Weak Local Residuals as Smoothness Indicators in Adaptive Mesh Methods for Shallow Water Flows

Sudi Mungkasi 1,* and Stephen Gwyn Roberts 2

1 Department of Mathematics, Faculty of Science and Technology, Sanata Dharma University, Yogyakarta, DIY 55281, Indonesia
2 Mathematical Sciences Institute, College of Science, The Australian National University, Canberra, ACT 2600, Australia; stephen.roberts@anu.edu.au
* Correspondence: sudi@usd.ac.id

Received: 29 January 2020; Accepted: 13 February 2020; Published: 1 March 2020

Abstract: This paper proposes some formulations of weak local residuals of shallow-water-type equations, namely, one-, one-and-a-half-, and two-dimensional shallow water equations. Smooth parts of numerical solutions have small absolute values of weak local residuals. Rougher parts of numerical solutions have larger absolute values of weak local residuals. This behaviour enables the weak local residuals to detect parts of numerical solutions which are smooth and rough (non-smooth). Weak local residuals that we formulate are implemented successfully as refinement or coarsening indicators for adaptive mesh finite volume methods used to solve shallow water equations.

Keywords: balance laws; conservation laws; finite volume; shallow water equations; smoothness indicator; weak local residual

1. Introduction

In the literature, fluid dynamics have been modelled into mathematical equations in the forms of partial differential equations. For example, Lee [1] used the forced KdV equation to model dynamics of trapped solitary waves. Shi et al. [2] studied the coupled time-fractional Boussinesq–Burgers system and provided some exact solutions to the system. Caputo et al. [3] reported some coupling conditions for water waves at forks. Paliathanasis [4] researched the two-dimensional rotating ideal gas dynamics and found a one-dimensional optimal system approximation. Castaings and Navon [5] proposed a numerical method for solving reaction–diffusion problems. These fluid dynamics often relate to the theory of conservation laws [6–9] with finite volume methods [10–13] as solving techniques. Overall, we can choose a fluid dynamics model in the literature that best suits the problem we want to solve.

This paper deals with fluid dynamics and focuses on solving the shallow water equations numerically using an adaptive mesh finite volume method with weak local residuals as the refinement indicators. Weak local residuals of the one-dimensional scalar conservation law were proved by Karni and Kurganov [14] to have lower degrees of accuracy at discontinuities than at smooth regions. Therefore, a weak local residual can be used as a smoothness indicator of numerical solutions of the one-dimensional scalar conservation law. Furthermore, even though the theory developed by Karni and Kurganov [14] is valid only for the one-dimensional scalar conservation law, the same behaviour of weak local residuals was seen to also be true for systems of conservation laws by numerical simulations [14–16].

A weak local residual formulation depends on the choice of test functions having a compact support on the computational domain. Karni et al. [15] chose to take quadratic B-splines as the test functions for both time and space variables. Constantin and Kurganov [16] chose linear B-splines for both time and space variables leading to a less expensive computational formulation of the weak
local residual. All weak local residuals of the aforementioned work of Kurganov et al. [15,16] assume uniform discretisation of spatial domain.

Our main contribution in this paper is the weak local residual formulation as refinement or coarsening indicators in adaptive mesh finite volume methods used to solve shallow-water-type equations in generic dimensions, including one, one and a half, and two dimensions. This generic formulation for different dimensions, and more importantly, for two dimensions in triangular grids, has never been published by the authors and any other researchers. By adaptive mesh we mean that the spatial domain discretisation can be non-uniform as the time evolves because of mesh refinement and mesh coarsening processes. We extend the work of Constantin and Kurganov [16] and so we choose linear B-splines as the test functions for both time and space variables in the formulation of weak local residuals.

Three shallow-water-type equations are considered, namely, one-, one-and-a-half-, and two-dimensional shallow water equations. Weak local residuals that we formulate are implemented in adaptive mesh finite volume methods to solve shallow water problems. Our numerical solutions show that weak local residuals are successful in detecting non-smooth regions of the solutions and lead to adaptive mesh methods which result in accurate numerical solutions.

There are several adaptive methods in the literature using other than weak local residuals as their refinement or coarsening indicator. One of them is the adaptive method based on the numerical entropy production, as proposed by Puppo and Semplice [17]. The theory of numerical entropy production was given by Puppo [18,19]. This method is tested for shallow water equations by Mungkasi [20,21]. The numerical entropy production is calculated using iterations of the discretised entropy inequality. Therefore, the computation of this refinement or coarsening indicator can be expensive. Another adaptive method is based on the gradient of the solution for the refinement and coarsening indicator, as used by Felcman and Kadrnka [22]. However, the gradient of the solution is actually not an error indicator, so the error indication may not be accurate.

