02$. Snapshots of the wave at the time $t^* = t \sqrt{g/h_0}$ are given in Figure 5. They show the wave propagating before breaking, at $t^* = 15$, the breaking phase, at $t^* = 20$, and the run-up phase, at $t^* = 25$ and $t^* = 30$. A good agreement between numerical simulation and experimental data is observed in all phases.

The breaking phase occurs approximately between $t^* = 18$ and $t^* = 23$. This is well predicted by our model and is illustrated in Figure 6.

In 2012 the US National Tsunami Hazard Mitigation Program (NTHMP) conducted a model benchmarking workshop [35]. The models that passed a
series of pre-established tests were approved for tsunami inundation modeling. One of the benchmark tests was based on the Synolakis (1987) experiment. The models had to achieve a normalized root-mean-squared-error (RMSE) smaller than 10% at the times shown in Figure 5. In Table 1 we compare the error of our approach with the average error of the approved models. As can be seen, the error of our Serre$_{\alpha,\beta}$ is below the average error

<table>
<thead>
<tr>
<th>Run-up Phase - Normalized RMSE (%)</th>
<th>$t^* = 15$</th>
<th>$t^* = 20$</th>
<th>$t^* = 25$</th>
<th>$t^* = 30$</th>
<th>Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serre$_{\alpha,\beta}$</td>
<td>3</td>
<td>8</td>
<td>6</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>Serre$_{M,\alpha,\beta}$</td>
<td>4</td>
<td>10</td>
<td>5</td>
<td>3</td>
<td>6</td>
</tr>
<tr>
<td>NTHMP</td>
<td>7</td>
<td>9</td>
<td>6</td>
<td>4</td>
<td>7</td>
</tr>
</tbody>
</table>

Table 1. Comparison of the normalized RMSE with respect to the Synolakis (1987) experiment; proposed Serre$_{\alpha,\beta}$, Serre$_{M,\alpha,\beta}$, and mean of the four models presented in NTHMP (2012).
at all times. The results with \( \text{Serre}_{\alpha,\beta}^M \) are also given, and they show that the Manning’s friction formula is less accurate.

Snapshots of the run-down phase are given in Figure 7. Again, we can observe a good agreement between numerical simulation and experimental data.

![Snapshots of the run-down phase](image)

**Figure 7.** Comparison between numerical (solid line) and experimental data (dots) at four different times for the run-down phase of the Synolakis (1987) experiment.

The normalized RMSE for the run-down phase is given in Table 2. For this phase \( \text{Serre}_{\alpha,\beta} \) and \( \text{Serre}_{\alpha,\beta}^M \) achieved a similar performance. We note that the NTHMP benchmark test did not include the run-down phase.

<table>
<thead>
<tr>
<th>( t^* )</th>
<th>( \text{Serre}_{\alpha,\beta} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>55</td>
<td>5</td>
</tr>
<tr>
<td>60</td>
<td>5</td>
</tr>
<tr>
<td>70</td>
<td>3</td>
</tr>
<tr>
<td>80</td>
<td>3</td>
</tr>
<tr>
<td>Mean</td>
<td>4</td>
</tr>
</tbody>
</table>

**Table 2.** Normalized RMSE for \( \text{Serre}_{\alpha,\beta} \) with respect to the Synolakis (1987) experiment.

In Table 3 we analyze the sensitivity of our implementation to the mesh size \( \Delta x \). The same mean of normalized RMSE for the run-up and run-down phases prove the robustness against this parameter.

### 5.2.2. CADAM Project [34]

- Dam break flow over a triangular hump.

The CADAM (Concerted Action on Dambreak Modelling), was a European project whose main goal was to push forward the numerical simulation of dam break events [34, 28]. During the project duration a database of tests cases was created. One of such tests cases is shown in the top left image of
Table 3. Mean of normalized RMSE in the Synolakis (1987) experiment in function of $\Delta x$.

<table>
<thead>
<tr>
<th>Phase</th>
<th>$\Delta x = 0.1$ (m)</th>
<th>$\Delta x = 0.05$ (m)</th>
<th>$\Delta x = 0.025$ (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Run-up</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Run-down</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
</tbody>
</table>

Figure 8. The figure in the top left corner shows a schematic representation of the setup used in the CADAM project (2000). The dots indicate the gauges positions. The remaining figures show a comparison between numerical (solid line) and experimental data (dots) at the five gauges.

Figure 8. It consists of a reservoir with a gate located at $x = 15.5$ (m). The initial water depth upstream was 0.75 m and downstream the channel was dry. At time equal zero the gate is open and the water floods the dry portion of the channel interacting with a triangular obstacle. The reservoir is closed at the upstream end and open at the downstream end. To collect data seven gauges were placed across the reservoir.
The results of our simulation are given in Figure 8, and they illustrate the good agreement between numerical and experimental data. In Figure 9 we show snapshots of the numerical wave at four different times. The first two images, at $t = 13.53$ (s) and $t = 20.01$ (s), show the wave after interacting with the obstacle. Part of the wave overtakes the obstacle and other part is reflected creating a breaking wave traveling back in the upstream direction. At $t = 30.50$ (s) the breaking wave is now moving in the opposite downstream direction after being reflected in the solid wall at the upstream end. Finally, at $t = 41.60$ (s), the wave is no longer breaking and it is overtaking the triangular obstacle for the second time. For this test we used $\Delta x = 0.1$ (m) and $n_k = 0.01$.

