Author’s response to the reviews

November 20, 2014

Author’s response to executive editor D. Lunt

We have contacted Prof. Lutz Gross, the editor handling this manuscript, and GMD publisher. We have prepared the codes of global SWE model described in our manuscript as requested.

Author’s response to reviewer #1

Thank you for carefully reading the paper. Your constructive comments are extremely helpful for use to improve the manuscript. We have fully taken your comments into account and accordingly modified the paper. Below, please find our point-to-point responses to your comments. (The comments from the reviewer are marked in red color.)

This paper reports on a very interesting method for solving conservation laws, with an application to global ocean modelling. The method is in some senses a hybrid between discontinuous Galerkin methods, Finite volume methods and Collocation methods. High order is obtained from the high order polynomial representation in the volumes, and robust high order approximation of the fluxes is obtained via an interesting combination of averaging the fluxes at the volume boundaries via (approx) Riemann solvers and interpolation in the interiors of the volumes. The numerical experiments and the linear analysis support the conclusion that the method produces a simple robust high order method.
The paper does need an edit to pick up slight problems in the language, for instance, there are a number of places where definite or indefinite articles are missing.

Thank you for your comments. We have made a thorough linguistic edit.

Some minor points or comments:

(1) As discussed on page 4255, the pointwise representation of the quantities and the fluxes are obtained via high order polynomial interpolation. A question: Is it always safe to do this, i.e., is it ever necessary to limit the interpolants or introduce some sort of artificial viscosity? Or does the use of the average fluxes at the boundaries stabilize the calculation of the derivative of the fluxes at the interior points?

At cell boundaries, we solve a Riemann problem to determine the values of the numerical fluxes. As a result, the proposed scheme is an upwind scheme with inherent numerical viscosity, which guarantees the computational stability as shown by the Fourier analysis of the semi-discrete system in the paper. However, in the presence of discontinuous solutions the non-physical oscillations might occur without effective limiters or artificial dissipations, which is a common issue of the high-order schemes. There are some existing works for this purpose. We added some explanations in the revised manuscript (line 122 to line 125 and line 467 to line 473 in the marked-up manuscript).

(2) Equation (12). It would be good to remind the reader of the exactness of Gaussian Quadrature for polynomials of degree 5.

Thank you for suggesting this. We added some words in the revised manuscript (line 133 in the marked-up manuscript).


Two types of explicit Runge-Kutta schemes have been adopted in present study. The third-order scheme has been widely used in the existing models. Third-order scheme is a good trade-off between numerical accuracy and computational cost. The fifth-order scheme was used here only for assuring the fifth-order accuracy in both time and spatial discretizations in numerical con-
vergence tests. The references for the Runge-Kutta method is added to the revised manuscript (line 144 and line 151 in the marked-up manuscript).

(4) Page 4260. Provide a very short overview of the methods DG3, MCV5 which you use for your numerical comparisons.

We conduct a comparison between DG3, MCV5 and the proposed scheme since these three schemes have fifth-order accuracy and can be derived by flux reconstruction framework using different constraint conditions for spatial reconstruction of flux functions. As detailed in Huynh(2007), the DG3 scheme uses the Radau polynomial as the correction functions to derive the flux reconstruction which assure the continuity of the numerical fluxes computed from Riemann solvers at the cell interfaces. MCV5 scheme can be derived by the general framework for flux reconstruction using multi moments proposed in Xiao et al. (2013). MCV5 uses constraint conditions on the point values, first- and second-order derivatives of flux functions at the cell interfaces where Riemann solvers in terms of derivatives of the flux function are required. We added a brief description in the revised manuscript as suggested (line 206 to line 215 in the marked-up manuscript).


We added the relationship between covariant and contravariant velocity components in the revised manuscript as suggested (Eq.(28)).

(6) Can your method work with wet/dry interfaces?

The numerical techniques designed for the traditional finite volume schemes, as well as other high order schemes, like DG, can be straightforwardly applied to the present scheme without substantial barriers. The calculation of wet/dry front in shallow flows has been extensively studied under such numerical frameworks, and can be adopted in our model.

(7) Can your method reproduce a still lake (ocean) i.e. is well balanced for this stationary solution?

The stationary state can be exactly preserved by the proposed model since the topography source term is formulated in the way that satisfies the "exact
C-property”. The numerical procedure for this was described in details in Chen & Xiao, JCP, 2008.

Author’s response to reviewer #2

Thank you for carefully reading the paper. Your constructive comments are extremely helpful for use to improve the manuscript. We have fully taken your comments into account and accordingly modified the paper. Below, please find our point-to-point responses to your comments. (The comments from the reviewer are marked in red color.)

This paper presents a flux reconstruction method for discontinuous elements applied to 2D shallow water equations. While this research would be interesting for the modeling community, I have many comments on the quality of the presentation (given by this paper). The main objection is that the paper does not provide enough explanation on spatial discretizations. Also, there are comments on the model and on English usage.

I recommend it for publication in GMD after major revisions.

Comments:

1. In the introduction, the paper claims using the flux reconstruction method as described in Huynh, 2007 and modified in Xiao, 2013. The FR technique is to redistribute flux to all elements nodes, but Eqns. (6), (7) and further only update end points. If the paper uses one of the schemes derived in Xiao, 2013, then it should be stated. Still, in Xiao, 2013 (http://arxiv.org/pdf/1206.4406.pdf) it seems that interior points are modified by the FR process. In summary, much more should be given on the scheme used. Also, describe what (if anything) is different from previously published works.

   Basically, the flux reconstruction (FR) includes the following steps:

   1. Define the unknowns as the local degrees of freedom, which are the nodal values at the solution points within each cell;

   2. Build a high-order spatial reconstruction for flux function which is a consistent approximation to the solution over each cell and satisfies the con-
continuity conditions using the Riemann solver at cell boundaries;

3. Evaluate the derivatives of flux function at the solution points to get the
time evolution equations to update the solutions.

The key is step 2, and different constraint conditions can used for flux re-
construction, which result in different numerical schemes.

In Huynh 2007, FR is formulated by two correction functions which assure
the continuity at the two cell boundaries and collocate with the so-called pri-
mary Lagrange reconstruction at their zero-points. So, the existing nodal type
schemes can be recast under the FR framework with different correction func-
tions.

In Xiao et al. 2013, a more general FR framework was proposed by introduc-
ing the multi-moment constraint conditions including nodal values, first-order
derivatives and even second-order derivatives to determine the flux reconstruc-
tion.

In this study, we present a more straightforward and simpler approach to
derive the FR formulation using the collocation method, which has not been
discussed in either Huynh (2007) or Xiao et al (2013). The resulting scheme,
GaussLegendre-point based conservative collocation (GLPCC) scheme, is new
and has not be reported by anyone else to our knowledge.

We added some explanations in the revised manuscript (line 86 to line 93 in
the marked-up manuscript).

2. Eqn. (12) and conservation: It seems that conservation is only achieved
for uniform meshes, because Eqn. (12) depends on the element length. So, in
general, the scheme does not conserve mass if non-uniform grids are used? Also,
cube-sphere meshes are only quasi-regular. Is scheme conservative on a sphere?

It would be useful to provide a plot for mass conservation from one of the
shallow water tests, similar to Figure 14.

The proposed scheme is conservative even for non-uniform grids. The to-
tal mass within each control volume, i.e. \( \Delta x_i \tilde{q}_i \) is exactly conserved, which
is in fact computed by a finite volume formulation with the numerical fluxes
calculated at the cell boundaries as shown in Eq. (12). Rather than volume-integrated average (VIA) itself, the product of VIA and the volume is conserved.