In contrast, the weak local residual is cheap to compute and it is an error indicator. In each iteration of the conserved quantities, the residual can be obtained using a very short algebraic formula. Due to this reason, we choose a weak local residual as the coarsening or refinement indicator in our adaptive method in the present paper.

This paper is organised as follows. We recall the formulations of the weak local residuals (WLR) of Kurganov et al. [14–16] in Section 2. Section 3 presents weak local residuals for shallow-water-type equations. In Section 4, we implement the weak local residuals as error indicator for adaptive mesh refinement or coarsening. Some concluding remarks are then drawn in Section 5.

2. Balance Laws and Their Weak Local Residuals

In this section, we provide our numerical method of weak local residuals for detecting the smoothness of the solutions to balance laws. We follow the work described by Mungkasi [20]. We note that Kurganov et al. [14–16] have formulated weak local residuals of conservation laws. Mungkasi and Roberts [23,24] have extended those weak local residual formulations for balance laws. In this section, we recall the weak local residual formulations for balance laws. The formulations following Mungkasi et al. [20,23–25] are indeed important for the completeness of our work in the present paper.

We consider the scalar balance law

\[
\begin{align*}
q_t + f(q)_x &= s, \quad -\infty < x < \infty \\
q(x, t) &= q_0(x), \quad t = 0.
\end{align*}
\]  

(1)

Here, \(x\) represents the one-dimensional spatial variable, \(t\) represents the temporal variable, \(q\) denotes the conserved quantity, \(s\) is the source function, and \(f\) denotes the flux function. In addition,
We follow the methods of Mungkasi [20]. The one-dimensional shallow water equations (1D SWE) are
\[ \int_{-\infty}^{\infty} [q(x,t)T_t(x,t) + f(q(x,t))T_x(x,t) + s(x,t)T(x,t)] \, dx \, dt + \int_{-\infty}^{\infty} q_0(x)T(x,0) \, dx = 0 \]  
where \( T(x, t) \) is an arbitrary test function. The derivation of this weak form can be found in the work of Knobel [26] and Smoller [27].

Let \( q^n \) be approximate values of \( q(x, t^n) \) computed by a conservative numerical method and suppose that we have uniform grids \( (x_i := i \Delta x, t^n := n \Delta t) \). Using the notations of Karni and Kurganov [14], the corresponding approximation \( q^\Delta(x, t) \) (which is piecewise constant) is defined as
\[ q^\Delta(x, t) := q^n, \quad \text{if} \quad (x, t) \in [x_{i-1/2}, x_{i+1/2}] \times [t^{n-1/2}, t^{n+1/2}] \]  
where \( x_{i+1/2} := x_i + \Delta x/2 \) and \( t^{n+1/2} := t^n + \Delta t/2 \). We choose localized linear B-splines as our test functions \( T_{j+1/2}(x, t) := B_{j+1/2}(x)B^{n-1/2}(t) \), where \( B_{j+1/2}(x) \) and \( B^{n-1/2}(t) \) are centered at \( x = x_{j+1/2} \) and \( t = t^{n-1/2} \) with support of size \( 2\Delta x \) and \( 2\Delta t \). These are
\[ B_{j+1/2}(x) = \begin{cases} \frac{x-x_{j-1/2}}{\Delta x}, & \text{if } x_{j-1/2} \leq x \leq x_{j+1/2}, \\ \frac{x_{j+1/2}-x}{\Delta x}, & \text{if } x_{j+1/2} \leq x \leq x_{j+3/2}, \\ 0, & \text{otherwise,} \end{cases} \]  
and
\[ B^{n-1/2}(t) = \begin{cases} \frac{t-t_{n-3/2}}{\Delta t}, & \text{if } t_{n-3/2} \leq t \leq t^{n-1/2}, \\ \frac{t^{n+1/2}-t}{\Delta t}, & \text{if } t^{n-1/2} \leq t \leq t^{n+1/2}, \\ 0, & \text{otherwise.} \end{cases} \]

This leads to the weak local residual
\[ E_{j+1/2}^{n-1/2} = - \int_{t_{n-3/2}}^{t_{n+1/2}} \int_{x_{j-1/2}}^{x_{j+3/2}} \left[ q^\Delta(x, t) \left[ T^{n-1/2}_{j+1/2}(t) \right] + f(q^\Delta(x, t)) \left[ T^{n-1/2}_{j+1/2}(t) \right] + s^\Delta(x, t)T^{n-1/2}_{j+1/2}(t) \right] \, dx \, dt, \]  
which can be expressed straightforwardly as
\[ E_{j+1/2}^{n-1/2} = \frac{\Delta x}{2} \left[ q^n_j - q^n_{j-1} + q^n_{j+1} - q^n_{j-1} \right] + \frac{\Delta t}{2} \left[ f \left( q^n_{j+1} \right) - f \left( q^n_{j-1} \right) \right] + \frac{\Delta x \Delta t}{4} \left[ s^n_j + s^n_{j+1} + s^n_{j-1} + s^n_{j+1} \right]. \]  

Constantin and Kurganov [16] proposed the use of linear B-splines in the construction of the test function for conservation laws. Therefore, the weak local residual Equation (7) is denoted by CK (Constantin–Kurganov) indicator.

