Discontinuous-Galerkin discretization of a new class of Green-Naghdi equations
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Abstract

We describe in this work a discontinuous-Galerkin Finite-Element method to approximate the solutions of a new family of 1d Green-Naghdi models. These new models are shown to be more computationally efficient, while being asymptotically equivalent to the initial formulation with regard to the shallowness parameter. Using the free surface instead of the water height as a conservative variable, the models are recasted under a pre-balanced formulation and discretized using a nodal expansion basis. Independently from the polynomial degree in the approximation space, the preservation of the motionless steady-states is automatically ensured, and the water height positivity is enforced. A simple numerical procedure devoted to stabilize the computations in the vicinity of broken waves is also described. The validity of the resulting model is assessed through extensive numerical validations.

Introduction

Depth-averaged equations are widely used in coastal engineering for the simulation of nonlinear waves propagation and transformations in nearshore areas. The full description of surface water waves in an incompressible, homogeneous, inviscid fluid, is provided by the free surface Euler (or water waves) equations but this problem remains mathematically and numerically challenging. As a consequence, the use of depth averaged equations helps to reduce the three-dimensional problem to a two-dimensional problem, while keeping a good level of accuracy in many configurations.

Many Boussinesq-like models are used nowadays and a detailed review can be found in [44] and the recent monograph [43]. Denoting by $L$ the typical horizontal scale of the flow and $h_0$ the typical depth, the shallow water regime usually corresponds to the configuration where $\mu := \frac{h_0^2}{L^2} \ll 1$. If approximations of order $O(\mu^2)$ of the free surface Euler equations are furnished by the Boussinesq-type (BT equations in the following) equations, see [54, 56, 60] for instance, an additional smallness amplitude assumption on the typical wave amplitude $a$ is classically performed: $\varepsilon := \frac{a}{h_0} = O(\mu)$. This assumption often appears as too restrictive for many applications in coastal oceanography.
Removing the small amplitude assumption while still keeping all the \(O(\mu)\) terms, we obtain the so-called Green-Naghdi equations (GN equations in the following) [36], also referred to as Serre equations [66] or fully non-linear Boussinesq equations [80].

A large number of numerical methods have been developed in the past few years for the BT equations. Let us mention for instance some Finite-Difference (FDM in the following) approaches[56, 79, 69, 51], Finite-Element methods (FEM in the following) [48, 77, 70, 62], Finite-Volume discretizations (FVM in the following) for 1d equations [23], hybrid FDM / FVM [26, 27, 68, 73, 57], or even a purely 2d FVM discretization on unstructured meshes [39], allowing for mesh refinement and flexibility for large scale simulations.

As far as flexibility is concerned, the use of discontinuous-Galerkin methods (dG methods in the following) would appear as a natural choice. Indeed, this class of method provides several appealing features, like compact discretization stencils and hp-adaptivity, flexibility with a natural handling of unstructured meshes, easy parallel computation and local conservation properties in the approximation of conservation laws. A general review of dG methods for convection dominated problems is performed in [16]. Concerning the approximation of more general problems, involving higher-order derivatives, several methods and important developments have been proposed in recent years, following [5] on Navier-Stokes equations and [17] on convection-diffusion systems. A recent review is performed in [84] and a unified analysis can be found in [4], and [28, 29], respectively for elliptic problems and both 1st and 2nd order problems in the framework of Friedrichs’ systems.

The application of dG methods to the Saint-Venant equations (also called Nonlinear Shallow Water equations, NSW in the following) has recently lead to several improvements, see for instance [30, 3, 83, 82] and the recent review [21]. However, dG methods for BT equations have been under-investigated. In [31], a hp/spectral element model is introduced for the 1d enhanced equations of Nwogu [56], while the 2d equations of Peregrine [59] are studied in [32], in the flat bottom case, relying on a scalar reformulation that allowed some computational savings. This formulation is further investigated in [33], accounting for variable depth, and in [34] with the study of the enhanced equations of Madsen and Sorensen [53]. In [24, 25], an arbitrary order nodal dG-FEM method is developed for the set of highly-dispersive BT equations introduced in [52], respectively in 1d and 2d on unstructured meshes. These equations have a larger range of validity and can theoretically model fully nonlinear waves transformation, but they are also more complex, introducing a dependence in the vertical velocity, and consequently additional degrees of freedoms in the discrete approximations. As stated by the authors themselves, the bottleneck of their approach lies in the need of reconstruction and resolution of the large associated linear systems: this process can rapidly become a drawback for large-scale simulations.

Surprisingly, the GN equations have received far less attention. In the 1d framework and formulated in terms of free-surface elevation above the still water depth \(\zeta\) and horizontal velocity \(u\), these equations as derived in [2, 9], read as follow:

\[
\begin{aligned}
\left\{ \begin{array}{l}
\partial_t \zeta + \partial_x (hu) = 0, \\
\left[ 1 + \alpha \mathbb{T}[b, h] \right] \left( \partial_t (hu) + \partial_x (hu^2) + \frac{\alpha - 1}{\alpha} gh \partial_x \zeta \right) + \frac{1}{\alpha} gh \partial_x \zeta + h \mathcal{Q}_1(u) = 0,
\end{array} \right.
\end{aligned}
\] (1)

with the nonlinear differential operators

\[
\mathcal{Q}_1(u) = 2h \partial_x h (\partial_x u)^2 + \frac{4}{3} h^2 \partial_x u \partial_x^3 u + h \partial_x b (\partial_x u)^2 + h \partial_x^2 b u \partial_x u + (\partial_x \zeta \partial_x^2 b + \frac{h}{2} \partial_x^3 b) u^2
\]

and

\[
\mathbb{T}[b, h] = h \mathcal{T}[b, h] \left[ \frac{1}{h} \right],
\]

with for any scalar valued function \(w\):

\[
\mathcal{T}[b, h] w = -\frac{h^2}{3} \partial_x^2 w - h \partial_x h \partial_x w + (\partial_x \zeta \partial_x b + \frac{h}{2} \partial_x^3 b) w
\]

(2)
where $b(x)$ is a parametrization of the bottom variations, $h = \zeta + h_0 - b$ refers to the water height (see Fig. 1) and $\alpha$ is an optimization parameter.

A compact FVM approach is proposed in [13] and a FVM for a particular 2d flat bottom system in [46]. A pseudospectral approach is introduced in [58] and FEM discretization in [55] for the 1d case on flat bottom. Let us also mention [12] and more recently [67] for 2D cartesian numerical models based on the formulation of [80], respectively relying on FDM and FVM methods. In [9, 11, 72], a high-order accurate hybrid FVM-FDM model is introduced, embedded in a splitting approach. A robust treatment of moving shoreline and well-balancing for motionless steady states are ensured. The 2d extension of the hybrid FVM-FDM method on cartesian meshes has recently been performed in [45].

To the authors knowledge, the only dG method for GN equations has been recently proposed in [47], for the 1d equations with uneven bottom, relying on a centered dG approach and applications with second-order polynomial approximations, but the issue of preserving the water height positivity is not addressed. Still to our knowledge, no arbitrary order dG discretization of the GN equations has been proposed yet, and this is one of the goal of this work.

Another challenging issue is that although the GN equations provide a very good description of waves up to the breaking point, they are not entirely satisfactory as a nearshore flows simulation model. They are indeed unable to handle broken waves. To amend this important limitation, several approaches have been proposed. The most common one is to add an artificial viscous term to the momentum equation, whose role is to account for the energy dissipation that occurs during wave breaking (see for instance [14, 41, 12] for a related approach). This is also the method chosen in [25], embedded in the nodal-dG method. One of the drawbacks of this approach, as mentioned in [15], is that the computation remains very sensitive to the calibrations of the dissipation parameters.

An alternative approach to handle broken waves, which is becoming very popular, is to switch from the GN to the NSW equations in the vicinity of breaking waves. Indeed, NSW equations actually provide a very good description of broken waves [8], which are regarded as shock waves, and can therefore be accurately computed using shock-capturing methods. This switching strategy can be performed in a very simple fashion, provided the dispersive BT or GN equations are written in conservative forms. This approach is used for instance in [9, 75, 57, 63, 40, 67] where the occurrences of shock waves are accounted for through the use of FVM. Naturally, the method requires the introduction of some criteria devoted to detect waves which are about to break, and eventually to follow broken waves to detect if a "switch-back" to dispersive equations is needed. In [73] a criterium relying on the wave height to water depth ratio is introduced and subsequently used in [74] and [67] for instance. Another criterium based on the momentum gradient is successfully used in [63]. In [72], a relevant combination of the local energy dissipation, front slope and Froude number is proposed. Let us also mention the recent 2d approach of [40].

In the present work, we describe a strategy to compute discontinuous-Galerkin approximations of the solutions of some one-dimensional GN equations, including an efficient wave-breaking method. Indeed, the use of such fully nonlinear equations appear as a reasonable compromise between the weakly non-linear equations studied in [32, 33] and the highly-dispersive (and computationally costly) three-variables equations investigated in [24, 25]. Additionally, it is easily possible to extend the range of validity of the GN equations to moderately deep water by the introduction of some optimization parameters, as shown in [11, 45].

A closer look to equations (1) highlights that, if this formulation has real advantages (i.e. it does not involve any third order derivatives, and the presence of the second-order operator $I + \alpha T[b, h]$ makes the model very stable with respect to high frequency perturbations, see [9]), its main drawback is the time-dependency of this operator, through the water height $h$. Indeed, from a computational point of view, the dG discretization of $I + \alpha T[b, h]$ with a Local Discontinuous Galerkin approach [17] for higher order derivatives, and the associated linear system resolution, implies a global assembly process, for gathering the local discrete operators into the global one. And this should be done at each time step, or even sub step if high order time marching algorithms are used, together with the
corresponding algebraic system resolution.

To overcome this, the strategy recently proposed in [45], and also applied in this work, consists in deriving a new class of GN equations, asymptotically equivalent to the formulation (1) but for which the time dependency of the operator $I + \alpha T[b, h]$ is removed. The resulting model shares the same level of accuracy with the original GN equations, with respect to the approximation of the full water waves equations, but allows to build the global discrete operator in a pre-processing step. This operator is then used throughout all the computation, leading to dramatic computational savings.