We have rewritten Eq. (12) in the revised manuscript to avoid possible misleading. The plots for mass conservation of case 5 and 6 are shown in Fig. 10 and Fig. 13 in the revised manuscript.

3. Eqn. (19) and spectral analysis: How was eqn. (19) derived and what are the coefficients? Also, it seems that the spectral problem is formulated globally because neighbor values are included. I found more details in paper Xiao, 2013 (http://arxiv.org/pdf/1206.4406.pdf) but at least if notations are used, they should be clarified.

We have revised this part in the revised manuscript with more details (line 160 to line 193 in the marked-up manuscript). The spectral analysis adopted here follows the procedure in Huynh (2007).

4. Super convergence: The authors mention super convergence a few times. I believe they mean that their method is an h-p method with corresponding convergence properties. It would be desirable to clarify terminology. Also, the authors state (p. 4253) that “The Fourier analysis and numerical tests show that the present scheme has the super convergence property same as the DG method.” First, they did not show this numerically because there are no tests for the p refinement. Second, which DG method are they referring to? Third, how exactly the Fourier analysis can be used for exponential convergence?

The discussion on the super convergence follows the context in Huynh (2007) where a scheme using $K$ solution points is said to be super-convergent if its order of accuracy is higher than $K$. As shown in Huynh (2007) that the nodal DG scheme has a convergence rate of $2K − 1$, we demonstrate that the present three-point GLPCC scheme has 5th-order convergence rate by the Fourier analysis used in Huynh(2007). Here, the DG scheme is referred to the nodal type DG defined in Huynh (2007) which uses the Radau polynomial as the correction function. A relevant theoretical work can be found in Guo et al. (W. Guo, X. Zhong and J. Qiu, J. Comput. Phys. Vol. 235, 458-485 (2013)).

5. “The parameter $a$ in Eq. (24) is determined by the contravariant veloc-
ity component and the water depth, which are exactly same on two adjacent patches.” A continuous velocity field in contravariant coordinates on an edge has two components, and one of them, corresponding to a basis vector perpendicular to the edge, is the same (up to the sign) for adjacent elements.

It has been clarified in the revised manuscript (line 330 to line 333 in the marked-up manuscript).

6. Figure 15 needs labels ((a), (b) ...) and captions for them.

Thank you for suggestion. We added the figure labels in the revised manuscript (Fig. 17 in the revised manuscript).


Thank you for your comments. We revised the sentence in the manuscript (line 281 in the marked-up manuscript).

8. The paper does not cover diffusive properties of the proposed method and possible applications of artificial diffusion. I believe this is a valid point for discussion. Shallow water models are often considered as preliminary studies for 3D models. In 3D models, diffusion mechanisms cannot be ignored.

Based on the Riemann solver at cell interfaces, the proposed scheme is essentially an upwind type method. As a result, the inherent numerical dissipation is included and stabilizes the numerical solutions. We did not use any extra artificial viscosity in the shallow water model for the numerical tests presented in the paper. We agree with you that additional dissipation or limiter projection might be necessary in other cases in 3D. Because of the algorithmic similarity, the existing works on high-order limiting projection and artificial dissipation devised for DG or spectral element methods should be applicable to GLPCC
without substantial difficulty. Some comments have been included in the revised manuscript (line 122 to line 125 and line 467 to line 473 in the marked-up manuscript).

Comments on English:

1. More attention should be given to articles.

2. p. 4262: Revise “… coordinate system (ξ, η) are shown in Fig.”, “…the governing equations is rewritten.”

3. p. 4264: Revise “… we solving …”

4. p. 4267: Revise “The conservation errors of total energy and enstrophy are interest for atmospheric modelling”

5. p. 4268: Revise “Two kinds of setup of this test are usually checked in literatures.”

We have made a thorough linguistic check. All your comments are reflected in the revised manuscript. Thank you.

Appendix

A marked-up version of the revised manuscript is included as follows, where the revisions are indicated by using latexdiff.
A high-order conservative collocation scheme and its application to global shallow water equations

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Abstract. An efficient and conservative collocation method is proposed and used to develop a global shallow water model in this paper. Being a nodal type high-order scheme, the present method solves the point-wise values of dependent variables as the unknowns within each control volume. The solution points are arranged as Gauss–Legendre points to achieve the high-order accuracy. The time evolution equations to update the unknowns are derived under the flux-reconstruction (FR) framework (Huynh, 2007). Constraint conditions used to build the spatial reconstruction for the flux function include the point-wise values of flux function at the solution points, which are computed directly from the dependent variables, as well as the numerical fluxes at the boundaries of the computational element, which are obtained as the Riemann solutions between the adjacent cells. Given the reconstructed flux function, the time tendencies of the unknowns can be obtained directly from the governing equations of differential form. The resulting schemes have super convergence and rigorous numerical conservativeness.

A three-point scheme of fifth-order accuracy is presented and analyzed in this paper. The proposed scheme is adopted to develop the global shallow-water model on the cubed-sphere grid where the local high-order reconstruction is very beneficial for the data communications between adjacent patches. We have used the standard benchmark tests to verify the numerical model, which reveals its great potential as a candidate formulation for developing high-performance general circulation models.

1 Introduction

A recent trend in developing global models for atmospheric and oceanic general circulations is the increasing use of the high-order schemes that make use of local reconstructions and have
the so-called spectral convergence. Among many others are those reported in Giraldo et al. (2002); Thomas and Loft (2005); Giraldo and Warburton (2005); Nair et al. (2005a,b); Taylor and Fournier (2010); Blaise and St-Cyr (2012). Two major advantages that make these models attractive are (1) they can reach the targeted numerical accuracy more quickly by increasing the number of degrees of freedom (or unknowns), and (2) they can be more computationally intensive with respect to the data communications in parallel processing (Dennis et al., 2012).

The discontinuous Galerkin (DG) (Cockburn et al., 2000; Hesthaven and Warburton, 2008) and spectral element(SE) (Patera, 1984; Karniadakis and Sherwin, 2005) methods are the widely used frameworks in this context. A more general formulation, so-called flux reconstruction (FR), was presented in Huynh (2007) which covers a wide spectrum of nodal type schemes, including the DG and SE as the special cases. A FR scheme solves the values at the solution points located within each grid element, and the volume-integrated value, which are the weighted summation of the solutions, can be numerically conserved. We recently proposed a class of local high-order schemes, named multi-moment schemes, which were used to develop the accurate shallow water models on different spherical grids (Chen and Xiao, 2008; Li et al., 2008; Li and Xiao, 2010; Chen et al., 2014b). By introducing multi-moment concept, we show in Xiao et al. (2013) that the flux reconstruction can be implemented in a more flexible way, and other new schemes can be generated by properly chosen different types of constraint conditions.

In this paper, we introduce a new scheme which is different from the existing nodal DG and SE methods under the FR framework. The scheme, so-called Gauss–Legendre-point based conservative collocation (GLPCC) method, is a kind of collocation method that solves the governing equations of differential form at the solution points, and is very simple and easy to follow. The Fourier analysis and the numerical tests show that the present scheme has the super convergence property same as the DG method. A global shallow water equation (SWE) model has been developed by implementing the three-point GLPCC scheme on a cubed-sphere grid. The model has been verified by the benchmark tests. The numerical results show the fifth-order accuracy of the present global SWE model. All the numerical outputs look favourably comparable to other existing methods.