3. Shallow Water Equations and Their Weak Local Residuals

In this section, we focus on formulating weak local residuals for shallow water equations. We follow the methods of Mungkasi [20]. The one-dimensional shallow water equations (1D SWE) are
\[ h_t + (hu)_x = 0, \]  
\[ (hu)_t + \left( hu^2 + \frac{1}{2}gh^2 \right)_x = -ghz, \]
which are the mass and the momentum equations, respectively. Here, \( x \) is a free variable denoting
the one-dimensional position, \( t \) is a free variable denoting the time, \( u = u(x, t) \) represents the fluid
velocity dependent on \( x \) and \( t \), \( h = h(x, t) \) represents the fluid height (depth) dependent on \( x \) and \( t \),
\( z = z(x) \) is the topography (bed elevation) dependent on \( x \) measured from the horizontal reference,
and \( g \) is constant representing the acceleration due to gravity. We introduce an additional variable
called the stage \( w = h + z \) representing the position of water surface from the horizontal reference.

In the computation of CK indicator in this case, we have three options. We can compute the CK
indicator based on either the equation of mass conservation, that of momentum conservation, or a
combination between equations of mass and momentum conservations. The simplest computation is
the one based on the mass equation

\[
E_{i+1/2}^{h,n-1/2} = \left\{ \begin{array}{ll}
1/2 \left\{ \Delta x_i \left[ h^n_i - h^{n-1}_i + h^n_{i+1} - h^{n-1}_{i+1} \right] + \Delta t^n \left[ h^{n-1}_{i+1} u^{n-1}_{i+1} - h^{n-1}_i u^{n-1}_i + h^n_0 h^n_{i+1} - h^n_i u^n_i \right] \right\} & \text{if } E_{i+1/2}^{h,n-1/2} \geq E_{i-1/2}^{h,n-1/2} \\
E_{i+1/2}^{h,n-1/2} & \text{otherwise.}
\end{array} \right.
\]  

(10)

defined at time \( t = t^n-1/2 \) and position \( x = x_{i+1/2} \), where the superscript \( h \) in \( E_{i+1/2}^{h,n-1/2} \) represents
the mass quantity (height). Here \( \Delta x_i \) is the width of the \( i \)th cell, and \( \Delta t^n \) is the time step of the \( n \)th iteration.
Computing the residuals of the mass equation is simpler than computing those of the momentum
equation, because we do not need to deal with any source term. This is a similar situation to that
in the computation of numerical entropy production as a smoothness indicator as in the work of
Mungkasi and Roberts [24]. If we choose to compute the weak local residuals of the the momentum
equation, then the source term must be accounted for, so that the computation of the weak local
residuals is well-balanced. Mungkasi and Roberts [24] have proposed a well-balanced computation of
weak local residuals for the the momentum equation. As in the work of Mungkasi and Roberts [24],
we observed that weak local residuals with respect to the momentum and mass equations have similar
behaviour [24].

We define \( \Delta x := \min \{ \Delta x_i \} \), for one-dimensional problems, as the minimum cell width over the
whole discretised spatial domain to be substituted in Equation (10) and \( \Delta t \) is the time step, which
varies from an iteration to another one. We conjecture that over the whole discretised spatial domain,
both \( \Delta t^n \) and \( \Delta x_i \) in Equation (10) can be taken as constants. The difference of the order between rough
and smooth regions makes \( E_{i+1/2}^{h,n-1/2} \) able to detect the smoothness of the numerical solution. This is
guaranteed as the order of \( E_{i+1/2}^{h,n-1/2} \) is \( O(1) \) around discontinuities and \( O \left( \Delta x^{\min(3,r+1)} \right) \) at smooth
regions [16] when the numerical method has the formal order \( r \).

Equation (10) is defined at each vertex of computational cells. We take one having the largest
magnitude of two available indicators at its cell vertices, and we take the absolute value of them in
order to define the CK indicator at the centroids of each cell. Therefore, the CK indicator at the centroid
of the \( i \)th cell is given by

\[
E_{i}^{h,n-1/2} = \left\{ \begin{array}{ll}
E_{i+1/2}^{h,n-1/2} & \text{if } E_{i+1/2}^{h,n-1/2} \geq E_{i-1/2}^{h,n-1/2} \\
E_{i-1/2}^{h,n-1/2} & \text{otherwise.}
\end{array} \right.
\]

(11)

We note some remarks as follows. The assignment of the maximum indicator to the cell centroid
is not the only choice. We can also use a linear interpolation to get the centroid value. However, we use
the maximum, because it is better for the indicator to pick a cell to be refined, so that we maintain
the solution to be more accurate. This is important when we are not sure that a cell is to be refined or
coarsened, especially if the indicators have values close to the tolerance for refinement.