As a consequence, the computational effort can be oriented towards a high order of accuracy in space and time, and the enforcement of some essential robustness properties. An arbitrary order of accuracy in space is obtained through the use of the Legendre polynomials hierarchical basis, and low storage strong-stability preserving Runge-Kutta methods (SSP-RK in the following) are used for the time discretization. The whole model is shown to exactly preserve the motionless steady states, thanks to the pre-balanced reformulation of the surface gradient term and suitable interface fluxes [22], and a robust treatment is implemented for the moving shoreline, based on the enforcement of an element-wise water height positivity preservation property, borrowing the recent accuracy-preserving method introduced for the dGM in [85, 83].

We lastly describe a very simple way to stabilize the computations in the vicinity of possibly occurring steep wave fronts, relying on the GN-NSW switching method. The strong super-convergence criteria introduced in [42] is used to detect such wave fronts and locally suppress the dispersive effects, which can be done in a natural fashion, as the dispersive part of the GN equations is regarded as an additional source term in the NSW equations. The corresponding shock waves are subsequently computed using the natural shock-capturing abilities of the dG approach, combined with an efficient local limiting process.

To summarize, the main features of the 1d dG-based GN numerical model described in this work are:

- an improved computational efficiency when compared to the original equations (1), due to the time-independency of the involved regularizing second order dispersive operator,
- an arbitrary order of spatial accuracy,
- a well-balanced property for the motionless steady states,
- a robust treatment of the shoreline motions,
- a simple way of stabilizing the computations when steep wave fronts appear, based on a GN/NSW equations switching strategy.

This paper is organized as follows: in a first part we detail the 1d derivation of a new family of GN model, following the strategy detailed above. The second part is devoted to the description of a robust arbitrary order of accuracy nodal dG approach for this family of model, ensuring the preservation of motionless steady states. The emphasize is put on the discretization of the higher order derivatives. Finally, the accuracy and robustness of the resulting numerical model are assessed through extensive 1d numerical validations. The main differences with [9, 72] are consequently the considered equations and the chosen numerical method. Two-dimensional validations are left for future works.

1 Derivation of the model

1.1 A new class of GN equations

In this section, we detail the simplified 1d derivation of a new family of GN equations for which the time dependency of the second-order differential operator $I + \alpha T[b, h]$ is amended. Choosing $h_0$ a reference water depth, we recall that $\zeta(x, t)$ stands for the free surface elevation, $b(x)$ the variation of the bottom with respect to the rest state, $u$
the horizontal velocity, and \( h(x, t) = \zeta(x, t) + h_0 - b(x) \) the water height (see Fig. 1).

\[
\zeta(x, t) = h_0 - b(x) - h(x, t)
\]

**Figure 1: 1D configuration**

Defining \( \varepsilon = a/h_0 \) the nonlinearity parameter, \( \mu = h_0^2/L^2 \) the shallowness parameter, \( \beta = a_b/h_0 \) the bottom variations parameter, with \( a_b \) the typical bottom deformation amplitude, and choosing \( \alpha = 1 \) for the sake of simplicity, the 1d GN equations (1) can be written in their non-dimensionalized formulation:

\[
\begin{align*}
\partial_t \zeta + \varepsilon \partial_x (hu) &= 0, \\
[1 + \mu \tilde{T}[b, h]] (\partial_t (hu) + \varepsilon \partial_x (hu^2)) + h \partial_x \zeta + \mu \varepsilon h \tilde{Q}_1(u) &= 0,
\end{align*}
\]

where the non-dimensionalized water depth is \( h(x, t) = 1 + \varepsilon \zeta(x, t) - \beta b(x) \) and the non-dimensionalized differential operators are written as follows:

\[
\tilde{Q}_1(u) = 2h \partial_x h(\partial_x u)^2 + \frac{4}{3} h^2 \partial_x u \partial_x^2 u + \\
\beta \left( h \partial_x b(\partial_x u)^2 + h \partial_x^2 b \partial_x u + (\varepsilon \partial_x \zeta \partial_x^2 b + \frac{h}{2} \partial_x^2 b) u^2 \right),
\]

\[
\tilde{T}[b, h] = h \tilde{T}[b, h] \frac{1}{h},
\]

with for any scalar valued function \( w \):

\[
\tilde{T}[b, h]w = -\frac{h^2}{3} \partial_x^2 w - h \partial_x h \partial_x w + \beta (\varepsilon \partial_x \zeta \partial_x b + \frac{h}{2} \partial_x^2 b) w.
\]

Using the fact that

\[
\partial_t (hu) + \varepsilon \partial_x (hu^2) = -h \partial_x \zeta + O(\mu),
\]

we introduce the following decomposition

\[
\tilde{T}[b, h](\partial_t (hu) + \varepsilon \partial_x (hu^2)) = \Xi[h](\partial_t (hu) + \varepsilon \partial_x (hu^2)) + \tilde{Q}_2(\zeta) + O(\mu)
\]

with

\[
\Xi[h]w = -\frac{h^3}{3} \partial_x^2 \left( \frac{w}{h} \right) - h^2 \partial_x h \partial_x \left( \frac{w}{h} \right),
\]

\[
\tilde{Q}_2(\zeta) = -\beta (\varepsilon \partial_x \zeta \partial_x b + \frac{h}{2} \partial_x^2 b) \partial_x \zeta.
\]
We can therefore write the second equation of (3) as follows:

\[
[1 + \mu \Xi[h]] \left( \partial_t (hu) + \varepsilon \partial_x (hu^2) \right) + h \partial_x \zeta + \varepsilon \mu h \dot{Q}_1(u) + \mu h \dot{Q}_2(\zeta) = O(\mu^2).
\]  

(7)

Now, introducing the still water depth \( h_b = h_0 - \beta b \) (which depends on \( x \) but not on \( t \)), we observe that \( \Xi[h] \) can be seen as a perturbation of \( \Xi[h_b] \). More precisely, we have for any arbitrary scalar valued function \( w \):

\[
\Xi[h]w = \Xi[h_b]w - \dot{Q}_3[h^2 - h_b^2]w,
\]

with

\[
\dot{Q}_3[h^2 - h_b^2]w = \frac{1}{6} \partial_x (h^2 - h_b^2) \partial_x w + \frac{h^2 - h_b^2}{3} \partial_x^2 w - \frac{1}{6} \partial_x^3 (h^2 - h_b^2) w.
\]  

(8)

As a consequence, equation (7) can be reformulated as:

\[
[1 + \mu \Xi[h_b]] \left( \partial_t (hu) + \varepsilon \partial_x (hu^2) \right) - \mu \dot{Q}_3[h^2 - h_b^2] \left( \partial_t (hu) + \varepsilon \partial_x (hu^2) \right) + h \partial_x \zeta + \varepsilon \mu h \dot{Q}_1(u) + \mu h \dot{Q}_2(\zeta) = O(\mu^2).
\]  

(9)

Denoting that

\[
\partial_t (hu) + \varepsilon \partial_x (hu^2) = - [1 + \mu \Xi[h_b]]^{-1} (h \partial_x \zeta) + O(\mu),
\]  

(10)

and injecting it into \( \dot{Q}_3 \), we obtain the following non-dimensionalized model:

\[
\left\{ \begin{array}{l}
\partial_t \zeta + \varepsilon \partial_x (hu) = 0, \\
[1 + \mu \Xi[h_b]] \left( \partial_t hu + \partial_x (hu^2) \right) + h \partial_x \zeta + \varepsilon \mu h \dot{Q}_1(u) + \mu h \dot{Q}_2(\zeta) + \mu \dot{Q}_3[1 + \mu \Xi[h_b]]^{-1} (h \partial_x \zeta) = 0,
\end{array} \right.
\]  

(11)

Switching back to dimensionalized quantities, introducing the optimization parameter \( \alpha \) again, and dropping the \( O(\mu^2) \), we obtain the following family of 1d Green-Naghdi equations with a time-independent operator:

\[
\left\{ \begin{array}{l}
\partial_t \zeta + \partial_x (hu) = 0, \\
[1 + \alpha \Xi[h_b]] \left( \partial_t hu + \partial_x (hu^2) \right) + \frac{1}{\alpha} gh \partial_x \zeta + \frac{1}{\alpha} \partial_x^2 \zeta + \partial_x^2 \zeta + \frac{1}{\alpha} \partial_x^3 \zeta + g \partial_x^2 \zeta + g Q_3 \left[ 1 + \alpha \Xi[h_b] \right]^{-1} (gh \partial_x \zeta) = 0,
\end{array} \right.
\]  

(11)

where \( h_b, Q_1, Q_2 \) and \( Q_3 \) are obtained from (24), (1.1), (6b) and (8) replacing \( \varepsilon, \beta \) and \( \mu \) by 1 and dropping the \( [h^2 - h_b^2] \) in \( Q_3[h^2 - h_b^2] \). As for (1), this new formulation does not involve any 3rd-order derivatives on the variables.