The rest of this paper is organized as follows. In Sect. 2, the numerical formulations in one dimensional case are described in detail. The extension of the proposed scheme to a global shallow water model on cubed-sphere grid is then discussed in Sect. 3. In Sect. 4, several widely used benchmark tests are solved by the proposed model to verify its performance in comparison with other existing models. Finally, a short conclusion is given in Sect. 5.
2 Numerical formulations

2.1 Scheme in one dimensional scalar case

The first order scalar hyperbolic conservation law in one dimension is solved in this subsection,
\[ \frac{\partial q}{\partial t} + \frac{\partial f(q)}{\partial x} = 0, \]  
(1)

where \( q \) is dependent variable and \( f \) is flux function.

The computational domain, \( x \in [x_I, x_R] \), is divided into \( I \) elements with the grid spacing of \( \Delta x_i = x_{i+\frac{1}{2}} - x_{i-\frac{1}{2}} \) for the \( i \)th element \( C_i = [x_{i-\frac{1}{2}}, x_{i+\frac{1}{2}}] \).

A class of schemes can be devised under the framework of the flux reconstruction (FR) (Huynh, 2007; Xiao et al., 2013). For each grid, the computational variables (unknowns) are defined at several solution points within each element, e.g. within element \( C_i \), the point values, \( q_{im} \) (\( m = 1, 2, \ldots, M \)), are defined at the solution points \( (x_{im}) \) which are located within the element, are treated as the computational variables (unknowns). High order schemes can be built by increasing the number of the solution points. In this paper, we describe the GLPCC scheme that has three solution points for each grid element \( (M = 3) \).

Three configuration of local degrees of freedom (DOFs) (unknowns), i.e. \( q_{im} \), \( m = 1 \) to 3, are point wisely defined at solution points \( x_{im} \) within each element as shown in Fig. 1 (hollow circles) by the hollow circles. To achieve the best accuracy, the DOFs are arranged at Gauss-Lagrange points in this study.

\[ x_{i1} = x_i - \frac{1}{2} \sqrt{\frac{3}{5}} \sqrt{2} \Delta x_i, \quad x_{i2} = x_i \quad \text{and} \quad x_{i3} = x_i + \frac{1}{2} \sqrt{\frac{3}{5}} \sqrt{3} \Delta x_i, \]  
(2)

where \( x_i \) is the center of the element \( x_i = (x_{i-\frac{1}{2}} + x_{i+\frac{1}{2}})/2 \).

The unknowns are updated by applying the differential-form governing equations Eq. (1) at solution points as

\[ \frac{\partial q_{im}}{\partial t} = - \left[ \frac{\partial f(q)}{\partial x} \right]_{im}. \]  
(3)

As a result, the key task left is to evaluate the derivatives of the flux function, which is realized by reconstructing the piecewise polynomial for flux function, \( F_i(x) \), over each element. Once the reconstructed flux function is obtained, the derivative of flux function is approximated by

\[ \left[ \frac{\partial f(q)}{\partial x} \right]_{im} \approx \left[ \frac{\partial F_i(x)}{\partial x} \right]_{im}. \]  
(4)

In this study, we Huynh (2007), FR is formulated by two correction functions which assure the continuity at the two cell boundaries and collocate with the so-called primary Lagrange reconstruction at their zero-points. So, the existing nodal type schemes can be recast under the FR framework with different correction functions. In Xiao et al. (2013), a more general FR
framework was proposed by introducing the multi-moment constraint conditions including nodal values, first-order derivatives and even second-order derivatives to determine the flux reconstruction. Here, we will develop a new method to reconstruct the flux function, which is more straightforward and simpler compared with the methods discussed in either Huynh (2007) or Xiao et al. (2013).

We assume that the reconstructed flux function over the \( i \)th element, \( \mathcal{F}_i(x) \), has the form of

\[
\mathcal{F}_i(x) = c_{10}^i + c_{11}^i (x - x_i) + c_{12}^i (x - x_i)^2 + c_{13}^i (x - x_i)^3 + c_{14}^i (x - x_i)^4, 
\]

(5)

where the coefficients, \( c_{10}^i, c_{11}^i, \ldots, c_{14}^i \), are determined by a collocation method, which meets five constrained constraint conditions specified at five constrained constraint points (shown in Fig. 1 by the solid circles) as

\[
\begin{align*}
\mathcal{F}_i(x_{im}) &= f(q_{im}), \ m = 1 \text{ to } 3 \\
\mathcal{F}_i(x_{i-\frac{1}{2}}) &= \tilde{f}_{i-\frac{1}{2}} \\
\mathcal{F}_i(x_{i+\frac{1}{2}}) &= \tilde{f}_{i+\frac{1}{2}}
\end{align*}
\]

(6)

where \( \tilde{f}_{i\pm\frac{1}{2}} \) are the values of flux function at the interfaces between different element cell boundaries.

In Eq. (6), \( f(q_{im}) \) are calculated by three known DOFs at solution points. The values of flux function at interfaces the boundaries are obtained by solving the Riemann problems with the values of dependent variables at the interfaces between two neighboring elements, which are interpolated separately from two adjacent elements. Considering the interface at \( x_{i-\frac{1}{2}} \), we get two values of flux function from elements \( C_{i-1} \) and \( C_i \) as

\[
\begin{align*}
\tilde{f}_{i-\frac{1}{2}} &= f\left(q_{i-\frac{1}{2}}^L\right) = f\left[Q_{i-1}\left(x_{i-\frac{1}{2}}\right)\right] \\
\tilde{f}_{i+\frac{1}{2}} &= f\left(q_{i+\frac{1}{2}}^R\right) = f\left[Q_i\left(x_{i+\frac{1}{2}}\right)\right],
\end{align*}
\]

(7)

where \( Q_i(x) \) is a spatial reconstruction for dependent variable based on local DOFs, having the form of

\[
Q_i(x) = \sum_{m=1}^{3} [\mathcal{L}_m(x)q_{im}],
\]

(8)

where the Lagrange basis function \( \mathcal{L}_m(x) = \prod_{s=1, s\neq m}^{3} \frac{x-x_s}{x_m-x_n} \).

Then the numerical flux \( \tilde{f}_{i-\frac{1}{2}} \) at the element interface boundary is obtained by an approximate Riemann solver as

\[
\tilde{f}_{i-\frac{1}{2}} = \frac{1}{2} \left[ f_{i-\frac{1}{2}}^L + f_{i-\frac{1}{2}}^R \right] + \frac{1}{2} a \left[ q_{i-\frac{1}{2}}^L - q_{i-\frac{1}{2}}^R \right],
\]

(9)

where \( a = |f'(q_{avg}^L)| \) with \( f'(q) = \frac{\partial f(q)}{\partial q} \) being the characteristic speed. A simple averaging \( q_{avg}^{i-\frac{1}{2}} = \frac{q_{i-\frac{1}{2}}^L + q_{i-\frac{1}{2}}^R}{2} \) is used in the present paper.

Based on the Riemann solver at cell boundaries, the proposed scheme is essentially an upwind type method. As a result, the inherent numerical dissipation is included and stabilizes the numerical
solutions. We did not use any extra artificial viscosity in the shallow water model for the numerical tests presented in the paper.