An extended application of the CK indicator for one-and-a-half-dimensional problems needs
special treatment. An available model for one-dimensional shallow water flows involving a passive
tracer is the one-and-a-half-dimensional shallow water equation. We can also consider it a model
of one-dimensional shallow water flows involving a shear or transverse velocity. We recall the one-and-a-half-dimensional shallow water equations (1.5 D SWE)

\[ h_t + (hu)_x = 0, \]

\[ (hu)_t + \left( h u^2 + \frac{1}{2} g h^2 \right)_x = \frac{g}{x} h z, \]

\[ (hv)_t + (huv)_x = 0. \]

Here, \( v(x) \) is the concentration of the passive tracer for the passive tracer model. Alternatively, we consider here \( v := v(x) \) as the horizontal velocity orthogonal to the \( x \)-axis, that is, the transverse velocity. Note that the variables, except the additional variable \( v \), are the same as those in the one-dimensional model.

Variable \( v(x) \) must be incorporated in the computation of WLR for the one-and-a-half dimensional shallow water model. This means that WLR is also able to detect the smoothness of the tracer solution. We propose to use the combination of both equations only will not detect the smoothness of the tracer solution. We choose one of four available indicators with the largest magnitude at its cell vertices in order to define the CK indicator at the centroids of each cell. Let

\[ M_i = \max \left\{ \left| E_{i-1/2}^{h,n-1/2} \right|, \left| E_{i+1/2}^{h,n-1/2} \right|, \left| E_{i-1/2}^{hv,n-1/2} \right|, \left| E_{i+1/2}^{hv,n-1/2} \right| \right\}. \]

Then CK indicator at the centroid of \( i \)th cell is

\[ E_i^{h,n-1/2} = \begin{cases} E_{i-1/2}^{h,n-1/2}, & \text{if } M_i = \left| E_{i-1/2}^{h,n-1/2} \right|, \\ E_{i+1/2}^{h,n-1/2}, & \text{if } M_i = \left| E_{i+1/2}^{h,n-1/2} \right|, \\ E_{i-1/2}^{hv,n-1/2}, & \text{if } M_i = \left| E_{i-1/2}^{hv,n-1/2} \right|, \\ E_{i+1/2}^{hv,n-1/2}, & \text{otherwise}. \end{cases} \]

As follows, we extend the discussion of the CK indicator for two-dimensional problems. The two-dimensional shallow water equations (2D SWE) have the forms of the following system of equations

\[ h_t + (hu)_x + (hv)_y = 0, \]
Here, \( t \) denotes the time variable, \((x, y)\) represents the coordinate in two dimensions, \( g \) is the acceleration due to gravity, \( z = z(x, y) \) is the topography (bed elevation), \( h = h(x, y, t) \) denotes the fluid height, \( u = u(x, y, t) \) denotes the fluid velocity along the \( x \)-axis, and \( v = v(x, y, t) \) denotes the fluid velocity along the \( y \)-axis. We discretise the spatial domain into triangular cells in order to solve the two-dimensional shallow water equations numerically. We can use the one-dimensional formulation Equation (10) locally in the computation of the CK indicator in order to define the CK indicator at the middle point of an edge. For one-dimensional problems, our conjecture is that over the whole discretised spatial domain, both \( \Delta t \) and \( \Delta x \) in Equation (10) can be taken as constants.

In the finite volume method evolution, we recall that \( \Delta t \) is the time step. We take \( \Delta x = \Delta t \) over the whole discretised spatial domain and substitute it into in Equation (10) in order to define the CK indicator for two-dimensional problems. Dependent on fluid velocity, wave propagation velocity, and the CFL (Courant-Friedrichs-Lewy) number, the values of \( \Delta t \) may vary from an iteration to another one. Then one having the largest magnitude of the available indicator values at its edges defines the CK indicator at the centroid of a cell.

Before we continue to the next section, we provide some remarks here.

**Remark 1.** The assignment of the maximum indicator to the cell centroid is actually not the only choice. We can also use a linear interpolation to get the centroid value of the indicator. However, we use the maximum, because it is better for the indicator to pick a cell to be refined, so that we maintain the solution to be more accurate. This is important when we are not sure that a cell is to be refined or coarsened, especially if some indicators have values close to or the same as the tolerance for refinement.