1.2 Frequency Dispersion

Let us now briefly investigate the linear behavior of a small perturbation (\( \zeta, \dot{u} \)) to a constant state solution (\( \zeta, u \)) for equations (11) with a flat bottom. Coming back to primitive variables, and defining \( b = h_0 + \zeta \), the linear equations that govern these perturbations are

\[
\left\{ \begin{array}{l}
\partial_t \dot{\zeta} + \mu \partial_x \dot{\zeta} + h \partial_x \dot{u} = 0, \\
[1 - \frac{\alpha}{3} h_0^2 \partial_x^2] \left( \partial_t \dot{u} + \mu \partial_x \dot{u} + \frac{1}{\alpha} g \partial_x \dot{\zeta} \right) + \frac{1}{\alpha} g \partial_x \dot{\zeta} + \frac{h^2 - h_b^2}{3} \partial_x^2 \left[ 1 - \frac{\alpha}{3} h_0^2 \partial_x^2 \right]^{-1} \partial_x \dot{\zeta} = 0.
\end{array} \right.
\]  

(12)
When compared with the original GN equations (1), we can observe an additional linear contribution coming from the new operator $Q_3$. As a consequence, the linearized system (12) for our new model (11) is the same as the linearized system derived from the standard GN equations when we consider the linearization around the rest state $(\zeta, u) = (0, 0)$. This can be also observed considering the associated dispersion relation, looking for plane wave solution of the form $(\zeta^0, u^0)e^{i(kx - \omega t)}$ to (12). One finds the following dispersion relation

$$(\omega - kw)^2 = ghk^2 \frac{1 + \frac{\alpha - 1}{3} \frac{h_0^2k^2}{h_0 - h_0^2} - \frac{h_0^2 - h_0^2}{3} \frac{k^2}{1 + \frac{\alpha h_0^2}{3} k^2}}{1 + \frac{\alpha h_0^2}{3} k^2},$$

which is different from the classical dispersion relation associated with the GN equations (1) when the linearization is performed around a non trivial constant state $(\zeta, u) \neq (0, 0)$, due to the additional term $-\frac{h_0^2}{3} \frac{k^2}{1 + \frac{\alpha h_0^2}{3} k^2}$ in the numerator of the right-hand-side of (13). A rapid study of the numerator shows that $h_0$ should be far greater than $h_0$ to allow complex roots for (13); this does not happen in practice. We refer to [9, 11] for the optimal computation of $\alpha$. We take $\alpha = 1.159$ throughout this article.

**Remark 1.1.** We recall that there is no particular assumption of the nonlinearity parameter $\epsilon$, for the GN equations. Our new model, as for the original GN equations, is a fully nonlinear model. Additionally, the study above shows that the frequency dispersion properties are similar to the original GN equations, see Fig. 2 of [9] for the phase and group velocity errors on the range $kh_0 \in [0, 4]$.

**Remark 1.2.** The simplest strategy to improve the computational efficiency in the approximation of GN equations is certainly to avoid the inversion of any linear system. This can be easily achieved, while keeping the $O(\mu^2)$ accuracy, by directly using (5) into (3) and write

$$\partial_t(hu) + \epsilon \partial_x(hu^2) + h\partial_x\zeta + \epsilon \mu hQ_1(u) - \mu \tilde{T}[b, h](h\partial_x\zeta) = O(\mu^2).$$

However, by doing so, we would loose the stabilizing effects associated to the inversion of $[1 + \mu \tilde{T}[b, h]].$ Indeed, the associated linear dispersion relation reads as follows

$$(\omega - kw)^2 = ghk^2(1 - \frac{1}{3} h_0^2 k^2),$$

which admits complex roots as soon as $h_0k$ is large enough.

**Remark 1.3.** We refer the reader to [45] for a generalized 2d derivation and further optimizations, with additional remarks on the linear stability issues, and in particular on the reason why we substitute relation (10) in (9) instead of (5).

### 1.3 Pre-balanced formulation and dry areas

Before describing in details the dG discretization of (11), let us reformulate the model under the pre-balanced formulation, in order to adapt the FVM well-balanced discretization introduced in [22, 50] and handle the occurrence of dry areas. The main ideas are, firstly, to use the free surface elevation instead of $h$ as a flow variable, and secondly to introduce an alternative splitting and redistributing of the free surface gradient term $gh\partial_x\zeta$, exploiting the deviations from the system’s unforced equilibrium. This method was first introduced in [64] for the NSW equations in the framework of FV methods. However, the total free surface $\zeta$ can be undefined in dry areas, in
the case of emerging topography. This approach is therefore modified to handle the occurrence of dry areas in [49, 50], using the free surface elevation above the datum \( \eta = h + b \) instead of \( \zeta \). Note that \( \eta \) is well defined in dry areas \( (h = 0) \) and simply reduces to \( b \). More precisely, the following splitting of the hydrostatic pressure term is introduced:

\[
g h \partial_x \zeta = \frac{1}{2} g \partial_x (\eta^2 - 2\eta b) + g \eta \partial_x b,
\]

leading to the following formulation of the NSW system:

\[
\partial_t w + \partial_x F(w, b) = \mathcal{B}(w, b),
\]

with \( w = t(\eta, q) \), \( q = hu \) and

\[
F(w, b) = \begin{pmatrix} F_q(w) \\ \mathcal{F}_q(w, b) \end{pmatrix} = \begin{pmatrix} \frac{q}{\eta - b} + p(\eta, b) \\ 0 \end{pmatrix},
\]

\[
\mathcal{B}(w, b) = \begin{pmatrix} \mathcal{B}_q(w) \\ \mathcal{B}_q(w, b) \end{pmatrix} = \begin{pmatrix} 0 \\ -g \eta \partial_x b \end{pmatrix},
\]

and

\[
p(\eta, b) = \frac{1}{2} g \eta (\eta - 2b).
\]

Simple and robust well-balanced schemes for the NSW equations relying on this pre-balanced formulation are proposed for instance in [78, 22]. Using dG approximations for the NSW equations, we show in the next section that this formulation can also provide a perfect balance between fluxes and the topography source term, providing that the corresponding integral terms are computed exactly. We notice that using (15), the second equation of (11) can be equivalently written as follows:

\[
\partial_t q + \partial_x \left( \frac{q^2}{\eta - z} + p(\eta, b) \right) + \mathcal{D}_c(\eta, u) = -g \eta \partial_x b,
\]

with

\[
\mathcal{D}_c(\eta, u) = \left[ 1 + \alpha T[h_b] \right]^{-1} \left( \frac{1}{\alpha} g h \partial_x \eta + h(\mathcal{Q}_1(u) + g \mathcal{Q}_2(\eta)) + g \mathcal{Q}_3\left( \left[ 1 + \alpha T[h_b] \right]^{-1} (g h \partial_x \eta) \right) - \frac{1}{\alpha} g h \partial_x \eta, \right.
\]

where \( u = \frac{q}{\eta - z} \) is kept in the writing of \( \mathcal{D}_c(\eta, u) \) for the sake of simplicity.

**Remark 1.4.** The original system (1) can of course also be written as follows:

\[
\partial_t q + \partial_x \left( \frac{q^2}{\eta - z} + p(\eta, b) \right) + \mathcal{D}_o(\eta, u) = -g \eta \partial_x b,
\]

with

\[
\mathcal{D}_o(\eta, u) = \left[ 1 + \alpha T[h_b] \right]^{-1} \left( \frac{1}{\alpha} g h \partial_x \eta + h \mathcal{Q}_1(u) - \frac{1}{\alpha} g h \partial_x \eta. \right.
\]

From this, equations (1) and (11) are nothing but the pre-balanced NSW equations (16) supplemented by a dispersive source term:

\[
\partial_t w + \partial_x F(w, z) = \mathcal{B}(w, z) + \mathcal{D}(w),
\]
with
\[
\text{model (1) : } \mathcal{D}(w) = \begin{pmatrix} 0 \\ -\mathcal{D}_c(\eta, u) \end{pmatrix}, \tag{23a}
\]
\[
\text{model (11) : } \mathcal{D}(w) = \begin{pmatrix} 0 \\ -\mathcal{D}_c(\eta, u) \end{pmatrix}. \tag{23b}
\]

**Remark 1.5.** We notice that the new model (11) is derived under the implicit assumption that \( h_b \geq 0 \). To obtain a model which is defined even with the occurrence of dry states, and is able to handle the case of emerging topography, we slightly modify the definition of \( h_b \) as follows:
\[
h_b = \max(h_0 - \beta_b, \epsilon_b), \tag{24}
\]
where \( \epsilon_b \) is a freely chosen threshold that ensures that the new quantity \( h_b \) remains positive in dry areas, preserving the regularizing effects of the elliptic operator \( 1 + \alpha T[h_b] \). We highlight that this manipulation does not create any additional consistency error in the model, since it is compensated by a similar change in the definition (8) of \( Q_3 \). In the validations of §3 we use \( \epsilon_b = 0.1 \).

## 2 Discontinuous Galerkin discretization

To implement a discontinuous Galerkin scheme, let \( \mathcal{T}_h \) be a partition of the computational domain \( \Omega = [L, R] \) into \( N_e \) non-overlapping elements, denoted \( E^j \), \( 1 \leq j \leq N_e \). The element \( E^j = [x^j_1, x^j_r] \) has a length \(|E^j|\), a boundary \( \partial E^j \) (reduced to the 2 boundary nodes \( x^j_1 \) and \( x^j_r \)) and a unit outward normal \( \hat{n}^j \) (reduced to +1 and −1 respectively at the right and the left boundary). Let \( x \) be the coordinate in \( \Omega \) and we denote \( h = \max_{1 \leq j \leq N_e} |E^j| \). We aim at computing an approximated solution, denoted \( w_h = (\epsilon, q_h) \), on this partition. Let us define
\[
\mathcal{V}_h := \{ v \in L_2(\Omega) \mid \forall E \in \mathcal{T}_h, v_{|E} \in \mathbb{P}^N(E) \}, \tag{25}
\]
where \( \mathbb{P}^N(E) \) denotes the space of polynomials in \( E \) of degree at most \( N \). We consider only the new model (11) but the subsequent weak formulation and discretization can of course be straightforwardly applied to the original model (1).