It is easy to show that the proposed scheme is conservative in terms of the volume-integrated average of each element,

\[ \eta_i = \sum_{m=1}^{3} (w_{im}q_{im}), \quad (10) \]

where the weights \( w_{im} \) are obtained by integrating the Lagrange basis function as

\[ w_{im} = \frac{1}{\Delta x_i} \int_{x_{i-1/2}}^{x_{i+1/2}} L_m(x)dx, \quad (11) \]

and exactly same as those in Guassian quadrature of degree 5.

A direct proof of this observation is obtained by integrating Eq. (3) over the grid element, yielding

\[ \frac{d\eta_i}{dt} = \frac{\partial}{\partial t} (\Delta x_i \eta_i) = \Delta x_i \sum_{m=1}^{3} \left( w_{im} \frac{dq_{im}}{dt} \right) = -\frac{1}{\Delta x} \left( \tilde{f}_{i+1/2} - \tilde{f}_{i-1/2} \right), \quad (12) \]

where \( \Delta x_i \eta_i \) is the total mass within the element \( C_i \).

With the above spatial discretization, Runge–Kutta method is used to solve the following semi-discrete equation (ODE),

\[ \frac{dq_{im}}{dt} = D(q^*), \quad (13) \]

where \( D \) represents the spatial discretisation and \( q^* \) is the dependent variables known at time \( t = t^* \).

A fifth-order Runge–Kutta scheme (Fehlberg, 1958) is adopted in the numerical tests to examine the convergence rate,

\[ q_{im}(t^* + \Delta t) = q_{im}^* + \Delta t \left( \frac{17}{144} d_1 + \frac{25}{36} d_3 + \frac{3}{16} d_5 \right) + \frac{25}{48} d_6, \quad (14) \]

where

\[
\begin{align*}
    d_1 &= D(q^*) \\
    d_2 &= D(q^* + \frac{1}{2}\Delta td_1) \\
    d_3 &= D(q^* + \frac{5}{9}\Delta td_2) \\
    d_4 &= D(q^* + \frac{13}{90}\Delta td_1 + \frac{11}{20}\Delta td_2 - 5\Delta td_3) \\
    d_5 &= D(q^* - \frac{63}{100}\Delta td_1 + \frac{9}{5}\Delta td_2 - \frac{19}{10}\Delta td_3 + \frac{2}{5}\Delta td_4) \\
    d_6 &= D(q^* - \frac{6}{25}\Delta td_1 + \frac{2}{5}\Delta td_2 + \frac{23}{50}\Delta td_3 + \frac{8}{75}\Delta td_4)
\end{align*}
\]

(15)

In other cases, a third-order scheme (Shu, 1988) is adopted to reduce the computational cost, which does not noticeably degrade the numerical accuracy since the truncation errors of the spatial
discretisation are usually dominant. It is written as

\[ q_{im}(t^* + \Delta t) = q_{m,im}^* + \Delta t \left( \frac{1}{6} d_1 + \frac{1}{6} d_2 + \frac{2}{3} d_3 \right), \]  

(16)

where

\[
\begin{aligned}
d_1 &= \mathcal{D}(q^*) \\
d_2 &= \mathcal{D}(q^* + \Delta t d_3) \\
d_3 &= \mathcal{D}(q^* + \frac{1}{3} \Delta t d_1 + \frac{1}{4} \Delta t d_2)
\end{aligned}
\]  

(17)

### 2.2 Spectral analysis and convergence test

We conduct the spectral analysis (Huynh, 2007; Xiao et al., 2013) to theoretically study the performance of GLPCC scheme by considering the following linear equation

\[
\frac{\partial q}{\partial t} + c \frac{\partial q}{\partial x} = 0 \quad (x \in [-\infty, +\infty]) \quad \text{and} \quad c = 1.
\]  

(18)

The linear equation is discretised on an uniform grid with \( \Delta x = 1 \). Since \( c > 0 \) the advection speed is positive, the spatial discretisation for the three DOFs defined in element \( C_i \) involves the six DOFs within elements \( C_i \) and \( C_{i-1} \). Using the proposed scheme, we have the semi-discrete equation as

\[
\begin{bmatrix}
\frac{d}{dt} q_1 \\
\frac{d}{dt} q_2 \\
\frac{d}{dt} q_3
\end{bmatrix} =
\begin{bmatrix}
c_{i-1,11} & c_{i-1,12} & c_{i-1,13} & c_{i,11} & c_{i,12} & c_{i,13} \\
c_{i-1,21} & c_{i-1,22} & c_{i-1,23} & c_{i,21} & c_{i,22} & c_{i,23} \\
c_{i-1,31} & c_{i-1,32} & c_{i-1,33} & c_{i,31} & c_{i,32} & c_{i,33}
\end{bmatrix}
\begin{bmatrix}
q_{i-1,1} \\
q_{i-1,2} \\
q_{i-1,3} \\
q_{i,1} \\
q_{i,2} \\
q_{i,3}
\end{bmatrix}.
\]

and can be written as the following linear combination as

\[
\frac{\partial q_{im}}{\partial t} = - \frac{\partial q}{\partial x}_{im} = \sum_{s=1}^{3} \left( \hat{b}_{i,m,s} q_{i-1,s} \right) + \sum_{s=1}^{3} (b_{i,m,s} q_{i,s}),
\]  

(19)

where the coefficients \( \hat{b}_{i,m,s} \) and \( b_{i,m,s} \) are the coefficients for the DOFs within elements \( C_{i-1} \) and \( C_i \) respectively, which can be obtained by applying the proposed scheme to governing equation Eq. (18) in element \( C_i \).

With a wave solution \( q(x,t) = e^{i\omega x + i\omega t} \), semi-discrete formulation Eq. (19) is written as

\[
q_{i-1,m} = e^{-i\omega \Delta x} q_{im} = e^{-i\omega \Delta x} q_{im},
\]  

(20)
Above spatial discretization can be simplified as

\[
\frac{dq_i}{dt} \frac{\partial q_{im}}{\partial t} = \mathcal{P} - \mathcal{B}_s q_i,
\]

where \( q_i = [q_{i1}, q_{i2}, q_{i3}]^T \) and the components of the \( 3 \times 3 \) matrix \( \mathcal{B}_s \) are coefficients \( B_{i,ms} \) \((m = 1, 2, 3)\), where \( q_i = [q_{i1}, q_{i2}, q_{i3}]^T \) and

\[
\mathcal{P} = \begin{bmatrix}
    c_{i-1,11}e^{-I\omega} + c_{i,11} & c_{i-1,12}e^{-I\omega} + c_{i,12} & c_{i-1,13}e^{-I\omega} + c_{i,13} \\
    c_{i-1,21}e^{-I\omega} + c_{i,21} & c_{i-1,22}e^{-I\omega} + c_{i,22} & c_{i-1,23}e^{-I\omega} + c_{i,23} \\
    c_{i-1,31}e^{-I\omega} + c_{i,31} & c_{i-1,32}e^{-I\omega} + c_{i,32} & c_{i-1,33}e^{-I\omega} + c_{i,33}
\end{bmatrix},
\]

The exact solution. With the wave solution, the exact expression for the spatial discretization of Eq. (18) is

\[
\frac{dq_i}{dt} \frac{\partial q_i}{\partial t} = -I - I\omega q_i.
\]