**Remark 2.** In this paper, we use the triangle centered finite volume scheme. The mesh is constructed using the uniform bisection of triangles. In iFEM [29,30], the procedure is called UNIFORMBISECT.

**Remark 3.** For details of our algorithm, we have uploaded our computer codes in GitHub https://github.com/stoiver/ifem-shallow-water. These computer codes are used to generate numerical results in the next section.

## 4. Numerical Tests

In this section, we simulate implementations of the adaptive mesh finite volume method based on weak local residuals. Numerical flux due to Kurganov, Noelle, and Petrova (KNP) [31] is used. Quantities are measured in SI units. We implement the well-balanced finite volume method [32]. The numerical flux is described as follows.

We consider the problem stated in Equation (1). The semi-discrete central-upwind scheme for this problem is

\[
\frac{d}{dt} q_j(t) = - \frac{F_{j+\frac{1}{2}}(t) - F_{j-\frac{1}{2}}(t)}{\Delta x} + s_j(t) \tag{21}
\]

where the numerical flux \( F_{j+\frac{1}{2}} \) is given by [31]

\[
F_{j+\frac{1}{2}}(t) = \frac{a_{j+\frac{1}{2}}^r f(q_{j+\frac{1}{2}}^-) - a_{j+\frac{1}{2}}^l f(q_{j+\frac{1}{2}}^+)}{a_{j+\frac{1}{2}}^r - a_{j+\frac{1}{2}}^l} + \frac{a_{j+\frac{1}{2}}^r a_{j+\frac{1}{2}}^l}{a_{j+\frac{1}{2}}^r - a_{j+\frac{1}{2}}^l} \left[ q_{j+\frac{1}{2}}^+ - q_{j+\frac{1}{2}}^- \right]. \tag{22}
\]

Here \( q_{j+\frac{1}{2}}^+ \) and \( q_{j+\frac{1}{2}}^- \) are the right and the left approximate interpolant values of the conserved quantities at \( x = x_{j+\frac{1}{2}} \), where the interpolant is reconstructed at each time step from the previously computed
cell averages, \( \tilde{q}_j(t) \). In order to have a stable method, it is necessary that the time step \( \Delta t \) satisfies the Courant–Friedrichs–Lewy (CFL) condition \[31\], that is,

\[
\Delta t \leq C_r \frac{L}{2M}
\]

where \( C_r \leq 1 \) is the CFL number and \( M = \max_j \left\{ \left| a^+_{j+\frac{1}{2}} \right|, \left| a^-_{j-\frac{1}{2}} \right| \right\} \). For SWE,

\[
a^+_{i+\frac{1}{2}} = \max \left\{ u^+_{i+\frac{1}{2}} + \sqrt{g \left( h^+_{i+\frac{1}{2}} \right)^*}, u^-_{i+\frac{1}{2}} + \sqrt{g \left( h^-_{i+\frac{1}{2}} \right)^*}, 0 \right\},
\]

\[
a^-_{i+\frac{1}{2}} = \min \left\{ u^-_{i+\frac{1}{2}} - \sqrt{g \left( h^-_{i+\frac{1}{2}} \right)^*}, u^+_{i+\frac{1}{2}} - \sqrt{g \left( h^+_{i+\frac{1}{2}} \right)^*}, 0 \right\},
\]

and the variables with star signs are obtained from the hydrostatic reconstruction \[32–34\]

\[
z^*_{i+\frac{1}{2}} := \max \{ z_{i,r}, z_{i+1,l} \},
\]

\[
h^*_{i,r} := \max \{ 0, h_{i,r} + z_{i,r} - z^*_{i+\frac{1}{2}} \},
\]

\[
h^*_{i+1,l} := \max \{ 0, h_{i+1,l} + z_{i+1,l} - z^*_{i+\frac{1}{2}} \}.
\]

In one dimension (1D) and one-and-a-half dimensions (1.5D), \( L \) is the minimum cell width among all cells in the whole spatial domain in the corresponding time step. In the two-dimensional (2D) case, \( L \) is the minimum incircle radius of triangle among all the triangular cells in the corresponding time step. As follows, we provide four numerical tests.

### 4.1. A One-Dimensional Problem with Topography

As the first numerical test, we consider a one-dimensional problem with topography as studied by Felcman and Kadrňka \[22\].

Let us assume that we have a channel of length 25 with topography

\[
z(x) = \begin{cases} 
0.2 - 0.05 \left( x - 10 \right)^2 & \text{if} \ 8 \leq x \leq 12, \\
0 & \text{otherwise},
\end{cases}
\]

(29)

together with the Dirichlet boundary conditions

\[
hu(0, t) = hu(25, t) = 1.53, \quad h(25, t) = 0.66,
\]

(30)

and the initial condition

\[
w(x, 0) = 0.66, \quad u(x, 0) = 0.
\]

(31)

The width of the coarsest cell allowed is 0.25. The maximum level of the binary refinement is 5. Cells with \( |CK| \leq 0.1CK_{\text{tol}} \) are coarsened, whereas those with \( |CK| > CK_{\text{tol}} \) are refined. Two neighbouring cells are coarsened as long as they are at the same level and have the same parent. The tolerance of the CK indicator is

\[
CK_{\text{tol}} = 0.05 \max \{|CK|\},
\]

(32)

where CK is defined as in Equation (11). The spatial domain is discretised initially into 100 cells.