### 2.1 Weak formulation

Due to the number of terms and derivatives involved in the dispersive source term \( \mathcal{D}^{pbc}(h_b)(\eta, u) \), trying to write equation (18) as a first-order system, with the use of several auxiliary variables, would only bring up some unnecessary complexity in the formulation. Instead, we reformulate (18) as follows:
\[
\partial_t \epsilon + \partial_x q = 0, \tag{26a}
\]
\[
\partial_t q + \partial_x \left( \frac{q^2}{\eta - z} + p \right) + \mathcal{H} = -g\eta \partial_x b, \tag{26b}
\]
with \( \mathcal{H} \) is an auxiliary scalar valued variable, obtained as the solution of the following auxiliary problem:
\[
[1 + \alpha \Xi[h_b]](\mathcal{H} + \frac{1}{\alpha} gh \partial_x \epsilon) = \mathcal{K}, \tag{27a}
\]
\[
\mathcal{K} = \frac{1}{\alpha} gh \partial_x \epsilon + h \left( Q_1(u) + g Q_2(\eta) \right) + g Q_3(\mathcal{L}), \tag{27b}
\]
\[
[1 + \alpha \Xi[h_b]] \mathcal{L} = gh \partial_x \epsilon. \tag{27c}
\]
From this, a weak formulation of the problem (26) is obtained by multiplying by test functions \((\phi, \pi) \in \mathcal{V}_h^2\) and integrating over a given element \(E^j\); the flux term is integrated by parts. We obtain: \(\text{find } \mathbf{w}_h = (\eta_h, q_h) \in (\mathcal{V}_h)^2\) such that, \(\forall (\phi, \pi) \in (\mathcal{V}_h)^2\), and \(\forall E^j \in \mathcal{P}_h\) we have:

\[
\int_{E^j} \partial_t \eta_h \phi_h \, dx - \int_{E^j} q_h \phi_h \, dx + \left[q_h \phi_h\right]_{x_l}^{x_r} = 0, \tag{28}
\]

\[
\int_{E^j} \partial_t q_h \pi_h \, dx - \int_{E^j} \mathcal{F}_q(\mathbf{w}_h, z_h) \pi_h \, dx + \left[\mathcal{F}_q(\mathbf{w}_h, z_h) \pi_h\right]_{x_l}^{x_r} = \int_{E^j} \mathcal{B}_q(\mathbf{w}_h, b_h) \pi_h \, dx - \int_{E^j} H_b \pi_h \, dx. \tag{29}
\]

### 2.2 Semi-discrete formulation

On each element \(E^j\), we write \(\forall x \in E^j, \forall t \in [0, t_{max}]\):

\[
\eta_{h|E^j}(x, t) = \sum_{i=1}^{N_d} \tilde{\eta}_i^j(t) \theta_i^j(x), \quad \text{and} \quad q_{h|E^j}(x, t) = \sum_{i=1}^{N_d} \tilde{q}_i^j(t) \theta_i^j(x), \tag{30}
\]

where \(N_d = N + 1\) is the number of freedom degrees per element, \(\{\theta_i^j\}_{i=1...N_d}\) is a polynomial expansion basis for \(\mathbb{P}^N(E^j)\), and

\[
\tilde{\eta}^j = 4(\tilde{\eta}_1^j(t), \ldots, \tilde{\eta}_{N_d}^j(t)), \quad \tilde{q}^j = 4(\tilde{q}_1^j(t), \ldots, \tilde{q}_{N_d}^j(t)),
\]

are the local expansion coefficients vectors. In the following, \(\{\theta_i^j\}_{i=1...N_d}\) refers to the local Lagrange interpolating polynomial basis on the element \(E^j\), defined on the Legendre-Gauss-Lobatto (LGL) set of \(N_d\) nodes (a nodal-element basis). We consider the local polynomial expansion for the topography parameterization \(b_{h|E^j}(x) = \sum_{i=1}^{N_d} \tilde{b}_i^j \theta_i^j(x)\), and we assume that a local expansion is available for \(H_b\), given by:

\[
\tilde{H}^j = 4(\tilde{H}_1^j(t), \ldots, \tilde{H}_{N_d}^j(t)).
\]

Injecting these expansions into the weak formulation (28)-(29), replacing the test functions \((\phi_h, \pi_h)\) by the local basis functions and classically substituting numerical flux functions \(\tilde{q}\) and \(\tilde{\mathcal{F}}\) to the discontinuous intercell flux functions arising in the interface terms, we obtain the semi-discrete formulation of our dG approximation:

\[
\sum_{i=1}^{N_d} d \frac{dt}{dx} \tilde{\eta}_i^j(t) \mathbf{M}_{ik}^j - \sum_{i=1}^{N_d} \tilde{q}_i^j(t) \mathbf{S}_{ik}^j + \left[\tilde{\mathcal{F}} \theta_k^j\right]_{x_l}^{x_r} = 0, \tag{31}
\]

\[
\sum_{i=1}^{N_d} d \frac{dt}{dx} \tilde{\eta}_i^j(t) \mathbf{M}_{ik}^j - \int_{E^j} \mathcal{F}_q(\mathbf{w}_h, z_h) \frac{d}{dx} \theta_k^j \, dx + \left[\mathcal{F}_{\theta_k^j}\right]_{x_l}^{x_r} = \int_{E^j} \mathcal{B}_q(\mathbf{w}_h, b_h) \theta_k^j \, dx - \sum_{i=1}^{N_d} \tilde{H}_i^j(t) \mathbf{M}_{ik}^j, \tag{32}
\]

for \(1 \leq k \leq N_d\), and \(1 \leq j \leq N_e\), where

\[
\mathbf{M}_{ik}^j = \int_{E^j} \theta_i^j(x) \theta_k^j(x) \, dx \quad \text{and} \quad \mathbf{S}_{ik}^j = \int_{E^j} \theta_i^j(x) \frac{d}{dx} \theta_k^j(x) \, dx
\]

are respectively the local mass and stiffness matrix coefficients on \(E^j\).
Remark 2.1. As \( \{ \theta_i^j \}_{i=1,...,N_d} \) refers to the Lagrange interpolating polynomials, the expansion coefficients represent the values of the considered approximated quantity at the Gauss-Legendre-Lobatto set of nodes. The local expansion for the topography parameterization \( \tilde{b}^j = (\tilde{b}_1^j, \ldots, \tilde{b}_{N_d}^j) \) is therefore obtained just by “reading” the values of \( b \) at the LGL nodes.

Remark 2.2. In (32), the integrals \( \int_{\mathcal{E}_j} F_\eta(w, b_h) \frac{d}{dx} \theta_i^j dx \) and \( \int_{\mathcal{E}_j} B_q(w_h, b_h) \theta_i^j dx \) have to be computed exactly when motionless steady states are reached, with a suitable quadrature formula, in order to obtain a well-balanced scheme, as shown in \( \S 2.5 \). On the other hand, the computation of \( \tilde{\mathcal{H}}^{j} \), which involves higher order derivatives, is done directly, in a collocation way. This is temporarily postponed to \( \S 2.3 \).

Remark 2.3. At a given interface, the computation of the exchanging flux term \( \tilde{q} \) and \( \tilde{\mathcal{F}} \) is classically performed with numerical flux functions \( F^0 \) and \( F^q \), which are Lipschitz continuous, monotone and consistent with the exact flux \( F(w, b) = (F_w(w), F_q(w, b)) \). To ensure the positivity of the water height, we also need to use a water height positivity preserving flux (see \( \S 2.7 \)). We use the global Lax-Friedrichs flux in the numerical validations of \( \S 3 \). These interface fluxes computations, carefully chosen to ensure the preservation of motionless steady-states, are detailed in \( \S 2.5 \).

2.3 High-order derivatives and dispersive terms computation

We are now left with the resolution of systems (27a)-(27c), to compute the expansion \( \tilde{\mathcal{H}} \). Although the semi-discrete formulation (31)-(32) could appear purely local, the computation of the dispersive term expansions \( \tilde{\mathcal{H}} \) requires a global assembly process, for gathering the local discrete operators into a global one, in the computation of the inverse of the operator \( 1 + \alpha \mathbb{T}[h_b] \). Additionally, this computation involves the discrete approximation of space derivatives up to \( 2^{nd} \) order. We use the Local Discontinuous Galerkin (LDG) approach [17] to compute these derivatives. To illustrate this procedure, let consider the following \( 2^{nd} \) order equation, for an arbitrary scalar valued function \( w \):

\[
\ell - \partial_x^2 w = 0 ,
\]

reformulated as a set of two coupled \( 1^{st} \) order equation using an auxiliary variable \( v \):

\[
v + \partial_x w = 0 , \quad \ell + \partial_x v = 0 .
\]

Then, multiplying by a test function \( \phi_h \in \Phi_h \) and integrating over an element \( \mathcal{E}_j \), we obtain the associated weak formulation:

\[
\int_{\mathcal{E}_j} v \phi_h - \int_{\mathcal{E}_j} w \phi_h + \tilde{w}_r \phi_h(x_j^l) - \tilde{w}_l \phi_h(x_j^l) = 0 , \\
\int_{\mathcal{E}_j} \ell \phi_h - \int_{\mathcal{E}_j} v \phi_h + \tilde{v}_r \phi_h(x_j^l) - \tilde{v}_l \phi_h(x_j^l) = 0 .
\]

To define the exchanging fluxes \( \tilde{w} \) and \( \tilde{v} \) at a given interface, we use the following fluxes [16]:

\[
\tilde{w} = w - \xi [w] , \\
\tilde{v} = v + \nu [v] + \lambda [w] ,
\]

with the interface average \( w = (w^+ + w^-)/2 \) and jump \( [w] = (w^+ - w^-)/2 \), \( w^- \) and \( w^+ \) respectively the right and left interface values of \( w \) (and similar notations for \( v \)). Taking \( \xi = \nu = \lambda = 0 \) gives the centered Bassi and Rebay (BR) fluxes [5] which usually provide sub-optimal convergence rates for odd values of \( N \), while the choice \( \xi = \nu = 1 \), corresponding to the Cockburn and Shu alternate upwind discretization (refers to as LDG flux in the following), may allow to recover the optimal convergence order.
Remark 2.4. Note that $\frac{\lambda}{b}[w]$ is a penalization term, with a $O(h^{-1})$ scaling. Taking for instance $\xi = \nu = 0$ and $\lambda \neq 0$ gives the stabilized centered fluxes (sBR), sometimes helping to recover optimal convergence rates. In the numerical validation of §3, we only investigate and compare the BR and LDG choices. Note that extensive comparisons of fluxes including their stabilized versions and the impact of the coefficient $\lambda$ are performed in [33] for weakly non-linear BT equations.

Let us now come back to (35). At the discrete level, starting from the $N_e$ expansion coefficients vectors $\{\tilde{w}^i\}_{i=1..N_e}$, which are gathered in a $N_d \times N_e$ vector $\tilde{W} = t(\tilde{w}^1, \ldots, \tilde{w}^{N_e})$, we aim at computing the $N_d \times N_e$ vector $\tilde{L} = t(\tilde{\ell}^1, \ldots, \tilde{\ell}^{N_e})$ of expansion coefficients for the $2^{nd}$ order derivative $\partial^2 w$.