The numerical property of the proposed scheme can be examined by analysing the eigenvalues of matrix \( \mathcal{P} - \mathcal{B}_s \) in Eq. (23). Truncation errors of the spatial discretization are computed by comparing the principal eigenvalues of matrix \( \mathcal{P} \) and the exact solution \( -I \omega \mathcal{B}_s \) and its exact solution \( -I \omega \mathcal{B}_s \mathcal{P} \). The convergence rate can be approximately estimated by errors for the errors at two different wavenumbers. The results are shown in Table 1 and the fifth-order accuracy is achieved. The spectrum of the eigen matrix of Eq. (23) \( \mathcal{B}_s \) is shown in Fig. 2. A scheme achieves better numerical performance when the hollow circles become closer to imaginary axis. And the maximum of spectral radius determines the largest available CFL number, i.e. a larger spectral radius corresponding to a smaller available CFL number. Numerical dispersion and dissipation relations dominated by the principal eigenvalues are shown in Fig. 3. Numerical properties of several schemes were analyzed in Xiao et al. (2013), shown in their Fig. 1 for spectra and Fig. 2 for numerical dispersion and dissipation relations. The proposed scheme has the following numerical properties as we conduct a comparison between DG3 (Huynh, 2007) and MCV5 (Ii and Xiao, 2009) schemes. Since these three schemes have the fifth-order accuracy and can be derived by FR framework using different constraint conditions for spatial reconstruction of flux functions. As detailed in Huynh (2007), the DG3 scheme uses the Radau polynomial as the correction functions to derive the flux reconstruction which assure the continuity of the numerical
fluxes computed from Riemann solvers at the cell boundaries. MCV5 scheme can be derived by a general framework for flux reconstruction using multi moments proposed in Xiao et al. (2013). MCV5 uses constraint conditions on the point values, first- and second-order derivatives of flux functions at the cell boundaries where Riemann solvers in terms of derivatives of the flux function are required. Compared with DG3 scheme, the proposed scheme is easier to be implemented and thus has less computational overheads. Though MCV5 scheme gives better spectra (eigenvalues are closer to imaginary) than DG3 scheme and the present scheme, it adopts more local DOFs under the same grid spacing, i.e. 4I + 1 DOFs for MCV5 and 3I DOFs for DG3 and the present scheme where I is the total number of elements. Both MCV5 and the present scheme are accelerating up to wavenumber 2π show slightly higher numerical frequency in the high wavenumber regime, which is commonly observed in other spectral-convergence schemes, like DG. Considering the results of the spectral analysis, the proposed scheme is a very competitive framework to build high-order schemes compared with existing advanced methods.

Advection of a smooth sine wave is then computed by GLPCC scheme on a series of refined uniform grids to numerically checking the converge rate. The test case is specified by solving Eq. (18) with initial condition \( q(x, 0) = \sin(2\pi x) \) and periodical boundary condition over \( x \in [0, 1] \).

CFL number of 0.1 is adopted in this example. Normalized \( l_1, l_2 \) and \( l_\infty \) errors and corresponding convergence rate are given in Table 2. Again, the fifth-order convergence is obtained, which agrees with the conclusion in the above spectral analysis.

### 2.3 Extension to system of equations

The proposed scheme is then extended to a hyperbolic system with \( L \) equations in one dimension, which is written as

\[
\frac{\partial q}{\partial t} + \frac{\partial f(q)}{\partial x} = 0, \quad (24)
\]

where \( q \) is the vector of dependent variables and \( f \) the vector of flux functions.

Above formulations can be directly applied to each equation of the hyperbolic system, except that the Riemann problem, which is required at interfaces the cell boundaries between different elements to determine the values of flux functions, is solved for a coupled system of equations.

For a hyperbolic system of equations, the approximate Riemann solver used at interface \( x_{i-\frac{1}{2}} \) is obtained by rewriting Eq. (9) as

\[
f_{i-\frac{1}{2}} = \frac{1}{2} \left[ f_{i-\frac{1}{2}}^L + f_{i-\frac{1}{2}}^R \right] + \frac{1}{2} \beta a \left[ q_{i-\frac{1}{2}}^L - q_{i-\frac{1}{2}}^R \right], \quad (25)
\]

where the vectors \( f_{i-\frac{1}{2}}^L, f_{i-\frac{1}{2}}^R, q_{i-\frac{1}{2}}^L \) and \( q_{i-\frac{1}{2}}^R \) are evaluated by applying the formulations designed for scalar case to each component of the vector. In this paper, we use a simple approximate Riemann solver, the local Lax–Friedrich (LLF) solver, where \( a \) is reduced to a positive real number as

\[
a = \max(|\lambda_1|, |\lambda_2|, \ldots, |\lambda_L|), \quad (26)
\]
\[ \lambda_l (l = 1 \text{ to } L) \] are eigenvalues of matrix \( A \left( q_{i-\frac{1}{2}}^{\text{avg}} \right) \) with \( A(q) = \frac{\partial f(q)}{\partial q} \) and \( q_{i-\frac{1}{2}}^{\text{avg}} = q_{i-\frac{1}{2}}^l + q_{i-\frac{1}{2}}^e. \)

3 Global shallow-water model on cubed-sphere grid

3.1 Cubed-sphere grid

The cubed-sphere grid (Sadourny, 1972), shown in Fig. 4, is obtained by projecting an inscribed cube onto a sphere. As a result, the surface of a sphere is divided into six identical patches and six identical curvilinear coordinates are then constructed. Two kinds of projections are adopted to construct the local curvilinear coordinates, i.e. gnomonic and conformal projections (Rancic et al., 1996). Considering the analytic projection relations and more uniform areas of the computational elements grid spacing, the equiangular gnomonic projection is adopted in the present study. The transformation laws and the projection relations can be referred to Nair et al. (2005a,b) for details. Whereas, a side-effect of this selection choice is that the discontinuous coordinates are found along the boundary edges between adjacent patches. In Chen and Xiao (2008), we have shown that the compact stencils for the spatial reconstructions through using local DOFs are beneficial to suppress the extra numerical errors due to the discontinuous coordinates.

3.2 Global shallow-water model

The local curvilinear coordinate system \((\xi, \eta)\) are shown in Fig. 5, where \(P\) is a point on sphere surface, and \(P'\) is corresponding point on the cube surface through a gnomonic projection. \(\lambda\) and \(\theta\) represent the longitude and latitude. \(\alpha\) and \(\beta\) are central angles spanning from \(-\frac{\pi}{4}\) to \(\frac{\pi}{4}\) for each patch. Local coordinates are defined by \(\xi = R\alpha\) and \(\eta = R\beta\) where \(R\) is the radius of the Earth.