Figure 1 shows the simulation results when the flow is at the steady state (time \( t = 100 \)). We observe that refinement is done by the adaptive method above the parabolic bump, as this region is rougher than other regions. In addition, refinement is also done close to the right boundary, because
there is a bore (a static shock) in this region. This means that the adaptive method is successful in simulating the problem.

Figure 1. Flow over a parabolic bump at time $t = 100$ using the adaptive method. For this figure, we enforce that the maximum level of refinement is 5.

4.2. A One-Dimensional Dam Break with Passive Tracer

In this simulation, a one-dimensional problem with horizontal topography is considered. We shall show that our proposed method produces an accurate solution around the rarefaction, the contact discontinuity, and the shock.

Suppose that we have a dam break problem involving a passive tracer (1.5D dam break problem) on horizontal topography

$$z(x) = 0, \quad \text{for } x \in [-2000, 2000].$$

The initial condition is

$$w(x, 0) = \begin{cases} 10 & \text{if } -2000 < x < 0, \\ 4 & \text{if } 0 < x < 2000, \end{cases}$$

$$u(x, 0) = 0, \quad v(x, 0) = \begin{cases} 3 & \text{if } -2000 < x < 0, \\ 0 & \text{if } 0 < x < 2000. \end{cases}$$

The width of the coarsest cell allowed is set to be 250. The maximum level of the binary refinement is set to be 10. We take

$$\text{CK}_{\text{tol}} = 0.01 \max \{|\text{CK}|\},$$

where CK is defined as in Equation (17). The spatial domain is discretised initially into 16 uniform cells.

We may use the same numerical flux formulation for all three (transport, momentum, and mass) equations in order to solve this problem numerically. However, we choose to use the standard flux formulation (Kurganov, Noelle, and Petrova (KNP) [31]) only to mass and momentum equations so that we obtain a more accurate solution at the contact discontinuity of the passive tracer solution. The passive transport flux is

$$F_{i+1/2}^{h} = \begin{cases} P_{i+1/2}^{h} & \text{if } P_{i+1/2}^{h} \geq 0, \\ P_{i+1/2}^{h} & \text{otherwise}, \end{cases}$$
where \( F^h_{i+1/2} \) is the mass flux at \( x = x_{i+1/2} \). This passive transport flux is the upwind flux based on \( v(x) \) as proposed by Bouchut [35]. Two neighbouring cells are coarsened as long as they are at the same level and have the same parent. Cells with \( |CK| \leq 0.1CK_{\text{tol}} \) are coarsened, whereas those with \( |CK| > CK_{\text{tol}} \) are refined.

\[ F^h_{i+1/2} = v(x_{i+1/2}) \left( f(x_{i+1/2}) - f(x_i) \right) \]

**Figure 2.** Flow with passive tracer at time \( t = 90 \) after dam break. CK is defined as in Equation (17). ‘Ana sol’ means the analytical solution. ‘Num sol’ is the numerical solution. Both solutions coincide.

A representative of our simulation is Figure 2, where at this time, the number of cells is 2748. This figure shows the flow with passive tracer at time \( t = 90 \). In these results, the contact discontinuity remains sharp. In addition, rarefaction and shock occurring in the water surface solution are well-resolved. This indicates that our adaptive method indeed leads to a very accurate solution over the whole region. Recall that contact discontinuities are much harder to resolve than shocks, and numerical solutions are diffusive around contact discontinuities generally [17,36].

### 4.3. A Planar Dam Break in Two Dimensions

Suppose that we have all \((x, y)\) in \([-1, 1] \times [-1, 1]\) as the spatial domain. The initial condition is given by

\[
w(x, y, 0) = \begin{cases} 
0.5 & \text{if } x < 0, \\
0.2 & \text{if otherwise,}
\end{cases}
\]

\[ v(x, y, 0) = 0, \quad u(x, y, 0) = 0. \]  