This is done globally, using (36) to build the differentiation matrices, that also account for exchanging interface fluxes. Injecting $\tilde{w}^j$ into (35) and replacing $\phi_b$ by the local basis functions $\{\theta^i\}_{i=1,N_e}$, for all elements $\{\mathcal{E}^j\}_{i=1..N_e}$, we obtain the global discrete formulation:

$$
\begin{align*}
\mathcal{M}\tilde{V} &= \mathcal{S}\tilde{W} - (\mathcal{E} - \xi\mathcal{F})\tilde{W}, \\
\mathcal{M}\tilde{L} &= \mathcal{S}\tilde{V} - (\mathcal{E} + \nu\mathcal{F})\tilde{V} - \frac{\lambda}{b}\tilde{W},
\end{align*}
$$

(37)

where the square $N_d \times N_e$ global mass and stiffness matrices $\mathcal{M}$ and $\mathcal{S}$ have a block-diagonal structure

$$
\mathcal{M} = \begin{pmatrix} M^1 & & \\
& \ddots & \\
& & M^{N_e} \end{pmatrix}, \quad \mathcal{S} = \begin{pmatrix} S^1 & & \\
& \ddots & \\
& & S^{N_e} \end{pmatrix},
$$

(38)

and the matrices $\mathcal{E}$ and $\mathcal{F}$ accounting for the inter-element exchanging fluxes have the following structure:

$$
\mathcal{E} = \begin{pmatrix} 1 & & \\
\frac{1}{2} & \frac{1}{2} & \\
-\frac{1}{2} & -\frac{1}{2} & \\
& \ddots & \\
\frac{1}{2} & \frac{1}{2} & \\
-\frac{1}{2} & -\frac{1}{2} & 1 \end{pmatrix}, \quad \mathcal{F} = \begin{pmatrix} 1 & & \\
-\frac{1}{2} & \frac{1}{2} & \\
\frac{1}{2} & -\frac{1}{2} & \\
& \ddots & \\
\frac{1}{2} & \frac{1}{2} & \\
\frac{1}{2} & -\frac{1}{2} & 1 \end{pmatrix}
$$

(39)

Note that the issue of boundary conditions is postponed to §2.8. We obtain a global discrete formulation of the first and second order derivative operators, based on the LDG approach (36):

$$
\tilde{V} = \mathbb{D}_x \tilde{W}, \quad \tilde{L} = \mathbb{D}_x^2 \tilde{W},
$$

(40)

with

$$
\mathbb{D}_x = \mathcal{M}^{-1}(\mathcal{S} - \mathcal{E} + \xi\mathcal{F}),
$$

(41)

$$
\mathbb{D}_x^2 = \mathcal{M}^{-1}\left((\mathcal{S} - \mathcal{E} - \nu\mathcal{F})\mathbb{D}_x - \frac{\lambda}{b}\mathcal{F}\right).
$$

(42)
Using these global differentiation matrices, we are now able to approximate all the derivatives occurring in (27a)-(27c). The nonlinear products are treated directly, in a collocation manner, inspired from [24]. We can also build the global square $N_d \times N_c$ matrix of the discrete version of $1 + \alpha \mathcal{X}[h_b]$. For instance, considering the simplified flat bottom case ($h_b = h_0$), we approximate the corresponding operator as follows:

$$1 + \alpha \mathcal{X}[h_0] = 1 - \alpha \frac{h_0^2}{3} \partial_x^2 \Rightarrow I - \alpha \frac{h_0^2}{3} \partial_x^2,$$

where $I$ is the $N_d \times N_c$ identity matrix. The locality of the LDG approach results in a sparse block-structure matrix, which is stored in a sparse format and LU-factorized at the beginning of the computation, in a pre-processing step. For the validations of $\mathcal{X}(3)$, the factorization and the resolution of the resulting triangular linear systems are performed using the unsymmetric multi-frontal method [19].

**Remark 2.5.** The same approach can be used for the original model (1), the only difference being that the $1 + \alpha \mathcal{X}[b, h]$ matrix has to be build and factorized at each time step (or substep).

**Remark 2.6.** The use of direct interpolation/collage methods for the computation of the nonlinear products in the dispersive source terms reduces the computational cost but can generate some aliasing which deteriorates the solution quality and can lead to instabilities. To amend this, we use a stabilization filtering method (mild nodal filter), as suggested in [24].

### 2.4 Time discretization

We use the method of lines, based on Strong Stability Preserving Runge-Kutta (RK-SSP) schemes [35], as time marching algorithms. Up to $N = 3$, RK-SSP schemes of order $N + 1$ are considered, to ensure equal orders of convergence in space and time. A $4^{th}$ order SSP-RK scheme is used for $N \geq 3$. At the fully discrete level, we denote $\Delta t^n$ the time step used to advance from discrete time value $t^n$ to $t^{n+1}$, and the local polynomial expansions at the discrete time $t^n$ as follows:

$$\eta_b^n(x) = \sum_{i=1}^{N_d} \tilde{\eta}_i^n \theta_i(x), \quad \text{and} \quad q_b^n(x) = \sum_{i=1}^{N_d} \tilde{q}_i^n \theta_i(x),$$

with the corresponding expansion vectors:

$$\tilde{\eta}_i^n = t(\tilde{\eta}_{1,n}^{i,n}, \ldots, \tilde{\eta}_{N_d,n}^{i,n}), \quad \tilde{q}_i^n = t(\tilde{q}_{1,n}^{i,n}, \ldots, \tilde{q}_{N_d,n}^{i,n}).$$

### 2.5 Preservation of motionless steady states

In this section, we detail the computation of the interface fluxes $[\mathcal{F}_b^j] \hat{\eta}_b^j$ appearing in the semi-discrete local discretization (32). To ensure the preservation of motionless steady states, we adapt to the dG framework the method introduced in[22, 49] for the FVM discretization of the pre-balanced SW equations. The following scheme can also be regarded as the adaptation to the pre-balanced formulation of the ideas introduced in [81].

Let us introduce the simplified notations:

$$w_r^- = w_b(x_r^i), \quad b_r^- = b_b(x_r^i), \quad w_r^+ = w_b(x_r^{i+1}), \quad b_r^+ = b_b(x_r^{i+1}),$$
$$w_l^- = w_b(x_l^i), \quad b_l^- = b_b(x_l^i), \quad w_l^+ = w_b(x_r^{i-1}), \quad b_l^+ = b_b(x_r^{i-1}).$$

13
(at the left and right boundaries of $\mathcal{E}^j$, the superscripts $-$ and $+$ stand respectively for the interior and exterior values). Dropping the subscripts, we define at each boundaries:

$$b^* = \max(b^-, b^+), \quad b = b^* - \max(0, b^+ - \eta^-),$$

$$\tilde{h}^* = \max(0, \eta^+ - b^*), \quad \tilde{\eta}^* = \tilde{h}^* + \tilde{b},$$

leading to new interior and exterior values:

$$\tilde{w}^- = (\tilde{\eta}^-, \frac{\tilde{h}^-}{\tilde{h}^- q^-}) \quad \text{and} \quad \tilde{w}^+ = (\tilde{\eta}^+, \frac{\tilde{h}^+}{\tilde{h}^+ q^+}),$$

with $h^\pm = \eta^\pm - b^\pm$. Then, the interface fluxes in (31)-(32) are computed as follows:

$$\tilde{q} = F_q(\tilde{w}^-, \tilde{w}^+),$$

$$\tilde{F} = F_q(\tilde{w}^-, \tilde{w}^+, \tilde{b}) - g\tilde{\eta}^-(\tilde{b}^- - \tilde{b}),$$

where $F = F_q(F_q, F_q)$ is the Lax-Friedrich flux, defined as follows for two arbitrary interface states $w_l, w_r$:

$$F_q(w_l, w_r) = \frac{1}{2} \left( (q_l + q_r) - \omega_0 (\eta_l - \eta_r) \hat{n} \right),$$

$$F_q(w_l, w_r, b_l, b_r) = \frac{1}{2} \left( (F_q(w_l, b_l) + F_q(w_r, b_r)) - \omega_0 (q_l - q_r) \hat{n} \right),$$

with $\omega_0 = \max_{1 \leq j \leq N_e} \left( \max(|\eta|, \sqrt{g(\eta - b)}) \right)$ and $\hat{n}$ the outward normal of the element. We point out that the modified fluxes (49)-(50) induce perturbations of order $N + 1$ when compared to the traditional interface fluxes $F_q(w^-, w^+)$ and $F_q(w^-, w^+, b^-, b^+)$ and do not modify the consistency and convergence.

**Remark 2.7.** $F$ can be replaced by any monotone and positive preserving numerical flux function, like the HLL flux [76] or the VFRoe-relaxation flux [7].

**Remark 2.8.** In (48), we implicitly assumed that we have $h^\pm > 0$. The case $h^\pm = 0$ (or numerically $h^\pm < \epsilon$) is classically handled by setting the corresponding values of $u^\pm$ and $q^\pm$ to 0. We set $\epsilon = 10^{-6}$ in the numerical validations of §3.

**Proposition 1.** The dG scheme (31)-(32), with the interface fluxes defined in (49)-(50), preserves the motionless steady states.