To build a high-order global model, the governing equations are rewritten onto the general curvilinear coordinates. As a result, the numerical schemes developed for Cartesian grid are straightforwardly applied in the computational space. The shallow-water equations are recast on each spherical patch in flux form as

\[
\frac{\partial q}{\partial t} + \frac{\partial e(q)}{\partial \xi} + \frac{\partial f(q)}{\partial \eta} = s(q),
\]

where dependent variables are \(q = \left[ \sqrt{G}h, u, v \right]^T\) with water depth \(h\), co-variant velocity vector \((u, v)\) and Jacobian of transformation \(\sqrt{G}\), flux vectors are \(e = \left[ \sqrt{G}h\hat{u}, g(h + h_s) + \frac{1}{2}(\hat{u}u + \hat{v}v), 0 \right]^T\) in \(\xi\) direction and \(f = \left[ \sqrt{G}h\hat{v}, 0, g(h + h_s) + \frac{1}{2}(\hat{u}u + \hat{v}v) \right]^T\) in \(\eta\) direction with gravitational acceleration \(g\), height of the bottom mountain \(h_s\) and contravariant velocity vector \((\hat{u}, \hat{v})\), source term is \(s = \left[ 0, \sqrt{G}\hat{v} (f + \zeta), -\sqrt{G}\hat{u} (f + \zeta) \right]^T\) with Coriolis parameter \(f = 2\Omega \sin\theta\), rotation speed of the Earth \(\Omega = 7.292 \times 10^{-5} \text{s}^{-1}\) and relative vorticity \(\zeta = \frac{1}{\sqrt{G}} \left( \frac{\partial v}{\partial \xi} - \frac{\partial u}{\partial \eta} \right)\).
The expression of metric tensor $G_{ij}$ can be found in Chen and Xiao (2008), Nair et al. (2005a,b). Jacobian of the transformation is $\sqrt{G} = \sqrt{\det(G_{ij})}$ and the covariant and the contravariant velocity components are connected through

$$
\begin{bmatrix}
\dot{u} \\
\dot{v}
\end{bmatrix} = G^{ij} \begin{bmatrix}
u \\
v
\end{bmatrix},
$$

(28)

where $G^{ij} = (G_{ij})^{-1}$.

Here, taking $\sqrt{G}h$ as the model variable assures the global conservation of total mass. And the total height is used in the flux term. Consequently, the proposed model can easily deal with the topographic source term in a conservative balanced way (Xing and Shu, 2005).

The numerical formulations for two dimensional schemes are easily obtained under the present framework by implementing the one-dimensional GLPCC formulations in $\xi$ and $\eta$ directions respectively as

$$
\left(\frac{\partial q}{\partial t}\right)^\xi = -\frac{\partial (q) (\xi)}{\partial \xi} + \frac{\partial (q) (\eta)}{\partial \eta} + s,
$$

(29)

where

$$
\left(\frac{\partial (q)}{\partial t}\right)^\xi = -\frac{\partial (q)}{\partial \xi} \text{ and } \left(\frac{\partial (q)}{\partial t}\right)^\eta = -\frac{\partial (q)}{\partial \eta},
$$

(30)

are discretised along the grid lines in $\xi$ and $\eta$ directions.

We describe the numerical procedure in $\xi$ direction here as follows. In $\eta$ direction, similar procedure is adopted for spatial discretisation by simply exchanging $\xi$ with $f$ and $\eta$. Considering three DOFs, i.e. $q_{ij1nk}, q_{ij2nk}$ and $q_{ij3nk}$, along the $n$th row ($n = 1$ to $3$) of element $C_{ijk} = \left[\xi_{i-\frac{1}{2}}, \xi_{i+\frac{1}{2}}\right] \times \left[\eta_{j-\frac{1}{2}}, \eta_{j+\frac{1}{2}}\right]$ on patch $k$ (defined at solution points denoted by the hollow circles in Fig. 6), we have the task to solve the following equations

$$
\left(\frac{\partial q_{ijmnk}}{\partial t}\right)^\xi = -\left(\frac{\partial e}{\partial \xi}\right)_{ijmnk}.
$$

(31)

As in one dimensional case, a fourth-order polynomial $E_{ijnk}(x)$ is built for spatial reconstructions of flux functions $e$ to calculate the derivative of $e$ with regard to $\xi$ as

$$
\left(\frac{\partial e}{\partial \xi}\right)_{ijmnk} = \left[\frac{\partial E_{ijnk}(\xi)}{\partial \xi}\right]_{ijmnk},
$$

(32)

where $E(\xi)$ can be obtained by applying the constrained conditions at five constrained points (solid circles in Fig. 6) along the $n$th row of element $C_{ijk}$, which are point-wise values of flux functions $e$ including three from DOFs directly and other two by solving Riemann problems along the $n$th lines of different columns of the adjacent elements.

The LLF approximate Riemann solver is adopted. It means that the parameter $a$ in Eq. (25) reads

$$a = |\tilde{u}| + \sqrt{G^{11}} gh.$$

Details of solving Riemann problem in global shallow water model using governing equations Eq. (27) can be referred to Nair et al. (2005b).
How to set up the boundary conditions along twelve patch boundaries is a key problem to construct a global model on cubed-sphere grid. With the enough information from the adjacent patch, above numerical formulations can be applied on each patch independently. In present study, the values of dependent variables are required to be interpolated from the grid lines in the adjacent patch, for example, as shown in Fig. 7 for the boundary edge between patch 1 and patch 4. When we solve the Riemann problem at point \( P \) on patch 1, \( q^R_P = \left[ \left( \sqrt{Gh} \right)^R_P, u^R_P, v^R_P \right]^T \) is obtained by interpolation along the grid line \( PP_1 \). Whereas, \( q^L_P = \left[ \left( \sqrt{Gh} \right)^L_P, u^L_P, v^L_P \right]^T \) need to be interpolated from the DOFs defined along grid line \( P_4 P \) on patch 4. Since the coordinates on patch 1 and patch 4 is discontinuous at point \( P \), the values of the covariant velocity vector on the coordinate system on patch 4 should be projected to coordinate system on patch 1 and the values of the scalar can be adopted directly. Different from our previous study (Chen and Xiao, 2008), we solve the Riemann problem at patch boundary only in the direction perpendicular to the edge in present study. The parameter \( a \) in Eq. (25) is determined by the contravariant velocity component perpendicular to the edge and the water depth, which are exactly same in two adjacent coordinate systems since the water depth is a scalar independent of coordinate system and the basis vector perpendicular to the edge is continuous between adjacent patches. As a result, solving Riemann problem obtains the same result wherever the numerical procedure is conducted on patch 1 or patch 4. So, no additional corrections are required and the global conservation is guaranteed automatically.

4 Numerical tests

Representative benchmark tests, three from Williamson’s standard test cases (Williamson et al., 1992) and one introduced in Galewsky et al. (2004), are checked in this section to verify the performance of the proposed global shallow water model. All measurements of errors are defined following Williamson et al. (1992).

4.1 Williamson’s standard case 2: steady-state geostrophic flow

A balanced initial condition is specified in case by using a height field as

\[
gh = gh_0 - \left( R \Omega u_0 + \frac{u_0^2}{2} \right) (- \cos \lambda \cos \theta \sin \gamma + \sin \theta \cos \gamma)^2
\]

(33)

where \( gh_0 = 2.94 \times 10^4 \), \( u_0 = 2\pi R/(12 \text{ days}) \) and the parameter \( \gamma \) represents the angle between the rotation axis and polar axis of the Earth, and a velocity field (velocity components in longitude/latitude grid \( u_\lambda \) and \( u_\theta \)) as

\[
\begin{align*}
  u_\lambda &= u_0 \left( \cos \theta \cos \gamma + \sin \theta \cos \lambda \sin \gamma \right) \\
  u_\theta &= -u_0 \sin \lambda \sin \gamma
\end{align*}
\]

(34)

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As a result, both height and velocity fields should keep unchanging during integration. Additionally, the height field in this test case is considerably smooth. Thus we run this test on a series of refined grids to check the convergence rate of GLPCC global model. The results of \( l_1, l_2 \) and \( l_\infty \) errors and convergence rates are given in Table 3. After extending the proposed high-order scheme to the spheric geometry through the application of the cubed-sphere grid, the original fifth-order accuracy as shown in one-dimensional simulations and spectral analysis preserved in this test. Numerical results of height fields and absolute errors are shown in Fig. 8 for tests on a series of refined grids are checked. As above case, to compare with our former fourth-order model this test case is checked on grid \( G_{32} \) (with similar number of DOFs, 256 DOFs along the equator) in Chen and Xiao (2008). The influence of patch boundaries on the numerical results can be found in the plots of the absolute errors. The distributions of absolute errors can reflect the locations of patch boundaries, especially in the flow with \( \gamma = 0 \).