In this example, as illustrated in Figure 10, $\text{Serre}_M^{\alpha,\beta}$ is slightly more accurate than $\text{Serre}_G^{\alpha,\beta}$. The most noticeable difference is at the last gauge, located after the obstacle at $x = 35.5$ (m).
The initial condition for $h$ is discontinuous across the dam location. To avoid numerical instabilities we use the expression

$$h(x, 0) = h_d + \frac{h_u - h_d}{2} \left(1 + \tanh \left(\frac{x_g - x}{\delta}\right)\right),$$

(28)

where $h_d$ and $h_u$ are the water depth downstream and upstream, respectively, $x_g$ is the gate location, and $\delta$ is a small positive parameter that we set equal to 0.1 [37].

5.2.3. Roeber et al. [40] - Solitary wave propagating over an exposed reef. In last example of this section we analyze the interaction of a solitary wave with an idealized fringing reef [40]. The experimental setup is shown in Figure 11. The outer reef has a slope of 1/12 and the still water depth of 2.5 (m) leaves the reef crest exposed by 6 (cm) and the reef flat submerged by 14 (cm) of water.

![Figure 11. Schematic representation of the setup used in the experiment of Roeber et al. (2010). The horizontal gray line represents the initial still water depth, $h_0 = 2.5$ (m).](image)

The solitary wave with amplitude $a/h_0 = 0.3$ and initially centered around $x_0 = 15$ (m), shoals over the outer reef and breaks just before the crest at around $t^* = 67$, with $t^* = t\sqrt{g/h_0}$. The flume is closed at both ends and a series of breaking and reflection events occur during the experiment. The comparison between the experimental data recorded at ten different gauges and the numerical simulation is depicted in Figure 12. The data starts at $t^* = 95.7$, after the first passage of the wave front by all the gauges. Overall, our model reproduces the experimental data with good accuracy. For later times, as the wave dynamics becomes more and more complex, some discrepancies start to appear. For this simulation we used $\Delta x = 0.1$ (m) and $n_k = 0.02$.

Wave profiles at four different times are given in Figure 13. At $t^* = 71.11$ we see the first interaction of the wave front with the still water in the reef flat. An hydraulic jump is originated and a downstream propagating bore is
seen at $t^* = 79.36$. At $t^* = 97.25$ the bore is already moving upstream after being reflected by the solid wall at the downstream end. By time $t^* = 120.02$ the bore has crossed the reef crest creating and hydraulic jump at the top of the outer reef. This wave dynamics is in accordance with the description presented in [40].

The comparison between $\text{Serre}^M_{\alpha,\beta}$ and $\text{Serre}_{\alpha,\beta}$ is given in Figure 14. Here, $\text{Serre}_{\alpha,\beta}$ is more accurate than the Manning based $\text{Serre}^M_{\alpha,\beta}$ at seven of the ten gauges. At the remaining three gauges $\text{Serre}^M_{\alpha,\beta}$ has a smaller RMSE in one of them, $x = 61.5$ (m), and the RMSE is identical at the other two, $x = 54.4$ (m) and $x = 65.2$ (m).
5.3. Highly dispersive waves. In this section we illustrate the ability of our approach to deal with wave breaking and highly dispersive effects simultaneously. The goal is to highlight the superiority of the improved Serre$\alpha,\beta$ over classical Serre. In the simulation of such type of processes, besides the breaking parameters, also the parameters $\alpha$ and $\beta$ become relevant. For the experiments of this section we used $F_f$ instead of $\phi_f$ for breaking termination.

5.3.1. Carmo et al. [18] - Dam break flow over a wet and flat channel. Our first test case is a laboratory dam break problem presented in [18]. As shown in the top left image of Figure 15, it consists of a reservoir with a gate located at $x = 3.8$ (m). The initial water depth upstream and downstream was 0.099 m and 0.051 m, respectively. The channel was open at the downstream end and closed at the upstream end. Three gauges were placed at the downstream portion of the channel.

To simulate this experiment we have used $\Delta x = 0.01$ (m) and the initial condition (28) with $\delta = 0.05$. The parameters $\alpha$ and $\beta$ were set equal to
Figure 15. The figure in the top left corner shows a schematic representation of the setup used in the experiment of Carmo et al. (1993). The dots indicate the gauges positions. The remaining figures show a comparison between numerical (solid line) and experimental data (dots) at the three gauges.

Figure 16. Snapshots of the simulated Carmo et al. (1993) experiment at two different times. The area between the dash lines corresponds to breaking regions.