**Proof.** We adapt the ideas of [81] to the pre-balanced formulation (22). We want to show that

$$\frac{d}{dt} \tilde{\eta}^j(t) = 0 \quad \text{and} \quad \frac{d}{dt} \tilde{\theta}^j(t) = 0, \quad \text{for} \quad 1 \leq j \leq N_e.$$

If a motionless steady state is reached, that is

$$w_b = \tilde{t}(\eta_0, 0),$$

we have $\tilde{q}^j(t) = 0$ on each element and using (31), $\frac{d}{dt} \tilde{q}^j(t) = 0$ obviously holds. For the second equation (32), we need the residue

$$R = -\int_{\mathcal{E}^j} F_q(w_b, b) \frac{d}{dx} \theta_k^j dx + [\tilde{F} \theta_k^j]_{x^j}^e + g \int_{\mathcal{E}^j} \eta_b \frac{d}{dx} b \theta_k^j dx + \sum_{i=1}^{N_e} \tilde{\mathcal{H}}_i^j(t) \tilde{M}_i^j,$$
to vanish when a motionless steady state is reached, for \(1 \leq k \leq N_d\) and \(1 \leq j \leq N_c\). We first notice that the definition of the differential operators \(\{Q_i\}_{1 \leq i \leq 3}\), combined with (53), directly leads to \(\mathcal{H}(w_h) = 0\) at the discrete level, as a solution of (27a)-(27b)-(27c).

When (53) holds, (46)-(47)-(48) leads to \(\hat{q}^- = \hat{q}^+ = \hat{\eta}_0\) for both interfaces, and using (50), the interface flux \(\hat{F}(x_j^i)\) reduces to

\[
\hat{F}(x_j^i) = \frac{g}{2} (\eta_0^2 - 2\eta_0 \hat{b}_r) - g \eta_0 (b^-_r - \hat{b}_r) = \frac{g}{2} (\eta_0^2 - 2\eta_0 b^-_r) = F_q(w^-_r, b^-_r).
\]

We show in a similar way that, at the left boundary, \(\hat{F}(x_j^i) = F_q(w^-_l, b^-_l)\). Consequently we have,

\[
R = \int_{\mathcal{E}^j} \partial_x(F_q(w_h, b_h)) \theta^j_k dx + g \int_{\mathcal{E}^j} \eta_h \frac{d}{dx} b_h \theta^j_k dx = 0
\]
as \(w_h\) is a steady state solution, leading to \(\partial_x(F_q(w_h, b_h)) = -g \eta_h \frac{d}{dx} b_h\), and the integral terms are computed exactly.

\[\square\]

### 2.6 Handling broken waves and limiting strategy

Let us now consider the issue of wave steepening and breaking, which can lead to numerical instabilities. As stated in the introductory lines, we can find several methods in the literature since the last few years, that allow to handle broken waves in BT equations. They all rely on an accurate detection of wave fronts, near the breaking point, in the introductory lines, we can find several methods in the literature since the last few years, that allow to handle broken waves in BT equations. They all rely on some suitable physical parameters, like for instance the critical wave front slope or the energy dissipation, we choose in this work to use a simple (and purely numerical) smoothness detection criterion to identify the potential instability regions. Indeed, our goal here is only to stabilize the computations and avoid numerical instabilities due to the occurrence of steep front waves, while keeping things as simple as possible and avoiding a wave-by-wave tracking strategy.

The chosen criteria uses the ideas of the popular discontinuity detector of [42], which is based on a strong super-convergence property of the dG method at the outflow boundary of each element in smooth regions of the flow: smooth solutions with polynomial approximations of degree \(N\) locally converge at an enhanced \(O((\mathcal{E}^j)^{2N+1})\) rate at outflow boundaries of \(\mathcal{E}^j\), see [1]. To use this property, the following detector is proposed in [42]:

\[
\mathcal{J}_j = \frac{\Delta^- |\hat{h}_1^j - \hat{h}_{N_d}^{j-1}| + \Delta^+ |\hat{h}_{N_d}^j - \hat{h}_1^{j+1}|}{|\mathcal{E}^j| \|\hat{h}^j\|_\infty},
\]

where

\[
\Delta^- = \begin{cases} 1 & \text{if } \tilde{q}_1^j \geq 0, \\ 0 & \text{otherwise}, \end{cases} \quad \Delta^+ = \begin{cases} 1 & \text{if } \tilde{q}_{N_d}^j \leq 0, \\ 0 & \text{otherwise}, \end{cases}
\]

and \(\hat{h}^j = \hat{\eta}^j - \hat{b}^j\). Then, as we have \(\mathcal{J}_j \to 0\) in smooth solution areas, and \(\mathcal{J}_j \to +\infty\) at discontinuities, the discontinuity detector reads as follows:

\[
\begin{align*}
\text{if } \mathcal{J}_j < 1, & \quad \text{the cell is identified as a regular cell,} \\
\text{if } \mathcal{J}_j > 1, & \quad \text{the cell is identified as a troubled cell.}
\end{align*}
\]

Basically, considering a wave traveling over a sloping topography, the shore facing side of the wave is gradually steepening, while the offshore facing side is stretching. As the free surface gradient in the steepening area is also automatically increasing, waves about to break are identified using the previous indicator, in elements such that
\( \mathcal{J}_j > 1 \) (called troubled elements, following [61]) and we locally suppress the dispersive term in such elements \((i.e.\) we locally switch to the NSW equations) to avoid possible instabilities. After breaking, the wave fronts are handled as shocks by the NSW equations and only the hyperbolic part of the equations is solved for the wave fronts. The breaking wave dissipation is represented by shock local energy dissipation [8]. Of course, waves about to break can’t be considered as true discontinuities yet, but our numerical investigations show that, at a certain point and when the mesh is fine enough, the free surface gradient become large enough to activate the criteria (57). The validity of this simple approach, at least in the 1d case, is confirmed on several test cases in §3. Again, we highlight that this approach is quite rough, especially when compared with elaborated physical-based strategies, such as [72, 40] for instance. There is no wave-by-wave analysis and no threshold allowing to switch back towards GN equations when enough energy has been dissipated. In this regard, our stabilization technic is more to be compared with the approach introduced in [63] for instance, based on the momentum gradient.

Remark 2.9. Our numerical investigations show that, for a given breaking wave, the switching areas may need to be slightly enlarged, to prevent the possible occurrence of spurious oscillations due to the model’s transition, especially when high order polynomials are used. To achieve this, some neighboring cells are added to the switching area to include the steepening shore facing side of the wave, and a part of the offshore facing side. This can be simply done with the help of a mask, defining the switching area as a band of length 2\( r \) centered to the elements verifying \( \mathcal{J}_j > 1 \) (see Fig. 2). In this work, to keep things simple and avoid additional wave’s analysis, an arbitrary number of five cells is used to define \( r \). Such a choice has satisfyingly lead to robust computations in the test cases of §3. Ultimately, some more elaborated approaches should be used, relying for instance on an estimation of the roller length, as done in [72].

Additionally, the use of a local stabilization technique is classically required in troubled elements, to avoid spurious oscillations near discontinuities. In this work, we choose to use local limitation/projection procedure \( w_h \leftarrow \Pi w_h \) which maintains the conservation of mass element by element. We have implemented the improved moment limiting strategy of [10], which is free of problem dependence as there is no parameters needed to be calibrated unlike in the widely used TVB limiter [16] in which a constant has to be calibrated to enforce the TVB property. Although the limiting approach of [10] can be generalizable to any order of approximation \( \mathcal{N} \), we choose, for the sake of simplicity in the implementation, to locally truncate the development of the numerical solution in troubled cell, following the classical implementation of the TVB generalized slope limiter based on piecewise linear approximations. The order of the approximation is locally reduced to \( \mathcal{N} = 1 \), we only reconstruct the first order moment following exactly the lines of [10], and the higher order moments are arbitrarily set to zero. The reader is referred to [21] for the detailed reconstruction of the slope in our NSW equations based numerical model.

2.7 Robustness

An additional limitation is performed to ensure the positivity of the water height in the vicinity of dry (or almost dry) areas during the computation. Let us consider \( h^n_{\mathcal{E}_j}(x) \) obtained at the discrete time \( t^n \) from the fully discrete previous dG method \((i.e.\) deduced from \( \eta^n_{\mathcal{E}_j}(x) \) and \( b_{\mathcal{E}_j}(x) \)) and let denote \( \tilde{h}_j^n \) its average. Practically, starting from positive mean values \( \tilde{h}_j^n \geq 0 \) on all element \( \mathcal{E}_j \)\( \{E\} \}_{1,...,\mathcal{N}_e} \), we want to ensure that \( \tilde{h}_j^{n+1} \geq 0 \) on all element \( \{E\} \iota_{1,...,\mathcal{N}_e} \), without destroying the order of accuracy.

Such a property is enforced adapting the ideas developed in [85, 83] to the pre-balanced formulation. We first note that the scheme satisfied by the cell averaged of the free surface in the dG approximation (31) is

\[ \tilde{\eta}_j^{n+1} = \tilde{\eta}_j^n - \frac{\Delta t}{\lvert \mathcal{E}_j \rvert} \left[ \tilde{h}_j^n \right]_j, \quad (58) \]
Proof. Proposition 2. We consider the following first-order scheme

\[ \eta_j^{n+1} = \eta_j^n - \Delta_n \left[ \frac{\beta_j}{2|E_j|} \left( \frac{\tilde{h}_j^-}{\tilde{h}_j^0} \tilde{q}_j^n + \frac{\tilde{h}_j^+}{\tilde{h}_j^0} \tilde{q}_{j+1}^n \right) - \alpha_0 (\tilde{\eta}_j^+ - \tilde{\eta}_j^-) - \left( \frac{\tilde{h}_j^-}{\tilde{h}_j^0} \tilde{q}_j^n + \frac{\tilde{h}_j^+}{\tilde{h}_j^0} \tilde{q}_{j-1}^n \right) + \alpha_0 (\tilde{\eta}_j^- - \tilde{\eta}_j^+) \right] \]

with \( \tilde{\eta}_j^+ \) and \( \tilde{\eta}_j^- \) defined at each interfaces following (46)-(47)-(48) and (59)-(60). If \( \eta_j^n - b_j \) and \( \eta_j^{n+1} - b_{j+1} \) are positive, then \( \eta_j^{n+1} - b_j \) is positive under the condition \( \frac{\Delta_n}{|E_j|} \alpha_0 \leq 1 \).