### 4.2 Williamson’s standard case 5: zonal flow over an isolated mountain

The total height and velocity field in this case is same as above case 2 with \( \gamma = 0 \), except \( h_0 = 5960 \text{ m} \) and \( u_0 = 20 \text{ m s}^{-1} \). A bottom mountain is specified as

\[
h_s = h_{s0} \left( 1 - \frac{r}{r_0} \right),
\]

where \( h_{s0} = 2000 \text{ m} \), \( r_0 = \frac{\pi}{2} \) and \( r = \min \left[ r_0, \sqrt{(\lambda - \lambda_c)^2 + (\theta - \theta_c)^2} \right] \).

This test is adopted to check the performance of a shallow water model to deal with a topographic source term. We run this test on a series of refined grid \( G_6, G_{12}, G_{24} \) and \( G_{48} \). Numerical results of height fields are shown in Fig. 9 for total height field of the test on grid \( G_{12} \) at day 5, 10 and 15, which agree well with the spectral transform solutions on T213 grid (Jakob-Chien et al., 1995). Furthermore, the oscillations occurring at boundary of bottom mountain observed in spectral transform solutions are completely removed through the conservative treatment of a numerical scheme to the spheric geometry through the application of the cubed-sphere grid, the original fifth-order accuracy as shown in one-dimensional simulations and spectral analysis preserved in this test. The conservation errors of total energy and enstrophy are shown in Fig. 10. The conservation errors for total energy (left panel) and potential enstrophy (right panel) of tests on a series of refined grids are checked. As above case, to compare with our former fourth-order model this test case is checked on grid \( G_{20} \) having the similar DOFs on former \( 32 \times 32 \times 6 \) grid. The conservation errors are \(-9.288 \times 10^{-7}\) for total energy and \(-1.388 \times 10^{-5}\) for potential energy.
enstrophy and much smaller than those by fourth-order model in Chen and Xiao (2008).

4.3 Williamson’s standard case 6: Rossy–Haurwitz wave

Rossby–Haurwitz wave case checks a flow field including the phenomena of a large range of scales. As a result, the high-order schemes are always preferred to better capture the evolution of small scales. The spectral transform solution on fine T213 grid given by Jakob-Chien et al. (1995) is widely accepted as the reference solution to this test due to the good capability of spectral method to reproduce the behaviour of small scales. Numerical results of height fields by GLPCC model are shown in Fig. 12 for tests on grids $G_{12}$ and $G_{24}$ at day 7 and 14. At day 7, no obvious difference is observed between the solutions on different grids and both agree well with the reference solution. At day 14, obvious differences are found on different grids. Eight circles of 8500 m exist in the result on coarser grid $G_{12}$, which are also found in the spectral transform solution on T42 grid, but not in the results on finer grid $G_{24}$ by GLPCC model and the spectral transform ones on T63 and T213 grids. Additionally, the contour lines of 8100 m exists in spectral transform solution on T213 grid, but not in present results and spectral transform ones on T42 and T63 grids. According to the analysis in Thuburn and Li (2000), this is due to the less inherent numerical viscosity on finer grid. As in case 5, total mass is conserved to the machine precision as shown in Fig. 13 and the conservation errors for total energy and potential enstrophy are given in Rossby–Haurwitz test are also shown in Fig. 14 for tests with different resolutions. Total energy error of $-6.131 \times 10^{-6}$ and potential enstrophy error of $-1.032 \times 10^{-5}$ are obtained by the present model running on grid $G_{20}$, which are smaller than those by our fourth-order model on $32 \times 32 \times 6$ grid (Chen and Xiao, 2008). This test was also checked in Chen et al. (2014a) by a third-order model (see their Fig. 19(c) and (d)), where much more DOFs (nine times than those on grid $G_{24}$) are adopted to obtain a result without eight circles of 8500 m at day 14. It reveals a well accepted observation that a model of higher order converges faster to the reference solution, and should be more desirable in the atmospheric modelling. High-order accuracy is very beneficial to simulating this test and the atmospheric dynamics.

4.4 Barotropic instability

A barotropic instability test was proposed in Galewsky et al. (2004). Two kinds of setup setups of this test are usually checked in literatures, i.e. the balanced setup and unbalanced setup. The balanced setup is same as Williamson’s standard case 2, except the water depth changes with much larger gradient within a very narrow belt zone. This test is of special interest for global models on the cubed-sphere grid, since that narrow belt zone is located along the boundary edges between patch 5 and patches 1, 2, 3, and 4. Extra numerical errors near boundary edges would easily pollute the numerical results. In practice, 4-wave four-wave pattern errors may dominate the simulations on the coarse grids. For this case, we run the proposed model on a series of refined grids. By
checking the convergence of the numerical results, we can figure out if the extra numerical errors generated by discontinuous coordinates can be suppressed by the proposed models with the increasing resolution. The unbalanced setup introduces a small perturbation to the height field. Thus, the balanced condition can not be preserved and the flow will evolve to a very complex pattern. Exact solution does not exist for unbalanced setup and a spectral transform solution on T341 grid to this case given in Galewsky et al. (2004) at day 6 is widely adopted as reference solution. The details of setup of this test can be referred to Galewsky et al. (2004).

4.4.1 Balanced setup

We test the balanced setup at first. The proposed model runs on two grids with different resolutions of $G_{24}$ and $G_{72}$. Numerical results of water depth after integrating for 5 days are shown in Fig. 15 and evolution of normalized $l_1$ errors of water depth of two simulations are depicted in Fig. 16. On a coarse grid with $G_{24}$, the numerical result is dominated by four-wave pattern errors and the balanced condition can not be preserved in simulation. The accuracy is obviously improved by increasing the resolution using grid $G_{72}$. The numerical result of height field at day 5 is visually identical to the initial condition. The improvement of the accuracy can be also proven by checking the velocity component $u_\theta$. Numerical results of $u_\theta$, which keeps zero in exact solution, vary within a range of ±31 $\text{m s}^{-1}$ on grid $G_{24}$ and are much smaller range of ±0.8 $\text{m s}^{-1}$ on grid $G_{72}$. This test is more challenging for cubed-sphere grid than other quasi-uniform spherical grids, e.g. Yin–Yang grid and icosahedral grid. As shown in Fig. 16, at very beginning of the simulation the $l_1$ errors increase to a magnitude of about $10^{-4}$ on coarse grid $G_{24}$ and this character does not change with the grid resolution on refined grid $G_{72}$. This evolution pattern of $l_1$ errors are different from those of models on Yin–Yang and icosahedral grids, where initial startup errors also decrease on fine grids as shown in Chen et al. (2014a, Fig. 23).