0.04 and 0.1, respectively. For breaking we used $\phi_i = 0.43$ and $F_f = 1.32$, and we found the friction term to be negligible. The images of Figure 15 show that the numerical results are in close agreement with the laboratory data. The wave height is well predicted and no significant phase distortion is observed. Two wave snapshots at time $t = 2.42$ (s) and $t = 2.66$ (s) are given in Figure 16. They show the front wave breaking as reported in [18].

In Figure 17 we give the numerical results obtained with classical Serre at the last two gauges. Here the dispersive effects are more relevant and a significant phase discrepancy is observed. Note that the height of the breaking front wave is well predicted. This suggests that the differences observed are in fact due to the weakly dispersive properties of Serre. At the first gauge the results are similar to the ones obtained with Serre$_{\alpha,\beta}$. 
5.4. Beji et al. [1] - Periodic wave propagation over a submerged bar. Our last experiment deals with the propagation of a sinusoidal wave over a submerged bar [1]. The setup is illustrated in the top left image of Figure 18. It shows the bar with a front slope of 1:20 and a back slope of 1:10. The initial water height is $h_0 = 0.4$ (m), leaving the 0.3 (m) high bar completed submerged. The incident wave height is 0.027 (m), the frequency is $1/2.5$ (s), and the wavelength is 4.7904 (m). This challenging case involves wave shoaling, wave breaking, and higher harmonic release.

To simulate the incident wave we used an input condition at the left boundary [16]. The mesh size was set to $\Delta x = 0.015$ (m) and the breaking parameters were set to $\phi_i = 0.5$ and $F_f = 1.08$. We also defined $\alpha = 0.04$, $\beta = 0.07$, and the friction term $n_k = 0.02$. In Figure 18 we present the comparison
between Serre_{n,\beta} and experimental data collected with three gauges placed at the bar crest. The results reveal that the wave shape is well captured by our model.

**Figure 19.** Snapshots of the flow propagation in the Beji et al. (1993) experiment; the area between the two dash lines corresponds to the breaking region.

Two snapshots of the wave are given in Figure 19. They show one of the first moments after a wave front begins to break, at \( t = 21.71 \) (s), and one of last moments before the wave front stops breaking, at \( t = 23.50 \) (s). These results reveal that our model correctly predicts the onset and duration of breaking.

**Figure 20.** Comparison between numerical (solid line) and experimental data (dots) at the last three gauges. From left to right: Serre_{n,\beta} and classical Serre.

For the comparison between Serre_{n,\beta} and classical Serre we use the data recorded with three gauges placed after the bar crest. The results of the simulations are given in Figure 20. Unlike Serre, the proposed Serre_{n,\beta} is
capable of reproduce the experimental data with good accuracy. The limitations of Serre are clearly evident at the last gauge located at $x = 17$ (m). During the shoaling phase the wave accumulates higher harmonics that are released as highly dispersive waves during the breaking phase and on the downward slope. Therefore, the last gauge is the most challenging since the higher harmonic are already fully released. At the gauges of Figure 18 Serre and Serre$_{\alpha,\beta}$ have a similar behavior.

![Figure 21](image)

**Figure 21.** Comparison of the RMSE at the six gauges of the Beji et al. (1993) experiment; Serre$_M^{\alpha,\beta}$ (black line) and Serre$_{\alpha,\beta}$ (grey line).

The comparison between Serre$_M^{\alpha,\beta}$ and Serre$_{\alpha,\beta}$ is given in Figure 21. It shows that Serre$_{\alpha,\beta}$ results in a somewhat lower RMSE at all gauges except the first one.

6. Conclusion

In this paper we propose, implement, and test a complete wave model for nearshore and dam break simulation. Our approach is based on a Serre system with improved dispersion properties. We complement this system with a wave breaking methodology and friction terms.

For breaking we follow a well established strategy that consists in ignoring all the dispersive terms in the neighborhood of a breaking wave. This strategy takes advantage of our splitting scheme that separates the model in a hyperbolic part and a dispersive part. A high order positivity preserving finite volume method is used to discretize the hyperbolic part and finite difference are used to discretize the dispersive part. For friction we propose a more physically based alternative to the usual Manning formula. The numerical results suggest that the proposed formulation is more accurate than the one based in Manning.

Our model is tested against several experimental problems. We separate the problems in two groups: weakly dispersive waves and highly dispersive waves. In the first group we focus on the breaking and friction methodologies.
The model proved to be efficient and robust. The second group is particularly challenging and innovating since it involves wave breaking and dispersive effects. These complex examples reveal the limitations of classical Serre system and highlight the advantage of our improved Serre system.

References


José S. Antunes do Carmo
IMAR, Dep. of Civil Engineering, University of Coimbra, Portugal
E-mail address: jsacarmo@dec.uc.pt

José A. Ferreira
CMUC, Dep. of Mathematics, University of Coimbra, Portugal
E-mail address: ferreira@mat.uc.pt

Luís Pinto
CMUC, Dep. of Mathematics, University of Coimbra, Portugal
E-mail address: luisp@mat.uc.pt