Proof. The scheme (61) can be written, using (51):

\[ \eta_j^{n+1} = \eta_j^n - \Delta_n \left[ \frac{\beta_j}{2|E_j|} \left( \frac{\tilde{h}_j^-}{\tilde{h}_j^0} \tilde{q}_j^n + \frac{\tilde{h}_j^+}{\tilde{h}_j^0} \tilde{q}_{j+1}^n \right) - \alpha_0 (\tilde{\eta}_j^+ - \tilde{\eta}_j^-) - \left( \frac{\tilde{h}_j^-}{\tilde{h}_j^0} \tilde{q}_j^n + \frac{\tilde{h}_j^+}{\tilde{h}_j^0} \tilde{q}_{j-1}^n \right) + \alpha_0 (\tilde{\eta}_j^- - \tilde{\eta}_j^+) \right] \]

where we set \( h_j^n = \eta_j^n - b_j \) for the sake of simplicity. Denoting that \( \tilde{\eta}_j^+ - \tilde{\eta}_j^- = \tilde{h}_j^+ - \tilde{h}_j^- \) and \( \tilde{\eta}_j^- - \tilde{\eta}_j^+ = \tilde{h}_j^- - \tilde{h}_j^+ \), and subtracting \( b_j \) at both sides, we can conclude as in [83], writing (61) as a linear combination of positive values:

\[ \eta_j^{n+1} - b_j = \left( 1 - \frac{\Delta_n}{2|E_j|} (\alpha_0 + \frac{q_j^n}{h_j^n} \tilde{h}_j^- \tilde{h}_j^0 - \frac{\Delta_n}{2|E_j|} (\alpha_0 - \frac{q_j^n}{h_j^n} \tilde{h}_j^+ \tilde{h}_j^0) h_j^n \right) h_j^n \]

\[ + \left( \frac{\Delta_n}{2|E_j|} (\alpha_0 + \frac{q_j^n}{h_j^n} \tilde{h}_j^- \tilde{h}_j^0) \frac{\tilde{h}_j^+}{\tilde{h}_j^0} \right) h_{j-1}^n + \left( \frac{\Delta_n}{2|E_j|} (\alpha_0 - \frac{q_j^n}{h_j^n} \tilde{h}_j^+ \tilde{h}_j^0) \frac{\tilde{h}_j^-}{\tilde{h}_j^0} \right) h_{j+1}^n. \]

Based on Proposition 2, the ideas of [83] can be directly applied. Let us briefly recall the main steps of the procedure for a 1st order time discretization. For each element \( E_j \), assuming that \( \tilde{h}_j^n \geq 0 \):
1. Let $S_j^i = \{r^i_k\}_{k=1,...,d}$ be the set of $d$ LGL points on the element $\mathcal{E}^j$, and $\bar{\omega}_k$ the corresponding quadrature weights. $d$ is chosen such that the associated quadrature rule is exact for polynomials of degree $N$ (i.e. $2d - 3 \geq N$). We compute $m^n_j = \min_{r^i_k \in S_j^i} h^n_{\mathcal{E}^j}(r^i_k)$.

2. We modify $h^n_{\mathcal{E}^j}(x)$ in order to ensure that it is positive at the previous set of $d$ LGL nodes. This is done using the following conservative accuracy-preserving linear scaling around the cell average:

$$\hat{h}^n_{\mathcal{E}^j}(x) = \theta^n_j (h^n_{\mathcal{E}^j}(x) - \bar{h}^n_j) + \bar{h}^n_j,$$

where

$$\theta^n_j = \min \left( \frac{\bar{h}^n_j}{\bar{h}^n_j - m^n_j}, 1 \right).$$

We deduce from (63) a modified $N$ order polynomial $\tilde{\eta}^n_{\mathcal{E}^j}(x)$, which is injected into (31)-(32) instead of (30). Thus, following [85, 83], the positivity of the mean water height $\bar{h}^{n+1}_j$ is ensured under the condition:

$$\alpha \frac{\Delta^n_j}{|\mathcal{E}^j|} \leq \bar{w}_1.$$ 

In practice, we have $\bar{w}_1 = \frac{1}{6}$ for $N = 2, 3$ and $\frac{1}{12}$ for $N = 4, 5$.

2.8 Boundary conditions

The boundary conditions are classically imposed weakly, by enforcing suitable reflecting relations at a virtual exterior nodes, at each boundaries, through the boundary interface fluxes. Periodic, Neumann and Dirichlet conditions can be enforced following this simple process. For the approximation of second order derivatives with the LDG strategy, we classically enforce Neumann boundary conditions for the auxiliary variable. We also use Neumann boundary conditions for the derivatives involved in the discrete version of the operator $1 + \alpha \mathcal{S}[h_b]$. The first and last rows of (39) should be adapted for Dirichlet conditions. As far as generation and absorption of waves are concerned, we use relaxation techniques, as in [11, 45]. The computational domain is slightly extended to include the sponge layers, which length has to be calibrated from the incoming waves (generally 2 wavelengths).

We enforce periodic waves generation with the use of a generation/relaxation zone, following [52]. We have implemented the relaxation functions proposed in [79] and at the inlet boundary, we progressively impose on a two-wavelength long generation layer the targeted wave train.

3 Numerical examples

In this section, we assess the ability of our dG approach to compute waves propagation and transformations. The two models/schemes given by (31)-(32) involving (23a) or (23b) will be referred to as GNO (for original) and GNC (for constant) respectively. Unless stated otherwise, we use Neumann boundary conditions at both boundaries, the optimized value $\alpha = 1.159$ (see [9]) is used and we set $\epsilon_b = 0.1$. We highlight that our numerical investigations have shown that the value of $\epsilon_b$ does not influence the numerical results. Unless stated otherwise, and accordingly with the robustness result of the previous section, we do not suppress the dispersive effects in the vicinity of dry areas.
3.1 Motionless steady states preservation

We consider a 100 m channel and start from an initial steady state at rest $\eta = 0$ and $q = 0$, over the following topography (see Fig. 3):

$$b(x) = -h_0 \exp\left( - \left( \frac{x - d}{l} \right)^2 \right) + 0.8.$$  \hspace{1cm} (64)

After 10 s of simulation, the steady state is still preserved up to the machine accuracy, regardless of the polynomials’ order in the expansion basis or the refinement of the mesh. As an example, for $N = 3$ and $N_e = 300$, the computations give an $L^1$-error of $1.56 e^{-16}$ for the free surface and $7.13 e^{-15}$ for the discharge.

3.2 Accuracy and convergence analysis in the presence of non-flat bottom

We now focus on the evolution of a smooth gaussian profile over the previous non-flat and smooth topography, to highlight the convergence properties of our approach. The initial water height is defined as $h(x) = ab(x)$, where $a/h_0 = 0.2$ and $b$ defined in the previous test case (64). No exact solution is available, so a reference solution is
computed at $t = 0.15s$ with $N = 7$, $N_e = 1280$. Computations for both models (22)-(23a) and (22)-(23b) are run on a sequence of regular meshes with increasing refinement and polynomial expansions of increasing orders, from 1 to 5. For the approximation of high order derivatives, we use successively LDG and BR fluxes. The numerical $L^1$-errors computed using the reference solution at $t = 0.15s$ are reported in Tab. 5 for the water height and plotted on Fig. 4 for both water height and discharge, in log-log scale.

As expected, we obtain similar orders of accuracy for both GNO and GNC models. Concerning the impact of the numerical flux choice on the convergence rates, we can observe on Table (5) that, for both models and as mentioned in previous studies [17, 24, 32, 33], the use of BR fluxes may lead to sub-optimal convergence rates for odd values of $N$, while the LDG fluxes lead to optimal convergence rates. The corresponding convergence orders are reported on the last column of Table (5). As shown in Table 6, a similar behaviour is observed for the $L^2$ norm, while an order of $N + 1/2$ is obtained with the $L^\infty$ error using the LDG fluxes.

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Figure 5: Accuracy analysis in the presence of non-flat bottom: $L^1$ error for the discharge.
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Figure 6: Accuracy analysis for the GNC model in the presence of non-flat bottom: convergence rates in $L^1$, $L^2$ and $L^\infty$ norms for increasing values of $N$.

Let us now investigate the computational improvements obtained with the new GNC model. To this purpose, an averaged cpu-time per time-step is measured for both models, for increasing values of polynomials’ order and number of elements. These cpu times are denoted respectively $\rho_o$ and $\rho_c$ for the original and constant models. We focus on the ratio $\tau := \rho_o/\rho_c$ and the corresponding values are reported in Tab. 7. These values confirm the computational savings provided by the GNC model, clearly more efficient for all given couple $(N, N_e)$. We observe that the ratio increases with respect to the polynomial’s order, confirming the benefit of this new approach for high order simulations. For polynomial expansions of order 6 the speed-up can reach 7.15.

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Figure 7: Accuracy analysis in the presence of non-flat bottom: time ratio $\tau = \rho_o/\rho_c$ for increasing values of $N$ and $N_e$.

Based on these investigations, all the following computations are performed with the GNC model.

### 3.3 Propagation of a solitary wave

In this test, we investigate the propagation of a solitary wave over a flat bottom. We recall that the original model (1) admits the following class of solitary waves solution:

\[
\begin{align*}
    h(x, t) &= h_0 + a \text{sech}^2 \left( \kappa (x - ct) \right), \\
    u(x, t) &= c \left(1 - \frac{h_0}{h(x, t)}\right),
\end{align*}
\]

where $\kappa = \sqrt{ \frac{3a}{4h_0^2(h_0 + a)} }$, and $c = \sqrt{g(h_0 + a)}$. We also recall that these solitary waves are only solutions up to $O(\mu^2)$ of the new GNC family of models.
Figure 8: Propagation of a solitary wave: water surface profiles at t=0, 80, 160, 240 and 320s.

Figure 9: Propagation of a solitary wave: head on collision: water surface profiles at several times during the propagation.

For this test, the reference water height is fixed at $h_0 = 10 \text{ m}$ and we use the new model (11), with $\alpha = 1$ to compute the propagation of a solitary wave initially centered at $x_0 = 200 \text{ m}$, with a relative amplitude set to $a = 0.2h_0$. The computational domain is a 4000m long channel, regularly meshed with 1000 elements. We investigate several orders of polynomial expansions and follow the flow motion along the channel until $t = 320s$.  