4.4.2 Unbalanced setup

We run the unbalanced setup on a series of refined grids to check if the numerical result will converge to the reference solution on fine grid refined grids. Numerical results for relative vorticity field after integrating the proposed model for 6 days are shown in Fig. 17. Shown are the results on four grids with gradually refined resolutions of $G_{24}$, $G_{48}$, $G_{72}$ and $G_{96}$. On grid $G_{24}$, the structure of numerical result is very different from the reference solution. After refining the grid resolution, the result is improved on grid $G_{48}$. Except the structure in top-left corner, it looks very similar to the reference solution. On grid $G_{72}$ and $G_{96}$, numerical results agree with the reference solution very well and there is no obvious difference between these two contour plots. Compared with the results of our former fourth-order model, the contour lines look slightly less smooth. Similar results are found in the spectral transform reference solution. Since this test contains more significant gradients in the solution, a high-order scheme might need some extra numerical dissipation to remove the
noise around the large-gradients. Increasing the grid solution can effectively reduce the magnitude of the oscillations as shown in the present simulation.

5 Conclusions

In this paper, a three-point high-order GLPCC scheme is proposed under the framework of flux reconstruction. Three local DOFs are defined within each element at Gauss–Legendre points and a super convergence of fifth order is achieved. This single-cell based method shares the advantages with the DG and SE methods, such as high-order accuracy, grid flexibility, global conservation and high scalability for parallel processing. Meanwhile, it is much simpler and easier to implement. With the application of the cubed-sphere grid, the global shallow water model has been constructed using GLPCC scheme. Benchmark tests are checked by using the present model, and promising results reveal that it is a potential framework to develop high-performance general circulation models for atmospheric and oceanic dynamics. As any high-order numerical scheme, additional dissipation or limiter projection might be needed in simulations of real case applications. Because of the algorithmic similarity, the existing works on high-order limiting projection and artificial dissipation devised for DG or SE methods are applicable to GLPCC without substantial difficulty. Also it is an important future study to design more reliable limiting projection formulations for GLPCC and other FR schemes, which are able to deal with discontinuities without losing the overall high-order accuracy.

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References


Fig. 1. Configuration of DOFs and constrained constraint conditions in one dimensional case.

Fig. 2. The spectrum of the semi-discrete scheme.

Fig. 3. Numerical dispersion (left) and dissipation (right) relations of the semi-discrete scheme.
Fig. 4. The cubed-sphere grid.

Fig. 5. The gnomonic projection.
Fig. 6. Configuration of DOFs and constrained conditions in two dimensional case.

Fig. 7. Riemann problem along patch boundary edge between patch 1 and 4.
Fig. 8. Numerical results and absolute errors of water depth for case 2 on grid $G_{12}$ at day 5. Shown are water depth (top-left) and absolute error (top-right) of the flow with $\gamma = 0$ and water depth (bottom-left) and absolute error (bottom-right) of the flow with $\gamma = \frac{\pi}{4}$.

Fig. 9. Numerical results of total height field for case 5 on grid $G_{12}$ at day 5 (top-left), day 10 (top-right) and day 15 (bottom).
Fig. 10. Normalized conservation error of total mass on grid $G_{12}$ for case 5.

Fig. 11. Normalized conservation errors of total energy and potential enstrophy on refined grids for case 5.
Fig. 12. Numerical results of water depth for Rossby–Haurwitz–wave test case 6 on grid $G_{12}$ at day 7 (top-left), day 14 (top-right) and on grid $G_{24}$ at day 7 (bottom-left) and day 14 (bottom-right).

Fig. 13. Normalized conservation error of total mass on grid $G_{12}$ for case 6.
Fig. 14. Normalized conservation errors of total energy and potential enstrophy on refined grids for Rossby–Haurwitz wave test case 6.

Fig. 15. Numerical results of water depth for balanced setup of barotropic instability test on two grids $G_{24}$ (left) and $G_{72}$ (right). Contour lines vary from 9000 m to 10 100 m.
Fig. 16. Normalized $l_1$ error of water depth for balanced setup of barotropic instability test on two grids.
Fig. 17. Numerical results of relative vorticity for unbalanced setup of barotropic instability test on a series of refined grids. Contour lines vary from $-1.1 \times 10^{-4}$ to $-0.1 \times 10^{-4}$ by dashed lines and $0.1 \times 10^{-4}$ to $1.5 \times 10^{-4}$ by solid lines.
**Table 1.** Numerical errors at two wavenumbers and corresponding convergence rate.

<table>
<thead>
<tr>
<th>Wavenumber</th>
<th>$\omega = \frac{\pi}{8}$</th>
<th>$\omega = \frac{\pi}{4}$</th>
<th>Order</th>
</tr>
</thead>
<tbody>
<tr>
<td>Error</td>
<td>$-3.1408 \times 10^{-5}$</td>
<td>$-1.0466 \times 10^{-7}$</td>
<td>4.97</td>
</tr>
</tbody>
</table>

**Table 2.** Numerical errors and convergence rates for advection of a sine wave.

<table>
<thead>
<tr>
<th>Resolution</th>
<th>$l_1$ error</th>
<th>$l_2$ error</th>
<th>$l_\infty$ error</th>
<th>Order</th>
</tr>
</thead>
<tbody>
<tr>
<td>$I = 4$</td>
<td>$3.9392 \times 10^{-3}$</td>
<td>$3.9623 \times 10^{-3}$</td>
<td>$3.9702 \times 10^{-3}$</td>
<td>$-5.9623 \times 10^{-3}$</td>
</tr>
<tr>
<td>$I = 8$</td>
<td>$1.5683 \times 10^{-4}$</td>
<td>$1.4841 \times 10^{-4}$</td>
<td>$1.3396 \times 10^{-4}$</td>
<td>$4.89$</td>
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<tr>
<td>$I = 16$</td>
<td>$5.3627 \times 10^{-6}$</td>
<td>$4.8431 \times 10^{-6}$</td>
<td>$4.1707 \times 10^{-6}$</td>
<td>$5.01$</td>
</tr>
<tr>
<td>$I = 32$</td>
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<td>$1.5327 \times 10^{-7}$</td>
<td>$1.3293 \times 10^{-7}$</td>
<td>$4.97$</td>
</tr>
<tr>
<td>$I = 64$</td>
<td>$5.3017 \times 10^{-9}$</td>
<td>$4.8092 \times 10^{-9}$</td>
<td>$4.1670 \times 10^{-9}$</td>
<td>$5.00$</td>
</tr>
</tbody>
</table>

**Table 3.** Numerical errors and convergence rates for case 2 with flow in north-east direction (with $\gamma = \frac{\pi}{4}$).

<table>
<thead>
<tr>
<th>Grid</th>
<th>$l_1$ error</th>
<th>$l_2$ error</th>
<th>$l_\infty$ error</th>
<th>Order</th>
</tr>
</thead>
<tbody>
<tr>
<td>$G_6$</td>
<td>$3.394 \times 10^{-5}$</td>
<td>$5.492 \times 10^{-5}$</td>
<td>$1.868 \times 10^{-4}$</td>
<td>$-4.79$</td>
</tr>
<tr>
<td>$G_{12}$</td>
<td>$1.440 \times 10^{-6}$</td>
<td>$2.321 \times 10^{-6}$</td>
<td>$8.924 \times 10^{-6}$</td>
<td>$4.39$</td>
</tr>
<tr>
<td>$G_{24}$</td>
<td>$5.367 \times 10^{-8}$</td>
<td>$8.317 \times 10^{-8}$</td>
<td>$3.457 \times 10^{-7}$</td>
<td>$4.69$</td>
</tr>
<tr>
<td>$G_{48}$</td>
<td>$1.942 \times 10^{-9}$</td>
<td>$2.957 \times 10^{-9}$</td>
<td>$1.487 \times 10^{-8}$</td>
<td>$4.54$</td>
</tr>
</tbody>
</table>