The domain is discretised initially into 2048 triangular cells. We simulate this problem with the time step being taken in such a way that the CFL condition is satisfied and the method is stable. We stop the simulation at time \( t = 0.2 \). We have five different areas of cells after coarsening or refinement, because the maximum level of refinement and coarsening is set to be 2. That is, an original initial triangular cell can be coarsened or refined up to two level coarser or two level finer. The CK tolerance is \( CK_{\text{tol}} = 0.5 \max \{|CK|\} \) (see Section 3 about how to compute CK indicator for a two-dimensional problem). Cells having CK smaller than \( CK_{\text{tol}} \) are candidates for coarsening; otherwise for refinement. We use the iFEM algorithm to do the coarsening and refinement. The data structure and description of iFEM can be found in a technical report authored by Chen [29] and a research paper written by Chen and Zhang [30].
Before we proceed, let us recall the mesh adaptivity algorithm in iFEM, where an illustrative example is given in Figure 3. Suppose that initially we have the nodes $N_1, N_2, N_3, N_4$ which are the coordinates $(0,0), (1,0), (1,1), (0,1)$, respectively. The elements are initially set to be two triangles with the nodes $[N_2, N_3, N_1]$ as triangle $T_1 = T_1[N_2, N_3, N_1]$ and the nodes $[N_4, N_1, N_3]$ as triangle $T_2[N_4, N_1, N_3]$, as shown in the left subfigure of Figure 3. The order of the nodes in a triangle is started from the node in front of the longest edge followed by other nodes in the counter-clockwise direction. Suppose that we want to refine the area around the node $N_5$ that is, the area around the coordinates $(1,1)$. Triangle $T_1[N_2, N_3, N_1]$ is bisected, so the node $N_5$ having coordinates $(0.5, 0.5)$ is created. The bisection of triangle $T_1[N_2, N_3, N_1]$ becomes the new $T_1[N_5, N_2, N_3]$ and $T_3[N_5, N_1, N_2]$. To maintain the conformity, triangle $T_2[N_4, N_1, N_3]$ is also bisected into the new $T_2[N_5, N_4, N_1]$ and $T_4[N_5, N_3, N_4]$. This lead to the subfigure shown in the middle of Figure 3. For the next refinement, we use the same procedure. As we want to refine the area around the node $N_5$, triangle $T_1[N_5, N_2, N_3]$ is bisected into new triangles $T_1[N_6, N_5, N_2]$ and $T_5[N_6, N_5, N_3]$; and triangle $T_4[N_5, N_3, N_4]$ is bisected into new triangles $T_4[N_7, N_5, N_3]$ and $T_6[N_7, N_4, N_5]$. (Note that the coarsening algorithm is the reverse of this refinement algorithm.)

Representatives of the results of our simulation are illustrated in Figures 4 and 5. Figure 4 shows results at time $t = 0.2$ the scaled CK indicator $|CK|/ \max \{|CK|\}$ and the water surface. At this time ($t = 0.2$), the number of triangles is 1656. Figure 5 depicts the corresponding momentum and mesh. We see from these results that in order to get an accurate solution, the numerical method achieves refinements around the rarefaction and the shock.

![Figure 3](image-url)  
**Figure 3.** Example of refining the mesh twice around the northeast corner node $N_5(1,1)$ of the domain.

![Figure 4](image-url)  
**Figure 4.** The left subfigure is the water surface and the right one is $|CK|/ \max \{|CK|\}$ at time $t = 0.2$ after the planar dam is broken.
4.4. A Radial Dam Break in Two Dimensions

Suppose that we have the spatial domain as all \((x, y)\) in \([-1, 1] \times [-1, 1]\). The initial condition is given by
\[
v(x, y, 0) = 0, \quad u(x, y, 0) = 0, \quad \text{(40)}
\]
\[
w(x, 0) = \begin{cases} 
1 & \text{if } x^2 + y^2 < 0.25, \\
0.5 & \text{if otherwise.} 
\end{cases} \quad \text{(41)}
\]

The time step is taken in such a way that the CFL condition is satisfied and the method is stable. The domain is discretised initially into 2048 triangular cells. We stop the simulation at time \(\Delta t = 0.05\). The maximum level of coarsening and refinement is 2. That is the original initial triangular cells can be refined or coarsen twice at the maximum. As a result, we have five different areas of cells. The NEP tolerance is \(\text{CK}_{\text{tol}} = 0.5 \max \{|\text{CK}|\} \). Cells having absolute CK smaller than \(\text{CK}_{\text{tol}}\) are candidates for coarsening, while otherwise are candidates for refinement.

Figures 6 and 7 show representatives of our simulation results. At time \(t = 0.05\), Figure 6 depicts the scaled CK indicator \(|\text{CK}|/ \max \{|\text{CK}|\}\) and the water surface. At this time \((t = 0.05)\), the number of triangles is 2168. Figure 7 shows the corresponding momentum and mesh. Like in the case of the planar dam simulation, we observe that refinements occur around the rarefaction and the shock leading to an accurate solution.
Before we conclude the paper, we provide some remarks as follows.