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We show on Fig. 8 some free surface profiles at several times obtained with 3rd order expansions and a fourth order SSP-RK time-marching scheme. We can observe an excellent agreement between numerical solutions provided by the new model (11) and formula (65), showing the negligible discrepancies introduced by the $O(\mu^2)$ approximation in the new family of models. The free surface profile is accurately preserved during the propagation. However, let us mention that even if we do not report results here, some convergence studies have been performed for this particular test case, and we obtain sub-optimal $N + \frac{1}{2}$ convergence rates, for both BR and LDG fluxes. Similar behavior are observed with various wave relative amplitudes.

To complete the picture, and have a deeper assessment of the influence of non-linearities and dispersive terms, we compute the interaction of two counter-propagating waves of equal amplitudes, initially located at $x=1500m$ and $x=2500m$. Same mesh and polynomials order are used. The overall behavior of the propagation can be assessed through the free surface snapshots proposed in Fig. 9. As we can observe in the last pictures, the head-on collision generates dispersive tails of small amplitude. A zoom on these non-linear interactions is lastly proposed at $t=200s$, exhibiting a very proper description of the phenomenon. Indeed, this typical behavior is also studied in [55], leading to very similar observations. The propagation and transformations of solitary waves of increased non-linearity are studied in the next test.

<table>
<thead>
<tr>
<th>Gauge Number</th>
<th>location(m)</th>
<th>location(m)</th>
<th>location(m)</th>
<th>location(m)</th>
</tr>
</thead>
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<tr>
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<td>5.75</td>
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<tr>
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<td>6.25</td>
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<td>5.00</td>
</tr>
<tr>
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<td>8.75</td>
<td>6.75</td>
<td>5.75</td>
<td>5.75</td>
</tr>
</tbody>
</table>

Figure 10: Guibourg test case : location of the wave gauges for the four experiments.

### 3.4 Shoaling of solitary waves ([37])

We now investigate the dispersive properties of the model and study the nonlinear shoaling, using the data issued from a laboratory study performed at the LEGI (Grenoble, France). In this test, we consider a 36m channel with constant bed slope and a train of solitary waves generated at the inflow boundary. Measurements of the free surface are available at several wave gauges in the vicinity of the breaking point during the simulation. We consider 4 series of experiments, involving an increasing relative amplitude, starting from $a/h_0 = 0.096$ and ending with $a/h_0 = 0.534$. 

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Figure 11: Guibourg test case: comparison between computed (solid lines) and experimental (dots) time series of total free surface at several gauges before the breaking point.

The displayed numerical results are obtained with \((N, N_e) = (3, 400)\). Numerical results are reported on Fig. 11, together with the experimental data available from [37]. The gauges locations for these experiments are specified on Table 10. We can observe a very good behavior of the numerical model, even for large wave’s amplitude. These results assess the good accuracy of our model in the reproduction of the shoaling process.

3.5 Periodic waves over a submerged bar ([6, 20])

Going ahead in the assessment of the dispersive properties of the GNC model, we investigate now the propagation of periodic waves over a submerged bar, following the set-up proposed in [20]. The topography set-up and wave gauges locations are shown in Fig. 12. We choose here to perform test A, in which the amplitude of the input waves is set to \(a = 0.01m\). The time period \(T\) is 2.02 s and the initial depth \(h_0\) is 0.4m. When the incident waves encounters the upward part of the bar, it shoals and steepens, which generates higher-harmonics as the nonlinearity increases. These higher-harmonics are then freely released on the downward slope, and become deep-water waves behind the bar. The domain is regularly meshed with \(N_e = 800\) elements, and computations are run with increasing orders of polynomial expansions. Incident waves do not encounter breaking during the propagation, so that the switching/limiting process is not activated in this test.
Figure 12: Periodic waves over a submerged bar: sketch of the basin and gauges location.

Figure 13: Propagation of highly dispersive waves: free surface evolution at the four first gauges. Experimental data appear in plain lines.
Figure 15: Beji and Battjes plunging breaking case: free surface evolution at gauge $x = 6\, \text{m}$ and seven gauges regularly spaced from $x = 11\, \text{m}$ to $x = 17\, \text{m}$. Experimental data appear in plain lines.

Figure 14: Propagation of highly dispersive waves: several profiles for increasing $N$ at the last gauge.

We show on Fig. 13 some time series of the free surface elevation at several wave gauges located along the channel, compared with the experimental data. These results are obtained with 3$^{rd}$ order polynomials. We obtain a very good agreement at the first gauges. Some discrepancies are noticed at the last gauge, mostly explained by the high non-linear interactions generated as the waves encounters the upward part of the bar. As exhibited in Fig. 14, these effects are not correctly reproduced up to $N = 3$ at the last gauge for this choice of space step and we tried to locally increase the model accuracy, leading to sensible improvements. However, in the area of the last three gauges, the dispersion properties of the current model are no more satisfactory enough to accurately reproduce the complete release of the higher-harmonics, which can be regarded as highly dispersive waves. These results can be improved with the use of some optimized GN models, as for instance the 3 parameters optimized Green-Naghdi.
models proposed in [11] and in the 2d case in [45].

Using the same mesh and polynomials order, the wave amplitude is increased to 0.054 $m$ and $T$ is set to 2.525 $s$, leading to the simulation of the long wave plunging case described in [6]. Wave breaking is observed on the top part of the bar. Time series of the free surface are plotted on Fig. 15 at several reference points located along the channel. They exhibit a good agreement between numerical results and data coming from the experiment, which can be compared with those obtained in [72] for example. In Fig. 16, we locally follow the breaking process during a period, indicating areas where dispersive terms are suppressed.

![Figure 16: Beji and Battjes plunging breaking case: focus on the breaking process during a period. The NSW equations are used in areas delimited by the red bands to allow wave breaking.](image)

3.6 Solitary wave breaking over a sloping beach ([71])

Let us now assess the ability of our numerical model to deal with breaking waves and dry areas. In this test, we study the propagation, shoaling, breaking and run-up processes of a solitary wave over a beach with constant slope $s=1/19.85$, following the experiments of [71]. The incident wave is supplied by formula (65), with a water level at rest $h_0 = 1m$ and an amplitude set to $a = 0.28m$. The couple $(N, N_e)$ is fixed to $(3, 600)$ for the computation and our numerical results are compared with the experimental data. We observe that the breaking wave method described in (§2.6) is able to identify the wave steepening, occurring approximatively around $t^* = 17s$. We emphasize the very satisfying agreement between the numerical results and the data, as shown on Fig. 17.

3.7 Cox’s experiment ([18])

To further investigate the breaking waves / switching strategy, we study in this test the evolution of an incoming train of regular waves over a beach with constant bed slope 1/35. Waves of relative amplitude $a/h_0 = 0.29$ and period $T = 2.2s$ are generated at the inflow boundary. We set $N = 2$ and $|\xi_j| = 0.1$. A sketch of the computational domain is available in Fig. 18, with the location of the wave gauges.
In this picture, we also point out the areas where the dispersive effects are turned off (in red) and the NSW equations are used, following the switching strategy detailed in §2.6. In agreement with experimental observations, the first wave to break is detected between gauges ♯2 and ♯3. We can observe on Fig. 19 the time series of the free surface elevation at the six wave gauges along the domain and the comparison with the data taken from the experiment. We observe a very good matching. Note that the limiter \([10]\) is not applied in the whole switching (red) areas but only on the few troubled elements in the vicinity of the discontinuities. Note that we don’t use very high order schemes in test involving wave breaking. Indeed, as the limiter is activated in the vicinity of the discontinuities, the overall order of convergence and accuracy is reduced, limiting the interest of using very high order. Ultimately, some h/p adaptivity strategies should be implemented.
Figure 18: Cox’s experiment: sketch of the basin and identification of breaking areas. The NSW equations are used in red areas to describe the breaking waves.

Figure 19: Cox’s experiment: comparison between computed and experimental time series of total free surface at the gauges.
3.8 Wave overtopping a seawall ([38])

Figure 20: Overtopping over fringing reefs: sketch of the basin and gauge locations.

Figure 21: Overtopping over fringing reefs: Free surface profiles at several times during the propagation.

In this last validation, we aim at demonstrating that our numerical model appear as a promising tool for more complex simulations. The following test is based on the experiments carried out at the Tainan Hydraulics Laboratory (THL), National Cheng Kung University. It implies a tsunami-like solitary wave collapsing on a seawall located on a 1:20 sloping beach. A cross-section of the 22 m flume is described in Fig. 20, together with the location of several gauges.
We consider an initial water depth of 0.2 m, and an incoming wave with a relative amplitude $a/h_0 = 0.35$. For this simulation we set $N = 2$ and to reduce the number of cells, we use two mesh sizes: $|\mathcal{E}_j| = 0.03$ for $9 < x < 12$ m and $|\mathcal{E}_j| = 0.09$ elsewhere, giving a total number of $N_e = 300$ elements. Indeed, the use of irregular and locally refined meshes is very natural within dG methods, in contrast with other approaches like WENO schemes for instance. Note also that we don’t add any friction terms.

In this case, the solitary wave encounters breaking before reaching the seawall. It follows an overtopping flow supplemented by a train of reflected waves subsequently generated after the impact on the wall. Time series of the free surface were recorded at several gauges along the computational domain, and are compared with experimental data (see Fig. 22). We can note a very good agreement, similar to those exhibited in [38] with a obtained with a Volume Of Fluid (VOF) method for the Reynolds averaged Navier-Stokes equations and [72] with a hybrid FVM discretization of the original GN equations.

Conclusion

In this work, an arbitrary order dG discretization is proposed for a new family of 1d Green-Naghdi equations for the simulation of fully non-linear and weakly dispersive waves propagation and transformation over uneven bottom. This approach provides robustness and preservation of motionless steady states. These new models are shown to be more computationally efficient than the original ones, especially for high order polynomial expansions. We also propose a simple way to stabilize the computations when steep wave fronts appear, which gives satisfying results.
The 2d models with time-independent operator are already derived in [45] and the extension of the present dG method is studied in an on-going work. In particular, the extension of the switching strategy to the 2d case is not a simple task.

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**References**


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