**Remark 4.** The tolerance for the weak local residuals is a matter of choice to the users of the proposed method. If the tolerance is too strict, then the refinement will be excessive, but if the tolerance is too loose then the mesh will be too coarse. We indicate that the tolerance is about 5% of the maximum weak local residual for the refinement and 0.5% of the maximum weak local residual for the refinement. That is, if a cell has the value of residual which is larger than 5% of the maximum weak local residual, then the cell must be refined as long as the cell area has not reached the minimum value of the allowed area. If a cell has the value of residual which is smaller than 0.5% of the maximum weak local residual, then the cell must be coarsened as long as the cell area has not reached the maximum value of the allowed area. However, once again, the tolerance is a matter of choice of the users.

**Remark 5.** For the first order method, if a cell is bisected, then each of the two new cells has the same quantity values as those of the original cell. That is, suppose that the triangle $T$ is bisected into two triangles $T_1$ and $T_2$. The quantity $q_{T_1}$ of the triangle $T_1$ and the quantity $q_{T_2}$ of the triangle $T_2$ are the same as the quantity $q_T$ of the original triangle $T$. In mathematical formulas

$$q_{T_1} = q_T \quad \text{and} \quad q_{T_2} = q_T. \quad (42)$$

For the second order method, if a cell is bisected, then quantities of each of the two new cells should be computed using proportional values of the linear reconstruction of the quantities of the original cell after implementing a slope limiter.

If two cells are coarsened, then the new cell has quantity values that are computed from the quantity values of the two old cells with respect to area-based weights. That is, supposed that two triangles $T_1$ and $T_2$ are coarsened to be the triangle $T$. The quantity of the triangle $T$ is computed as

$$q_T = \frac{\|T_1\|}{\|T\|} q_{T_1} + \frac{\|T_2\|}{\|T\|} q_{T_2} \quad (43)$$

where $\|T\|$ means the area of the triangle $T$. In all of our 1D, 1.5D, and 2D cases in this paper,

$$\frac{\|T_1\|}{\|T\|} = \frac{\|T_2\|}{\|T\|} = \frac{1}{2}. \quad (44)$$
5. Conclusions

We have developed weak local residual formulations according to Constantin and Kurganov, so that they can be implemented as smoothness or refinement indicator to adaptive mesh finite volume methods used to solve shallow-water-type equations. Our formulation has been shown to be successful in the numerical tests for one-, one-and-a-half-, and two-dimensional problems. In the present work, the tolerance of the indicator for refinement and coarsening are chosen manually without a fixed rule. Future work would probably examine how to set this tolerance based on a fixed rule which can be applied for general problems. It could also investigate our conjecture that both the time step and the cell width in the formulation of CK indicator can be taken as constants.

Solving the 2D SWE involving topography using the adaptive method with the WLR indicators could be a big continuation topic of research. This is because we have to deal with balancing the discrete SWE as well as balancing the WLR. In this paper, we have limited our scope to solving the shallow water equations using the proposed adaptive method for all 1D, 1.5D, and 2D cases, where for the 1.5D and 2D cases, we have not included the topography. The inclusion of topography in the 1.5D and 2D cases could be a future research direction. Research results in this paper are limited to first order methods. Extending our adaptive method to second order methods is also a possible future research direction. In addition, optimising the computer programming codes of the adaptive method could be investigated further.

Author Contributions: Both authors have contributed in the conceptualization, methodology, software, validation, formal analysis, investigation, resources, data curation, writing—original draft preparation, writing—review and editing, visualization, supervision, project administration, and funding acquisition. All authors have read and agreed to the published version of the manuscript.

Funding: This work was conducted in Australia and Indonesia. The work in Australia was financially supported by The Australian National University. The work in Indonesia was financially supported by the Ministry of Education and Culture as well as the Ministry of Research and Technology (Badan Riset dan Inovasi Nasional) of The Republic of Indonesia in the form of Hibah Penelitian Dasar Unggulan Perguruan Tinggi for the years 2018, 2019, and 2020.

Acknowledgments: Some parts of this work were conducted as a portion of the unpublished Ph.D. thesis [20] of Sudi Mungkasi (the first author) under the supervision of Stephen Gwyn Roberts (the second author) at The Australian National University, Canberra, Australia. Other parts were conducted at Sanata Dharma University, Yogyakarta, Indonesia after the first author received the Ph.D. degree.

Conflicts of Interest: The authors declare no conflict of interest.

Abbreviations

The following abbreviations are used in this manuscript:

1D one dimension
1.5D one and a half dimensions
2D two dimensions
CFL Courant–Friedrichs–Lewy
CK Constantin–Kurganov
iFEM integrated finite element method
KNP Kurganov–Noelle–Petrova
SI Systéme International (International System)
SWE shallow water equations
WLR weak local residual

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